

ACCEPTED MANUSCRIPT • OPEN ACCESS

Roadmap on Advancements of the FHI-aims Software Package

To cite this article before publication: Volker Blum *et al* 2026 *Electron. Struct.* in press <https://doi.org/10.1088/2516-1075/ae8067>

Manuscript version: Accepted Manuscript

Accepted Manuscript is “the version of the article accepted for publication including all changes made as a result of the peer review process, and which may also include the addition to the article by IOP Publishing of a header, an article ID, a cover sheet and/or an ‘Accepted Manuscript’ watermark, but excluding any other editing, typesetting or other changes made by IOP Publishing and/or its licensors”

This Accepted Manuscript is © 2026 The Author(s). Published by IOP Publishing Ltd.



As the Version of Record of this article is going to be / has been published on a gold open access basis under a CC BY 4.0 licence, this Accepted Manuscript is available for reuse under a CC BY 4.0 licence immediately.

Everyone is permitted to use all or part of the original content in this article, provided that they adhere to all the terms of the licence <https://creativecommons.org/licenses/by/4.0>

Although reasonable endeavours have been taken to obtain all necessary permissions from third parties to include their copyrighted content within this article, their full citation and copyright line may not be present in this Accepted Manuscript version. Before using any content from this article, please refer to the Version of Record on IOPscience once published for full citation and copyright details, as permissions may be required. All third party content is fully copyright protected and is not published on a gold open access basis under a CC BY licence, unless that is specifically stated in the figure caption in the Version of Record.

View the [article online](#) for updates and enhancements.

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

Roadmap on Advancements of the
FHI-aims Software Package

May 29, 2026

Accepted Manuscript

*Roadmap on Advancements of the FHI-aims Software Package***Authors:**

Joseph W. Abbott, Carlos Mera Acosta, Alaa Akkoush, Alberto Ambrosetti, Viktor Atalla, Alexej Bagrets, Jörg Behler, Daniel Berger, Hannah Bertschi, Björn Bieniek, Jonas Björk, **Volker Blum***, Saeed Bohloul, Connor L. Box, Nicholas Boyer, Danilo Simoes Brambila, Gabriel A. Bramley, Kyle R. Bryenton, María Camarasa-Gómez, Christian Carbogno, Fabio Caruso, Sucismita Chutia, Michele Ceriotti, Gábor Csányi, William Dawson, Francisco A. Delesma, Fabio Della Sala, Bernard Delley, Robert A. DiStasio Jr., Maria Dragoumi, Sander Driessen, Marc Dvorak, Simon Erker, Ferdinand Evers, Eduardo Fabiano, Matthew R. Farrow, Florian Fiebig, Jakob Filser, Lucas Foppa, Lukas Gallandi, Alberto Garcia, Ralf Gehrke, Simiam Ghan, Luca M. Ghiringhelli, Mark Glass, Stefan Goedecker, Dorothea Golze, Matthias Gramzow, James A. Green, Andrea Grisafi, Andreas Grüneis, Jan Günzl, Stefan Gutzeit, Samuel J. Hall, Felix Hanke, Ville Havu, Xingtao He, Joscha Hekele, Olle Hellman, Uthpala Herath, Jan Hermann, Daniel Hernangómez-Pérez, Oliver T. Hofmann, Johannes Hoja, Simon Hollweger, Lukas Hörmann, Ben Hourahine, Wei Bin How, William P. Huhn, Marcel Hülsberg, Timo Jacob, Sara Panahian Jand, Hong Jiang, Erin R. Johnson, Werner Jürgens, J. Matthias Kahk, Yosuke Kanai, Kisung Kang, Petr Karpov, Elisabeth Keller, Roman Kempt, Danish Khan, Matthias Kick, Benedikt P. Klein, Jan Kloppenburg, Alexander Knoll, Florian Knoop, Franz Knuth, Simone S. Köcher, Jannis Kockläuner, **Sebastian Kokott***, Thomas Körzdörfer, Hagen-Henrik Kowalski, Peter Kratzer, Pavel Kůs, Raul Laasner, Bruno Lang, Björn Lange, Marcel F. Langer, Ask Hjorth Larsen, Hermann Lederer, Susi Lehtola, Maja-Olivia Lenz-Himmer, Moritz Leucke, Sergey Levchenko, Alan Lewis, O. Anatole von Lilienfeld, Konstantin Lion, Werner Lipsunen, Johannes Lischner, Yair Litman, Chi Liu, Qing-Long Liu, Songrui Liu, Andrew J. Logsdail, Michael Lorke, Zekun Lou, Iuliia Mandzhieva, Andreas Marek, Johannes T. Margraf, Reinhard J. Maurer, Tobias Melson, Florian Merz, Jörg Meyer, Georg S. Michelitsch, Teruyasu Mizoguchi, Evgeny Moerman, Dylan Morgan, Jack Morgenstein, Jonathan Moussa, Akhil S. Nair, Lydia Nemeč, Harald Oberhofer, Alberto Otero-de-la-Roza, Ramón L. Panadés-Barrueta, Thanush Patlolla, Mariia Pogodaeva, Alexander Pöpl, Alastair J. A. Price, Thomas A. R. Purcell, Jingkai Quan, Nathaniel Raimbault, Markus Rampp, Karsten Rasim, Ronald Redmer, Xinguo Ren, Karsten Reuter, Norina A. Richter, Stefan Ringe, Patrick Rinke, Simon P. Rittmeyer, Herzain I. Rivera-Arrieta, Matti Ropo, **Mariana Rossi***, Victor Ruiz, Nikita Rybin, Andrea Sanfilippo, **Matthias Scheffler***, Christoph Scheurer, Christoph Schober, Franziska Schubert, Tonghao Shen, Christopher Shepard, Honghui Shang, Kiyoo Shibata, Andrei Sobolev, Ruyi Song, Aloysius Soon, Daniel T. Speckhard, Pavel V. Stishenko, Elia Stocco, Muhammad N. Tahir, Izumi Takahara, Jun Tang, Zechen Tang, Thomas Theis, Franziska Theiss, Alexandre Tkatchenko, Milica Todorović, George Trenins, Oliver T. Unke, Álvaro Vázquez-Mayagoitia, Oscar van Vuren, Daniel Waldschmidt, Han Wang, Yanyong Wang, Jürgen Wieferink, Jan Wilhelm, Scott Woodley, Jianhang Xu, Yong Xu, Yi Yao, Yingyu Yao, Mina Yoon, Victor Wen-zhe Yu, Zhenkun Yuan, Marios Zacharias, Igor Ying Zhang, Min-Ye Zhang, Wentao Zhang, Xingchen Zhang, Rundong Zhao, Shuo Zhao, Ruiyi Zhou, Yuanyuan Zhou, Tong Zhu

(*) journal guest editor

Contents

1	Introduction	5
1.1	Introduction and Overview	6
2	Basic Concepts and Numerics	11
2.1	Numeric Atom-Centered Orbital Basis Sets for DFT and Beyond	11
2.2	Technical Underpinnings for All-Electron Density Functional Theory with Numeric Atom-centered Orbitals	22
2.3	Relativistic Effects: Scalar Relativity, Spin-Orbit Coupling, and Four-Component Relativity	32
2.4	Solvers for Large-Scale Electronic Structure Theory: ELPA and ELSI	40
2.5	Electronic-Structure Analysis Tools in Numeric Atomic Orbital Framework	49
2.6	Explicit and Implicit Embedding Approaches	57
3	Exchange and Correlation	63
3.1	Exchange and Correlation for Density Functional Theory and the DFT+U Approach	64
3.2	Hybrid Density Functionals for Large Scale Simulations	71
3.3	Density-Functional Based Methods for Dispersion Interactions	78
3.4	Random Phase Approximation and Beyond for Total Energies and Forces	83
4	Electronic Excited States	91
4.1	Coupled-Cluster Theory for the Ground State and for Excitations	92
4.2	All-Electron GW for Finite Systems	98
4.3	GW Calculations for Periodic Systems	106
4.4	Neutral Excited States: Casida Equation for Linear-Response Time-Dependent Density Functional Theory and Bethe-Salpeter Equation	112
4.5	Real-Time Time-Dependent Density Functional Theory	118
4.6	All-Electron Δ -SCF Calculations in Molecular and Periodic Systems	124

Roadmap on Advancements of the FHI-aims Software Package

CONTENTS

5	Linear Response Methods	129
5.1	Density-Functional Perturbation Theory with Numeric Atom-Centered Orbitals	130
5.2	Nuclear Magnetic Resonance with Numeric Atom-Centered Basis Sets	136
5.3	Polarization, Born Effective Charges, and Topological Invariants	143
6	Vibrations and Dynamics	149
6.1	Phonons, Anharmonicity Quantification, and Strongly Anharmonic Vibrations	150
6.2	Performing Advanced Ab Initio Molecular Dynamics	157
6.3	Electronic Structure Calculations with Quantum Nuclei	164
6.4	Electron-Phonon Coupling and Electronic Friction	171
6.5	Temperature-Dependent Electronic Spectral Functions from Band-Structure Unfolding	175
7	Transport	181
7.1	Vibrational Heat Transport in Weakly, Moderately, and Strongly Anharmonic Solids	182
7.2	Electrical Transport for Anharmonic Materials	187
7.3	Molecular Transport	193
7.4	Computing Diabatic Couplings	198
8	Workflows and AI	203
8.1	Code-Agnostic Materials Science Libraries, Workflow Managers, and Visualization Tools	204
8.2	Workflows for Artificial Intelligence	209
8.3	Machine Learning the Electronic Structure in a Real-Space and Atom-Centered Framework	215
9	Appendix	223
9.1	Summary of the Delta-DFT test for the FHI-aims species defaults	223

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

1 Introduction

Accepted Manuscript

1.1 Introduction and Overview

Volker Blum^{1,2}, Sebastian Kokott^{3,4}, Mariana Rossi⁵, and Matthias Scheffler⁴

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin Germany

⁴The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

Abstract

Electronic-structure theory is the foundation of the description of materials including multiscale modeling of their properties and functions. Obviously, without sufficient accuracy at the base, reliable predictions are unlikely at any level that follows. The software package FHI-aims has proven to be a game changer for accurate free-energy calculations because of its scalability, numerical precision, and its efficient handling of density functional theory (DFT) with hybrid functionals and van der Waals interactions. It treats molecules, clusters, and extended systems (solids and liquids) on an equal footing. Besides DFT, FHI-aims also includes quantum-chemistry methods, descriptions for excited states and vibrations, and calculations of various types of transport. Recent advancements address the integration of FHI-aims into an increasing number of workflows and various artificial intelligence (AI) methods. This *Roadmap* describes the state-of-the-art of FHI-aims and advancements that are currently ongoing or planned.

Introduction

The Fritz Haber Institute *ab initio* materials simulation (FHI-aims) software project is an accurate and precise electronic-structure and molecular dynamics software that treats molecules, clusters, and extended systems (solids and liquids) on an equal footing. Here the terms "accurate" and "precise" refer to the basic theoretical concepts and employed numerical methods, respectively. The FHI-aims project was conceived in 2003 at the Fritz Haber Institute of the Max Planck Society. FHI-aims has since grown into a large global community. The software is being developed and advanced by numerous researchers without whom it would not exist the way it does. This is exemplified by the 32 topical contributions and the 201 authors featured in this *Roadmap*, highlighting the breadth and diversity of expertise within the FHI-aims ecosystem.

This *Roadmap* describes the state-of-the-art of the FHI-aims software package and its ongoing advancements and the plans for the near future. Detailed discussions on the background mathematics and physics are referred to the original publications or to other, topically focused roadmaps [1-3]. In addition to descriptions of the FHI-aims capabilities, most contributions also sketch and provide links to practical tutorials. In short, the FHI-aims software is described in terms of three topical bullets:

- Versatile: FHI-aims serves for the description of molecules, clusters, nanostructures, surfaces, solids, and liquids. It addresses the electronic structure as well as vibrations and (long-time) molecular dynamics. It is also links to state-of-the-art AI tools.

Roadmap on Advancements of the FHI-aims Software Package

1.1. INTRODUCTION AND OVERVIEW

- Precise and Accurate: FHI-aims employs DFT with highly efficient implementations of advanced exchange-correlation functionals, many-body methods (GW approximation and Bethe-Salpeter equation) and quantum chemistry methods, such as Møller-Plesset perturbation and coupled-cluster theories. The latter also includes the equation-of-motion approach for excited states.
- Scalable: FHI-aims runs on laptops, and for larger systems and long-time molecular dynamics it runs efficiently on highest-performance computers.

The *Roadmap* covers all these aspects, and more. Besides this 1-Introduction, it is organized into seven further chapters: 2-Basic Concepts and Numerics, 3-Exchange and Correlation, 4-Electronic Excited States, 5-Linear Response Methods, 6-Vibrations and Dynamics, 7-Transport, and 8-Workflows and AI. Each of these chapters contains between 3 and 6 contributions, covering different subtopics. Let give a brief summary of the various chapters.

Chapter 2 introduces the foundational methodological concepts employed in FHI-aims. Numerically tabulated atom-centered orbital (NAO) basis sets are employed to mathematically express the electronic structure. The first roadmap segment covers the details of this choice, which is central to the code's ability to achieve high numerical precision while remaining affordable up to very large system sizes. The technical choices (integration grids, electrostatic potential, forces, stresses, structure optimization) necessary to implement electronic structure theory based on NAOs efficiently are covered in a separate segment. Next, approaches to include relativistic effects (scalar, spin-orbit coupled or four-component) are central for the physical accuracy of computational predictions by electronic structure theory. For computational efficiency, linear-algebra based solutions of the self-consistent field eigenvalue problem or the density matrix are essential and are handled by open-source libraries, including the Eigenvalue solvers for Petaflop Applications (ELPA) library and the ELectronic Structure Infrastructure (ELSI) interface, which integrates this and other approaches into FHI-aims as a dedicated software layer. Finally, the wealth of data acquired in a single electronic structure calculation is immense and appropriate tools to analyze and understand this data are discussed. DFT represents a most important practical concept. However, it only exists in terms of approximations to the exchange and correlation (xc) functional. [4]

Chapter 3 addresses the various levels of approximations, starting from the local-density approximation and Perdew's generalized gradient approximation, which is the currently widely favored approach, and comes in several flavors. For several applications the self-interaction error of these functionals can be critical. Here the DFT+U method can heal part of the problem, in some cases. A significant advancement of FHI-aims concerns the recently improved implementation of the next level of the treatment of xc, namely hybrid functionals. Here FHI-aims achieves top-notch efficiency. All the mentioned functionals miss the long-range dispersion (van der Waals) interactions which can be critical for a reliable description, e.g. of organic systems. FHI-aims offers several options for adding vdW contributions to the standard xc functionals. The highest level xc treatment in DFT, close to "heaven of xc functionals" is the random-phase approximation (RPA), for which FHI-aims offers total energies and also forces. In fact, FHI-aims also offers quantum-chemistry methods such as coupled-cluster theory. However, this is discussed in the next chapter because it also includes the coupled-cluster equation-of-motion treatment to describe excited states. A special contribution of this chapter deals with explicit and implicit embedding approaches that allows the use of high-level first-principles methods in a reduced region, which is embedded in a coarse-grained environment. This technique is especially relevant for applications in biochemistry and homo- and heterogeneous catalysis.

Chapter 4 reports numerous approaches for calculating electronic excited states. It covers coupled-cluster theory (including the equation of motion), the GW approach, time-dependent DFT (also in its real-time

Roadmap on Advancements of the FHI-aims Software Package

1.1. INTRODUCTION AND OVERVIEW

formulation), and the Delta-SCF method. These implementations cover neutral excitations as well as ionization and electron affinities, which means charged excitations. As in essentially all FHI-aims contributions, the methods enable the treatment of core levels and valence bands and they apply to finite systems as well as to extended solids and liquids. Let us just mention real-time time-dependent DFT, because this is a less common concept. Such simulations can address important questions, especially pertaining to non-equilibrium electron dynamic phenomena, including interfacial charge transfer, atom-cluster collisions, topological quantum matter, etc. Furthermore, Ehrenfest dynamics can be performed on top of real-time time-dependent DFT so that non-adiabatic effects of electrons can be taken into account in first-principles molecular dynamics simulations.

Chapter 5 covers the implementation of methods that are capable of describing the response of the electronic density to external stimuli, in a linear-response regime. Contribution 5.1 summarizes the implementation of density-functional perturbation theory (DFPT) for electric-field and nuclear displacement perturbations. This gives access to polarizabilities and optical dielectric constants of materials, which enter the calculation of several kinds of vibrational spectroscopy (Raman, sum-frequency generation, second-harmonic generation, etc.), as well as the quantities needed for the calculation of electron-phonon and non-adiabatic coupling terms, further detailed in Chapter 6. Contribution 5.2 details the implementation of DFPT for a magnetic-field perturbation, which provides the necessary ingredients for the calculation of NMR shifts. Finally, contribution 5.3 describes the implementation and gives practical guidelines for the calculation of the polarization in periodic systems, including derived quantities such as Born-effective charges and topological invariants. With the methods presented in this chapter, it is possible to assess a wide range of material response properties.

Complementing and enhancing the topics discussed in the previous chapters, Chapter 6 deals with nuclear motion and its impact on electronic properties. Contribution 6.1 centers on the description of FHI-vibes, a package associated with FHI-aims that interfaces to several other packages and workflows for vibrational motion and ab initio molecular dynamics (aiMD). It allows the computation of harmonic, perturbative, and fully anharmonic nuclear motion in several contexts. Contribution 6.2 describes several ways of performing aiMD with FHI-aims and highlights its use in the grand-canonical replica exchange method, which allows one to simulate material phases with fluctuating density or stoichiometry. Contribution 6.3 describes two efficient methods to include the quantum nature of nuclei in atomistic simulations. The nuclear-electronic orbital method, which is implemented natively and can approximate adiabatic and non-adiabatic regimes, and the path-integral molecular dynamics method, which is exact in the adiabatic approximation and can be performed through the connection of FHI-aims with i-PI. Contribution 6.4 looks at the coupling between atomic motion and electronic properties. It describes the computation of electron-phonon coupling and its use in the computation of nuclear motion in metallic environments, where dissipative dynamics that can be described in an electronic friction formalism is possible. In Contribution 6.5, electron-phonon coupling in the adiabatic regime is also addressed by showing how to perform electronic band-structure unfolding with numeric atom-centered orbitals and calculating temperature-dependent band-structures from thermally-displaced structures.

Chapter 7 deals with various aspects and forms of transport: vibrational heat transport, electrical transport of electrons or holes in semiconductors, and transport through single molecules and nanoscale systems where the molecule is positioned between two electrodes. With respect to the description of vibrational heat transport and electrical transport, FHI-aims features the most accurate methods considering the full statistical mechanics and all degrees of anharmonicities. The chapter also discusses perturbative methods that start from the phonon picture and are limited to largely harmonic materials. The more advanced methods employed by FHI-aims (Green-Kubo and Kubo-Greenwood) are particularly relevant for thermal

Roadmap on Advancements of the FHI-aims Software Package

1.1. INTRODUCTION AND OVERVIEW

insulators and thermoelectric materials. The method for molecular and nanoscale transport studies enables the treatment of a wide range of quantum phenomena, including quantum interference, non-equilibrium spin-crossover, and much more. Electronic couplings between electronic states localized in different components of a system, often also called transfer-integrals, are a useful tool to understand the efficiency of charge transport. The implementation which allows their calculation is described in the last contribution of this chapter.

Chapter 8 covers Workflows and AI. It starts with a description of efficient tools and infrastructure for creating and managing large datasets, visualizing and analyzing diverse data, and assessing the quality of simulation results. A wide array of tools is described, covering developments targeted mostly at FHI-aims, such as the Graphical Interface for Materials Simulations (GIMS), as well as more general community tools that can be efficiently used with FHI-aims. Important recent developments concern workflows that automate the AI guided acquisition of data, that perform AI-guided exploration of materials space, and that offer an AI-guided physico-chemical analysis of data. For example, the creation of machine-learning interatomic potentials is discussed, including active learning strategies to efficiently improve the quality of potentials. Because the structure space can be prohibitively high-dimensional, a strategy to reduce the dimensionality and efficiently sample conformational space is represented by the Bayesian Optimization Structure Search (BOSS) method. To understand structure-property relationships, the AI Sure Independence Screening and Sparsifying Operator (SISSO) method is integrated with FHI-aims. This enables the creation of maps of materials properties and functions to identify regions in materials space where high-performance can be obtained for a certain goal and/or where more data are needed. Finally, Chapter 8 describes several machine-learning methods that target the electronic-structure directly and aim to learn basic ingredients of DFT, such as the electronic density or the Kohn-Sham Hamiltonian, based solely on a description of the atomic structure. These methods, which are all interfaced with FHI-aims, can be tightly integrated to electronic-structure calculations and offer a route to completely bypass the costly DFT evaluation of all material properties while retaining accuracy.

In conclusion of this Introduction, let us mention that the lists of authors of the whole paper and of most (but not all) contributions are ordered alphabetically, and the coordinators of the respective contributions are typeset in bold.

We expect that the summaries of features and implementations presented in these contributions will raise the interest of the electronic structure community, and we are happy to receive further questions or comments by email to aims-coordinators@ms1p.org.

Acknowledgment

We are deeply grateful to Anna Lutz for her dedication in coordinating and incorporating all change requests, and for her thoughtful communication with each contributor to the Roadmap article. Her efforts were instrumental in bringing it all together.

The Roadmap editors and FHI-aims coordinators:
Volker Blum, Sebastian Kokott, Mariana Rossi, and Matthias Scheffler

*Roadmap on Advancements of the FHI-aims Software Package**1.1. INTRODUCTION AND OVERVIEW*

Accepted Manuscript

2 Basic Concepts and Numerics

2.1 Numeric Atom-Centered Orbital Basis Sets for DFT and Beyond

Jörg Behler^{1,2}, *Volker Blum^{3,4}, Nicholas Boyer⁵, Bernard Delley⁶, Maria Dragoumi⁷, Stefan Goedecker⁸, William Huhn³, Erin R. Johnson^{9,10,11}, Elisabeth Keller¹², *Sebastian Kokott^{7,13}, Raul Laasner³, Susi Lehtola¹⁴, Johannes Margraf¹², Jack Morgenstein³, Thanush Patlolla¹⁵, Xinguo Ren^{16,7a}, Karsten Reuter¹², *Mariana Rossi¹⁷, *Matthias Scheffler⁷, Yi Yao^{3,13}, Igor Ying Zhang^{7,18,b}, Rundong Zhao^{3,19,c}, and Tong Zhu³

¹Lehrstuhl für Theoretische Chemie II, Ruhr-Universität Bochum, 44780 Bochum, Germany

²Research Center Chemical Sciences and Sustainability, Research Alliance Ruhr, 44780 Bochum, Germany

³Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

⁴Department of Chemistry, Duke University, Durham, NC 27708, USA

⁵Department of Chemistry, University of North Carolina at Chapel Hill, Chapel Hill, NC, USA

⁶LTC, Paul Scherrer Institut, Forschungsstrasse 111, 5232 Villigen-PSI, Switzerland

⁷The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁸Department of Physics, University of Basel, Klingelbergstrasse 82, CH-4056 Basel, Switzerland

⁹Department of Chemistry, Dalhousie University, 6243 Alumni Crescent, Halifax, Nova Scotia, B3H 4R2, Canada

¹⁰Department of Physics and Atmospheric Science, Dalhousie University, 6310 Coburg Road, Halifax, Nova Scotia, B3H 4R2, Canada

¹¹Yusuf Hamied Department of Chemistry, University of Cambridge, Lensfield Road, Cambridge, UK, CB2 1EW

¹²Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹³Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

¹⁴Department of Chemistry, University of Helsinki, P.O. Box 55, FI-00014 University of Helsinki, Finland

¹⁵William G. Enloe High School, Raleigh, NC 27610, USA

¹⁶Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

¹⁷Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

¹⁸Department of Chemistry, Fudan University, Shanghai 200433, People's Republic of China

¹⁹School of Physics, Beihang University, Beijing 102206, China

(*) Coordinator of this contribution

^aCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

^bCurrent Address: Department of Chemistry, Fudan University, Shanghai 200433, People's Republic of China

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

^cCurrent Address: School of Physics, Beihang University, Beijing 102206, China

Summary

Density functional theory (DFT) – especially semilocal Kohn-Sham DFT and hybrid generalized Kohn-Sham DFT, which are based on an orthonormal set of effective single-particle wave function to construct the density – is arguably the most productive method for quantum-mechanical simulations of materials and molecules. The central numerical choice in implementing DFT is how to discretize the effective single-particle orbitals using a finite set of basis functions. The chosen basis set determines the numerical precision of a particular simulation and also the time to solution.

FHI-aims employs numerically tabulated atom-centered orbital (NAO) basis functions, exploiting several key advantages for precision and efficiency: (1) The rapidly varying shape of the orbitals and ground-state density towards the nucleus is captured practically exactly by including numerically calculated orbitals of free atoms in the basis set. Thus, accurate simulations including all electrons become straightforward, even for large systems. (2) Precise ground-state densities and total energies for molecules and for extended systems can be obtained with rather few additional basis functions beyond the minimal basis, i.e., the overall basis set size remains efficient even for high-precision calculations. (3) NAO basis functions can be strictly localized to a spatially limited region around their atomic center, \mathbf{R}_I , with a smooth decay to zero at a certain distance. Thus, different regions in space are treated by different subsets of basis functions, allowing one to implement almost all operations in standard DFT in a way that allows the computational effort to scale linearly with system size, N , in the limit of large systems. The sole exception is the solution of the matrix-based eigenvalue problem of DFT, which formally scales as $O(N^3)$ and dominates for the largest system sizes, even when employing highly efficient solvers [5, 6]. In the limit of a large number of atoms, this cost can often be circumvented by lower-scaling, direct solutions for the density matrix [7]. However, the efficiency of these solvers strongly depends on the sparsity patterns (e.g., atom density) in the simulated system. For 3D bulk materials, diagonalization is still often the most efficient (or, at least, most practical) approach even for the largest systems simulated today.

For ground-state DFT, FHI-aims possesses a complete, easy-to-use and well tested NAO basis set library for elements 1-102, which can be used without any further editing and which allows users to balance the desired precision with computational efficiency. Fast calculations are enabled by small but physically reliable “light” basis sets, while progressively larger basis sets (“intermediate”, “tight”, and beyond) allow one to achieve benchmark-quality numerical precision for targeted observables in production calculations (though of course dependent on the observable in question). The general form of NAO basis functions also allows one to use a wide range of other standard atom-centered orbital basis sets, including pre-constructed NAO basis sets for applications beyond ground-state DFT, as well as Gaussian- or Slater-type orbital (GTO or STO) basis sets.

Current Status of the Implementation

As noted above, the practical handling of FHI-aims’ basis sets is simple in routine simulations. Three well-tested levels of numerical precision, “light,” “intermediate,” and “tight,” cover ground-state DFT and many other tasks across the periodic table. In a practical FHI-aims run, simply copy the desired predefined defaults into FHI-aims’ input file `control.in` prior to running a calculation.

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

The remainder of this section reviews the mathematical definition of the basis sets inside the code, as well as the origins of appropriate basis sets for ground-state DFT and beyond. Throughout this section, the term “precision” is used to indicate the numerical quality of the result for a given set of physical approximations to the electronic structure problem (e.g., a particular density functional) as opposed to the accuracy of the approximate electronic structure method(s) themselves.

Numerical definition of basis functions in FHI-aims

In electronic structure theory, a set of N_b basis functions $\{\varphi_i(\mathbf{r})\}$ is employed to construct the electron density, $n(\mathbf{r})$, and other observables from effective, single-particle orbitals, $\psi_l(\mathbf{r})$, where l is the orbital index:

$$\psi_l(\mathbf{r}) = \sum_{i=1}^{N_b} c_{il} \varphi_i(\mathbf{r}). \quad (2.1)$$

FHI-aims uses numerically tabulated, atom-centered basis functions of the form [8]

$$\varphi_i(\mathbf{r}) = \frac{u_i(r)}{r} \cdot Y_L(\Omega). \quad (2.2)$$

Here, $u_i(r)$ is a radial function, usually centered at an atomic position \mathbf{R}_I (but can also be placed on user-defined empty sites). I denotes the center, $Y_L(\Omega)$ is a real-valued spherical harmonic function, $r = |\mathbf{r} - \mathbf{R}_I|$ denotes the distance to the atomic position, and $\Omega = (\theta, \phi)$ are the angles defined by the unit vector $(\vec{r} - \vec{R}_I)/r$. L indicates the combination of angular and magnetic quantum numbers, which are implicitly defined by the overall basis function index i (i.e., $L = L(i)$ in practice). The use of numerically tabulated radial functions, $u_i(r)$, in Eq. (2.2) was pioneered by Averill, Ellis, Zunger, Freeman, Delley, te Velde, Baerends, Koepernik, Eschrig, and others [9–14]. In FHI-aims, the radial functions, $u_i(r)$, are defined via smooth natural cubic spline functions on dense, logarithmically spaced grids and, thus, allow for a highly precise representation of any radial function shape. Calculations of periodic systems can be performed on exactly the same mathematical footing as non-periodic calculations by using linear combinations of the localized basis functions of Eq. (2.2) in every unit cell, with unit-cell-dependent phase factors accounting for reciprocal-space sums [8].

FHI-aims supports five types of radial functions, $u_i(r)$: (i) radial functions of spherical, self-consistent free atoms; (ii) radial functions of hydrogen-like atoms (i.e., one-electron atoms with a spherical nuclear potential, Z/r , in which Z is an effective parameter); (iii) radial functions of spherical, self-consistent free ions; (iv) GTO functions; and (v) STO functions. Radial functions, densities, and potentials for (i) spherical free atoms and (iii) free ions are generated on the fly using the atomic solvers originally included in the fhi98pp code [15], or in the atom_sphere [16] or four-component (Dirac) DFTatom codes [17], while hydrogen-like functions (ii) can be generated either from their analytical shape or by numerically solving the corresponding radial equation. The NAO radial function types (i-iii) in FHI-aims are strictly localized, i.e., they are non-zero inside a specified radius and they decay smoothly to zero when approaching the boundary. The smooth decay of the radial functions is achieved by applying a confinement potential during the construction of the basis functions [8, 12, 18] (i.e., also for numerically computed, hydrogen-like radial functions). This confinement potential is zero within a given radius, r_c , and then smoothly increases to a large value (formally, infinity) over a width, w . The intention of this localization is to avoid extended regions in which a basis function is close to, but not quite, zero, while keeping the impact on the overall attainable precision of a calculation minimal. Typical extents of the basis functions are thus still rather large: $r_c + w = 5 \text{ \AA}$ for light elements

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

and “light” computational settings and up to 8 Å for the largest atoms (Cs) and “tight” settings. As a result, products of NAO basis functions are only non-zero if they are located on atoms in reasonable proximity to one another, facilitating $O(N)$ scaling of the computational effort with increasing system size (where N is a measure of the system’s extent) for calculating Hamiltonian and overlap matrix integrals in semilocal DFT [19], the non-local exchange operator [6], and the electron density update. Depending on the exact mathematical context, the measure N might refer to the number of atoms, basis functions, electrons or similar quantities related to the system size.

Construction principle of the NAO basis set library for ground-state DFT

Following the same basic principles as other atom-centered basis sets in quantum chemistry, the preconstructed standard basis sets in FHI-aims consist of two parts:

- A “minimal basis” of atomic basis functions that can accommodate all electrons of each individual atom in a chemical system (non-periodic or periodic). The minimal basis can be constructed to represent the ground-state free-atom limit with high accuracy, including the structure of the orbitals and density near the nucleus, but it is not on its own sufficient to capture properties in extended systems.
- Additional radial functions that provide flexibility to represent the orbitals, electron density, and other observables in the full system. Consistent with the experience from other atom-centered orbital approaches (GTOs or STOs), only a relatively small set of specifically selected basis functions in addition to the minimal basis is needed to achieve physically accurate solutions for the orbitals, density, and total ground-state energy [12].

The fact that the shape of the radial functions, $u_i(r)$, in Eq. (2.2) is flexible enables the creation of basis sets that can represent different chemical structures and environments in DFT with high precision, while keeping the overall number of basis functions low (in the range of single digits to several tens per atom, depending on chemical element and desired precision; see below for some examples), i.e., remaining computationally efficient.

The minimal basis in FHI-aims consists of numerically calculated radial functions of self-consistent, spherical and non-spinpolarized free atoms in their ground states. This minimal basis captures the shape of the orbitals and density close to each atomic nucleus nearly exactly in a straightforward way, since the electron-nuclear potential is very large and dominates the solution close to the nucleus. Thus, shape approximations to the near-nuclear potential, orbitals, or density, such as pseudopotentials, effective core potentials, or projector-augmented waves are not needed. Likewise, the unphysical behavior of the kinetic energy operator that can arise for linear combinations of analytical functions [20], such as for contracted Gaussian functions near a nucleus, is avoided (cf. Figure A.2 and associated discussion in Ref. [20]). As an additional advantage, physically correct finite-nucleus potentials for different isotopes can be substituted for the unphysical Coulomb singularity, $-Z_I/r$ [21], if desired (FHI-aims has an implementation of finite-nucleus potentials from experimentally determined charge density distributions [22–24] as compiled in Ref. [25]), and fully relativistic four-component atomic basis functions can also be incorporated into the formalism in the quasi-four-component (Q4C) approach [26].

Beyond the minimal basis definition, FHI-aims provides a basis-set library of additional radial functions for each atom type (elements 1-102), enabling calculations of increasing precision as more functions are included. This basis set library was created in an automated fashion, sidestepping any potentially limiting

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

“human intuition” in the initial basis set construction steps [8]. The simple recipe that proved successful was to select specific basis functions one by one from a large pool of candidate functions, using a stepwise variational minimization of non-self-consistently computed total energies of atomic dimers at several different bond distances. The pool of candidate functions included orbitals of hydrogen-like single-electron atoms, as well as doubly charged free ions, considering angular momenta up to g for elements 1-18 and up to h for all elements beyond.

In practice, the sequences of basis functions resulting from this minimization procedure turn out to be transferable for use as basis functions in more complex structures as well. With few exceptions, the basis functions follow one another in groups of different angular momenta (e.g., a p -, d - and s -type function following the minimal basis for oxygen, see below). These groups are denoted as “tiers” or “levels” of basis functions for each element. The basis sets in the “light”, “intermediate”, and “tight” numerical defaults of FHI-aims are constructed by selecting basis functions from these “tiers” to achieve basis set convergence commensurate with the respective defaults. However, we note that convergence of an observable with the basis set size may be dependent on the system type (i.e., free atoms, molecules, surface or bulk systems), the chosen method (i.e., GGA, meta-GGA, or hybrid functional vs. correlated methods), or the observable itself.

As an example, Figure 2.1 illustrates the basis functions used in “light” settings for the H and O atoms in the H₂O molecule. The hydrogen atom has only a single minimal basis function ($1s$, Figure 2.1a). The next selected basis function is a parameterized, hydrogen-like $2s$ function (Figure 2.1b) followed by a $2p$ hydrogen-like function (Figure 2.1c). In multi-atom calculations, this initial group of basis functions builds on the atomic solution but provides variational flexibility both regarding the radial character and the angular momentum character of the density around the H atom. The minimal $1s$, hydrogen-like $2s$ and $2p$ functions shown in Figures 2.1a-c (the first “tier”) are enabled as part of FHI-aims’ “light” defaults for H. Considering that there are three angular momentum functions for a p orbital, this adds up to five radial functions for H with “light” settings.

The light elements B, C, N, O, and F all possess minimal basis sets composed of $1s$, $2s$, and $2p$ functions (shown for O in Figures 2.1d-e). Consistent with typical STO or GTO basis sets, the next “tier” for each of these elements contains one s , p , and d function, respectively (Figures 2.1g-i), amounting to a total of 14 basis functions per atom. As in the case of H, this group of functions again provides flexibility both regarding the radial and the angular momentum character of the density around each atom. This first tier of basis functions is used as “light” settings in FHI-aims. It yields rather reliable molecular geometries [8] and closely matches the design principles of so-called “double zeta plus polarization” STO and GTO basis sets.

Figure 2.1j shows the resulting electron density of a H₂O molecule for the PBE density functional [27]. Figure 2.1k displays the corresponding difference between the self-consistent electron density, $n(\mathbf{r})$, and a simple superposition of spherical free-atom densities without any chemical bond formation, showing the physically expected shift of charge from the red region near the H atoms to the green region closer to the O atom). As shown, e.g., in Refs. [8, 28], the numerical precision can be systematically increased to very high precision (well below the chemical accuracy threshold of 1kcal/mol for total energies and atomization energies) as more basis functions are added beyond “light” settings. For example, Figure 2 in Ref. [28] reports 0.8 kcal/mol Median Absolute Deviation (MAD) and 1.4 kcal/mol maximum absolute deviation (max AD) for tight species defaults, and 0.1 kcal/mol MAD and 0.8 kcal/mol max AD for tier4 species defaults for 100+ molecules, per molecule with respect to highly precise values obtained using the multiresolution wavelet code MRCHEM [29]. The deviations for atomization energies are correspondingly less, i.e., 0.3 kcal/mol

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

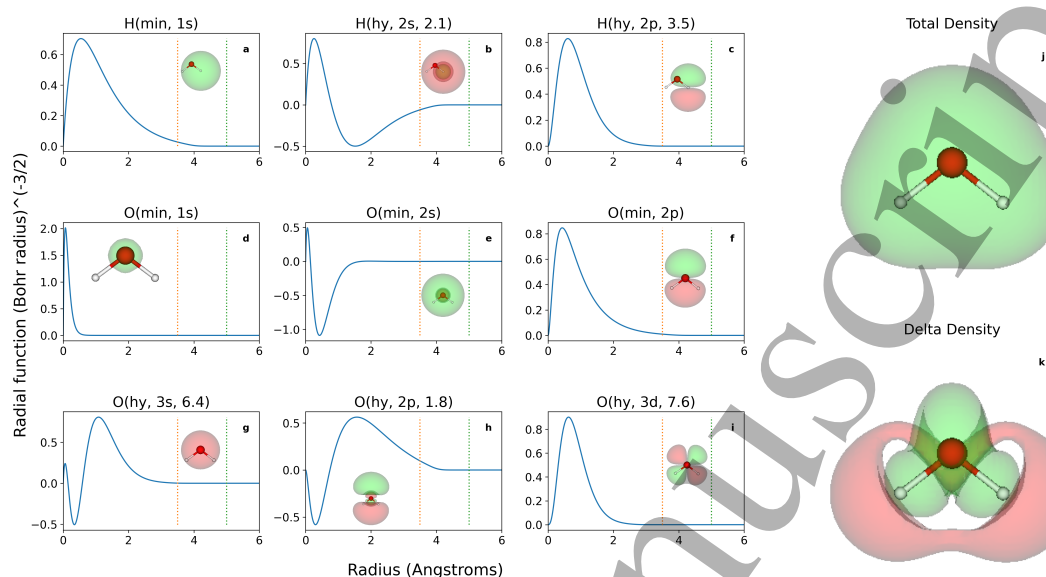


Figure 2.1: (a-i) Visualization of the NAO basis functions associated with “light” settings for a H_2O molecule. Each plot shows a radial function, $u_i(r)$, with an isosurface of the corresponding three-dimensional basis function, $\varphi_i(\mathbf{r})$, as an inset. The numerical definition of each basis function is indicated at the top of each plot. “min” indicates a minimal basis function, “hy” denotes a basis function from a hydrogen-like atom, for which the final number is the effective nuclear charge, Z , used in the construction of that function (e.g., $Z = 2.1$ for the hydrogen-like $2s$ function in b). Vertical red lines show the onset of the confinement potential (3.5 \AA) and vertical green lines show the radial distance at which the confinement potential approaches infinity (5 \AA). For larger distances, each radial function is strictly zero. (j) Density isosurface of the self-consistent total density of the H_2O molecule, computed using FHI-aims’ “light” settings and the PBE density functional. (k) Difference between the self-consistent density of the H_2O molecule and a simple addition of spherical free-atom densities. The isosurfaces of basis functions are shown for a value of $\pm 0.03 \text{ a.u.}$, with green denoting positive and red negative isosurfaces. Similarly, total and difference densities are plotted with positive (green) and negative (red) isosurfaces for values of $\pm 0.03 \text{ a.u.}$

MAD and 1.3 kcal/mol for tight species defaults, and 0.08 kcal/mol MAD and 0.5 kcal/mol max AD for tier4 species defaults, because some errors cancel.

The computer-generated basis set library in FHI-aims can also safeguard against potential omissions in human intuition. For example, the Zn atom ($Z=30$) has a closed $3d$ electron shell but not yet a p -type valence electron shell in the free atom. The first automatically selected additional basis function is indeed a p -type function, reflecting its importance in chemical bonds, then followed by another set of s , p , d and f functions. This group of 33 basis functions is retained in the the first tier of basis functions for Zn in FHI-aims, corresponding to “light” settings.

In relation to other basis function types than NAOs (e.g., plane waves), it is important to note that comparing just the number of basis functions on their own is not strictly meaningful, since the numerical effort per basis function will vary greatly with the basis function type, desired numerical precision, and other approximations made. Furthermore, precise points of comparison in the literature can be difficult to find. As one literature example, consider the augmented plane wave calculations showcased in Ref. [30]. In this work, for the CO molecule in a periodic box, 1,400 basis functions per atom were used. For a bulk Sb_2S_3 crystal, 120

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

basis functions per atom were employed. In contrast, FHI-aims' "tight" settings require 39 basis functions per atom for the CO molecule (an advantage brought about because no NAO basis functions are needed to represent the vacuum in a DFT total-energy calculation). In the Sb_2S_3 case, 44.2 NAO basis functions per atom would be used in FHI-aims ("tight" settings). Towards the largest systems, the relatively compact basis sets (i.e., relatively low ratio of number of basis functions to number of electrons in the system) will reduce the computational effort for matrix based eigenvalue or density matrix solutions (Sec. 2.4).

NAO basis sets beyond ground-state DFT

As with any other type of basis set, basis sets with different characteristics can be required for applications beyond ground-state DFT, e.g., to accurately represent explicit two-electron correlation (not part of ground-state semilocal or hybrid DFT) or to reflect response properties that arise in special spatial regions, such as the response of the density to a core level excitation or to the presence of nuclear spins.[31] For some scenarios, specialized NAO basis sets have been developed and benchmarked, as described in other sections of this review. Examples include:

- *Correlated methods for light-element molecules* (e.g., second-order Møller-Plesset theory, MP2, the random phase approximation, RPA, and *GW*): The valence correlation consistent (VCC) n -zeta (nZ) NAO basis sets ("NAO-VCC- nZ ") for the elements H-Ar introduced in Ref. [20] allow for a systematic extrapolation of ground- and excited-state energies to the complete basis-set limit, using basis sets of increasing order n ($n=2-5$). The principle follows the established scheme for GTO-based correlation-consistent basis sets by Dunning and others [32], but using a minimal basis set of free-atom NAO basis functions and additional hydrogen-like basis functions, all of which are numerically confined to a finite extent as described above.
- *Low-energy neutral excitations in light-element molecules* (e.g., optical absorption and excitons using the Bethe-Salpeter equation (BSE), linear-response or real-time time-dependent DFT): As shown in Ref. [33], numerically precise neutral excitation energies can be achieved by adding a few spatially extended "augmentation" basis functions to FHI-aims' standard basis sets. This strategy is analogous to Dunning's augmentation of standard GTO basis sets.[34] In practice, Ref. [33] shows that adding one s -like and one p -like augmentation function from Dunning's basis sets to FHI-aims' "tier 2" basis sets allows one to compute low-lying excitation energies of small molecules with a numerical precision of a few hundredths of an eV. These basis sets are referred to as "tier2_aug2" in FHI-aims.
- *Neutral excitation energies of light-element molecules involving core levels*: For excitations that promote a core electron to a valence level (e.g., in X-ray absorption spectroscopy), additional basis set flexibility in the near-nuclear region is needed. As shown in Ref. [35], precise core-level excitation energies can be obtained by adding a few highly localized STO basis functions to the "tier2_aug2" basis sets.
- *GW calculations of energy band structures of periodic solids*: FHI-aims' standard "tier 2" basis sets show qualitatively reliable results [36]. In line with other basis set types (e.g., plane waves, linearized augmented plane waves or GTOs), key parameters such as the fundamental gap reach a precision of several hundredths or tenths of eV for dense solids. Compared to experiment, the resulting band gaps are drastically improved over values extracted from single-particle eigenvalues of semilocal DFT. As explained in more detail in Ref. [36], additional, highly localized STO basis functions can be added

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

to further improve numerical precision. Similarly, optical properties of solids can be computed using the BSE based on GW quasiparticle calculations and FHI-aims' standard "tier" basis sets.[37]

- *Nuclear magnetic resonance observables for light-element molecules:* As shown in Ref. [31], nuclear shieldings can be computed with high numerical precision using the NAO-VCC- nZ basis sets. J -couplings, i.e., the effective coupling strength between two nuclear spins, require basis sets that include additional, highly localized functions around each nucleus, in line with experience from other basis set types. For J -couplings (elements 1-18), FHI-aims offers a series of basis sets called NAO- J - n . These basis sets combine the NAO-VCC- nZ basis sets with highly confined, primitive s -type GTOs derived from Jensen's GTO-based pc- J basis sets[38, 39].

In addition to NAO basis sets, FHI-aims can also employ any other basis set prescription based on STOs or GTOs from the standard quantum chemistry literature, since these basis functions have the same numerical form as Eq. (2.2). A broad range of GTO basis sets has been collected at the Molecular Sciences Software Institute (MolSSI) basis set exchange [40] and can be exported in a format suitable for FHI-aims. Thus, direct comparisons and benchmarks with standard quantum-chemistry approaches are possible on equal footing in FHI-aims, e.g., in the GW100 benchmark of molecular excitation energies[41].

Usability and Tutorials

As noted earlier, FHI-aims' standard basis sets are pre-tabulated within "light", "intermediate", or "tight" species defaults. Additionally, a set of "really_tight" species defaults is provided as a starting point for customizable convergence tests, with the same initial basis-set selection as "tight" but even more stringent numerical parameters otherwise. The user either appends these defaults for the given elements to the `control.in` input file of FHI-aims or uses an external package, such as GIMS [42] or ASE [43], to automatically build an input file with the specified defaults. Their use is documented across all tutorials published for the code (<https://fhi-aims-club.gitlab.io/tutorials/tutorials-overview/>)

The tabulated defaults that are distributed with the code have been benchmarked for reliable performance. If desired, all defaults can be trivially customized. In addition to other numerical settings (e.g., the real-space grids and the electrostatic potential expansion order[8]), they contain specifications of the NAO confining parameters, the NAO minimal basis, and the list of all further basis functions to be used in the calculation. This selection can be used as is or (if desired) can be adjusted – either by simply commenting / uncommenting suggested basis functions or by editing directly, allowing one to add any supported basis-function type with custom parameters. Thus, the modification of the parameters in the species defaults is straightforward for any user. The meanings of all these parameters are documented in the FHI-aims manual. Furthermore, standard GTO and STO basis sets from other quantum chemistry codes can be used, e.g., imported from the MolSSI Basis Set Exchange.[40]

For DFT, the high numerical precision of the pre-tabulated basis sets in FHI-aims has been established in a host of tests against other codes over time. Perhaps most prominently, in the widely cited 2016 "Delta test" of computationally predicted equations of state $E(V)$ for 70 elemental solids,[44] FHI-aims' "really_tight" settings with the first two angular momentum groups or "tiers" enabled, provided essentially indistinguishable numerical precision compared to other benchmark-quality all-electron codes (Wien2k[45] and Exciting[46]). FHI-aims' current "default_2020" light, intermediate, and tight production basis-set choices reflect selections that were assessed using the same Delta test data to ensure reliable precision at each level. A full compilation of the Delta test results is included in Appendix 9.1 and a set of corresponding input

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

and output files has been deposited at the NOMAD repository [47].

For total energies, atomization energies and dipole moments of isolated systems, a comparison to high-precision multiresolution wavelet calculations for 211 molecules demonstrated similarly high and systematically increasing precision of FHI-aims' NAO basis sets, down to few-meV total energy differences per molecule on average. [28] Energy band structures for 103 elemental and compound solids, covering 66 chemical elements, were later compared at the DFT level between FHI-aims and the high-precision all-electron code Wien2k. [48] For scalar-relativistic electronic valence states, differences remain below 10 meV on average for the band structures when using FHI-aims' "tier 2" NAO basis sets. For conduction bands up to 5 meV above the conduction band minimum or Fermi level, this difference increases to 50 meV for some elements; this is still remarkably close given the limits imposed by finite basis set size on the representation of high-lying unoccupied states. The same comparison was later carried out for self-consistent, spin-orbit-coupled band structures from the Wien2k code, compared to Q4C results from FHI-aims, also including spin-orbit coupling self-consistently. Despite the completely different implementations and overall different methodological choices, both codes agreed to within 50 meV on average even for the largest deviation (metallic Ir, for which FHI-aims' Q4C implementation is expected to be more accurate) for valence energy bands.

A standard benchmark for main-group quantum chemistry is the GMTKN55 database [49], which consists of 55 individual benchmark sets that can be grouped into five categories, as shown in Table 2.1. Since the data sets in each category have very different energy scales, their contributions to the total error metrics are weighted by the average magnitude of their component energy differences. The excellent performance of FHI-aims' NAO basis sets is demonstrated by their differences from the GTO values, $|\Delta|$ in Table 2.1, giving rather close results to those reported in Ref. [49] using quadruple- ζ GTO basis sets, while including far fewer basis functions. Since this comparison focuses primarily on numerical precision of the different basis set prescriptions, the PBE functional was selected for testing in this work due to its unambiguous definition. As shown by the relatively high values of the weighted mean absolute deviation, compared to the high-level reference data of Ref. [49], the physical accuracy of the PBE functional itself is of course still limited due to neglect of dispersion and inherent delocalization error.

Future Plans and Challenges

FHI-aims' standard NAO basis sets for DFT are highly robust and flexible, such that the fundamental basis-set selection has not needed to be changed since its original 2009 publication [8]. However, several additions are nevertheless desirable and on the way.

Methods that improve over the speed of "light" settings yet remain reliable have been a frequently desired wishlist item. The challenge is that the minimal basis of free-atom functions on its own is needed to guarantee the numerical representation of free atoms, but is itself not sufficient to guarantee physically correct results for bonded structures. Recent work by Keller *et al.* [51] provides a pathway forward, by employing almost minimal basis sets, to which only a single s function is added. Additionally, a parameterized correction energy is introduced that accounts for the some of the missing precision. This correction currently only supports the widely used PBE density functional, but could be extended to other functionals as well. The energy correction results in an average error in unit-cell volume of less than 5% across a wide range of different materials spanning from perovskites to noble gases and metalloids. In comparison with the commonly used FHI-aims-09 "light" and "tight" basis sets, this approach was found to offer average speedups of 1.92 and 4.76, respectively, for the cases considered in Ref. [51]. This decrease in computational time makes the

Roadmap on Advancements of the FHI-aims Software Package

2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND

Table 2.1: Weighted mean absolute deviations (WTMAD2, defined in Ref. [49], in kcal/mol) from high-level reference data for each category of the GMTKN55 benchmark. Results are shown for the PBE functional using either literature data [49] for Gaussian-type orbitals (GTOs) or FHI-aims calculations (NAOs). $|\Delta|$ denotes the difference between the calculated GTO and NAO WTMAD2 values, validating their numerical precision. The GTO calculations used the aug-def2-QZVP basis set, while the NAO calculations used tight species defaults in most cases. The exceptions are for the AHB21, BH76, BH76RC, G21EA, and IL16 benchmarks, which contain weakly bound anions. Here, the nonstandard "tier2.aug2" basis was used as it includes additional diffuse functions. The other exception is for WATER27 where, due to the large system sizes, tier2.aug2 was only applied to oxygen atoms for the specific reactions involving anions. The FHI-aims inputs are freely available on github [50].

Dataset description	Number of subsets	PBE		
		GTOs	NAOs	$ \Delta $
Reaction energies for small systems	18	7.93	7.99	0.06
Reaction energies for large systems	9	16.23	16.27	0.04
Barrier heights	7	16.72	16.42	0.30
Intermolecular non-covalent interactions	12	15.68	16.01	0.33
Intramolecular non-covalent interactions	9	19.70	19.93	0.23
Total data set	55	13.88	13.95	0.07

simulation of systems up to tens of thousands of atoms, or beyond, significantly more accessible.

The minimal basis sets of FHI-aims contribute greatly to the precision of electron densities and total energies near each nucleus. One remaining restriction is that numerically exact free-atom solutions in the atomic solver libraries employed in FHI-aims are not available for meta-GGA density functionals or hybrid DFT. Thus, the minimal basis sets for these functionals are currently typically constructed by the local-density approximation or the semilocal PBE functional. While the resulting difference is insignificant for energy differences in chemically bonded systems, a completely consistent treatment of the minimal basis for each density functional would be desirable in order to completely leverage the all-electron precision in FHI-aims across all functionals.

Finally, for speed reasons, the introduction of pseudopotentials in FHI-aims has often been desired, since this change would reduce computational cost to the attainable minimum by outrightly eliminating all core electrons. Indeed, this is the approach taken by the Siesta code [52] and others. NAOs are perfectly compatible with pseudopotentials. However, a new optimization of all basis sets, grids, and a complete pseudopotential set would be required. Norm-conserving pseudopotentials are already partially supported in FHI-aims in the context of embedding calculations. This functionality could easily be extended to the fully self-consistent case. In order to increase the speed of calculations even further, full support for norm-conserving pseudopotentials could certainly be envisioned, but has not yet been approached because of the considerable human time investment.

As an alternative approach to minimize the computational cost associated with core electrons, a frozen-core approximation has been implemented for the eigenvalue solver in the ELSI infrastructure. This approach requires no adjustments to the underlying potential and it allows one to seamlessly re-introduce the core electrons if desired. Carrying the frozen-core approach through the remaining FHI-aims code (beyond the eigensolver stage) is a promising route to achieve similar speedups to an outright pseudopotential variant of FHI-aims.

*Roadmap on Advancements of the FHI-aims Software Package**2.1. NUMERIC ATOM-CENTERED ORBITAL BASIS SETS FOR DFT AND BEYOND*

Acknowledgements

We acknowledge Dr. Axel Becke for his assistance with generating the GMTKN55 inputs. ERJ thanks the Atlantic Computational Excellence Network (ACENET) for computing resources, the Natural Sciences and Engineering Research Council (NSERC) of Canada for financial support, and the Royal Society for a Wolfson Visiting Fellowship. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC
ATOM-CENTERED ORBITALS

2.2 Technical Underpinnings for All-Electron Density Functional Theory with Numeric Atom-centered Orbitals

Viktor Atalla¹, Daniel Berger², Björn Bieniek¹, *Volker Blum^{3,4}, Saeed Bohloul⁵, Christian Carbogno^{6,7,a}, Sucismita Chutia¹, Ralf Gehrke¹, Mark Glass³, Dorothea Golze⁸, Jan Günzl⁸, Ville Havu⁹, Olle Hellman¹⁰, Uthpala Herath⁴, William Huhn⁴, Werner Jürgens^{1,b}, Florian Knoop^{6,11}, Franz Knuth⁶, *Sebastian Kokott^{6,5}, Björn Lange³, Maja Olivia Lenz-Himmer⁶, Konstantin Lion^{6,5}, Werner Lipsunen⁹, Florian Merz¹², Herzain I. Rivera-Arrieta⁶, Matti Ropo¹, Mariana Rossi¹³, Andrea Sanfilippo¹, Matthias Scheffler⁶, Franziska Schubert¹, Álvaro Vázquez-Mayagoitia¹⁴, Jürgen Wieferink¹, Scott Woodley¹⁵, Yi Yao^{4,5}, Yingyu Yao¹, Mina Yoon¹⁶, and Victor Wen-zhe Yu³

¹Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Chair for Theoretical Chemistry and Catalysis Research Center, Technische Universität München, Lichtenbergstr. 4, D-85747 Garching, Germany

³Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

⁴Department of Chemistry, Duke University, Durham, NC 27708, USA

⁵Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁶The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁸Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁹Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

¹⁰Weizmann Institute of Science, Rehovot, Israel

¹¹Department of Physics, Chemistry and Biology (IFM), Linköping University, SE-581 83, Linköping, Sweden

¹²Lenovo HPC Innovation Center, Stuttgart, Germany

¹³Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

¹⁴Computational Science Division, Argonne National Laboratory, 9700 South Cass Avenue, Lemont, Illinois 60439, United States

¹⁵University College London, Department of Chemistry, Gower Street, London WC1E 6BT, United Kingdom

¹⁶Center for Nanophase Materials Sciences, Oak Ridge National Laboratory, Oak Ridge, TN 37830, USA

(*) Coordinator of this contribution

^aCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

^bCurrent Address: Technische Hochschule Mittelhessen (THM), University of Applied Sciences, Fachbereich MND, Wilhelm-Leuschner-Straße 13, D-61169 Friedberg, Germany

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

Summary

Numeric atom-centered orbitals (NAOs) are the fundamental discretization choice for the electronic structure problem in FHI-aims. The remaining code is built around this foundation. This contribution summarizes the technical choices made to deliver a code that is efficient and convenient to use. We outline the implementation of the integration scheme and the underlying integration grids, as well as the evaluation of the electrostatic potential using an atom-centered multipole decomposition of the density. The two-electron Coulomb operator is treated in a separate scheme, using an efficient resolution of identity technique. The available choices for the k-grid are introduced. Furthermore, symmetry usage is considered, implemented as an option for periodic systems to reduce the computational effort for real-space and k-space operations. Beyond the basic algorithmic choices, a key element in FHI-aims is the handling of the message-passing interface (MPI) based parallelization. Two schemes exist in distributing the real-space Hamiltonian and overlap matrices: a globally sparse index representation and a locally-indexed dense index representation, where the latter is especially efficient for large-scale simulations. From the spatial derivatives of the NAOs, expressions for analytical forces and stresses can be derived, enabling efficient structure relaxation and molecular dynamics simulations. The technical details of the trust-radius-method relaxation algorithm, as well as constrained structure relaxation will be discussed. We lay out how the integration, force and stress evaluation can be further sped up by using graphics processing units (GPUs). Finally, we explain software engineering aspects, including how new developments are integrated into the software project and core functionalities of FHI-aims are continuously regression tested.

Current Status of the Implementation

Real-space grids. A central question for a code based on numerically tabulated functions is how to represent integrands for matrix elements, densities, potentials, and other objects relevant to electronic structure theory. A central constraint is that an all-electron method with sharply peaked potentials and rapidly varying orbitals near the nucleus cannot simply be implemented using a uniform, even spaced real-space grid of points in three dimensions – such a grid would need to be far too dense to resolve each nucleus correctly. Instead, FHI-aims successfully employs the methods of non-uniform, overlapping atom-centered real-space grids pioneered by Becke [53] and Delley [12]. In this formalism, real-space integrals are computed numerically on an integration grid that combines atom-centered spheres of grid points around each nucleus (at). Using this integration grid, integrals can be approximated as a summation:

$$\int d^3r f(\mathbf{r}) \approx \sum_{i=1}^{N_{\text{grid}}} w_i f(\mathbf{r}_i) = \sum_{\text{at}=1}^{N_{\text{at}}} \left[\sum_{s=1}^{N_{\text{rad}}(\text{at})} w_{\text{rad}}(\text{at}, s) \sum_{t=1}^{N_{\text{ang}}(\text{at}, s)} w_{\text{ang}}(\text{at}, s, t) p_{\text{at}}(\mathbf{r}_{\text{at}, s, t}) f(\mathbf{r}_{\text{at}, s, t}) \right], \quad (2.3)$$

where $w_{\text{rad}}(\text{at}, s)$ represents radial integration grid shell weights. The specific choice of radial integration grid shells is detailed in [8], Appendix A of [20] and in Appendix D.1 of Knuth et al. [54], where a variable radial multiplier is used to uniformly increase the number of radial integration grid shells s . The $w_{\text{ang}}(\text{at}, s, t)$ are the weights for the spherical integration grid, which typically use the grids by Lebedev and coworkers [55–57] (FHI-aims uses the variants created by Delley [58]). The Lebedev grids possesses octahedral, i.e., O_h symmetry. We further provide angular grids with hexagonal symmetry of D_{6h} and, additionally

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

non-symmetric grids “efficient spherical t-design” (ESTD) grids [59]. They can be triggered by the keyword `force_lebedev_d6hgrid` or `force_lebedev_estd`, respectively. The function $p_{\text{at}}(\mathbf{r}_{\text{at},s,t})$ is the partition function, which smoothly transitions from zero to one based on its distances to each nucleus. For the recommended and default partition function used in FHI-aims, we refer to Appendix C in Knuth *et al.* [54]. This partition function is based on the Stratmann’s partition function [60] with our modifications to enhance performance in periodic systems.

For an efficient evaluation of the integrals of the form of Eq. (2.3), the set of integration points P is divided into mutually distinct groups or “batches” of points $B_\nu \subset P$. The batches B_ν are distributed across the MPI tasks using a grid adapted cut-plane-method [19] demonstrating linear scaling behavior. As described in the same paper, the density and its gradients are evaluated in a linear-scaling fashion, using the same formalism and the localized nature of NAO basis functions. For periodic systems and for large non-periodic systems, the density is constructed using a density-matrix based formalism rather than by summing over molecular orbitals individually.

Electrostatic (Hartree) Energy. The average Coulomb energy, E_{es} , of electrons and nuclei, approximated by a continuous probability density $n(\mathbf{r})$ for the location of electrons, and fixed positions \mathbf{R}_I for the nuclei Z_I ($I=1, \dots, N_{\text{atoms}}$), is a key element of any electronic structure calculation. Importantly, and as laid out in Ref. [8], this group of terms must be treated together:

$$E_{\text{es}} = - \int d^3r \left(n(\mathbf{r}) \sum_I \frac{Z_I}{|\mathbf{r} - \mathbf{R}_I|} \right) + \frac{1}{2} \iint d^3r d^3r' \left(\frac{n(\mathbf{r})n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} \right) + \frac{1}{2} \sum_{I,J \neq I} \frac{Z_I Z_J}{|\mathbf{R}_J - \mathbf{R}_I|}. \quad (2.4)$$

In the limit of infinitely large or periodic systems, each term in Eq. (2.4) would diverge towards infinity on its own, even if taken per atom or per unit cell, and even for an overall neutral system, since each term amounts to the interaction of a finite charge with another, overall finite charge. Taken together, the overall system of electrons and nuclei with which each charge interacts is neutral, i.e., a combined treatment as laid out in Ref. [8] is tractable.

In practice [8], FHI-aims first subtracts a sum of spherical free-atom densities from the overall density $n(r)$. The free-atom densities and their electrostatic potentials are large but they can be calculated precisely on dense logarithmic grids, thus removing a large piece from the overall computation of electrostatics.

The remaining density difference between the full density and the sum of free-atom densities, $\delta n(\mathbf{r})$, is then evaluated numerically on the overlapping atom-centered integration grid. To obtain its electrostatic potential, FHI-aims follows Becke’s [53] and Delley’s [12] approach to first partition $\delta n(\mathbf{r})$ into localized density pieces around each atom and then multipole-decomposing these densities. Each multipole component can be treated separately, using the Green’s function of the Coulomb operator, to arrive at a corresponding multipole potential component of the electrostatic potential, which can then be summed up to give the overall electrostatic potential $\delta v_{\text{es}}(\mathbf{r})$ associated with $\delta n(\mathbf{r})$. Some care is required to ensure that each multipole-decomposed density component has the exact same charge or multipole moment when represented on different, dense or sparse integration grids during the procedure. Furthermore, for periodic systems, an Ewald summation technique is applied to each multipole component [58], ensuring that long-range tails of the potential can be summed up efficiently and at once in reciprocal space.

Charged periodic systems can be treated but require an artificial opposing compensating background charge to give a finite energy per unit cell. This background charge can either be a constant density or it can be distributed as small pieces over the nuclei in the system. In the case of a localized charged defect and a

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

constant background charge, additional techniques such as the Freysoldt-Neugebauer-Vandewalle correction [61] can then be employed to approximately restore the energy of an isolated charged defect in an otherwise neutral crystal.

Overall, the electrostatic potential evaluation described above is efficient and, in periodic systems, scales as $O(N) \log N$ with system size N . Recently, we redesigned the computation to evaluate the contribution of each multipole component to the Hartree potential across a batch of grid points simultaneously [6]. This restructuring eliminates branching in the innermost loop, reduces subroutine calls by two orders of magnitude, improves memory access efficiency and cache usage, enables compiler vectorization, and ensures that the Hartree energy evaluation is not a bottleneck even for extremely large periodic computations.

Two-electron Coulomb operator Unlike for the overall density and average electrostatic potential, a piecewise evaluation of the Coulomb operator over orbitals or basis functions cannot be avoided if the exchange operator or explicitly correlated method's ground- or excited-state energies are required. The two-electron Coulomb operator in terms of NAO basis functions $\varphi_i(\mathbf{r})$,

$$(ij|kl) = \iint d^3r d^3r' \left(\frac{\varphi_i^*(\mathbf{r})\varphi_j^*(\mathbf{r}')\varphi_k^*(\mathbf{r}')\varphi_l(\mathbf{r})}{|\mathbf{r} - \mathbf{r}'|} \right), \quad (2.5)$$

poses no particular mathematical difficulty, except that the sheer number of terms initially grows as N_{basis}^4 in finite systems and, in periodic systems, can never be cut off even for very large distances because of the slow decay of the $\frac{1}{|\mathbf{r} - \mathbf{r}'|}$ potential. In FHI-aims, any techniques that require the actual two-electron Coulomb operator can be treated efficiently using resolution of identity (RI) techniques, which expand products of two basis functions $\varphi_i(\mathbf{r})\varphi_j(\mathbf{r})$ in terms of simpler individual auxiliary basis function sets $\{P_\mu(\mathbf{r})\}$. The construction of these product basis sets and associated operations for NAOs is described in Ref. [62] (RI-V) and, in a localized form, in Ref. [63] (RI-LVL). Especially the latter, localized RI technique RI-LVL enables linear-scaling evaluations of the exchange operator even for extreme system sizes and efficient periodic implementations of exact exchange [6, 64] as well as many-body methods [36, 37]. The errors associated with the corresponding RI techniques are discussed in Ref. [63].

Importantly, the explicit form of the two-electron Coulomb operator is never involved in the overall electrostatic potential calculation in FHI-aims, which rather employs the density-based multipole decomposition formalism described following Eq. (2.4). Thus, the precision of the auxiliary basis set and associated RI technique have no impact on the Coulomb potential term, usually the biggest term in complex systems. The remaining, usually smaller exchange or explicitly correlated energy pieces can be computed stably and efficiently with reasonably sized and well benchmarked auxiliary NAO basis sets for RI.

k-point grids. By default, for periodic calculations, FHI-aims relies on Γ -centered regular k-space grids for Brillouin zone integrations, with fractional coordinates $k_i = (i_1/N_1, i_2/N_2, i_3/N_3)$, where $i_x = 0, \dots, N_x - 1$ for each reciprocal lattice vector direction $x = 1, 2, 3$. The number of k-points N_1, N_2, N_3 along their corresponding reciprocal lattice vector can be either specified directly by the keyword `k_grid N_1 N_2 N_3` or is computed from the k-point line density `n_k k_grid_density n_k` via $N_x = n_k |\mathbf{L}_x^*|$, where \mathbf{L}_x^* is the reciprocal lattice vector corresponding to lattice vector \mathbf{L}_x . Additionally, the user can specify a rigid shift of the k-grid by setting `k_offset d_1 d_2 d_3`, where (d_1, d_2, d_3) is the shift in fractional coordinates. The use of shifted k-grids is only supported for LDA, GGA, and meta-GGA functionals and does not work with any functional that requires exact exchange contributions. Simple sanity checks are performed to determine if the user input is meaningful, e.g., for 1D systems and surface models the corresponding vacuum

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

directions should have only one k-point. For LDA, GGA, and meta-GGA functionals, arbitrary k-points can be generated by a user-defined external list of k-points and weights. It is also possible to automatically generate generalized regular k-grids by using the autoGR [65] code, which is interfaced with FHI-aims. In the case of autoGR, generalized regular k-grids are constructed from rotated Born-von Karman cells that preserve the space-group symmetry of the system. They allow a more efficient sampling of the k-space for oblique crystals. Generalized regular k-grids are supported for all density functionals in FHI-aims.

For any nonrelativistic or scalar-relativistic DFT simulation, the code assumes time-reversal symmetry, which reduces the number of k-point by about half. In addition, it is possible to only compute the irreducible set of k-points based on the space-group symmetry of the input structure by specifying `symmetry_reduced_k_grid.spg . true .` in the `control.in` file. In this case, the symmetry matrices obtained from the library `spglib` [66] are used to reduce the k-grid.

Indexing the real-space matrices and load balancing in real-space operations. In evaluating Eq. 2.3 on an even-spaced grid for a group of matrix elements (e.g, for $N_{\text{basis}} \times N_{\text{basis}}$ Hamiltonian matrix elements h_{ij} , with $N_{\text{basis}} \times N_{\text{basis}}$ corresponding integrands $f(\mathbf{r})$ on each grid point \mathbf{r}), nearly ideal, communication-free parallel execution can in principle be achieved by distributing different segments of the integration grid to different MPI tasks. A problem arises with respect to re-assembling the full integrals h_{ij} from the partial integrals calculated on different MPI tasks, since the memory footprint of h_{ij} grows large towards large systems, making it necessary to distribute h_{ij} between different MPI tasks as well. The distribution of non-zero elements of h_{ij} , however, will not be the same as the distribution of grid segments in which those matrix elements are computed, leading to a non-trivial communication pattern during execution.

Optimizing these parallel distributions and the associated communication patterns creates an integration framework that scales efficiently towards extremely large systems and towards extremely large numbers of processor cores. We here introduce how real-space matrices like the Hamiltonian or overlap matrix are indexed and stored over MPI tasks after the integration is carried out. Since the approach is similar for all kind of real-space matrices, we will focus our notation on the Hamiltonian matrix. A detailed explanation can be found in the paper by Blum et al. [8] and Huhn et al. [67].

In brief, the real-space Hamiltonian is evaluated for the disjoint batches of grid points B_v as [67]

$$h_{ij} = \sum_v h_{ij}[B_v], \quad (2.6)$$

where

$$h_{ij}[B_v] = \sum_{\mathbf{r} \in B_v} w(\mathbf{r}) \varphi_i^*(\mathbf{r}) \hat{h}_{KS} \varphi_j(\mathbf{r}) \quad (2.7)$$

is the contribution of batch B_v to the real-space Hamiltonian element h_{ij} .

Once integrals of the real-space Hamiltonian (and overlap) are evaluated for each batch $h_{ij}[B_v]$, a local matrix copy of h_{ij} is updated on each MPI task. Within FHI-aims, different storage formats are used for this local copy depending on the calculation setup (see also the explanation in [67]).

For small clusters, every MPI task has a full dense copy of h_{ij} by default (i.e. `packed_matrix_format none`). This storage pattern becomes inefficient in terms of memory usage for even moderately sized systems and is thus not used as a default choice for any larger clusters or periodic systems. The default for smaller periodic systems or cluster systems is a globally-indexed sparse format (i.e. `packed_matrix_format index`). Here, only the non-zero elements of h_{ij} are stored in a compressed

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

sparse row (CSR) format along with two indexing arrays. By default, all matrix elements that could potentially be non-zero according to the extent of the basis functions involved are kept – the strict localization of the basis functions themselves still leads to a natural sparsity of large matrices that is already sufficient for efficient execution. While the sparse matrix storage in this format is efficient for smaller system sizes, its remaining disadvantage is that the sparse storage of h_{ij} on each MPI task still scales linearly with system size as $O(N_{\text{atom}})$, independent of the number of MPI tasks being used. This scaling behavior of per-processor memory will thus become the memory bottleneck in truly large-scale calculations.

In order to achieve constant, rather than linear per-processor memory of all matrix storage arrays towards large system sizes and correspondingly large CPU core counts, FHI-aims implements a so-called locally-indexed sparse or dense real-space matrix storage format [67] (activated by `use_local_index .true.` or automatically above a certain system size, typically 100 atoms, when supported by all requested observables in a calculation). In this approach, each processor works on batches of spatially adjacent grid points and stores only the subset of matrix elements (e.g., h_{ij}) that are non-zero on those grid points. Thus, individual matrix elements can have non-zero contributions from different processors (i.e., different MPI tasks), but in the limit of large systems, no single MPI task needs to store anything even close to the full matrix size - the actual, locally non-zero matrix segments to be stored are far smaller. The synchronization step to gather the full matrix from its local, distributed pieces is then more complex compared to a single global format. First, the locally-indexed matrix elements are distributed in an overlapping fashion across MPI tasks. Second, the fully integrated matrix is also stored in a distributed format, but normally a dense, block-cyclic format across MPI tasks that is not simply related to the parallelization of the underlying integration grid. The distributed format avoids the memory bottleneck of the global format for larger systems, provided that sufficiently many MPI tasks are available, but incurs a small performance overhead. To mitigate this overhead, one can set the keyword `load_balancing .true.`, where the code tries to achieve an even integration load across all cores. Enabling the load balancing also changes the matrix storage format from locally-indexed sparse format to a local dense matrix format (see Ref. [67] for details).

In general, computations that use quantities built from basis functions on the real-space grid can take advantage of sparsity in much the same way as for the Hamiltonian and overlap matrices. This includes the evaluation of the density and its derivatives, as well as of LDA, GGA and mGGA exchange-correlation terms, since they are subject to the same locality limitations within each grid-point batch.

MPI-3 shared memory. As a next step towards optimizing memory and communication, sharing common arrays between processors located on a physical node in FHI-aims is accomplished following the MPI-3 standard. Compared to the more frequently found OpenMP paradigm, the MPI-3 approach can be quite readily controlled from within a given run, without asking for any specific instructions for shared-memory layout to be provided by a user. At the time of writing, MPI-3 shared memory use is largely restricted to the computation of the exact exchange contribution.[6] A module for handling MPI-3 internode shared-memory arrays has been added to FHI-aims. In principle, this module makes the use of the shared memory arrays in FHI-aims almost as easy as using non-shared arrays. For reading from such arrays, there is no difference to conventional arrays. Only in case of writing to the shared arrays special care is needed as only a single MPI task is allowed write to an array element at the same time. After writing to the shared arrays, a special synchronisation step is needed. The shared memory modules set up the needed infrastructure, e.g., intranode communicators, which in turn enable the allocation and deallocation, as well as synchronisation of the shared memory arrays.

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

Forces and stress tensor. In an all-electron formalism, nearly complete basis set convergence can typically be achieved for total energies associated with fixed atomic positions, but the same fixed basis set would not describe a structure completely if the atom in question, but not its basis functions, were moved. As a result, when calculating analytical derivatives of the total energy for a basis set associated with fixed atomic positions, a number of terms arise that must be accounted for. In short (a detailed description is included in Ref. [8]), in addition to Hellmann-Feynman forces, i.e., the derivative of the Hamiltonian itself, so-called Pulay forces (associated with immobile basis functions), additional derivatives associated with the multipole decomposition of the electrostatic potential, and terms associated with the exchange-correlation functional (generalized gradient functional derivatives, derivatives associated with exact exchange [64], the random phase approximation [68]) must be computed. Recently, a correction of the Hellmann-Feynman forces that accounts for residual non-selfconsistency (cf. Eq. (6.5)) has been implemented, reducing the number of SCF iterations needed for precise forces. In addition, the branching inside of the main loops of the Pulay force evaluation has been simplified and the computation to aggregate or avoid unnecessary computations has been restructured.

Furthermore, the evaluation of stress, i.e., of the derivatives of the total energy E under a symmetric strain deformation $\varepsilon_{\alpha\beta}$ of the unit cell with volume V along the the Cartesian axes α, β ,

$$\sigma_{\alpha\beta} = \frac{1}{V} \frac{\partial E}{\partial \varepsilon_{\alpha\beta}} \quad (2.8)$$

is supported. A finite-difference approach is available for testing purposes, covering all total-energy methods. For production purposes, all semilocal (LDA, GGA, and mGGA) and hybrid density functionals are supported by analytical stress derivatives. A detailed account of the terms needed for GGA and hybrid functionals is given in Ref. [54]. These implementations enable structural relaxations of lattice degrees of freedom, as discussed below, enable *ab initio* molecular dynamics simulations in isobaric or isostress ensembles (see Contrib. 6.2), and play a pivotal role in fully anharmonic simulations of heat transport (see Contrib. 7.1).

Structure optimization The task of structure optimization is to find a local minimum configuration x_{\min} of the DFT total energy $E(x)$, where [69]

$$x = (R_0^x, R_0^y, \dots, R_N^z; a_1^x, a_2^x, \dots, a_3^z) , \quad (2.9)$$

with $3N$ atomic coordinates $R_I^{x,y,z}$, and three lattice vectors $a_L^{x,y,z}$ in the case of periodic solids. The degrees of freedom can be constrained in various ways, leading to a modified total energy function $\tilde{E}(x)$ without implications for the following discussion. For constraining the structure optimization either the space group symmetries can be used or, more generally, parametric constraints can be utilized that maintain local distortions or global symmetries of the structure [70]. In the quasi-Newton method, a step s_x toward the minimum is predicted by

$$s_x = B_{xx'}^{-1} f_{x'} . \quad (2.10)$$

where f_x denotes the generalized force, $f_x = -\partial_x E(x)$, and B is an approximation to the true Hessian matrix, $H_{xx'} = \partial_x \partial_{x'} E(x)$. The minimum configuration is characterized by $f_x = 0$. FHI-aims uses a quasi-Newton descent optimization with a hessian approximant B obtained via the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm [71-74], and a trust radius method (TRM) to limit the predicted step size. The initial guess for the approximate hessian, B^0 , is obtained in block-diagonal form via a Lindh model for the

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

atomic degrees of freedom [75], and

$$B_a^0 = c(a^{-1})^T a^{-1} \quad (2.11)$$

for the lattice degrees of freedom, where c is a numerical constant, a is the lattice matrix defined in Eq. (2.9), and T denotes the matrix transpose. The choice in Eq. (2.11) leads to strain-like relaxation steps [76, 77], improving the relaxation both in terms of the number of relaxation steps and the final outcome, compared to the original choice of a diagonal matrix B_a^0 in Ref. [54]. Furthermore, this choice can preserve space-group symmetry when used together with symmetrized forces and stress (cf. discussion below). More details can be found in appendix D of Ref. [78].

For the case where the predicted step length exceeds the trust radius r_{trust} , the step length is adjusted. We note that the adjustment of step length to the trust radius in the default TRM algorithm can lead to symmetry-breaking steps even when symmetric forces and stress are used. This is due to the adjustment

$$s'_x = (B + \lambda)_{xx'}^{-1} f_{x'} \quad (2.12)$$

where λ is chosen such that $s'_x = r_{\text{trust}}$, which can lead to space-group symmetry breaking in some cases, but is usually a good choice in particular for complex potential energy surfaces. To enforce a symmetry-preserving relaxation, the keyword LATTICE_TRM can be used, where the adjustment is performed via rescaling

$$s'_x = B_{xx'}^{-1} f_{x'} / \lambda \quad (2.13)$$

which preserves symmetry. Finally, FHI-aims monitors the consistency of forces, stresses, and total energies during a structure optimization, i.e., the total energy is not allowed to increase beyond a certain target tolerance (typically, 3 meV) over multiple relaxation steps.

In addition to structure optimization to a local minimum-energy structure, a python package “aimsChain” provides the ability to find transition states and minimum energy paths for chemical reactions. This package supports various flavours of the chain of states methods for finding the minimum energy path. Currently the nudged elastic band method (NEB) [79, 80], the string method [81], and the growing string method [82] are included. A tutorial is available at <https://fhi-aims-club.gitlab.io/tutorials/finding-transition-states-with-aimschain>.

Symmetry for periodic structures. Several ways to use symmetry for the simulation of periodic structures exist in FHI-aims. As noted before, FHI-aims assumes time-reversal symmetry for DFT calculations for the k-space grid in non-relativistic or scalar-relativistic computations by default. In addition, space-group symmetry can be used to further reduce the number of k-space grid points; here, the software package spglib[66] is used. Space-group symmetry can be employed as well to reduce the number of real-space grid points during the integration of the real-space matrix elements as well as for the computation of the density and of its gradients. This is especially beneficial for high-symmetry systems if force or stress evaluations are needed. If space-group symmetry is requested by the user, the input structure either needs to have the numerically correct atomic positions and lattice parameters, or the structure can be refined on the fly to the closest spacegroup that could be found by spglib. If the system has a space-group number larger than 1, the symmetry of the system will be preserved along relaxation trajectories.

Separately, FHI-aims has a complete framework for parametrically constrained relaxation of certain struc-

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

tures (i.e., user-imposed structural constraints beyond simply preserving the symmetry of a given system).

GPU acceleration. FHI-aims supports the use of graphics processing units (GPUs) for acceleration of DFT simulations. Currently, the integration of the Hamiltonian matrix and overlap matrix, the evaluation of forces and stress components, as well as the solution of the generalized eigenvalue problem can be executed on GPUs [67]. GPUs with support for Nvidia's CUDA and AMD's HIP can be used for the aforementioned parts of the computation. The solution of the generalized eigenvalue problem can be accelerated by the ELPA library (see **Contribution 2.5**).

Restarting calculations. Often it is useful to restart simulations from already converged density matrices, e.g. for executing post-processing tasks (DOS, band structure, etc.) or to split long calculations into several parts. The current recommended restart functionality is provided by ELSI (see **Contribution 2.5**) via writing the density matrix to disk. This enables the restart of the calculations for the same number of k-points and basis functions, while not restricting the number of MPI tasks.

Recently, a different type of restart from the resolution-of-identity decomposed density has been added [83]. While this functionality is mostly used for machine-learning-the-density approaches, it can be also used in general for restarting calculations, especially also for different numbers of k-points. More details are given and applications are demonstrated in **Contribution 8.4**. It is also possible to write checkpoints during the computation of the self-energy for the GW approximation. This feature is introduced in **Contribution 4.3**.

Wave function export to TREXIO. FHI-aims provides an interface to the TREXIO [84] software ecosystem, which allows one to export a wave function as a TREXIO file. TREXIO files store wave functions in a general format, enabling seamless transfer between different quantum chemistry codes. Several codes can read wave functions from TREXIO, Figure 2.2 provides an overview of codes for which interoperability with the NAO-based TREXIO wave function from FHI-aims has been verified. This includes the codes Quantum Package [85], NECI [86] and QMC=Chem [87, 88]. Quantum Package supports methods like coupled cluster (CCSD), configuration interaction singles and doubles (CISD) or CIPSI (Configuration Interaction using a Perturbative Selection made Iteratively). The NECI software suite enables full configuration interaction quantum Monte Carlo (FCIQMC), while QMC=Chem performs highly accurate variational and diffusion quantum Monte Carlo (VMC, DMC) calculations. Additionally, TREXIO files can be converted into other formats, such as FCIDUMP [89], to enhance interoperability with other codes.

Software Engineering: Regression testing and continuous integration. FHI-aims includes a suite of regression tests containing over 100 small calculations that test a broad range of FHI-aims' functionality across several different primary code branches (e.g. periodic/non-periodic systems, MPI parallel/serial code execution). To ensure integrity of the code, FHI-aims employs a range of continuous integration pipelines to run the suite of regression tests for different compilers (nvhpc, GNU, Intel oneAPI) on different architectures (CPU and GPU nodes) and for different MPI libraries. The continuous integration pipelines are implemented using GitLab continuous integration/continuous delivery, through a `gitlab-runner` instance installed on a small local cluster. The continuous integration process is initiated whenever a developer pushes code changes to the FHI-aims GitLab repository. The push event creates several continuous integration pipelines on the GitLab server, which in turn are executed through the `gitlab-runner` instance on the small local cluster. Every pipeline has a dedicated container (that is, a virtualization of the operation system with pre-installed dependencies). In this container, the current version of FHI-aims is built and the regression tests

2.2. TECHNICAL UNDERPINNINGS FOR ALL-ELECTRON DENSITY FUNCTIONAL THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

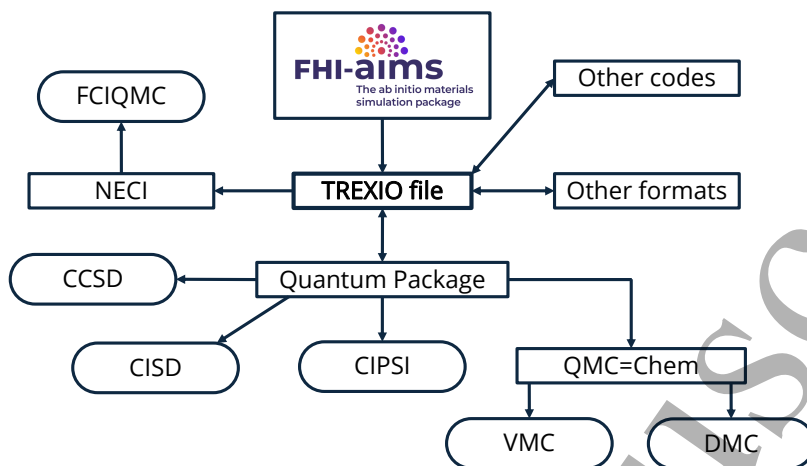


Figure 2.2: Scheme illustrating how an NAO wave function calculated by FHI-aims can be processed by other quantum chemistry codes after being exported to a TREXIO file. Rectangular boxes represent codes, while rounded boxes indicate the possible quantum chemistry methods. Refer to the text for explanations of the abbreviations.

are executed. Once the tests have successfully passed, the code changes can be merged into the main line of the code.

Future Plans and Challenges

FHI-aims in its present form supports some of the most demanding electronic structure calculations ever performed, especially for more demanding hybrid density functional theory calculations [6]. With the advent of exascale computing, the biggest challenge will be to keep FHI-aims current and functional on the most relevant platforms and architectures, which are themselves continually evolving. Even for the existing GPU infrastructure in FHI-aims more opportunities are possible to accelerate the code execution, especially porting the evaluation of the Hartree Potential and the exact exchange contribution to GPUs. Furthermore, new types of GPUs, such as Intel's current generation, are not yet supported at the time of writing and will require developer attention.

Acknowledgements

The authors acknowledge Arvid Ihrig for numerous contributions to the FHI-aims core functionalities.

We acknowledge the computational resources provided by the Aalto Science-IT project and CSC – IT Center for Science (Finland).

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

2.3 Relativistic Effects: Scalar Relativity, Spin-Orbit Coupling, and Four-Component Relativity

***Volker Blum**^{1,2}, **Sander Driessen**³, **Dorothea Golze**⁴, **Matthias Gramzow**⁵, **William P. Huhn**¹, **Timo Jacob**^{5,a}, **Sebastian Kokott**^{6,7}, **Qing-Long Liu**⁴, **Karsten Reuter**⁸, **Matthias Scheffler**⁶, **Han Wang**⁹, **Wentao Zhang**¹, and ***Rundong Zhao**⁹

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Materials Simulation and Modelling, Department of Applied Physics, Eindhoven University of Technology, PO Box 513, 5600 MB Eindhoven, The Netherlands

⁴Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁵Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁸Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁹School of Physics, Beihang University, Beijing 102206, China

(*) Coordinator of this contribution

^aCurrent Address: Institute of Electrochemistry, Ulm University, 89081 Ulm, Germany

Summary

While quantum mechanics is often introduced via the non-relativistic Schrödinger equation, this is actually not the correct equation to capture chemical and materials properties across the periodic table. In chemistry and materials science, relativistic effects can have large impacts and must be accounted for. Dirac's equation incorporates relativity more adequately and approximations derived from it are therefore used in practical simulations to ensure qualitatively correct results. The non-relativistic limit is often viewed as sufficient for chemical bonding characteristics of the lightest elements (atomic numbers $Z \leq 20$), but relativistic effects can have a considerable effect even on the properties of this limited group, e.g., on their core level energies[90]. The impact of relativity grows rapidly with increasing atomic number Z of the elements involved in a molecule or compound[26, 48]. Predicted non-relativistic energy band gaps are off by ~ 0.3 eV already for the semiconductors ZnO and ZnS ($Z=30$ for Zn). For semiconductors including the heavy $6p$ main group elements Pb and Bi ($Z=82$ and 83 , respectively), energy bands derived from the heavy elements are qualitatively incorrect without including the effects of spin-orbit coupling (SOC). Simply put, our world is a product of relativistic, not non-relativistic underlying interactions, including with electromagnetic fields.

This chapter focuses on relativistic approaches implemented in FHI-aims[8, 26, 48], not a general review of relativistic electronic structure theory, which can be found elsewhere[91]. NAOs offer a unique opportunity

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

for relativistic calculations since their functional form in the important region near the nucleus can be captured practically exactly, while their computational properties (compact basis sets, localization) make them amenable to treating large, complex systems. FHI-aims can include relativistic effects in density-functional theory (DFT) at four different levels of increasing accuracy, i.e., (1) non-relativistic, (2) scalar-relativistic (scalar Kohn-Sham orbitals with relativistic effects included in the kinetic energy operator)[8], (3) scalar-relativistic but with SOC effects included non-selfconsistently in orbitals and band structures [48], and (4) four-component orbitals in quasi-four-component (Q4C) Dirac-Kohn-Sham theory for total energies[26]. Scalar-relativistic effects are included using the atomic zero-order regular approximation (atomic ZORA) at no extra cost and complexity compared to the non-relativistic case and should therefore be included in all simulations. Non-selfconsistent SOC yields energy band structures that are quantitatively accurate at least up to $Z=48$ (Cd) and cover the properties of heavy-element compounds qualitatively correctly up to Pb and Bi. The Q4C approach provides an excellent foundation for future developments that cover spin, currents, and the interaction of the electrons with external fields.

Current Status of the Implementation

The Dirac-Kohn-Sham (DKS) DFT equations are built upon single-particle wave functions that are vectors of four scalar functions or “components”, typically denoted $(\phi_1(\mathbf{r}), \phi_2(\mathbf{r}), \chi_1(\mathbf{r}), \chi_2(\mathbf{r}))$. As a starting point, we consider the time-independent version of the four-component Dirac equation,

$$\begin{pmatrix} \begin{pmatrix} V & 0 \\ 0 & V \end{pmatrix} & c\boldsymbol{\sigma} \cdot \mathbf{p} \\ c\boldsymbol{\sigma} \cdot \mathbf{p} & \begin{pmatrix} -2mc^2 + V & 0 \\ 0 & -2mc^2 + V \end{pmatrix} \end{pmatrix} \begin{pmatrix} \phi_1 \\ \phi_2 \\ \chi_1 \\ \chi_2 \end{pmatrix} = \epsilon \begin{pmatrix} \phi_1 \\ \phi_2 \\ \chi_1 \\ \chi_2 \end{pmatrix}. \quad (2.14)$$

ϵ is the corresponding eigenvalue, m is the electron rest mass, c the speed of light, $\mathbf{p} = -i\hbar\nabla$ is the momentum operator, $V(\mathbf{r})$ is the effective DKS potential and $\boldsymbol{\sigma}$ is a 3-vector with the three individual (2×2) Pauli matrices $\sigma_x, \sigma_y, \sigma_z$ as its entries. As written, Eq. (2.14) already omits explicit magnetic fields and current terms that appear in the full Dirac equation [92]. Additionally, relativistic corrections to the Coulomb potential itself (which governs the electron-electron and electron-nuclear interactions) are neglected. In this form, Eq. (2.14) provides a practical starting point to connect systematically to the more approximate forms of Schrödinger-like scalar-relativistic equations with or without SOC [48]. The mathematical steps to derive these expressions are simple and are outlined next.

As formatted above, the matrix equation (2.14) emphasizes its structure in a form of four (2×2) blocks. One commonly combines ϕ_1 and ϕ_2 into a two-component vector called the “large component” $\phi(\mathbf{r}) = (\phi_1(\mathbf{r}), \phi_2(\mathbf{r}))$, and χ_1 and χ_2 into another two-component vector called the “small component” $\chi(\mathbf{r}) = (\chi_1(\mathbf{r}), \chi_2(\mathbf{r}))$. With this substitution, the lower two lines of Eq. 2.14 can be used to express the small component χ in terms of the large component:

$$\chi = \frac{c}{2mc^2 + \epsilon - V} \boldsymbol{\sigma} \cdot \mathbf{p} \phi. \quad (2.15)$$

Substituting Eq. 2.15 back into the upper two lines of Eq. 2.14 results in a single expression for the large component ϕ :

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

$$\boldsymbol{\sigma} \cdot \mathbf{p} \frac{c^2}{2mc^2 + \epsilon - V} \boldsymbol{\sigma} \cdot \mathbf{p} \phi + (V \cdot \mathbb{1}) \phi = \epsilon \phi. \quad (2.16)$$

Eq. (2.16) is a two-component equation for ϕ only, where $V \cdot \mathbb{1}$ denotes the scalar potential term V multiplied by the (2×2) unit matrix $\mathbb{1}$ to preserve the (2×2) structure of the equation. However, unlike Schrödinger's non-relativistic equation, Eq. (2.16) features an ϵ -dependent Hamiltonian on the left hand side and therefore is not solvable as a linear system with orthogonal solutions. Furthermore, based on the original equation (2.14), ϕ is not normalized on its own but only together with the small component χ . Nevertheless, Eq. (2.16) is a practical approximate equation for ϕ alone.

Using the explicit form of the Pauli matrices, one may further rewrite the kinetic energy operator in Eq.(2.16) as the sum of a spin-free part and a spin-containing part,[48] laying the foundation for the aforementioned scalar-relativistic and SOC approaches:

$$\left(\mathbf{p} \frac{c^2}{2mc^2 + \epsilon - V} \cdot \mathbf{p} + i \mathbf{p} \frac{c^2}{2mc^2 + \epsilon - V} \times \mathbf{p} \cdot \boldsymbol{\sigma} \right) \phi + (V \cdot \mathbb{1}) \phi = \epsilon \phi. \quad (2.17)$$

By writing down Eqs. (2.14)-(2.17) explicitly, the different levels of relativistic treatments available in FHI-aims can be rationalized straightforwardly.

Non-relativistic limit: Schrödinger equation. In regions of space where $\epsilon - V \ll 2mc^2 \approx 1 \text{ MeV}$, $\epsilon - V$ is negligible in the first two terms. By neglecting $\epsilon - V$ indiscriminately in those terms, the second term in Eq. (2.17) vanishes due to the cross product and the standard non-relativistic Schrödinger equation results:

$$\frac{\mathbf{p}^2}{2m} \phi + (V \cdot \mathbb{1}) \phi = \epsilon \phi. \quad (2.18)$$

In spin-polarized Kohn-Sham DFT, ϕ_1 and ϕ_2 in ϕ assume the roles of spin-up and spin-down electrons, coupled only implicitly through the exchange-correlation potential, which is part of V . However, the assumption $\epsilon - V \ll 2mc^2$ is never fully justified in all regions of space, since V assumes very large negative values close to a nucleus. As noted initially, the non-relativistic Schrödinger equation Eq. (2.18) is nevertheless frequently considered to be sufficient to capture properties associated with the valence electrons, away from the nucleus, in light-element molecules and solids ($Z=1-20$), e.g., chemical bonding energies or optical excitations. However, this assumption is far less appropriate for core electrons, which are localized close to the nucleus. Therefore, the most important use-case of the non-relativistic approximation in FHI-aims are benchmark comparisons of calculated observables to results from other codes in which the exact same, simple kinetic energy operator of Eq. (2.18) is used in both implementations. For production calculations, using scalar-relativistic theory (discussed next) has the same computational cost and is always preferable.

Scalar relativistic approach: Atomic ZORA. Three separate observations allow one to turn Eq. (2.17) into a Schrödinger-like equation that provides significantly broader physical accuracy than the non-relativistic limit.

1. The SOC term (which includes $\boldsymbol{\sigma}$) is primarily relevant in regions in which the gradient of V , via the momentum operator \mathbf{p} applied to $c^2/(2mc^2 + \epsilon - V)$, is large, i.e., near the nucleus. The SOC term is therefore frequently neglected for properties derived from valence electrons.
2. The potential $V(\mathbf{r})$ is only appreciably large compared to $2mc^2 + \epsilon$ in regions close to a nucleus. In those regions, $V(\mathbf{r})$ is very similar to the potential associated with a free atom A containing the same

Roadmap on Advancements of the FHI-aims Software Package

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

nucleus, V_A , which can therefore be used instead of V .

3. The energies of valence electrons (order $\sim eV$), which are primarily located away from the nucleus, always satisfy $\epsilon \ll 2mc^2$. The Hamiltonian therefore remains correct for valence electron properties even if setting ϵ to zero on the left side of Eq. (2.17).

By combining observations (1)-(3), Equation (2.17) can be transformed into a simpler form[8], which is nevertheless accurate for all valence electron properties in the absence of strong SOC[44]:

$$\mathbf{p} \frac{c^2}{2mc^2 - V_A} \cdot \mathbf{p} \phi + V \phi = \epsilon \phi \quad (2.19)$$

In a basis set representation, V_A is chosen to be the atomic potential associated with the basis function to its right and the operator is then symmetrized, as described in Eq. (2.20) and associated discussion. If the self-consistent V were used in the denominator instead of V_A , this would be the original zero-order regular approximation[93], which suffers from a gauge invariance problem[94]. Using V_A , which is determined once and does not change in a multi-atom self-consistent calculation, removes the gauge invariance problem.

In practice, Eq. (2.19) is never applied in real-space form in FHI-aims, but rather in the form of Hamiltonian matrix elements between basis functions,

$$h_{ij} = \langle \varphi_i | \mathbf{p} \frac{c^2}{2mc^2 - V_A} \cdot \mathbf{p} + V | \varphi_j \rangle. \quad (2.20)$$

In this form, the operator can be evaluated to the right, for the free atom potential V_A associated with the nucleus at which φ_j is centered, followed by symmetrization between matrix elements h_{ij} and h_{ji} . For a given matrix element, the resulting scalar-relativistic atomic ZORA kinetic energy operator is exactly analogous to the non-relativistic one, since V_A is independent of the three-dimensional structure of the molecule or solid under consideration and there is no dependence of h_{ij} on ϵ . For properties associated with valence electrons, e.g., the structure of solids, the results derived from FHI-aims' atomic ZORA scalar relativity are numerically indistinguishable from scalar-relativistic treatments in other benchmark-quality codes[44]. The remaining approximation is restricted to core level energies, for which $\epsilon \ll 2mc^2$ is not exact. However, this effect can be corrected by a simple correction that depends only on the chemical element in question, not on the structure within which an atom is embedded [90].

As a note, FHI-aims also implements a second, separate scalar-relativistic approach, "scaled ZORA" [8, 94]. Scaled ZORA addresses the gauge invariance problem of the original ZORA approximately, but not entirely. It is implemented non-selfconsistently, i.e., as a post-processing step. Scaled ZORA can be useful if one is interested in scalar-relativistic core level eigenvalues with absolute positions closer to their spin-orbit averaged positions in the Dirac equation. However, in general the atomic ZORA approach should be used since it is free of any gauge-invariance problems and is broadly supported for practical simulations with FHI-aims.

Scalar relativistic approach with second-variational SOC correction: atomic ZORA+SOC. For single-particle energy levels (i.e., energy band structures), SOC effects can become qualitatively important even if they do not yet affect total energy differences drastically. One widely used approach to restore SOC in energy band structures is therefore to first carry out a self-consistent scalar-relativistic calculation without SOC. In a second step, the SOC operator is evaluated for a subset of the scalar-relativistic, self-consistent orbitals. The resulting, spin-orbit coupled Kohn-Sham single-particle Hamiltonian is then inverted once to obtain approximate spin-orbit coupled eigenvalues ϵ and two-component orbitals ϕ . This approach is called a

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

“second-variational” SOC correction. The specific Hamiltonian used in FHI-aims for this purpose is [48]

$$\left(\mathbf{p} \frac{c^2}{2mc^2 + \epsilon - V_A} \cdot \mathbf{p} + \frac{i}{4mc^2} \mathbf{p} V \times \mathbf{p} \cdot \boldsymbol{\sigma} \right) \phi + (V \cdot \mathbb{1}) \phi = \epsilon \phi, \quad (2.21)$$

where the second term results from a Taylor expansion in V of the earlier SOC term in Eq. 2.17. As discussed in more detail below, this non-selfconsistent approach is remarkably precise for band structures across much of the periodic table [48] while remaining computationally efficient, since a full two-component self-consistent field cycle is avoided.

Self-consistent four-component relativity: Q4C approach. For self-consistent relativistic calculations of all four components, which includes self-consistent SOC, FHI-aims implements the quasi-four-component (Q4C) approach [26, 95]. This method (closely related to the exact two-component, X2C approach of quantum chemistry [96]) relies on the idea that the small component χ is most relevant in the near-nuclear regions of a structure, where relativistic effects are strong, so that χ can be handled by an approximate, fixed relation with the large component ϕ , which is based on the free atom. In practice, this is accomplished by expanding the four-component wave functions into atom-centered, four-component basis functions instead of expanding each component individually:

$$\begin{pmatrix} \phi_i \\ \chi_i \end{pmatrix} = \sum_{\mu} C_{\mu i} \cdot \begin{pmatrix} \varphi_{\mu}^L \\ \varphi_{\mu}^S \end{pmatrix} \quad (2.22)$$

For each of the 4C basis functions here, there is a fixed relation between its large and small components, i.e.,

$$\varphi_{\mu}^S = \hat{K} \varphi_{\mu}^L; \quad \hat{K} = \left[\frac{c}{2mc^2 + \epsilon_{A,\mu} - V_A} \right] (\boldsymbol{\sigma} \cdot \mathbf{p}). \quad (2.23)$$

Here, A denotes the atom type where this basis function is centered, and $\epsilon_{A,\mu}$ and V_A represent the corresponding orbital energy and atomic radial potential, respectively. For the minimal basis, from which the majority of the density is derived, $\epsilon_{A,\mu}$ is the corresponding eigenvalue of each basis functions, i.e., \hat{K} is the exact free-atom relation Eq. (2.15) between the atomic χ and ϕ . For other basis functions, FHI-aims employs $\epsilon_{A,\mu}=0$ as in the atomic ZORA approximation [26]. In FHI-aims, a set of fully-relativistic atomic-orbital basis functions for the minimal basis can be precomputed by solving the radial Dirac equation for free atoms using the open-source atom-solver code DFTATOM [97].

Usability and Tutorials

Since different relativistic approaches are implemented directly within FHI-aims, using them is as simple as setting a keyword, `relativistic`, to an appropriate value. This keyword is used and explained in multiple basic FHI-aims tutorials at <https://fhi-aims-club.gitlab.io/tutorials/tutorials-overview>.

- Non-relativistic theory and scalar-relativistic atomic ZORA are supported for essentially all functionality in the code due to their simplicity and mathematically simple structure.
- Second-variational, spin-orbit coupled atomic ZORA is supported for energy eigenvalues and energy band structures in semilocal and hybrid DFT, as well as many derived quantities such as element-resolved densities of states, spin textures of energy bands, or absorption spectra.

Roadmap on Advancements of the FHI-aims Software Package

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

- At the time of writing, the Q4C method in FHI-aims is implemented to support system sizes up to and above 100 atoms. It does not yet include support for exchange-correlation beyond non-spinpolarized, semilocal DFT and it does not yet support forces or geometry optimization.

As mentioned above, the scalar-relativistic atomic ZORA approach achieves essentially identical results with other codes, indicating that the valence electron treatment in this approach is basically the exact one within the limits of a scalar relativistic approach.[44] However, energy band structures from scalar relativistic computations deviate from fully relativistic ones already for compounds containing elements beyond Zn ($Z=30$),[26] i.e., SOC is essential for band structure calculations involving even moderately heavy elements.

For energy band structures and (in the case of the Q4C method) total energies, the spin-orbit coupled atomic ZORA and Q4C methods have been extensively benchmarked in Refs. [48] and [26]. Ref. [48] defined a comprehensive benchmark of the band structures of 103 metallic, semiconducting and ionic solids for scalar-relativistic and second-variational spin-orbit coupled methods, covering chemical elements from Li to Po. Compared to self-consistent spin-orbit coupled energy band structures derived from the Wien2k code,[45] the second-variational atomic ZORA + SOC approach remains quantitatively exact up to at least Cd ($Z=48$). The following $5p$ and $6s$ elements (up to Ba) show band structures that agree within 50 meV on average, whereas the energy band structures of $6p$ elements (up to Po) show deviations up to 0.3 eV. In Ref. [26], the same group of compounds was used to benchmark other methods against the Q4C approach, showing close agreement with Wien2k's self-consistent SOC approach except for the energy band structures of the late $5d$ metals Os, Ir, and Pt, for which the Q4C approach is presumably more accurate. All corresponding Q4C band structure benchmark data can be accessed via the NOMAD database at (<https://doi.org/10.17172/NOMAD/2025.03.09-1>).

Figure 2.3 shows the impact of relativistic effects on the band structure of an organic-inorganic semiconductor, the chiral hybrid perovskite S-1-(1-naphthyl)ethylammonium lead bromide (S-NPB) [98] in a 118-atom structure (Figure 2.3a), computed using tight settings along the reciprocal space path shown in Figure 2.3b. The density functional used is DFT-PBE, since the objective is a comparison of different relativistic effects on equal footing, rather than a more accurate treatment of the overall band structure using a higher-level approach. Energy bands highlighted in red are those primarily associated with lead ($Z=82$), for which SOC effects are much stronger than for all other elements in the structure. The evolution of energy band structures from (c) non-relativistic to (d) scalar-relativistic (atomic ZORA) shows drastic shifts both of the band gap and of the energy band alignments between Pb and other components in the valence band. Similarly strong effects occur when introducing second-variational SOC (e). Without SOC, a group of flat, molecular-derived bands would form the conduction band minimum, whereas the physically correct inclusion of SOC shifts the conduction bands to the Pb-derived bands. Physically, this shift would have a drastic effect, since the spatial location of carriers in a semiconductor impacts both the electronic and the optical properties of the material. Finally, the transition to (f) with the Q4C approach (including SOC self-consistently, as well as the small component) shows further changes, but much smaller in scale. While the Pb-derived bands are not separately highlighted in this figure (computing a formal decomposition is not yet implemented for the four-component wave function), their character at the conduction band edge is clearly apparent in comparison to (e), with a shift of a few tenths of eV compared to the organic-derived flat bands immediately above. This shift is of the same order as the inherent uncertainty even of higher-level functionals (such as hybrid DFT or the G_0W_0 method with different starting points) [99]. Compared to Q4C, the band structure features of this closed-shell system are already captured well at the simpler, non-selfconsistent atomic ZORA + SOC approach. However, a higher level of relativistic treatment would be needed for non-collinear spins in open-shell systems or if the p -derived conduction bands were partially filled, in which case self-consistency

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

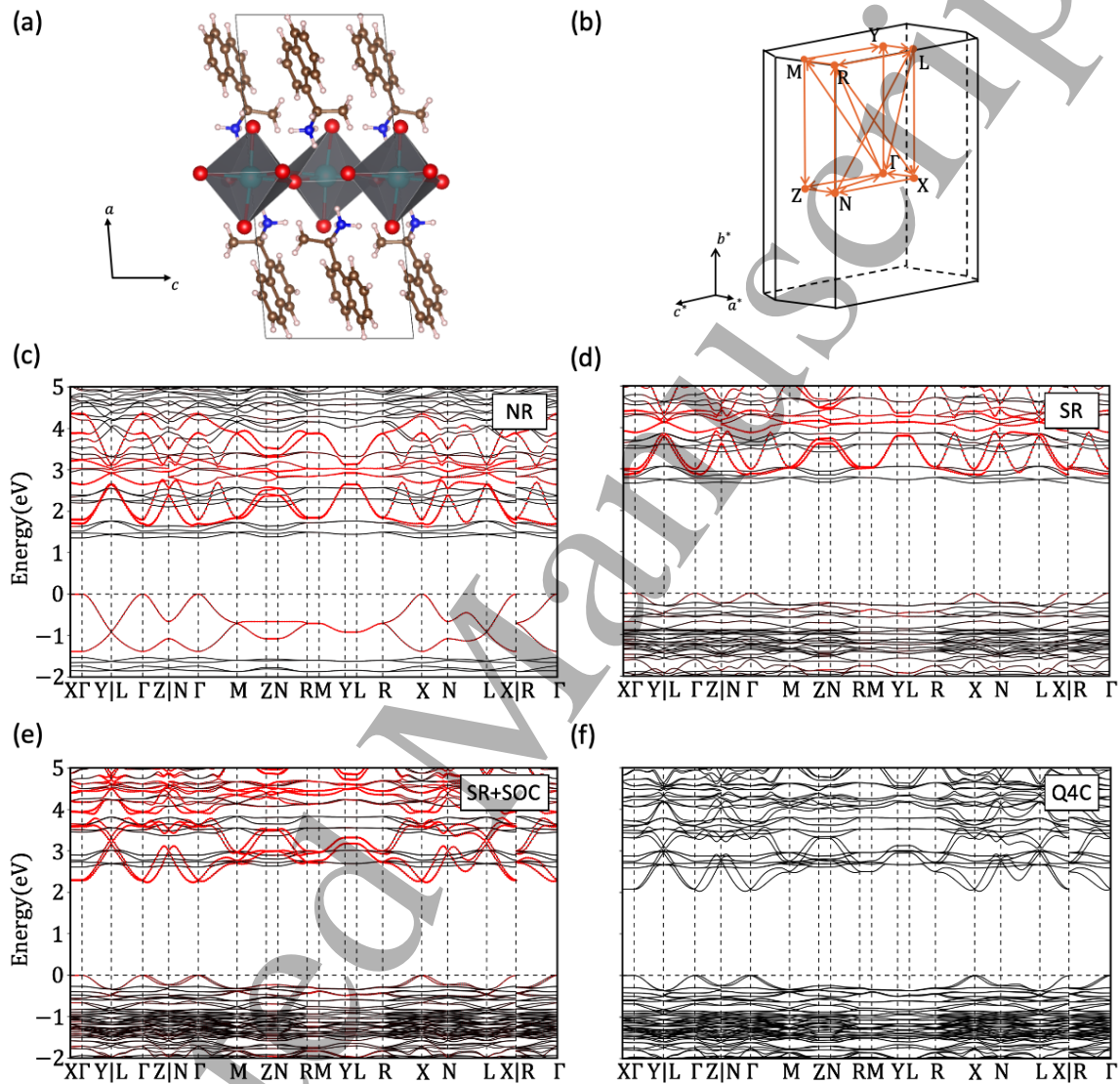


Figure 2.3: Structural and electronic properties of the layered hybrid perovskite S-1-(1-naphthyl)ethylammonium lead bromide (S-NPB). (a) A representative unit cell of the relaxed S-NPB structure in real space consists of 118 atoms, including 2 lead atoms, 8 bromine atoms, 4 nitrogen atoms, 32 carbon atoms, and 72 hydrogen atoms in a unit cell. (b) The corresponding Brillouin zone in reciprocal space, highlighting the k-path employed for band structure calculations. (c-f) Electronic band structure comparisons under different computational methods: (c) Nonrelativistic (NR), (d) Atomic Zero-Order Regular Approximation Scalar Relativistic (SR), (e) SR with non-selfconsistent Spin-Orbit Coupling (SR+SOC), and (f) Quasi-Four-Component (Q4C) method. The contributions for lead (Pb) in the bands (c-e) are highlighted in red.

2.3. RELATIVISTIC EFFECTS: SCALAR RELATIVITY, SPIN-ORBIT COUPLING, AND FOUR-COMPONENT RELATIVITY

would matter.

Future Plans and Challenges

For production DFT calculations across most of the periodic table, at least up to heavy elements such as Pb ($Z=82$), relativistic effects are reliably captured by scalar relativity (atomic ZORA) and, for band structures, enhanced by second-variational spin-orbit coupling (atomic ZORA + SOC).

For phenomena in which relativistic effects near the nucleus or the coupling between spin and orbital degrees of freedom play a particular role, future developments in FHI-aims will build on the Q4C approach, which has both self-consistent spin orbit coupling, the correct four-component wave function near the nucleus, and which can incorporate other effects such as the correct potential of a finite nucleus if needed. At the time of writing, ongoing developments include an extension of Q4C to system sizes well beyond the current reach, by adapting the method to FHI-aims' distributed-parallel, "locally-indexed" [67] infrastructure for large-scale integrals. Similarly, an extension of the Q4C-supported exchange-correlation functionals to DFT+U and support for spin-polarized exchange-correlation functionals, making non-collinear spin systems accessible in open-shell Q4C, are under active development.

Further important future Q4C developments include forces and stresses for geometry optimization and dynamics, support for hybrid density functionals, as well as extensions of the Q4C approach to many-body methods such as *GW* or the Bethe-Salpeter equation. Similarly, core-hole spectroscopies, nuclear magnetic resonance observables, and other physical observables which are impacted by nuclear and/or core electron properties are highly desirable targets for a fully relativistic implementation. Overall, work towards a complete, NAO based four-component treatment offers much promise for direct simulations of phenomena that are currently not within easy range of standard electronic structure methods.

Acknowledgements

We thank Dr. Ondřej Čertík for his technical support on the open source Dirac atom-solver code DFTATOM he authored. We are grateful to Levi Keller, who would have been a coauthor, for his contributions; Levi Keller passed away on January 28, 2023. For the application of the Q4C approach to large hybrid perovskite systems, V.B. and W.Z. were supported by NSF Award DMR-2323803.

2.4 Solvers for Large-Scale Electronic Structure Theory: ELPA and ELSI

***Petr Karpov**¹, ***Andreas Marek**¹, Tobias Melson¹, Pavel Kůs¹, Hermann Lederer¹, Bruno Lang², Alexander Pöppel³, Jun Tang⁴, Danilo Simoes Brambila⁶, Hagen-Henrik Kowalski⁵, Lydia Nemeč^{6,a}, Simone S. Köcher^{7,8,9,b}, Christian Carbogno^{5,8,c}, Christoph Scheurer^{7,8}, Matthias Scheffler⁵, Karsten Reuter^{7,8}, Victor Wen-zhe Yu¹⁰, Ben Hourahine¹¹, Alberto Garcia¹², William Dawson¹³, Yi Yao^{10,14}, William Huhn¹⁵, Jonathan Moussa¹⁶, Björn Lange¹⁰, Álvaro Vázquez-Mayagoitia¹⁷, Mina Yoon¹⁸, Samuel J. Hall¹⁹, Reinhard J. Maurer^{19,20,21}, Uthpala Herath¹⁰, Konstantin Lion^{5,14}, Sebastian Kokott^{5,14}, Ville Havu²², and ***Volker Blum**^{10,23}

¹Max Planck Computing and Data Facility, Garching, Germany

²University of Wuppertal, Mathematics and Natural Sciences

³Intel Corporation, Santa Clara, CA, USA

⁴Annapurna Labs, AWS

⁵The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Theoretical Chemistry and Catalysis Research Center, Technische Universität München, D-85747 Garching, Germany

⁸Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁹Institute of Energy Technologies (IET-1), Forschungszentrum Jülich GmbH, Wilhelm-Johnen-Str., 52425 Jülich, Germany

¹⁰Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

¹¹Department of Physics, SUPA, University of Strathclyde, John Anderson Building, 107 Rottenrow, Glasgow G4 0NG, UK

¹²Institut de Ciència de Materials de Barcelona, ICMA-B-CSIC, Campus UAB, 08193 Bellaterra, Spain

¹³RIKEN Center for Computational Science, Kobe 650-0047, Japan

¹⁴Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

¹⁵Intel Corporation, 500 Beaver Brook Road, Boxborough, MA, 01719, USA

¹⁶Molecular Sciences Software Institute, Blacksburg, Virginia 24060, USA

¹⁷Computational Science Division, Argonne National Laboratory, 9700 South Cass Avenue, Lemont, Illinois 60439, United States

¹⁸Center for Nanophase Materials Sciences, Oak Ridge National Laboratory, Oak Ridge, TN 37830, USA

¹⁹Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

²⁰Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

²¹Faculty of Physics, University of Vienna, Vienna A-1090, Austria

²²Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

²³Department of Chemistry, Duke University, Durham, NC 27708, USA

(*) Coordinator of this contribution

^aCurrent Address: Infineon Technologies AG, Am Campeon 1-15, 85579 Neubiberg, Germany

^bCurrent Address: Institute of Energy Technologies (IET-1), Forschungszentrum Jülich GmbH, Wilhelm-Johnen-

Str., 52425 Jülich, Germany

^cCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

In this contribution, we give an overview of the ELPA library and ELSI interface, which are crucial elements for large-scale electronic structure calculations in FHI-aims. ELPA is a key solver library that provides efficient solutions for both standard and generalized eigenproblems, which are central to the Kohn-Sham formalism in density functional theory (DFT). It supports CPU and GPU architectures, with full support for NVIDIA and AMD GPUs, and ongoing development for Intel GPUs. Here we also report the results of recent optimizations, leading to significant improvements in GPU performance for the *generalized* eigenproblem.

ELSI is an open-source software interface layer that creates a well-defined connection between “user” electronic structure codes and “solver” libraries for the Kohn-Sham problem, abstracting the step between Hamilton and overlap matrices (as input to ELSI and the respective solvers) and eigenvalues and eigenvectors or density matrix solutions (as output to be passed back to the “user” electronic structure code). In addition to ELPA, ELSI supports solvers including LAPACK and MAGMA, the PEXSI and NTPoly libraries (which bypass an explicit eigenvalue solution), and several others.

ELSI, ELPA, and other solver libraries supported in ELSI are mature, well-tested software, ensuring efficient support for large-scale simulations on current HPC systems. Future plans for ELPA include further optimization of GPU routines, particularly integrating the GPU collective communication libraries (NVIDIA’s NCCL and AMD’s RCCL) into the solution of tridiagonal the matrix problem and backtransformation of the eigenvectors in standard eigenproblem solvers. Similarly expanding support for Intel GPUs through deeper integration with Intel’s oneAPI libraries, specifically oneCCL, is planned. These advancements aim to ensure that ELPA, accessible either with its own Fortran/C/C++ APIs or via ELSI, remains at the forefront of large-scale computations, offering researchers powerful tools for addressing increasingly complex systems in electronic structure simulations. Future developments in ELSI will also continue to focus on GPU support and exascale readiness, continued support for a broad range of solvers, as well as extensions to solvers, e.g., for constrained density functional theory.

Current Status of the Implementation

Introduction: From effective single-particle equations to generalized eigenproblem and density

Using the Kohn-Sham (KS) [100] or generalized Kohn-Sham (gKS) [101] formalism, a full problem of n_{el} interacting electrons can be transformed to n_{el} or more auxiliary single-particle equations, $\hat{H}\psi_l = \varepsilon_l\psi_l$, e.g., for the scalar-relativistic or non-relativistic kinetic energy operator \hat{t} :

$$\left(\hat{t} + V_{\text{ext}}(\mathbf{r}) + \int \frac{n(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|} d\mathbf{r}' + v_{\text{xc}}(\mathbf{r}) \right) \psi_l(\mathbf{r}) = \varepsilon_l \psi_l(\mathbf{r}), \quad (2.24)$$

where $V_{\text{ext}}(\mathbf{r})$ is the external potential, $n(\mathbf{r}) = \sum_{l=1}^N f_l |\psi_l(\mathbf{r})|^2$ is the electron density, ψ_l are the effective single-particle orbitals of DFT with occupation numbers f_l and eigenvalues ε_l . N here denotes the number of single-particle states considered in the calculation. The exchange-correlation potential v_{xc} is the only un-

Roadmap on Advancements of the FHI-aims Software Package

2.4. SOLVERS FOR LARGE-SCALE ELECTRONIC STRUCTURE THEORY: ELPA AND ELSI

known term and can be treated using various density functional approximations [1]. To solve the equations (2.24) numerically, one has to choose a convenient set of N basis functions and approximate the KS state $\psi_l(\mathbf{r}) = \sum_{i=1}^N c_{li} \varphi_i(\mathbf{r})$. The set of equations (2.24) then transforms to the generalized eigenvalue problem

$$HC = SC\varepsilon \quad (2.25)$$

where H and S are the Hamiltonian and the overlap matrices, respectively, with elements $h_{ij} = \int \varphi_i(\mathbf{r})^* \hat{H} \varphi_j(\mathbf{r}) d\mathbf{r}$, $s_{ij} = \int \varphi_i(\mathbf{r})^* \varphi_j(\mathbf{r}) d\mathbf{r}$; C is the matrix of eigenvectors (stored as matrix columns); and $\varepsilon = \text{diag}(\varepsilon_1, \dots, \varepsilon_N)$. Given C , one may obtain the total energy $E[n(\mathbf{r})]$ via the ground-state density $n(\mathbf{r})$.

Solving the generalized eigenproblem

Since the basis functions $\{\varphi_i(\mathbf{r})\}$ are not orthonormal, S is not an identity matrix. In order to solve the generalized eigenproblem, $HC = \varepsilon SC$, for real symmetric/complex Hermitian matrices H , S , and positive-definite matrix S , one usually reduces it to the standard eigenproblem via the following steps:

1. Cholesky decomposition of $S = U^T U$, where U is an upper-triangular matrix.
2. Inversion of the upper triangular matrix $U \rightarrow U^{-1}$.
3. Calculation of two matrix products: $\tilde{H} = (U^{-1})^T H U^{-1}$.

Here, A^T corresponds to the transpose or conjugate-transpose of a real- or complex-valued matrix A , respectively. This reduces the *generalized* eigenproblem to the *standard* one, $\tilde{H}\tilde{C} = \varepsilon\tilde{C}$, which has the same set of eigenvalues. If the eigenvectors are needed, one has to perform the backtransformation, which requires another matrix-matrix multiplication: $C = U^{-1}\tilde{C}$.

A strict condition for the applicability of steps 1 to 3 is that the positive-definite matrix S must not be numerically singular, i.e., the overlap matrix S itself must not have eigenvalues that are zero; otherwise, Eq. (2.25) could have multiple different solutions C that correspond to the same physical eigenfunctions $\{\psi_l(\mathbf{r})\}$. As a consequence, in practical computations, S should not be ill-conditioned, i.e., the ratio between the largest and the smallest eigenvalue of S should normally not exceed the numerical range (around 10^{12}) accessible by double precision numbers.

The standard FHI-aims basis sets are compact enough to lead to well-conditioned overlap matrices S even for well-converged DFT calculations and steps 1 to 3 are therefore routinely used. Prior to any calculation, FHI-aims computes the eigenvalues σ_i of the overlap matrix and alerts the user if a dense basis set with a high condition number of the overlap matrix is detected. In the latter case (high condition number of S), electronic structure calculations can then be carried out by transforming the eigenproblem (2.25) to the basis of eigenvectors D_i of the overlap matrix, but omitting any D_i for which σ_i is smaller than a small positive number δ (typically $\delta \leq 10^{-5}$). This procedure is standard in the community and is implemented in ELSI.

Circumventing the eigenvalue problem: Density-matrix based solutions

In the limit of large systems, the computational effort for solving the eigenproblem Eq. (2.25) scales cubically with system size and becomes the computational bottleneck for Kohn-Sham DFT. Solving the eigenvalue problem (2.25) is one option to obtain $n(\mathbf{r})$. However, since the actual targets of DFT are $n(\mathbf{r})$ and $E[n(\mathbf{r})]$, the eigenvalues and eigenvectors are technically not required. Alternative strategies therefore target finding the single-particle density matrix $P_{ij} = \sum_{l=1}^N f_l c_{li}^* c_{lj}$ without an explicit eigenvalue solution. The density

Roadmap on Advancements of the FHI-aims Software Package

2.4. SOLVERS FOR LARGE-SCALE ELECTRONIC STRUCTURE THEORY: ELPA AND ELSI

can then be computed as

$$n(\mathbf{r}) = \sum_{ij} \int \varphi_i(\mathbf{r})^* P_{ij} \varphi_j(\mathbf{r}) d\mathbf{r}. \quad (2.26)$$

Especially for sparse, localized basis sets, the computational effort for solvers that target the density matrix without an explicit eigenvalue/eigenvector solution can scale quadratically or linearly with the system size. For sufficiently large systems, such solvers can therefore outperform the cubic-scaling eigenvalue solution. The crossover point, i.e., the system size beyond which the eigenvector-free solution becomes more favorable, depends on the specifics of the system, level of theory, and computer hardware used. Thus, maintaining access to different solver types within a single electronic structure code is desirable.

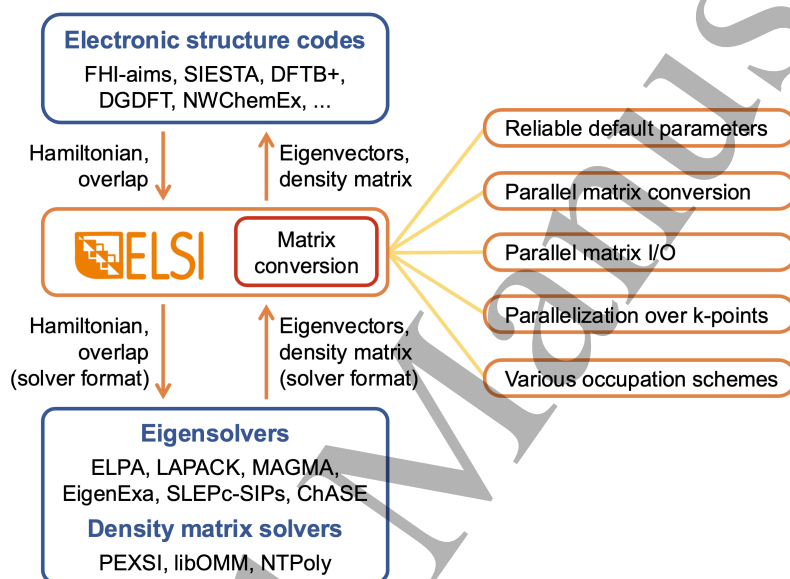


Figure 2.4: Tasks performed by the ELSI interface software, connecting different eigenvalue and density matrix solvers to electronic structure codes including FHI-aims, Siesta, DFTB+, NWChemEx and others. ELSI provides a uniform interface that is callable in Fortran, C, C++, and Python, and handles matrix format conversion between user codes and solver libraries. The solvers presented here include eigenvalue and density matrix solvers connected to ELSI at the time of writing. The ELSI interface provides functionality for tasks common to eigenvalue and density matrix operations, such as calculating occupation numbers or trivially parallel handling of independent eigenvalue problems for different k -points and spin channels.

ELSI

The ELSI interface software [7, 102, 103] (Figure 2.4) provides a uniform code layer that handles eigenvalue problems or density matrix calculations, supporting several electronic structure codes (FHI-aims [8], DFTB+ [104], Siesta [52], NWChemEx [105]) and integrating ten different solvers. The solvers have different APIs, making their direct integration into an electronic structure software a non-trivial task. Moreover, different electronic structure codes would all need to reimplement essentially the same connections to different solvers. ELSI simplifies the integration and use of multiple solver libraries by providing a unified interface, allowing users to access various eigensolvers and density matrix solvers optimized for various problem sizes

Roadmap on Advancements of the FHI-aims Software Package

2.4. SOLVERS FOR LARGE-SCALE ELECTRONIC STRUCTURE THEORY: ELPA AND ELSI

and types. The interface is designed around a derived type, the `elsi_handle`, to pass data between the user code that calls ELSI, the ELSI interface layer itself, and the solver codes. This construct enables programs written in Fortran, C, C++ or Python to connect to the ELSI interface with equal ease. Additionally, conversions between different distributed-parallel matrix formats are handled efficiently in ELSI.

ELSI supports cubic scaling eigensolvers like LAPACK [106, 107] and ELPA [5, 108, 109], as well as reduced scaling methods such as PEXSI [110, 111] or the linear-scaling NTPoly [112] density matrix solvers. Further supported solvers include the dense eigensolvers EigenExa [113] and MAGMA [114], the iterative eigensolvers ChASE [115, 116], SLEPc-SIPS [117], and libOMM [118], as well as BSEPACK [119] and DLA-Future [120, 121]. Although implemented in ELSI, ChASE and DLA-Future are not currently supported in FHI-aims. Given the plethora of solvers supported, we refer to the respective original publications for their in-depth mathematical definitions. ELSI returns either eigenvalues and eigenvectors or the density matrix and its energy-weighted counterpart. Since the density matrix includes the occupation numbers of the Kohn-Sham orbitals, ELSI also provides an implementation of numerically precise occupation numbers. For FHI-aims, ELSI is the designated source of standardized computed occupation numbers, f_i , including several non-Aufbau occupation approaches to perform occupation-constrained DFT calculations (e.g., the Δ SCF method).

ELPA

ELPA is an open-source massively parallel direct eigensolver library for real symmetric or complex Hermitian eigenvalue problems [5, 108, 109] that is supported by most major DFT software packages [8, 45, 52, 104, 122-132]. ELPA is capable of solving both *standard* and *generalized* eigenproblems: users are not required to perform the transformation from the *generalized* problem manually, although interfaces for each individual transformation step are also provided. Moreover, ELPA is ported to GPU accelerators (including NVIDIA, AMD, and Intel) [133, 134]. To the best of our knowledge, ELPA has recently set a new record [135] of a $3.2 \cdot 10^6 \times 3.2 \cdot 10^6$ dense matrix diagonalization for all eigenvalues and eigenvectors of the standard real-valued eigenproblem on the LUMI supercomputer, which is equipped with AMD MI250X GPUs.

Usability and Tutorials

Solver selection in FHI-aims

The numerical atomic orbital (NAO) basis sets in FHI-aims [8] are rather compact, i.e., the number of basis functions N required to represent the single-particle states with high precision is typically not larger than a single-digit multiple of n_{el} . Thus, since the number of basis functions N determines the dimension of H and S , typically a large fraction of their eigenpairs is needed. For this scenario and for small and mid-sized systems, direct eigensolvers, such as LAPACK, ScaLAPACK [136, 137] or ELPA are the solvers of choice.

One advantage of ELSI is its ability to enable direct, comparative benchmarks of different solvers within a single software framework, allowing one to make a choice that is specific to system and hardware characteristics of a particular computational campaign. Beyond the default serial (LAPACK) and parallel (ELPA) solvers available in FHI-aims, a range of additional solvers can be accessed through ELSI and are supported by FHI-aims directly by choosing appropriate keywords. At the time of writing, these include PEXSI, NTPoly, EigenExa, SLEPc-SIPS, libOMM, and MAGMA.

Past results for the NAO basis sets of FHI-aims showed that the crossover point beyond which different density-matrix-based solvers outperform ELPA lies beyond several hundred to several thousand atoms [7]. On parallel and/or GPU hardware, performing a direct eigenvalue solution using ELPA is therefore preferable for the majority of current production calculations in FHI-aims. For larger systems and for semilocal

Roadmap on Advancements of the FHI-aims Software Package

2.4. SOLVERS FOR LARGE-SCALE ELECTRONIC STRUCTURE THEORY: ELPA AND ELSI

DFT, the solution effort for the pole expansion and selected inversion (PEXSI) [110, 111] method scales at most quadratically with system size for three-dimensional systems, and even subquadratically for two- or one-dimensional geometries. Benchmarks in Ref. [7] showed that, for 1D systems, PEXSI outperformed a direct diagonalization for as few as 400 atoms. For 2D systems, the crossover was found around 1,000 atoms. Alternatively, for non-metallic systems, density matrix purification allows one to directly solve for the density matrix with a solution effort that grows linearly with the system size, independent of the density functional, as implemented, e.g., in the NTPoly [112] library for sparse matrix function calculations. Benchmarks in Ref. [7], for the DFTB+ code with significantly more sparse matrices than FHI-aims, show that the performance crossover depends heavily on the system type and matrix sparsity, and should be tested individually prior to production calculations for the largest systems.

ELPA - Usability

Within FHI-aims, ELSI and ELPA (as well as LAPACK calls for serial linear algebra steps) are automatically employed with parallelization strategies that leverage trivial parallelism (e.g., for separate k -points or spin channels) whenever possible. Since the performance of solvers other than ELPA depends on system, problem and hardware characteristics, their selection is a user choice by setting specific keywords.

One key consideration pertains to high-performance hardware beyond CPUs. Specifically, utilizing GPUs is essential for leveraging the computational power of modern supercomputers. This not only accelerates matrix diagonalization but also allows for the handling of larger matrices, enabling the investigation of physical systems with more atoms.

Much effort has been spent porting ELPA to GPUs. Nowadays ELPA fully supports both NVIDIA and AMD GPUs, including support for their respective GPU collective communication libraries, NCCL [138] and RCCL [139], respectively, for the most computationally intensive parts of the algorithm. Partial support is available for Intel GPUs using SYCL [140] and oneMKL [141], though this is currently limited to solving the standard eigenproblem.

For the core solver of the *standard* eigenproblem, ELPA provides implementations of the conventional one-stage diagonalization method (“ELPA1”) and the two-stage diagonalization (“ELPA2”) [142–144]. On CPUs, as a rule of thumb, the ELPA2 solver is preferable and it is also typically up to 1.5-2 times faster than MKL’s ScaLAPACK [145]. However, on GPUs, the ELPA1 solver is typically faster, providing a speedup of three times over the best setting on the CPU-only nodes if the local matrices per GPU are not extremely small [133]. If only a subset of all eigenvectors is computed (as is often the case in DFT), then ELPA2-GPU becomes more competitive compared to ELPA1-GPU as the number of targeted eigenvectors decreases.

For the *generalized* eigenproblem, until recently ELPA-GPU faced a significant bottleneck in the matrix-matrix multiplication step. ELPA relied on ScaLAPACK or its own implementations of the SUMMA [146, 147] and Cannon’s [148] algorithms for parallel matrix multiplication, none of which were fully GPU-ported until the 2024.05 release, leaving a substantial part of the computation to CPUs. Table 2.2 compares the ELPA 2023.11 and 2024.05 releases, with the latter including a full GPU port for the matrix-matrix multiplication (‘Multiply’). This is especially important for ‘NCCL’ setup (1 MPI task per GPU), where the total ‘Multiply’ time for forward and backward transformation is reduced from 559 to 25.6 seconds.

GPU porting of the matrix-matrix multiplication brings the performance of the ELPA-GPU *generalized* eigen-solver in line with that of the core *standard* eigenvalue solver, now achieving a similar speedup of approximately three to four times over the best-performing ELPA-CPU configuration. It is important to note that the CPU side of the comparison (last row of Table 2.2) involves all CPU cores on a given computational node. For instance, as shown in Table 2.2, we compare a GPU setup with 4 GPUs and tuned from 1 to 18 CPU cores

Roadmap on Advancements of the FHI-aims Software Package

2.4. SOLVERS FOR LARGE-SCALE ELECTRONIC STRUCTURE THEORY: ELPA AND ELSI

ELPA version	Forward transformation			Standard EVP			Backtr.	Total
	Choles.	Invert	Multiply	Tridiag.	Solve	Back	Mult.	
2023.11, MPS, 18 MPI per GPU	6.5	5.7	34.7	55.3	11.7	18.0	24.5	156.4
2024.05, MPS, 18 MPI per GPU	6.1	4.9	21.4	54.5	12.1	17.6	12.1	128.9
2024.05, MPS, 4 MPI per GPU	5.9	5.1	8.0	51.3	11.3	12.6	3.3	97.8
2023.11, NCCL, 1 MPI per GPU	2.8	2.1	294	49.7	40.4	15.3	265	669
2024.05, NCCL, 1 MPI per GPU	2.3	1.8	18.8	49.5	40.1	14.6	6.8	134
2024.05, CPU	27.3	23.4	72.7	54.4	42.9	141	30.3	393

Table 2.2: Comparison of the wall-clock times (in seconds) of the ELPA 2023.11 vs 2024.05 releases in different GPU and CPU setups for the individual steps of the *generalized* eigenproblem. The matrix size is 40960×40960 , and the diagonalization is performed to obtain all eigenvalues and eigenvectors on a single node on MPCDF's system *Raven* [150]. For the GPU calculations the ELPA1 solver with four NVIDIA A100 (40GB, SXM) GPUs was used. The 'MPS' configuration uses NVIDIA's Multi-Process Service [149] and several MPI processes per GPU. For the ELPA 2023.11 release, 18 MPI processes per GPU (full *Raven* node) show the best performance, while for ELPA 2024.05, the best performance is achieved with 4 MPI processes per GPU. The 'NCCL' configuration uses the NVIDIA Collective Communication Library [138], which is constrained to 1 MPI process per GPU. For the CPU calculations the ELPA2 solver with 72 MPI processes on two 36-core Intel Xeon Platinum 8360Y processors was used.

per GPU (using ELPA1) against the CPU setup with all 72 CPU cores (using ELPA2). At the moment, FHI-aims is best tested with NVIDIA MPS [149], as it allows multiple MPI processes per GPU to be used efficiently, which can be particularly beneficial for steps like 'Solve'. NVIDIA MPS using all CPU cores of a node seems still to provide a very good speedup compared to the CPU-only ELPA (cf. Table 2.2). In cases where ELPA dominates the FHI-aims runtime, it is worthwhile to tune for the best MPI process per GPU ratio. For the future development of ELPA, as more of its parts being ported to GPU over time, we expect that the 'NCCL' code path will become more performant than the 'MPS' one. This is because NCCL enables direct data transfer between GPU devices, eliminating the need for costly GPU-CPU memory copies required for MPI operations.

Overall, we recommend using ELPA-GPU when possible: it can provide up to a 4x speedup for the complete solution of standard and generalized eigenproblems, and even up to a 10x speedup for individual solution steps. Despite this drastic improvement provided by the ELPA 2024.05 release, the 'Multiply' step still dominates the time for forward and back transformations from *generalized* to *standard* eigenproblem, motivating its further optimization.

ELSI supports ELPA-GPU (NVIDIA only) directly in ELPA's 2020 release, and as an externally compiled library for later versions and other GPU types. Since newer ELPA versions offer significant performance enhancements on GPUs, it is recommended to use ELSI with externally linked ELPA.

Tutorials

Within the FHI-aims ecosystem, ELPA and ELSI are automatically used in standard tutorials, however, ELSI itself comes with its own extensive manual in its repository. Within the standard build process of FHI-aims, the 2020.05.001 version of ELPA is included as a default build step. A newer version of ELPA can be included first by compiling an external version of the ELPA library and then linking to this version in a subsequent build of FHI-aims; this process is described as a dedicated tutorial accessible at <https://fhi-aims-club.org/>

b.gitlab.io/tutorials/tutorials-overview/.

Since the 2024.05 release, ELPA provides a comprehensive and self-contained *ELPA Manual: User's Guide and Best Practices* [109]. This manual offers ELPA's quick-start guide and code examples, as well as detailed instructions on installation, how to use ELPA in applications, and troubleshooting. It is an essential resource for both new and experienced users, ensuring optimal performance and integration of ELPA in applications. Additionally, we provide a GitHub repository containing teaching materials from recent tutorials on ScaLAPACK and ELPA [151]. The repository includes presentation slides and various code examples that demonstrate the practical use of ELPA, covering both the CPU and GPU versions of the library.

Future Plans and Challenges

Several key optimizations are planned for ELPA to further improve its performance on modern HPC systems. The primary focus is on enhancing the GPU solvers for the standard eigenproblem, particularly focusing on the ELPA2 tridiagonalization, the solve step of the tridiagonal matrix, and the backtransformation steps, which are not yet utilizing the GPU collective-communication libraries. This should allow the code to keep all the data on the GPU memory, avoiding the costly data transfers between the CPU and GPU. Overall, we expect that with more and more parts of ELPA being ported to GPU, using NCCL/RCCL/oneCCL collective communication libraries allowing direct GPU-GPU communication will become more advantageous; in particular, we expect that ELPA's 'NCCL' codebranch will outperform the 'MPS' one in the near future. However, the use of NCCL and alike will require a reorganization of the management of the GPU use in FHI-aims: it is necessary to set up the dense matrices directly in the GPU memory corresponding to a given MPI task, instead of first distributing them in parallel across all CPUs.

Additionally, work is underway to optimize ELPA's parallel matrix-matrix multiplication routine, which is used for the transformation of the generalized to the standard eigenproblem. Although the GPU implementation has already reduced the bottleneck as we reported in this contribution, the matrix multiplication remains the most time-consuming part of the process.

Expanding support for Intel GPUs is another major goal. Currently, ELPA's functionality on Intel GPU hardware is limited to solving the standard eigenproblem. We plan to utilize oneCCL and further oneMKL operations to achieve feature parity with the other GPU implementations in ELPA.

Generally, ELPA aims to support and be optimized for all modern and emerging hardware architectures, including Accelerated Processing Units (APUs) [152] and Vector Processing Units (VPUs) [153]. Notably, there is an ongoing effort to port ELPA kernels to the RISC-V architecture [154].

ELSI will need similar architecture-specific optimizations. Additionally, support for a broader ecosystem of new and emerging solvers is a constant target, with recent, community-supported additions to ELSI of the Chebyshev eigensolver ChaSE [115, 116] and a new, distributed linear algebra eigensolver DLA-Future [120, 121] require implementation and testing in FHI-aims. For example, demonstration calculations of extremely large DFT calculations were performed on Google's proprietary Tensor Processing Units (TPUs) [155], based on an adaptation of the FHI-aims/ELSI software stack. This adaptation also made use of an efficient combination of single- and double-precision solutions during the self-consistent field cycle. Similar use of faster single-precision solutions in ELPA is possible using a reduced, frozen-core eigenvalue solution via ELSI and FHI-aims, offering further optimization potential in the future [156].

Acknowledgements

We are grateful to Marc Torrent for sharing his preliminary ELPA benchmarks with ABINIT and for fruitful discussions, which have triggered the work on optimizing the generalized eigenproblem solver in ELPA-GPU. We gratefully acknowledge the support provided by Markus Hrywniak from NVIDIA. We also thank Markus Rampp for the useful comments on the initial draft of the manuscript. This project was supported by NOMAD Center of Excellence (European Union's Horizon 2020 research and innovation program, Grant Agreement No. 951786) and the embedded CSE programme of the ARCHER2 UK National Supercomputing Service (<http://www.archer2.ac.uk>). We acknowledge CSC for awarding us access to LUMI supercomputer (Kajaani, Finland). ELSI was supported by the National Science Foundation (NSF), USA under Award No. 1450280. This research used resources of the Argonne Leadership Computing Facility, a U.S. Department of Energy (DOE) Office of Science user facility at Argonne National Laboratory and is based on research supported by the U.S. DOE Office of Science-Advanced Scientific Computing Research Program, under Contract No. DE-AC02-06CH11357. We also thank all the colleagues who contributed to ELSI/ELPA in the past and/or in the context of other projects, specifically: Jianfeng Lu, Lin Lin, Fabiano Corsetti, Weile Jia, Raul Laasner, Yingzhou Li, David B. Williams-Young, Xinzhe Wu, Edoardo di Napoli, Rocco Meli, Carolin Penke. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

2.5 Electronic-Structure Analysis Tools in Numeric Atomic Orbital Framework

Volker Blum^{1,2}, Simon Erker³, James A. Green⁴, Oliver T. Hofmann^{3,5,a}, Simon Hollweger³, Lukas Hörmann⁷, *Sebastian Kokott^{4,6}, Hagen-Henrik Kowalski⁶, Sergey V. Levchenko^{5,8,b}, Chi Liu², Teruyasu Mizoguchi⁹, Mariia Pogodaeva⁸, Norina A. Richter⁵, Mariana Rossi¹⁰, Matthias Scheffler⁶, Kiyoo Shibata⁹, Ruyi Song², Aloysius Soon^{5,c}, Izumi Takahara⁹, Yi Yao^{1,4}, and Rundong Zhao¹¹

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Institute of Solid State Physics, Graz University of Technology, Petersgasse 16/II, 8010 Graz, Austria

⁴Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁵Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Department of Chemistry and Department of Physics, University of Warwick, Coventry, CV3 5AB, UK

⁸Skolkovo Institute of Science and Technology, Moscow 121205, Russia

⁹Institute of Industrial Science, The University of Tokyo, Tokyo 153-8505, Japan

¹⁰Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

¹¹School of Physics, Beihang University, Beijing 102206, China

*Coordinator of this contribution.

^aCurrent Address: Institute of Solid State Physics, Graz University of Technology, Petersgasse 16/II, 8010 Graz, Austria

^bCurrent Address: Skolkovo Institute of Science and Technology, Moscow 121205, Russia

^cCurrent Address: Department of Materials Science & Engineering, Yonsei University, Seoul 03722, Republic of Korea

Summary

Modern first-principles electronic-structure methods allow us to calculate the wavefunctions and energies for ground and excited electronic states. In principle, this provides a complete description of system's physical and chemical properties in a wide range of temperature and pressure conditions. However, the information contained in a many-electron wavefunction, or even in the set of one-electron states and energies obtained from effective-potential theories (such as Kohn-Sham density-functional theory or the Hartree-Fock approximation) is too vast and redundant to be useful for physical understanding without further processing. Therefore, a set of electronic-structure analysis tools have been developed, including density of states (total and projected), band structure, charge and spin partitioning, visualization of electronic density and density differences, isosurfaces of one-electron states and their densities, and electron-localization functions. In this Chapter, these tools and some particularities of their implementation in a numeric atomic

2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK

orbital (NAO) framework are described.

Current Status of the Implementation

Density of States (DOS) and Band Structure (BS)

Total DOS is calculated from one-electron energies $\epsilon_{n\mathbf{k}}$ as follows:

$$g(\epsilon) = \frac{1}{F} \sum_{n\mathbf{k}} w_{\mathbf{k}} \delta(\epsilon - \epsilon_{n\mathbf{k}}), \quad (2.27)$$

where the \mathbf{k} -point grid is taken in the first Brillouin zone (BZ), $w_{\mathbf{k}}$ are \mathbf{k} -point weights (without any spatial symmetry $w_{\mathbf{k}} = 1/N$, where N is the total number of \mathbf{k} -points in the grid), and F is a normalization factor. This factor reflects physical scaling of the number of states per energy unit with the system size. By default F is set to 1, implying normalization per unit cell. When comparing DOS for different unit cells, however, it should be set to the cell volume or another factor that scales with the cell size.

Projected DOS (PDOS) in FHI-aims is based on Mulliken partitioning, and is calculated as follows:

$$g_{\nu}(\epsilon) = \frac{1}{F} \sum_{\mu} \sum_{n\mathbf{k}} w_{\mathbf{k}} \langle \psi_{n\mathbf{k}} | \phi_{\mu} \rangle S_{\mu\nu}^{-1}(\mathbf{k}) \langle \phi_{\nu} | \psi_{n\mathbf{k}} \rangle \delta(\epsilon - \epsilon_{n\mathbf{k}}), \quad (2.28)$$

where $\psi_{n\mathbf{k}}$ are one-electron wavefunctions, ϕ are a set of localized functions, ν, μ are indices of localized orbitals, and $S_{\mu\nu}(\mathbf{k})$ is the overlap matrix. By construction, $\sum_{\nu} g_{\nu}(\epsilon) = g(\epsilon)$. In FHI-aims, NAOs are used to calculate the projections $\langle \phi_A | \psi_{n\mathbf{k}} \rangle$. These orbitals are in general non-orthogonal, and interpretation of PDOS in terms of NAO contributions becomes more ambiguous with increasing basis set size. Therefore, PDOS should be used only as a qualitative analysis tool. In practice, the equation 2.28 is evaluated via Kohn-Sham coefficients $c_{n\mathbf{k}}^{\alpha}$ in wavefunctions $\psi_{n\mathbf{k}} = \sum_{\alpha} c_{n\mathbf{k}}^{\alpha} \sum_{\mathbf{R}} \phi_{\alpha}(\mathbf{r} + \mathbf{R}) e^{i\mathbf{k}\mathbf{R}}$ (\mathbf{R} is a lattice vector):

$$g_{\nu}(\epsilon) = \frac{1}{F} \sum_{\mu} \sum_{n\mathbf{k}} w_{\mathbf{k}} (c_{n\mathbf{k}}^{\nu})^* S_{\nu\mu}(\mathbf{k}) c_{n\mathbf{k}}^{\mu} \delta(\epsilon - \epsilon_{n\mathbf{k}}), \quad (2.29)$$

One can also use molecular or atomic-cluster orbitals as the projection functions ϕ , resulting in molecular orbital (MO) projected DOS (MODOS).

In crystal/MO overlap population (COOP/MOOP) analysis[157], orbital overlap is quantified to characterize chemical bonds (bonding, antibonding, or non-bonding). This is achieved by omitting the sum over basis functions μ in equation 2.28 and analyzing the obtained pair-contributions to DOS[158].

By convention, BS (the dependence of $\epsilon_{n\mathbf{k}}$ on \mathbf{k}) is plotted along straight lines connecting special points in the first BZ. The special points depend on the crystal symmetry, and are tabulated for most common crystal structures[159]. Similar to DOS, BS can be resolved in terms of NAO contributions by projecting $\epsilon_{n\mathbf{k}}$ onto NAOs. The projection is part of a Mulliken charge partitioning[160, 161].

In supercell calculations, for example for modelling defects or thermally perturbed crystal structures, bands with a \mathbf{k} -vector from the larger first BZ of the primitive cell fold into the smaller BZ of the supercell. A special procedure called band unfolding is required to recover important information about topology of electronic bands. Band unfolding is implemented in FHI-aims and is discussed in detail in Chapter 6.5.

*Roadmap on Advancements of the FHI-aims Software Package***2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK****Charge Partitioning (CP)**

CP allows one to assign a certain charge or spin to a particular orbital and/or atom. This is a useful tool for analyzing charge/spin redistribution in a system compared to, for example, free atoms or another system. However, there is no unique way to perform charge partitioning, since electrons in a many-electron system are often shared by several atoms. One can distinguish orbital-based (Mulliken[160, 161]), density-based (Hirshfeld[162, 163] and Bader[164]), and potential-based (by fitting an electrostatic potential, ESP) partitioning schemes. The ESP charges can be quite useful as they correctly reproduce electrostatics of the system. All of these methods except Bader partitioning are implemented in FHI-aims for both periodic and non-periodic systems. Bader partitioning and electron localizability indicator (ELI-D, a flavor of electron localization function, ELF, discussed in the next section) are accessible via the FHI-aims interface with the DGrid program package[165].

Berry-phase polarization[166] allows one to calculate (up to an integer number of quanta) polarization \mathbf{P} of an infinite periodic crystal with a band gap. Polarization can be used to calculate Born effective charges for any atom A as $Z_{A,ij} = \frac{V}{e} \frac{\partial P_i}{\partial R_{A,j}}$, where V is the unit cell volume, P_i is i th component of vector \mathbf{P} , and $R_{A,j}$ is the j th coordinate of atom A . Berry-phase polarization and Born effective charges are useful tools for understanding properties of systems with broken inversion symmetry, such as ferroelectrics, or polarization changes during dynamics. The details of implementation in FHI-aims can be found in Ref.[167].

Electron Localization

Solution of the electronic problem in a mean-field approximation usually results in delocalized one-electron states. However, observables such as total energy or electron density are invariant with respect to any unitary transformation among states, provided occupied and unoccupied states are not mixed. This freedom is used in FHI-aims to find MOs that are more localized and therefore easier to interpret in the context of chemical bonding.

An interesting way to analyze localization of electrons is the ELF[168]. Different from the orbital transformations, which can be cast into transformations of a one-particle density matrix, ELF is a way of understanding a two-particle density matrix. Topological analysis of the ELF yields regions of local pairing of electrons, where the ELF is maximal. ELF gives additional information to the widely used Bader topological analysis[164], as demonstrated in [168] by revealing the shell structure of heavy atoms using the ELF. Flavors[169, 170] of the original formulation[168] are also available in FHI-aims. Since the ELF is a function in three dimensions, software that can plot volumetric data is necessary to visualize it. FHI-aims can output a volumetric data file (in CUBE format[171]) for such a visualization.

Volumetric Data (electron density, electrostatic potential, etc.)

Three-dimensional visualization can provide vast amounts of information for understanding electronic structure. Currently FHI-aims can output data for visualizing objects listed in Fig. 2.5.

Usability and Tutorials

The necessary input parameters and methods for plotting (P)DOS and band structure are shown in Fig. 2.6. In practice, the δ -function in equations 2.27 and 2.28 is replaced by a Gaussian function at each \mathbf{k} -

Roadmap on Advancements of the FHI-aims Software Package

2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK

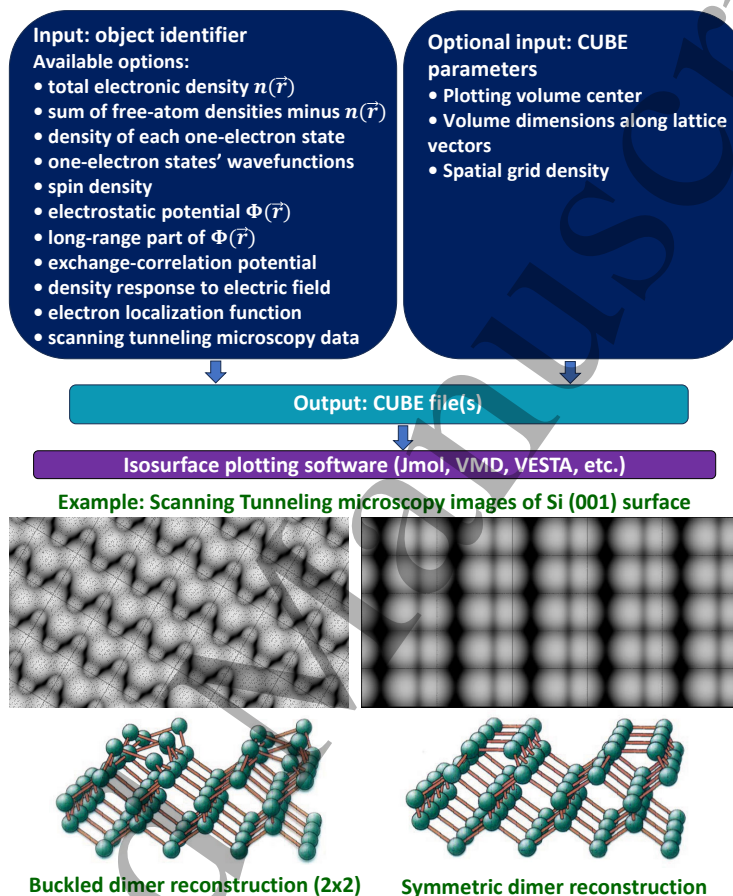


Figure 2.5: CUBE^[171] output infrastructure in FHI-aims. The density of each one-electron state can be plotted for different k -points and spin-channels, when relevant. Both real and imaginary parts of electronic wavefunctions can be plotted, if available. Spin density is defined as the difference between spin-majority and spin-minority densities. The long-range part of the electrostatic potential is defined by Ewald partitioning. Density response to an applied electric field needs a perturbation theory calculation. ELF can be calculated in different flavors^[168–170]. Scanning tunneling microscopy (STM) data are calculated in the Tersoff-Hamann approximation^[172]. Simulated STM images of the Si(001) surface with (left) and without (right) buckled dimer reconstruction, calculated with DFT-PBE functional, are shown as an example. The STM images were produced with the VMD software package (see https://fhi-aims-club.gitlab.io/tutorials/stm_visualization/ for a tutorial).

Roadmap on Advancements of the FHI-aims Software Package

2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK

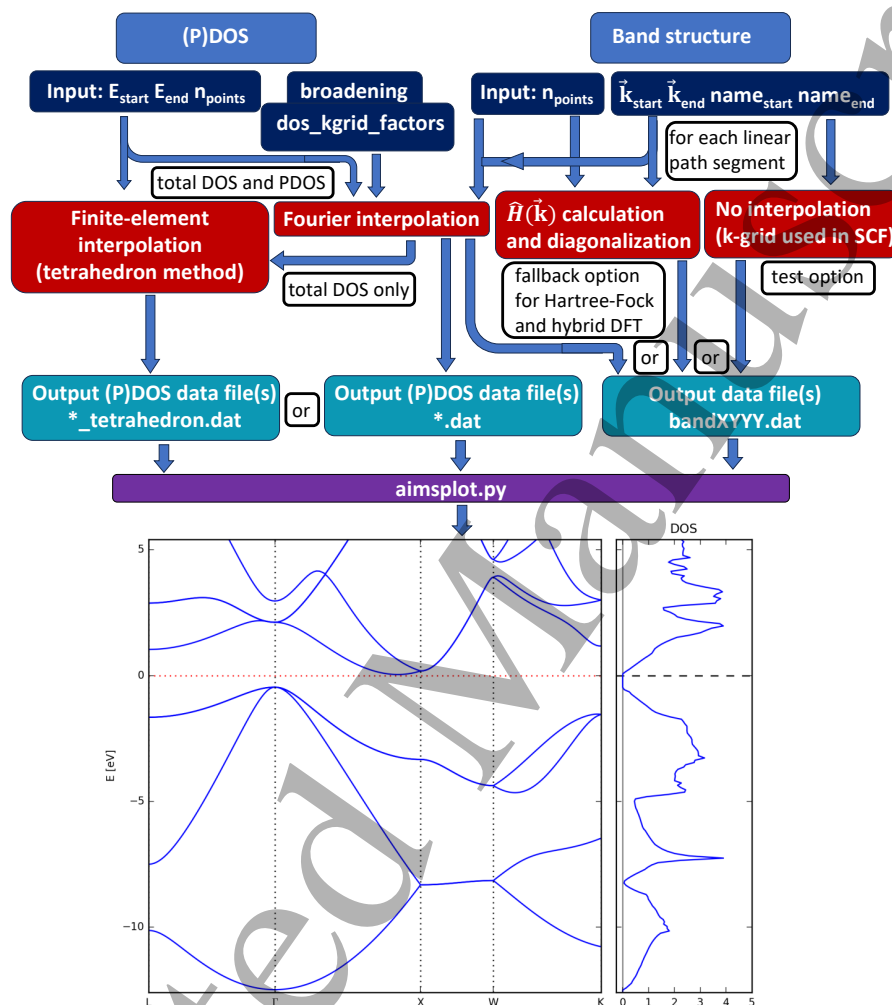


Figure 2.6: Available functionality for DOS and band structure calculations and plotting. E_{start} and E_{end} are starting and ending points, respectively, on the energy axis for DOS plotting (in eV), n_{points} is the number of points on the energy axis for DOS and along a linear path segment in k -space for band structure plotting. \vec{k}_{start} and \vec{k}_{end} are coordinates of starting and ending points of a linear path segment in k -space in the basis of reciprocal lattice vectors. Names of starting and ending points of each path segment are given by $name_{start}$ and $name_{end}$, respectively. An example combined band-structure/DOS plot obtained with python script `aimsplot.py` is also shown. The script is part of FHI-aims distribution.

Roadmap on Advancements of the FHI-aims Software Package

2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK

point. Each Gaussian is normalized to 1, and its width is a parameter (denoted "broadening" in Fig. 2.6). To get a converged DOS, one needs to increase the k -grid density and decrease the width. In order to achieve this in a reasonable computational time, an interpolation can be used to calculate ϵ_{nk} on a k -grid denser than the one used in the actual SCF calculation. One such interpolation scheme is the tetrahedron method[173, 174]. With this method, the Gaussian width is not requested, since it is calculated automatically. Another interpolation scheme exploits the locality of the NAO basis set. In this basis set, one can calculate the Hamiltonian matrix in real space for a set of lattice vectors \mathbf{R} . Then the Hamiltonian matrix on a dense k -point grid (defined by the parameter "dos_kgrid_factors" in Fig. 2.6) can be calculated by a Fourier transform of the real-space matrix (Fourier interpolation), and then diagonalized to yield new ϵ_{nk} without self-consistency. This approach is sometimes called perturbative. Both the tetrahedron (the recommended way to calculate (P)DOS) and perturbative approaches are implemented in FHI-aims. Moreover, they can be combined to improve total DOS (but currently not PDOS) convergence even further. For PDOS, there are three choices for the sets of ϕ_A is FHI-aims: atom, species, and MO. For atom sets, PDOS is output for every atom separately, while for species sets, sum of PDOS for all atoms of the same species are calculated. An example of species PDOS is shown in Fig. 2.7.

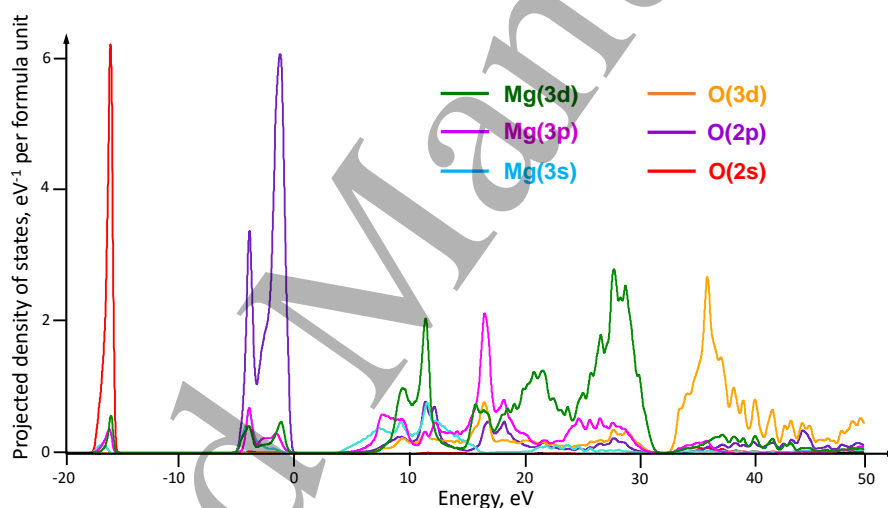


Figure 2.7: Species PDOS for MgO calculated with DFT-PBE, tight settings.

In the case of spin-polarized calculations, DOS for the different spin-channels can be calculated and analyzed separately. Moreover, DOS can be calculated with spin-orbit coupling (SOC) post-SCF correction[48]. For calculations without periodic boundary conditions, DOS becomes a set of separate states (some of which can be degenerate), since there are no k -points and no bands. PDOS/MODOS can be used to analyze the orbital character of states in such systems as well.

Just as for DOS, a k -space interpolation scheme is needed to converge BS. In FHI-aims this can be done with the perturbative approach. Bands can be also calculated with SOC post-SCF correction [48]. FHI-aims allows one to characterize bands in terms of spin texture, i.e., expectation values of Pauli matrices for spinors at every nk [98, 175] in calculations with SOC.

There is a difficulty in perturbative interpolation from real to reciprocal space of electronic structure when

*Roadmap on Advancements of the FHI-aims Software Package***2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK**

using non-local potentials, as, for example, is the case for exact exchange in hybrid functionals, or the Hartree-Fock approximation. For a non-local potential the Hamiltonian decays slowly with distance. For a given \mathbf{k} -grid the electronic problem for non-local potential can be solved efficiently by folding in the long-range effects into a \mathbf{R} -dependent matrix for a moderate number of lattice vectors \mathbf{R} (forming a so-called Born-von Karman supercell). However, this matrix may not be good enough for the Fourier interpolation on a denser \mathbf{k} -grid. This problem can be addressed by either increasing the \mathbf{k} -grid density in the SCF calculation, or by using a much less computationally efficient \mathbf{k} -space implementation of exact exchange. This is relevant for both DOS and BS calculations.

Tutorials are freely available online on how to compute BS and (P)DOS in FHI-aims. In particular, Part 3 and the Appendix of the “Basics of Running FHI-aims” (<https://fhi-aims-club.gitlab.io/tutorials/basics-of-running-fhi-aims/>) details the keywords required, output produced, and plotting tools available in FHI-aims, and compares the interpolative approaches to computing the total DOS.

For molecular systems, ESP charges are found by minimizing the difference between the actual ESP (from the electronic-structure calculation) and the ESP due to charges, under the condition that the sum of all charges is equal to the total charge of the system. The minimization is performed on a grid of points in space that fit between spheres around each atom, with the radii of these spheres a multiple of van der Waals radii for corresponding atoms. For periodic systems, two different methods are implemented[176, 177]. For the method in Ref.[176] one needs to provide electronegativity, self-Coulomb interaction, and weighting factors for each atom to ensure physically reasonable resulting charges. In the second method[177] charges for constraining ESP charges have to be provided directly, for example from Mulliken or Hirshfeld partitioning schemes.

For localization of MOs in non-periodic systems, the unitary optimization of localized MOs, developed by Lehtola and Jónsson[178], with the Pipek-Mizey localization method[179] is implemented in FHI-aims. The localization method requires a state-projected population matrix on each atom, and thus relies on a partitioning scheme. Both Mulliken and Hirshfeld partitioning schemes can be used in the localization procedure in FHI-aims. In addition, maximally localized Boys MOs can be calculated, following the procedure of finding the localizing unitary transformation described in Ref. [180].

FHI-aims can be requested to output volumetric data for any of the quantities listed in Fig. 2.5 in the Gaussian CUBE file format [171]. Several visualization software packages (e.g., Jmol, VMD, VESTA) can plot isosurfaces using the volumetric data from CUBE files. An example of such a plot is shown in Fig. 2.8.

Future Plans and Challenges

As was demonstrated above, the NAO basis and its efficient implementation in FHI-aims provides a powerful platform for electronic-structure analysis. There are many more possibilities for implementation of useful analysis tools in the future. Among ongoing developments are Wannier functions for periodic systems, Bader charge and magnetic moment analysis, and a combined perturbative/tetrahedron interpolation scheme for PDOS. An improved version of the tetrahedron method[181] is also desired.

In general, the FHI-aims development team is listening attentively to the requests from community regarding electronic-structure analysis tools. Do not hesitate to contact the team if you need a particular functionality, or if you are willing to implement the functionality in FHI-aims yourself. The only challenges here are the labor and computational costs. The latter is mainly associated with the computational cost of the underlying electronic-structure theory. For example, as discussed above, band-structure calculations can

Roadmap on Advancements of the FHI-aims Software Package

2.5. ELECTRONIC-STRUCTURE ANALYSIS TOOLS IN NUMERIC ATOMIC ORBITAL FRAMEWORK

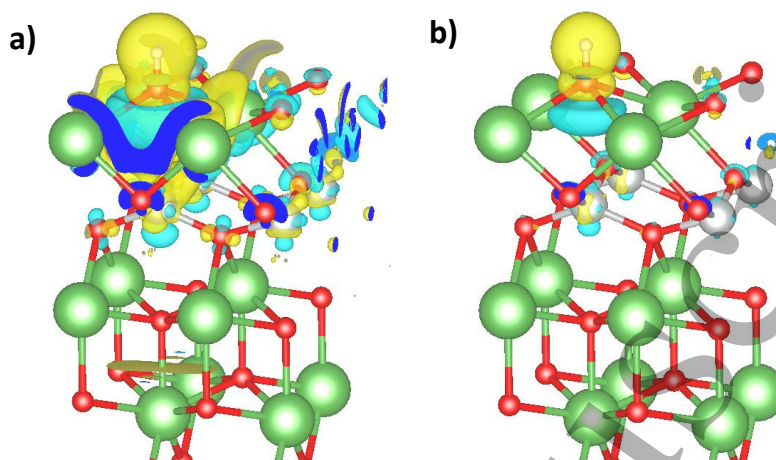


Figure 2.8: Electron density change upon adsorption of a hydrogen atom on a (001) surface (2×2 surface supercell) of La_2NiO_4 Ruddlesden-Popper oxide, calculated with the DFT-RPBE functional, tight settings. Oxygen atoms are shown as red spheres, Ni - grey, La - green, H - white. The yellow-colored isosurface shows electron depletion, cyan - electron accumulation. The isosurface value is $0.01 \text{ |e|/\text{\AA}^3}$ in panel a), and $0.03 \text{ |e|/\text{\AA}^3}$ in panel b). The image was obtained using the VESTA program and a CUBE file, obtained from FHI-aims-generated CUBE files for a surface with adsorbed H, a clean surface, and a free H atom. The positions of atoms in all systems are the same as in the relaxed surface with adsorbed H.

be quite expensive with hybrid functionals. Beyond-DFT approaches, such as GW and other many-body perturbation theory methods, are even more expensive, and the automatic functionality for DOS and band-structure calculations with these methods was not implemented for this reason. Development of robust interpolation schemes, as well as increasing efficiency of the methods themselves, e.g., by using graphical accelerators, will resolve these issues.

Acknowledgements

We acknowledge contributions from Frank Wagner and Alim Ormeci to the development of the interface between FHI-aims and DGrid. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

2.6 Explicit and Implicit Embedding Approaches

Daniel Berger¹, Volker Blum^{2,3}, Gabriel A. Bramley⁴, Matthew R. Farrow⁴, Jakob Filser⁵, Matthias Kick⁵, *Andrew J. Logsdail⁶, Harald Oberhofer⁷, Karsten Reuter⁵, Stefan Ringe⁸, Pavel V. Stishenko⁶, Oscar van Vuren⁶, Daniel Waldschmidt⁹, and Yi Yao^{2,10}

¹Chair for Theoretical Chemistry and Catalysis Research Center, Technische Universität München, Lichtenbergstr. 4, D-85747 Garching, Germany

²Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

³Department of Chemistry, Duke University, Durham, NC 27708, USA

⁴Department of Chemistry, Kathleen Lonsdale Materials Chemistry, University College London, London, United Kingdom

⁵Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶Cardiff Catalysis Institute, School of Chemistry, Cardiff University, Park Place, Cardiff CF10 3AT, United Kingdom

⁷Department of Physics and Bavarian Center for Battery Technologies, University of Bayreuth, Bayreuth, Germany

⁸Department of Chemistry, Korea University, Seoul 02841, Republic of Korea

⁹Technische Universität München (TUM), Munich, Germany

¹⁰Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

*Coordinator of this contribution.

Summary

Applied electronic structure calculations often benefit from approaches that simplify the system of interest, as approximate models reduce the number of atoms and electrons in a system to make the calculations more tractable; for example, the simulation of molecular adsorption on an extended surface is often performed with fixed-size periodic cells, finite-thickness surface slab models, a vacuum environment, and without thermal effects, which are all simplifications of the laboratory reality. However, models of real chemical systems can become inaccurate with such an approach, and this motivates efforts towards hierarchical representations where the all-electron site or species of interest is embedded within a coarser representation of the environment. Such multiscale embedding approaches typically ensure that the highest accuracy is maintained on the sites or species of interest, whilst the environment response is sufficiently captured and its energy contributions are included. The most common field of application for multiscale embedding is biological chemistry [182, 183], though there is demonstrable value for translation to homogeneous and heterogeneous systems, and application in specific fields such as electrochemistry, where long-range interactions can play a crucial role [184–186].

Embedding approaches apply either an explicit or implicit effective representation of the environment, as schematically shown in Fig. 2.9. In **explicit embedding**, basis centres (typically atomic sites) in the embedding environment can be coarsened, by removing electronic degrees of freedom and instead centering an effective embedding potential at the same point in the form of a monopole, dipole, or higher order multi-

Roadmap on Advancements of the FHI-aims Software Package

2.6. EXPLICIT AND IMPLICIT EMBEDDING APPROACHES

pole. Alternatively, hierarchical basis representations and/or approximate treatment of core electrons (such as freezing or pseudoising of electrons) can be applied in the embedding environment, providing greater subtlety in the form of the applied embedding potential [156, 187]. The latter approaches are particularly important where directed bonding interactions influence the active site. In contrast, **implicit embedding** considers the holistic dielectric response to/from an encapsulating medium for the active site or species. Often, the embedding medium is considered to be liquid, and response to any stimulus is represented in the all-electron calculation; directional bonding is therefore not considered, but rather the correct energetics of the system in its surroundings are approximated [184].

Current Status of the Implementation

FHI-aims has infrastructure that supports both implicit and explicit embedding approaches, including connectivity to external packages that can act as drivers for calculation workflows [188]. The FHI-aims functionality allows for the energy and forces of a system to be calculated self-consistently under the influence of an embedding environment.

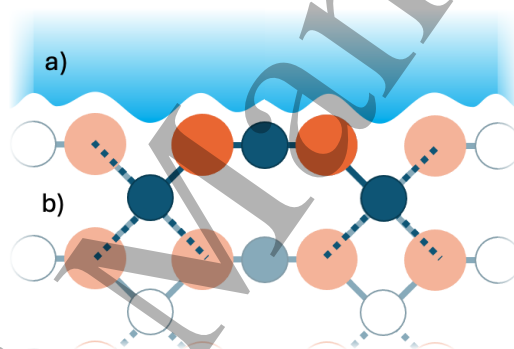


Figure 2.9: Schematic representations of: a) **Implicit embedding** environment; and b) **Explicit embedding** environment. The fully visible species represent quantum mechanical atoms of interest, with blue and orange spheres representing cation and anion species, respectively. For implicit embedding, the blue medium represents the surrounding environment with a dielectric, ϵ ; in the explicit embedding model, the partially transparent species represent pseudoised near-neighbours, and white spheres represent the long-range embedding environment that may be included as, *e.g.*, multipolar charges.

Explicit embedding is built on the all-electron approach used in a standard FHI-aims calculation, with the effective embedding potential from the surrounding environment captured through surrogate models centred at atomic sites. The embedding environments can take a coarse multipolar form, which is trivially included in the one-electron contributions to the Fock matrix in a manner similar to nuclei; however, this representation typically lacks the subtlety of the density distribution on a real atomic species, and therefore pseudopotential-type embedding is also available. The pseudopotential implementation in FHI-aims takes a Kleinman-Bylander form [187], which is separable into local and non-local components with additional terms for the non-linear core correction. The pseudopotentials can be particularly valuable for simulations of solid materials with ionic bonding [189–191]. For effective management of explicit embedding simulations, coupling FHI-aims with external packages that are designed for multiscale simulations is valuable for

Roadmap on Advancements of the FHI-aims Software Package

2.6. EXPLICIT AND IMPLICIT EMBEDDING APPROACHES

clear partitioning of the subsystems that require all-electron representation. FHI-aims presently supports an interface to the QM/MM wrapper, ChemShell [188, 192], which handles the appropriate labelling and geometry specification of pseudopotential centres (when close to the active site of interest), and/or multipolar charges (when at further distance). The infrastructure in FHI-aims is currently suitable for molecular calculations of energies, forces, and other observables.

Implicit embedding takes a contrasting holistic approach by representing the surrounding solvent environment as an effective embedding medium. From a classical electrostatic perspective, the integration of the environment's degrees of freedom is achieved by modifying the Poisson equation to the generalised form:

$$\nabla[\epsilon_0\epsilon(\mathbf{r})\nabla\Phi(\mathbf{r})] = -4\pi(n(\mathbf{r}) + n_{\text{ion}}[n(\mathbf{r})]) \quad (2.30)$$

The solution of Eq. (2.30) provides the electrostatic potential, $\Phi(\mathbf{r})$, given the all-electron density representation of the solute, $n(\mathbf{r})$, the possible charge density of solvent ions, n_{ion} , and the dielectric response of the solvent, $\epsilon(\mathbf{r})$, which is commonly functionalised based on the electron density tail of the solute [184].

FHI-aims offers three implicit embedding approaches based on the generalised Poisson equation. An additional interface with the Environ package provides further diversity of options for implicit embedding [193, 194].

The first approach is the *multipole-expansion* (MPE)[195] method, where the dielectric function is represented by a sharp step function of the electron density, leading to a discrete interface between the environment and the embedded system. The sharp interface in MPE reduces Eq. (2.30) to a problem that can be cast into an overdetermined system of linear equations (SLE), and the least squares solution can be solved efficiently using standard algorithms. The electrostatic potential in MPE is represented in a finite basis-set expansion, and computational efficiencies can be obtained by separating the regions in the solute 'cavity' space into atom-centered sub-regions to create multiple sub-cavities (MPE-*nc*), limiting the number of basis functions inside each sub-cavity to a size-independent finite number. The outcome is greatly reduced cost when preparing the SLE, as well as opportunity to apply a sparse solver algorithm.

The second implicit embedding approach is the *Stern layer modified Poisson-Boltzmann* (MPB) method, where a smooth dielectric permittivity function of the electron density is applied, which then requires Eq. (2.30) to be solved over the whole computational domain. MPB allows for higher flexibility in the model, and also allows the introduction of arbitrary forms for the ionic charge density in the solvent, n_{ion} , parameterised as a function of the electron density. The increased complexity of the interface leads inherently to higher computational cost, with MPE requiring negligible computation time compared to MPB.

The third implicit embedding approach is the *conductor-like screening model* (COSMO).[196]. Instead of solving the Poisson equation with a modified $\epsilon(\mathbf{r})$, COSMO uses $\epsilon = \infty$ in the medium, independent of its dielectric constant. This choice of boundary condition enables the solution of the Poisson equation through a more tractable boundary value problem of electric field inside a conductor. To account for the effect of a finite dielectric constant, an empirical scaling function is used to adjust the surface charge, given by:

$$f(\epsilon) = \frac{\epsilon - 1}{\epsilon + k}, \quad (2.31)$$

where k is an empirical parameter, typically set to 0.5 or 0. We implemented the smooth version of COSMO by York and Karplus[197], which eliminates discontinuities with respect to the atomic positions present in the original COSMO formulation.

Roadmap on Advancements of the FHI-aims Software Package

2.6. EXPLICIT AND IMPLICIT EMBEDDING APPROACHES

Usability and Tutorials

Explicit embedding approaches using multipolar charges and pseudoised atoms are realised via inclusion in the model geometry file (`geometry.in`) using the definitions `multipole` or `pseudocore`, respectively. For the pseudocode species, a standard species definition is also required with the calculation settings, including integration grids and a minimal basis, and with specific direction to a suitable `.cp1` format file containing the pseudopotential details [15, 198]. If the pseudopotentials are being used as a replacement for an all-electron species, the basis functions must be removed for subsequent applied calculations and a representative charge must be set appropriately, as detailed in the software manual. FHI-aims also supports the non-linear core correction, which remedies the non-physical linearisation of the exchange-correlation density functional [187]. Multipole species are simpler to define, requiring definition of location, multipole order, and charge in the system geometry, with no contribution in the `control.in` file. Forces are available via the Hellmann-Feynman formalism, with the `qmmm` keyword necessary in the `control.in` file for forces on multipoles. For all-electron species, spurious Coulomb singularities have been observed when multipoles spatially overlap with an integration grid point (*i.e.*, around real atoms); these singularities can be avoided by ensuring reasonable distance separation ($> 5 \text{ \AA}$) between real and embedding species. A full tutorial showcasing how explicit embedding calculations are performed with FHI-aims and ChemShell is given in the software tutorials [199].

Implicit embedding is available via a range of input keywords that manage the solvent model, and respective subsettings. The `solvent` keyword selects between the different methods. The options available for MPE include single cavity, piece-wise cavity (MPE-*nc*), and piece-wise solvent representation (MPE-*ncps*), with the latter supporting both molecular and periodic system representations [195, 200]. For MPE-*nc* and MPE-*ncps*, default settings are optimised for water as a solvent, but physical model parameters can also be set by the user. These parameters include the solvent dielectric constant, ϵ ; the choice of the isosurface that defines the interface between the solvent and solute; and whether non-electrostatic contributions (such as surface tension) are required in the total energy.

For the MPB approach, energies and analytical forces are supported for non-periodic systems [201]. After the appropriate choice for `solvent` in the calculation input, the dielectric function must be specified. At the current stage, only the dielectric function from the self-consistent continuum solvation (SCCS) method [193] has been extensively tested with the MPB package. The SCCS dielectric function has been popular in the community, and parameters for ionic [202] and neutral molecular solutes in aqueous [193], and non-aqueous [203] solutions (with generalization to any non-aqueous solvent), have been proposed. Implicit embedding methods rely critically on parameterizations, and therefore careful review of these references is encouraged for users. A notable additional feature of the MPB implementation is the possibility for modelling additional continuum charge distributions in the embedded region, which could for example represent salts in electrolytes. This is commonly achieved by modelling the ions as an ideal gas (Poisson-Boltzmann equation) with possibility for additional repulsion between the ions, to account for hydrated ion-ion interactions, and the ion-solute interactions that form a Stern layer [184, 201, 204]. Parameterisations for various dissolved salts are available and have been well-tested for molecular solutes [205]. Full details of all settings are available in the FHI-aims manual.

For the COSMO approach, both energies and analytical forces are supported for non-periodic systems. To enable a COSMO calculation, one must add the command `solvent cosmo` into the `control.in` file. Two additional inputs are required: the dielectric constant of the medium, specified with `cosmo_epsilon <value>`, and a file that represents the grid points on the dielectric surface. Full details on the format of the grid points file can be found in the FHI-aims manual.

Roadmap on Advancements of the FHI-aims Software Package

2.6. EXPLICIT AND IMPLICIT EMBEDDING APPROACHES

The external package Environ offers a wide range of cavity definitions, electrolyte models and additional energy contributions. Its interface with FHI-aims is kept as generic as possible, thereby making new developments in the Environ package immediately available to users of FHI-aims.[193, 194]

Future Plans and Challenges

The development and deployment of embedding approaches remains an active domain, with increasing need to understand chemical systems at large scale. Explicit atomistic and electronic embedding can therefore benefit from additional options for defining the environment, *via* either density [206] or wavefunction-based embedding [207], and these are fields of active on-going research. These developments will allow better representation of short-range interatomic interactions, and the coarser point charge and pseudopotential representations can be deployed at a greater distance from the active centre. Care will be required to manage the calculation model and coupling of energy landscapes accurately and coherently, and work is on-going in this domain within FHI-aims and complementary software packages, such as ChemShell. Furthermore, accessibility is as desirable as functionality, with a need for simpler workflows and transferability of approaches across the physical, chemical, biological, and material domains. Complementary to these aspects is the opportunity to integrate machine-learning representations of surrogate landscapes, as discussed elsewhere in this Roadmap.

Implicit environment embedding is equally important in future workflows, and will benefit from the forthcoming implementation of forces for the MPE model. These forces are in active development for molecular models but further ambitions exist for this functionality in periodic modelling approaches. The MPB model is complementary and offers further a high degree of flexibility for future implementations of advanced solvation techniques, such as non-linear or non-local dielectric response [184], and similar goals exist for extension to periodic modelling. The combined outcome will be access to models for solid/liquid interfaces relevant to the most timely challenges in environmentally-relevant chemistry, such as renewable energy and green catalysis.

Work also continues overall to improve the scalability of the implicit embedding approaches, by systematically constructing basis sets of limited size for piecewise solvent sub-regions, and seeking and implementing efficiently scaling sparse solvers such that hierarchical models are not a bottleneck when deployed [200]. Aspirations to couple implicit and explicit embedding approaches exist, but further development of the underlying framework is required before any attempts to realise this complete embedding model.

Acknowledgements

We acknowledge fruitful discussions with Oliviero Andreussi, Reinhard J. Maurer, Christoph Scheurer, Alexey Sokol and Scott Woodley. AJL, PVS, and GAB acknowledge funding by the UKRI Future Leaders Fellowship program (MR/To18372/1, MR/YO34279/1). OvV acknowledges funding of a PhD scholarship by Cardiff University. AJL, JF, HO, GAB, and PVS acknowledge funding from the ARCHER2 eCSE Programme (eCSE08-13). HO acknowledges support from the German Science Foundation (DFG) under grant number OB 425/9-1. SR additionally acknowledges financial support from the National Research Foundation of Korea (NRF), funded by the Ministry of Science and ICT (grant no. 2021R1C1C1008776).

*Roadmap on Advancements of the FHI-aims Software Package**2.6. EXPLICIT AND IMPLICIT EMBEDDING APPROACHES*

Accepted Manuscript

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

3 Exchange and Correlation

Accepted Manuscript

3.1 Exchange and Correlation for Density Functional Theory and the DFT+U Approach

Volker Blum^{1,2}, Fabio Della Sala^{3,4}, Eduardo Fabiano^{3,4}, Jan Hermann^{5,6,a}, J. Matthias Kahk⁷, Matthias Kick⁸, Sebastian Kokott^{5,9}, Susi Lehtola¹⁰, Johannes Lischner¹¹, *Andrew J. Logsdail¹², Harald Oberhofer¹³, Karsten Reuter⁸, Mariana Rossi¹⁴, *Matthias Scheffler⁵, and Alexandre Tkatchenko⁶

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Institute for Microelectronics and Microsystems (CNR-IMM), Via Monteroni, Campus Unisalento, 73100 Lecce, Italy

⁴Center for Biomolecular Nanotechnologies, Istituto Italiano di Tecnologia, Via Barsanti 14, 73010 Arnesano (LE), Italy

⁵The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶Department of Physics and Materials Science, University of Luxembourg, L-1511 Luxembourg City, Luxembourg

⁷Institute of Physics, University of Tartu, W. Ostwaldi 1, 50411 Tartu, Estonia

⁸Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁹Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

¹⁰Department of Chemistry, University of Helsinki, P.O. Box 55 (A. I. Virtasen aukio 1), FI-00014 University of Helsinki, Finland

¹¹Department of Physics, Department of Materials, and the Thomas Young Centre for Theory and Simulation of Materials, Imperial College London, London SW7 2AZ, United Kingdom

¹²Cardiff Catalysis Institute, School of Chemistry, Cardiff University, Park Place, Cardiff CF10 3AT, United Kingdom

¹³Department of Physics and Bavarian Center for Battery Technologies, University of Bayreuth, Bayreuth, Germany

¹⁴Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

*Coordinator of this contribution.

^aCurrent Address: Microsoft Research AI for Science, Karl-Liebknecht-Str 32, 10178 Berlin, Germany

Summary

In density functional theory (DFT), the ground state total energy is written as a function of the density, n , as [4, 8, 100, 208]:

$$E_{\text{DFT}}[n] = T_{\text{s}}[n] + \int d^3\mathbf{r} v_{\text{ext}}(\mathbf{r})n(\mathbf{r}) + E_{\text{H}}[n] + E_{\text{xc}}[n], \quad (3.1)$$

Roadmap on Advancements of the FHI-aims Software Package

3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH

where \mathbf{r} is a spatial coordinate. In Eq. (3.1), the first term ($T_s[n]$) is the kinetic energy of non-interacting electrons; the second term is the interaction energy of the electron density with an external potential, v_{ext} , generated by, e.g., the nuclei; the third term ($E_{\text{H}}[n]$) is the Hartree energy, which describes the electrostatic electron-electron interaction; and the fourth term ($E_{\text{xc}}[n]$) accounts for the many-electron exchange and correlation, also correcting the spurious self-interaction contained in the Hartree term.

To calculate $E_{\text{DFT}}[n]$, $T_s[n]$ can be evaluated exactly only when expressed in terms of single-particle wave functions. Furthermore, there is no proof that the functional for evaluating $E_{\text{xc}}[n]$ can be written down in terms of a closed mathematical form; indeed, $E_{\text{xc}}[n]$ is likely just an algorithm that, in its exact implementation, requires the solution of the many-body Schrödinger equation [4]. Importantly, Kohn-Sham DFT enables a route towards approximate functionals that can be solved.

Approximations to E_{xc} are central to DFT and they may be complex and intricate. The increasing intricacy is captured in the so-called Jacob's ladder [209], which stands on the ground (The Hartree approximation) and reaches towards heaven (The exact solution). The first rung of the ladder is the local-density approximation (LDA) where the density functional approximation (DFA) depends only on the electron density, n , and the exact solution is known numerically. The second rung of the ladder refers to the family of generalised gradient approximations (GGAs), which depend on n and ∇n . Then, on the third rung, meta-GGAs are introduced with dependence on the kinetic energy density (τ) or higher order derivatives of the density (e.g., $\nabla^2 n$). On rungs further up the ladder, DFAs also consider occupied Kohn-Sham states (e.g. hybrid-DFT) and then even unoccupied Kohn-Sham states. In principle, higher rungs of the ladder, and therefore more complex DFAs, offer greater mathematical flexibility and improved accuracy. For example, a recent development direction is represented by the Hartree-Fock adiabatic connection (HFAC) methods [210–212], which can be considered as non-linear double-hybrid functionals. HFAC functionals merge the MP2 term with density functionals from the strong-correlation limit [213], so that they can be applied also to systems with vanishing gap, and with very accurate results for non-covalent interactions [211, 212]. The reader is directed to the wider literature for more complete documentation of DFA formalisms and discussion of applicability for different chemical systems [209, 214–216].

Alongside DFA choices, corrective approaches also exist to improve system representation. DFT+ U is particularly successful at addressing the self-interaction error (SIE) for extended transition metal oxides, lanthanides, and actinide compounds with partially filled d- or f-shells, with outcomes similar to hybrid-DFT after careful parameterisation [217]. The Hubbard-corrected energy, $E_{\text{DFT}+U}$, is obtained by applying corrections to states that show a high degree of electron localization, described as correlated subspace, where the SIE can be large [191]:

$$E_{\text{DFT}+U}[n] = E_{\text{DFT}}[n] + E_U[\mathbf{n}_I] - E_{\text{dc}}[\mathbf{n}_I]. \quad (3.2)$$

Here, \mathbf{n}_I refers to the occupation number matrix of a correlated state for atom site I , and E_{dc} addresses double-counting of the Coulomb interaction. The DFT+ U formalism is simple and can provide improved accuracy with limited additional computational cost on the underlying DFAs.

In this Section of the Roadmap, the LDA, GGA, and meta-GGA **exchange-correlation DFAs** are considered, and we will also consider the **Hubbard-corrected DFT+ U** [191, 218, 219]. More complex DFAs that depend on the single-particle wave function, and methods for including correlated effects will be discussed in upcoming Sections. Importantly, these methods frequently build upon the basic DFAs discussed in the present section. Hybrid DFAs, which add a fraction of nonlocal exchange to E_{xc} , will be covered in Section 3.2. Van der Waals corrections, a correlation effect within E_{xc} , will be addressed in Section 3.3. The fifth-rung random phase approximation (RPA), discussed in Section 3.4, also includes van der Waals interactions, along with other

Roadmap on Advancements of the FHI-aims Software Package

3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH

aspects of correlation in chemical bonding. Finally, coupled cluster theory for ground and excited states is addressed in Section 4.1.

Current Status of the Implementation

Exchange-correlation DFAs

FHI-aims contains internal implementations of the energy density and its derivatives for various DFAs, which are accessible as separate subroutines. Complementary to the bundled implementations, FHI-aims can also interface to two libraries that provide auto-generated code for DFAs, substantially extending the list of available DFAs, and enabling access to high order derivatives. An interface to the LibXC library [214], which has become the standard implementation in >40 electronic structure packages, provides FHI-aims with >600 fully tested DFAs. FHI-aims is currently distributed with LibXC version 6.1.0. Access to such a standard implementation is beneficial for the reproducibility of computational results [220]; note that differences in bundled DFA implementations can lead to different programs giving vastly different results for nominally the same functional [221, 222]. Additionally, an interface to the `dfauto` package [223] provides a numerically robust framework for developing and testing new DFAs, noting that new implementations require developer interaction with the FHI-aims codebase. A functional form for `dfauto` is specified in the Maple language, and the program then automatically derives the functional derivatives and generates the Fortran code.

Both LibXC and `dfauto` have been extensively tested against the internal implementations of various DFAs, and the libraries have been found to be numerically accurate for the LDA, GGA, and meta-GGA methods outlined. The bundled implementations, as well as the DFAs from LibXC and `dfauto`, are also compatible with hybrid-DFT methods. For dispersion-corrected DFT, correlation energy is typically evaluated as an additive term to the total energy, as discussed in Section 3.3 in this Roadmap, although an implementation also exists for the dispersion-corrected DFA of Dion *et al.* that is evaluated during the SCF cycle [224, 225].

The energy density is evaluated discretely over a radial grid in real space, allowing batch-wise parallelisation on distributed computing architectures. The quantities that are necessary for estimating the energy at a given spatial point can include the density (n), gradient of the density (∇n), and the local kinetic energy density (τ), for which the respective DFA families are called the local density approximation (LDA), generalised gradient approximation (GGA), and meta-GGA approaches. The implementations calculate and return partial derivatives, allowing for calculation of a potential for the self-consistent field (SCF) cycle. For the GGA, the potential is calculated as [8]:

$$v_{xc}^{GGA} = \frac{\delta E_{xc}[n, \gamma]}{\delta n} = \frac{\partial f_{xc}}{\partial n} - 2\nabla \cdot \left[\frac{\partial f_{xc}}{\partial \gamma} \nabla n \right] \quad (3.3)$$

where γ is the scalar product of the spin-density gradients, *i.e.*, $|\nabla n|^2$, and f_{xc} is the spatially dependent exchange correlation energy density; a rearrangement is applied in the implementation of this term, in order to avoid an expensive evaluation of $\nabla^2 n$ [8, 226]. For the meta-GGA approach, the required derivative $\delta E_{xc}[n, \gamma, \tau]/\delta n$ cannot be easily obtained as the derivative requires knowledge of $\delta\tau/\delta n$ but τ is not an explicit functional of the electron density, n ; it is a functional of the occupied electronic states:

$$\tau = \frac{1}{2} \sum_i f_i |\nabla \psi_i|^2 \quad (3.4)$$

Roadmap on Advancements of the FHI-aims Software Package

3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH

where ψ_i is the i th molecular orbital and f_i is the occupation of this orbital. It is possible to calculate $\delta\tau/\delta n$ via the optimised effective potential (OEP) method as implemented by Sharp and Horton [227] or in the Krieger-Li-Iafrate approximation [228], but this approach is cumbersome and expensive, making it restrictive for practical applications [229]. Therefore, the potential is calculated by evaluating the functional derivative with respect to ψ , as first suggested by Neumann, Nobes and Handy (NNH) [230]:

$$v_{xc}^{mGGA} = \frac{\delta E_{xc}[n, \gamma, \tau]}{\delta n} \leftarrow \frac{1}{2} \frac{\delta E_{xc}[n, \gamma, \tau]}{\delta \psi} \quad (3.5)$$

and thus we can evaluate the derivatives with respect to the orbitals instead of the density [231]:

$$v_{xc}^{mGGA}\psi = v_{xc}^{GGA}\psi - \frac{1}{2} \nabla \cdot \left[\frac{\partial f_{xc}}{\partial \tau} \nabla \right] \psi. \quad (3.6)$$

The SCF cycle should be converged to obtain ground-state energies (Fig. 3.1). The framework is robust and computationally highly efficient, capable of performing DFT calculations on >10,000 cores [232]. Details of the configuration and convergence of the SCF cycle are available in Section 3.10 of the FHI-aims manual [233].

Hubbard-corrected DFT+U

The popular spherically averaged form of DFT+U is available in FHI-aims for addressing SIE in DFAs [219]. A scalar and site-dependent Hubbard parameter (U_I) is applied to determine the interaction strength of electrons localised at atom site I . The user is required to determine U via careful parameterisation to yield an experimentally known observable, such as a band structure [234]. The U parameters depend on the atom sites and their surroundings; the specifics of the projection operator used to determine the occupation of the correlated subspace; and on the employed basis set to form the projection operator.

The projector functions (Φ) for an atomic site I and a magnetic quantum number m are written in FHI-aims as a linear combination of the numerical atomic orbital (NAO) basis functions, $\Phi_{Im} = \sum_i c^i \phi_{Im}^i$, where c^i corresponds to the coefficient of an individual NAO basis function ϕ^i . NAOs are advantageous for this task, because they are the DFT solutions of non-spin-polarised free atoms and thus reflect the character of localised states well, especially when a minimal basis is used. The NAO formalism can result in equivalent charge localization with U values that are slightly lower, by roughly 1-2 eV, than those reported in other formalisms [191].

Usability and Tutorials

Exchange-correlation DFAs

FHI-aims currently provides access to ~40 LDA, GGA, and meta-GGA DFAs, as well as hybrid-DFT, via bundled implementations; full details of the available choices are in the software manual [233], and the reader is directed to wider literature for discussion of most appropriate DFAs for specific chemical systems [215, 216]. The implementations support spin-paired and spin-unpaired calculations. Choosing a DFA for a DFT calculation requires a simple input line: xc <name>, where <name> is an alphanumeric label as defined

Roadmap on Advancements of the FHI-aims Software Package

3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH

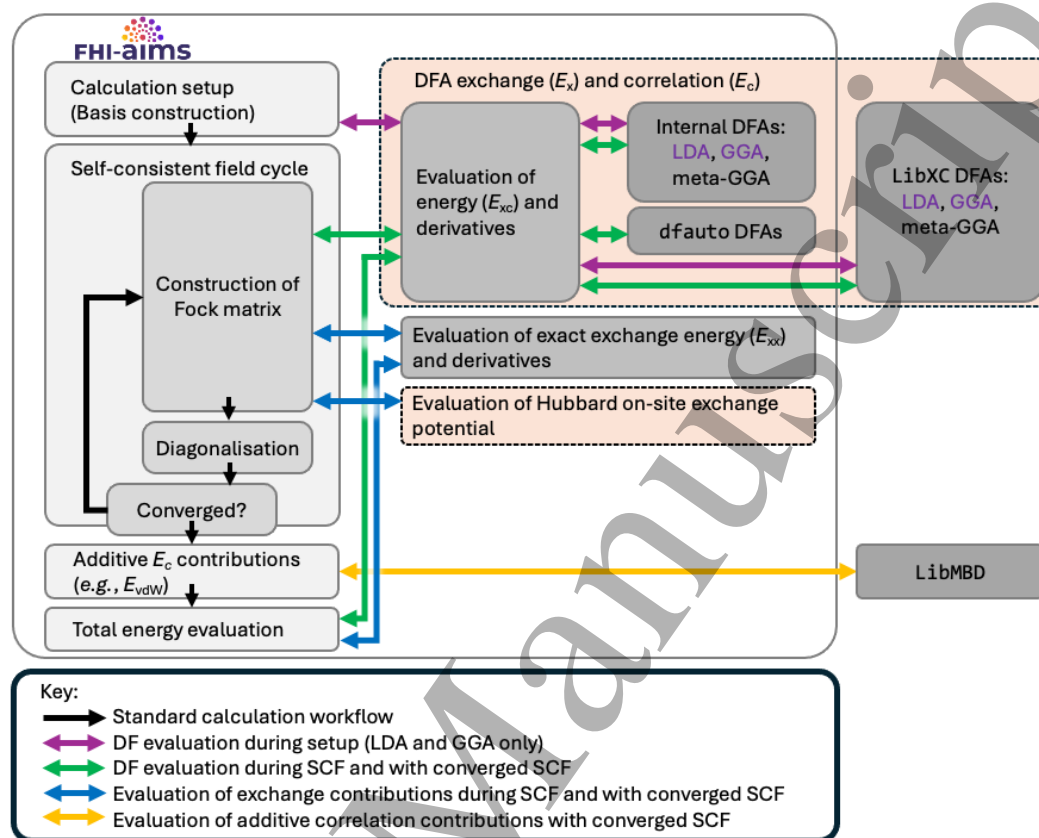


Figure 3.1: Schematic representation of the FHI-aims infrastructure for evaluation of exchange and correlation contributions. Highlighted in dashed boxes with an orange background are the aspects discussed in this contribution, namely the evaluation of density functional approximations (DFAs) and the evaluation of Hubbard on-site potentials. Additional energy components discussed elsewhere in this Roadmap include exact exchange, perturbative correlation, and additive van der Waals corrections. A key is provided for the calculation workflows during calculation setup, during the self-consistent field (SCF) cycle, and during post-convergence energy evaluation.

in the manual. For the LibXC and dfauto implementations, an additional keyword on the input line defines that these packages are being used: `xc libxc <name>` and `xc dfauto <name>`, respectively. For the LibXC DFAs, `<name>` is as listed in the LibXC documentation [235], with capacity to couple distinct exchange and correlation options.

The flexible framework developed for DFA implementation means that different DFAs can be used at different points of the calculation workflow. Any DFA can be used to perform x initial SCF iterations, for instance, to quickly determine a qualitative electronic structure before switching to a more expensive (or harder to converge) DFA, with the command `xc_pre <x> <name>`. A different DFA can then be used to converge the SCF cycle. Any DFA can also be used for estimating the total energy on top of a converged wave function from another method (e.g., density-corrected methods [236], for example), using the

Roadmap on Advancements of the FHI-aims Software Package

3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH

command `total_energy_method <name>`. The same command also provides access to contemporary double-hybrids [237, 238].

Any LDA and GGA DFAs in LibXC, or a subset of internally implemented LDA and GGA DFAs, can be used for free-atom calculations during the initialisation phase of FHI-aims, which defines the free-atom minimal basis set. The default DFA for free atom calculations has been well-tested, but it can also be changed in the input; this impacts absolute energies, but changes in relative energies are typically minimal.

Meta-GGA DFAs are known to require the use of larger radial grids than LDAs or GGAs, and a more thorough radial sampling can be necessary to obtain accurate derivatives [239–242]. The radial grids are a numerically convergent parameter within a DFT software package, and in FHI-aims a denser radial grid can be obtained by changing the overall FHI-aims basis accuracy option (e.g., “light” → “intermediate”), or by increasing the values of flags that control the number of radial grid points, such as the `radial_multiplier` parameter.

Hubbard-corrected DFT+ U

The DFT+ U functionality is compatible with all major FHI-aims features and can be easily evoked by defining species-specific U values for the desired angular momentum channel, as per the FHI-aims software manual. Default settings are optimised for systems with rather localised electrons, and there is high flexibility in the choice of double-counting corrections and the calculation of the occupation matrix [191]; for the latter, the options are a pure on-site electron counting or a Mulliken-like description. The features for occupation matrix control [243] allow fixing of the occupation matrix during an SCF cycle. The constraint acts like a bias potential, enabling convergence of the electron density towards a desired configuration, such as excess electrons localised on specific sites, a specific magnetic ordering, or a transition state [243, 244]. The subsequent electronic solution can be used to restart an unconstrained self-consistent calculation. The occupation matrix control shows great promise for challenging electronic structure situations and hard-to-converge systems.[191, 245–247]

Future Plans and Challenges

The DFT framework in FHI-aims is robust and complete, allowing one to evaluate the total energy and perform self-consistent field calculations for a range of LDA, GGA, and meta-GGA approaches to determine the ground state electronic structure. A thorough and extensive library of contemporary DFAs is obtained by interfacing with external packages, particularly LibXC. The LibXC interface in FHI-aims currently does not support meta-GGAs that depend on the Laplacian of the density ($\nabla^2 n$); such support will hopefully be added in the future. We also aim to ensure the completeness of implementations that depend on an external evaluation of correlation (i.e., dispersion-corrected methods). These challenges are within FHI-aims, where the specific system observable and/or energy contribution must be calculated, and not in LibXC.

With respect to future opportunities, work is on-going towards orbital-free approaches for evaluation of the kinetic energy density, including contemporary orbital-free meta-GGAs [248], as well as more established approaches that use kinetic energy functionals to address non-additive errors in hierarchical density embedding [249]. Future developments of the DFT+ U method in FHI-aims will focus on implementing the linear response method of computing DFT+ U [250], the site-to-site coupling term generally denoted as DFT+ U + V [251], and further optimisation of the employed projectors.

*Roadmap on Advancements of the FHI-aims Software Package**3.1. EXCHANGE AND CORRELATION FOR DENSITY FUNCTIONAL THEORY AND THE DFT+U APPROACH*

Acknowledgements

We acknowledge fruitful discussions with Albert P. Bartók and Sheng Bi. AJL acknowledges funding by the UKRI Future Leaders Fellowship program (MR/TO18372/1, MR/YO34279/1). SL thanks the Academy of Finland for financial support under project numbers 350282 and 353749. HO acknowledges support from the German Science Foundation (DFG) under grant numbers OB 425/9-1 and, together with MK, OB 425/3-2. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

3.2 Hybrid Density Functionals for Large Scale Simulations

Volker Blum^{1,2}, Christian Carbogno^{10,3,a}, Lukas Galland⁴, James A. Green⁵, Stefan Gutzeit^{6,7}, Felix Hanke^{7,8,b}, Danish Khan⁹, O. Anatole von Lilienfeld⁹, *Sebastian Kokott^{5,10}, Thomas Körzdörfer⁴, Sergey Levchenko^{7,11,c}, Michael Lorke¹², Florian Merz¹³, Alastair J. A. Price⁹, Markus Rampp¹⁴, Ronald Redmer⁶, Xinguo Ren^{15,10,d}, Mariana Rossi¹⁶, Matthias Scheffler¹⁰, Andrei Sobolev^{5,10}, Jürgen Wieferink⁷, and Yi Yao^{5,10}

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, North Carolina 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Institut für Chemie, Universität Potsdam, D-14476 Potsdam, Germany

⁵Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁶Institut für Physik, Universität Rostock, D-18051 Rostock, Germany

⁷Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁸Surface Science Research Centre and Department of Chemistry, University of Liverpool, Oxford Street, Liverpool L69 3BX, United Kingdom

⁹Chemical Physics Theory Group, Department of Chemistry, University of Toronto, St. George Campus, Toronto, ON, Canada.

¹⁰The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹¹Skolkovo Institute of Science and Technology, Moscow 121205, Russia

¹²Faculty of Physics, University of Duisburg-Essen, D-47057 Duisburg, Germany

¹³Lenovo HPC Innovation Center, Stuttgart, Germany

¹⁴Max Planck Computing and Data Facility, D-85748 Garching, Germany

¹⁵Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

¹⁶Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

*Coordinator of this contribution.

^aCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

^bCurrent Address: CuspAI Ltd, 20 Station Road, Cambridge, CB1 2JD, United Kingdom

^cCurrent Address: Skolkovo Institute of Science and Technology, Moscow 121205, Russia

^dCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

Summary

Hybrid density functionals (hybrids), which incorporate a fraction of nonlocal exchange into E_{xc} of Eq. (3.1), have emerged as a practical reference method for *ab initio* electronic-structure-based simulations. They address several accuracy issues inherent in lower levels of density functional approximations (DFAs) while re-

Roadmap on Advancements of the FHI-aims Software Package

3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS

maining computationally feasible on current high-performance computers. The main computational bottleneck for atomistic simulations using hybrid density functional theory (DFT) is the evaluation of the non-local exact exchange (EXX). The localized resolution-of-identity-based real-space implementation RI-LVL [62, 63] of the exact exchange algorithm in FHI-aims [64] enables the computation of the EXX operator with linear scaling by the number of atoms in the system. The RI-LVL algorithm was recently optimized to allow for much improved exploitation of sparsity and load balancing across ten thousands of parallel computational tasks. All technical details are summarized in Ref. [6]. The results demonstrate drastically improved memory and runtime performance, scalability, and workload distribution on CPU clusters. The current reach of the hybrids is documented by run times and scaling of hybrid DFT simulations for several challenging materials, including hybrid organic/inorganic perovskites [252] and organic crystals, with up to 30,576 atoms (101,920 electrons described by 244,608 basis functions) in the simulation cell [6] (cf. Figure 3.2). Remarkably, despite the scale of these systems, the simulations can be conducted with modest computational resources. Finally, optimizations have been implemented in the computation of band structures and density of states for periodic hybrid functional simulations, alongside the support for new hybrid functionals.

Current Status of the Implementation

The key concept for the efficient evaluation of the exact exchange is the RI-LVL real space formalism. [62, 63] The EXX operator $X_{ij}(\mathbf{R})$ in real space then reads:

$$X_{ij,\sigma}(\mathbf{R}) = \sum_{k\mathbf{R}'} \sum_{\mathbf{R}''} \sum_{\mu\mathbf{Q}'} \sum_{\nu\mathbf{Q}''} \tilde{C}_{ik(\mathbf{R}')}^{\mu(\mathbf{Q}')} V_{\mu\nu(\mathbf{R}+\mathbf{Q}''-\mathbf{Q}')} \tilde{C}_{jl(\mathbf{R}'')}^{\nu(\mathbf{Q}'')} \times D_{kl,\sigma}(\mathbf{R}+\mathbf{R}''-\mathbf{R}') \quad (3.7)$$

where $\tilde{C}_{ik(\mathbf{R}')}^{\mu(\mathbf{Q}')}$ are the RI expansion coefficients in the flavor of the RI-LVL [63], $V_{\mu\nu(\mathbf{R})}$ the Coulomb matrix (cf. Eq. (3.17)), and

$$D_{kl,\sigma}(\mathbf{R}) = \frac{1}{N_{\mathbf{k}}} \sum_{\mathbf{k}} \sum_m f_{m,\sigma}(\mathbf{k}) c_{km,\sigma}(\mathbf{k}) c_{lm,\sigma}^*(\mathbf{k}) e^{i\mathbf{k}\mathbf{R}} \quad (3.8)$$

is the Fourier transform of the density matrix, with the occupation numbers $f_{m,\sigma}(\mathbf{k})$, the Kohn-Sham eigenvectors $c_{km,\sigma}(\mathbf{k})$ from the solution of KS eigenvalue problem of the previous SCF iteration, and the number of k-points in the Brillouin zone $N_{\mathbf{k}}$. \mathbf{R} and \mathbf{Q} denote linear combinations of lattice vectors in the Born-von Karman cell; the sum over them is not restricted to the extent of the Born-von Karman cell, but solely by the overlap of the basis functions. The Latin symbols i, j, k , and l denote indices of the numerical atomic orbital (NAO) basis functions, m the eigenstate index, the Greek symbols μ and ν are the indices of the product basis functions, and σ is the spin index. All basis functions and auxiliary basis functions are labeled with a real-space lattice vector. Dropping that index refers to basis functions at an atom in the cell $\mathbf{R} = \mathbf{0}$, e.g., $i = i(\mathbf{0})$. The RI-LVL expansion is constrained so that products of basis functions $\phi_i(r - R)$ on atom I and $\phi_j(r - R')$ on atom J are expressed using auxiliary basis functions $P_\mu(r - R)$, which must be assigned to the same atom pair $\mathcal{P}(IJ)$,

$$\phi_i(\mathbf{r} - \mathbf{R})\phi_j(\mathbf{r} - \mathbf{R}') = \sum_{\mu(\mathbf{R}) \in \mathcal{P}(IJ)} \left(\tilde{C}_{i(\mathbf{R})j(\mathbf{R}')}^{\mu(\mathbf{R})} P_\mu(\mathbf{r} - \mathbf{R}) + \tilde{C}_{i(\mathbf{R})j(\mathbf{R}')}^{\mu(\mathbf{R}')} P_\mu(\mathbf{r} - \mathbf{R}') \right). \quad (3.9)$$

Roadmap on Advancements of the FHI-aims Software Package

3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS

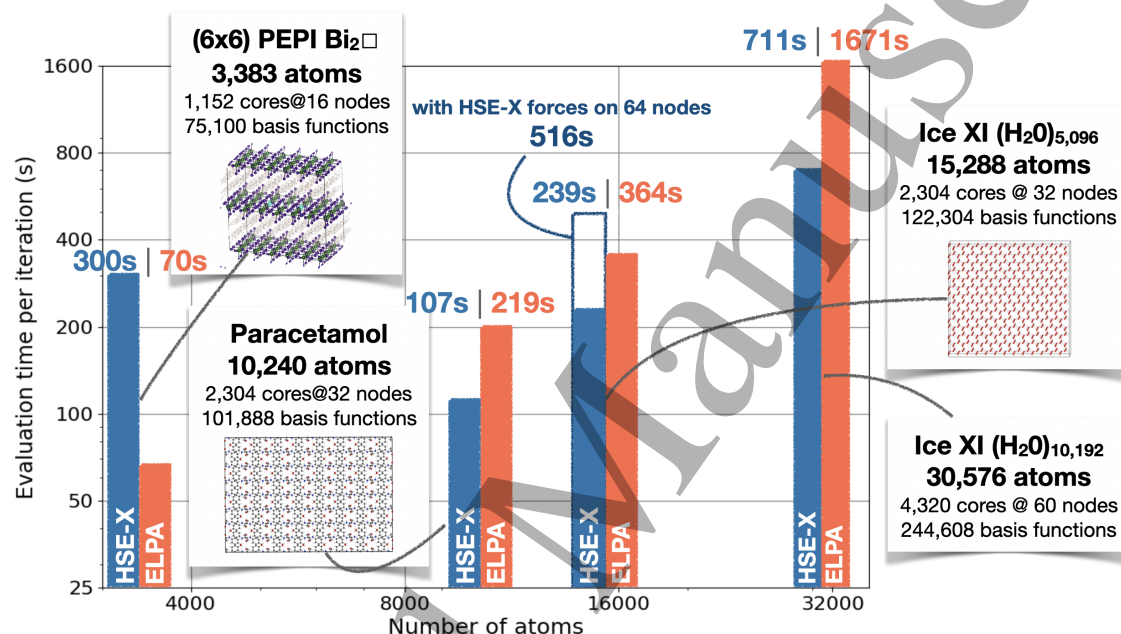


Figure 3.2: Benchmark results for the largest periodic structures analyzed in Ref. [6]. Average runtimes per self-consistent field iteration are shown for the HSE06 exchange operator (blue bars) and the ELPA two-stage eigenvalue solver (red bars). Simulations were performed using the HSE06 hybrid functional for the following systems (from left to right): phenylethylammonium lead iodide (PEPI) with a defect complex (indicated by a square in the chemical formula) [252], a $4 \times 4 \times 4$ paracetamol supercell, a 15,288-atom Ice XI supercell (including force evaluation), and a 30,576-atom Ice XI supercell. Calculations were conducted on the Raven HPC cluster at MPCDF, using Intel Xeon IceLake (Platinum 8360Y) nodes with 72 cores per node. Figure adapted from Ref. [6].

Roadmap on Advancements of the FHI-aims Software Package

3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS

All detailed expressions involved in Eq. (3.7) are given in Ref. [64].

The optimal distribution and storage of the RI coefficients $\tilde{\mathbf{C}}$, the Coulomb matrix \mathbf{V} , and the density matrix \mathbf{D} , as well as intermediate matrices of the multiplications in Eq. (3.7) is a hard task and strongly depends on the simulated system. The code tries to utilize the resources in an optimal fashion: Several rows of $X_{ij,\sigma}$ are aggregated into batches (so called Fock matrix blocks). The aggregation of rows of $X_{ij,\sigma}$ into larger blocks enables efficient CPU cache usage. If enough memory per node is available, the code duplicates all the data (that is, $\tilde{\mathbf{C}}$, \mathbf{V} , \mathbf{D} , index lists storing sparsity patterns, and communication patterns) that is needed to compute a row of the real-space Fock matrix $X_{ij,\sigma}$ (so called instances). This duplication allows a significant reduction of the Message Passing Interface (MPI) communication and enables almost perfect strong scaling for a wide range of core counts.[6] Both $\tilde{\mathbf{C}}$ and \mathbf{V} are sparse matrices and only computed and compressed once per self-consistent-field (SCF) cycle. The compression scheme removes columns and rows of the corresponding matrix, wherever the norm of the column and row is below 10^{-10} . \mathbf{D} is computed every SCF iteration and, formally, is the only input quantity for an EXX SCF evaluation. The real-space EXX operator $X_{ij}(\mathbf{r})$ is linked to the corresponding k-space operator $K_{ij,\sigma}(\mathbf{k})$ via the following Fourier transformation

$$K_{ij,\sigma}(\mathbf{k}) = \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} X_{ij,\sigma}(\mathbf{R}). \quad (3.10)$$

During a SCF iteration, $K_{ij,\sigma}(\mathbf{k})$ undergoes the same mixing scheme as used for the the density matrix. By default in FHI-aims, this is the Pulay mixing, which ensures a faster convergence of the SCF cycle. Then, the mixed EXX operator $\mathbf{K}_{\text{mix}}(\mathbf{k})$ (or a fraction thereof - depending on the used hybrid functional) is added to the Hamiltonian for solving the generalized eigenvalue problem.

The EXX contribution $K_{ij,\sigma}(\mathbf{k})$ can be computed also on generalized regular k-grids [253], which are k-grids constructed from rotated supercells to allow for a more efficient sampling of the Brillouin zone. This helps to reduce the needed compute resources for systems with a skewed cell, e.g. trigonal crystals. The generalized regular k-grid can be either specified by manual user input or automatically generated by autoGR [65]. Additionally, $K_{ij,\sigma}(\mathbf{k})$ can be computed at an arbitrary k-point in the Brillouin zone by a Fourier interpolation, e.g., needed for the computation of the band structure or the density of states. The Fourier interpolation requires a sufficiently dense k-grid. Empirically, our tests show that the Born-von Karman (BvK) cell should be large enough to contain all unique pairs of basis functions which have at least 25% of overlap measured by the maximal extent of the basis functions. The code checks at runtime whether this requirement is fulfilled. If the density of the k-grid is too low, the code stops and suggests a minimal needed k-grid.

Usability and Tutorials

The exact exchange code implementation has support for a large number of features in FHI-aims as summarized in Figure 3.3. It can perform computation of energy and forces for periodic and non-periodic systems, and, in addition, stress [54] for periodic systems. The key concept of any hybrid functional is the mixing of a fraction of exact exchange with fraction(s) of LDA, GGA, or meta-GGA exchange. Often, a range-separation function is introduced to divide the Coulomb potential v into long- and short-range parts. In FHI-aims, the error function (erf) is available as a range-separation function:

$$v(r) = \underbrace{\frac{1 - \text{erf}(\omega r)}{r}}_{v_{\text{SR}}(r)} + \underbrace{\frac{\text{erf}(\omega r)}{r}}_{v_{\text{LR}}(r)}, \quad (3.11)$$

Roadmap on Advancements of the FHI-aims Software Package

3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS

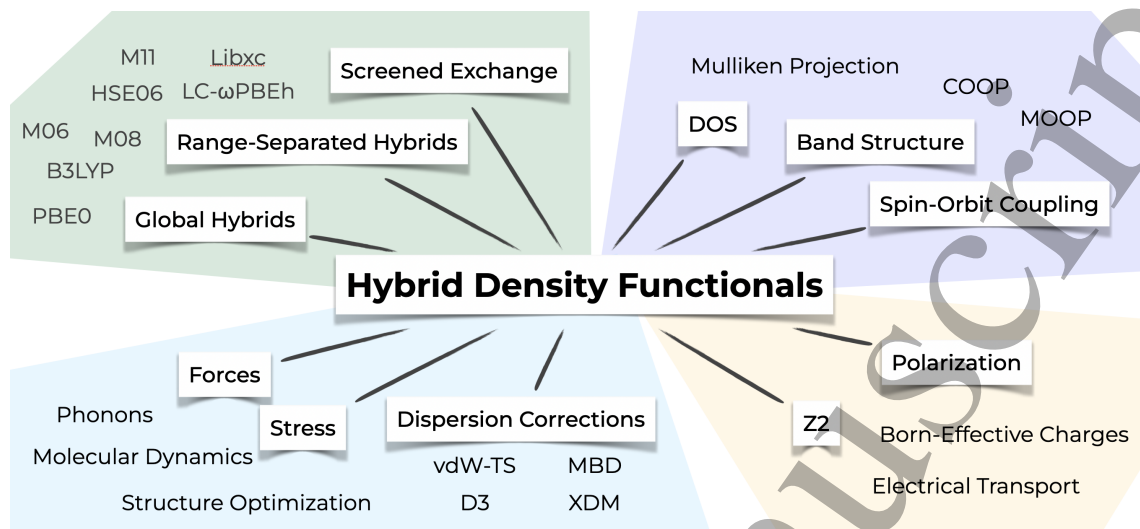


Figure 3.3: Overview of features implemented in FHI-aims with support for hybrid density functionals.

where $r = |\mathbf{r} - \mathbf{r}'|$, ω (an adjustable inverse length) is the range-separation parameter, and $v_{\text{SR}}(r)$ and $v_{\text{LR}}(r)$ are the short- and long-range Coulomb potential, respectively. As an alternative choice for the range-separation function, the exponential function $\exp(-\omega r)$ was enabled, recently. In general, the following contributions to the exchange energy E_x are obtained:

$$E_x(\alpha, \beta, \omega) = \alpha E_{\text{EXX}} + \beta E_{\text{EXX}}^{\text{SR}}(\omega) + (1 - \alpha) E_{\text{x-DFA}} - \beta E_{\text{x-DFA}}^{\text{SR}}(\omega), \quad (3.12)$$

where $\alpha + \beta \leq 1$ is required. E_{EXX} is the EXX energy using the full Coulomb potential and $E_{\text{EXX}}^{\text{SR}}(\omega)$ is the short-range EXX energy, where the short-range Coulomb potential $\frac{1 - \text{erf}(\omega r)}{r}$ of Eq. (3.11) is used. Breaking down the EXX contributions into long- and short-range part, we find that the parameter $\alpha + \beta$ governs the short-range EXX contribution and α the long-range EXX contribution. Similarly, $E_{\text{x-DFA}}$ is the semilocal density functional approximation (DFA) exchange energy for the full-range Coulomb operator and $E_{\text{x-DFA}}^{\text{SR}}(\omega)$ is the short-range semilocal DFA exchange energy. Choosing a functional via the `xc` flag in the `control.in` file will automatically set the default mixing parameters. An exception is the parameter ω , which the user needs to specify for range-separated hybrids directly implemented in FHI-aims, since ω can vary depending on the chosen reference publication. However, the most common choices for ω can be used through LibXC [214] by specifying the corresponding LibXC flag for the hybrid functional. The Hartree-Fock method and different hybrid functionals are implemented in FHI-aims, such as, global hybrids (PBE0 [254, 255], PBEsolo [256], B3LYP [257]), range-separated hybrids (HSE06 [258, 259], LC- ω PBEh [260]), hybrid meta-generalized gradient functionals (MO6 suite [261], MO8 suite [262], M11 [263]), and the Koopmans-compliant screened exchange [264]. Many more hybrid functionals are supported through the XC functional library LibXC [214]. In addition, for materials including heavy elements, perturbative spin-orbit coupling can be combined with the hybrid functionals [48]. Moreover, the hybrid functionals can be supplemented by dispersion corrections vdW-TS, D3, MBD, and XDM, and optimized dispersion parameters are provide alongside.

FHI-aims has been extended to support adaptive hybrid functionals through the methodology of aPBE0 [265] using on-the-fly machine learning models to estimate EXX admixtures for improved accuracy. A tutorial is

Roadmap on Advancements of the FHI-aims Software Package

3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS

provided in Ref. [266] on how to get the optimal fraction of EXX using the Atomic Simulation Environment (ASE). When applicable (i.e. query system is sufficiently similar to training instances), this integration allows for enhanced predictions of energetics, electron densities, and spin-state properties while maintaining computational efficiency. When not applicable, aPBE0 will revert to the default PBE0 admixture ratio of 0.25.

The current EXX code is production-ready and has been tested on several high-performance compute platforms and CPU architectures, i.e., x86_64 and arm64 architectures. When a hybrid functional is specified in the `control.in` file, the code tries to find the best setup for the resources specified by the user with the primary goal to avoid out-of-memory scenarios. However, it is also possible to manually tune the setup via keywords in the `control.in` file. The manual tuning sometimes enables additional speed-ups of 20-30%. The most important parameters used for the memory estimation are printed in the main output file and can be used by the user to find a better memory distribution. The performance of the code also varies for different hybrid functionals: Hybrid functionals that use a screened Coulomb potential (e.g. HSE06) are significantly faster (especially for dense systems, like bulk materials) than functionals that use a bare Coulomb potential, e.g., runtimes and required CPU resources increase by a factor of six when using the PBE0 functional instead of the HSE06 functional for the hematite crystal. Recently, a screening for the Coulomb potential has been implemented approximately accounting for long-range exact exchange contributions [267]. Using this screening function for the PBE0 functional gave results close to the original PBE0 functional, but come with significantly reduced computational cost and memory.

Future Plans and Challenges

The above-described improvements have become part of the standalone, open-source library libxc-X [268]. Writing and maintaining efficient HPC software, will be a big challenge in the future. Thus, making the code implementation open source will enable future research and development within the scientific community.

Graphics processing unit (GPU) strategies will be needed for the exact exchange algorithm, but are not yet exploited, as the porting of CPU code to GPU architecture is not at all straightforward. In the CPU implementation, the inherent sparsity of real-space approach keeps the size of matrices used for dense matrix-matrix operations moderate. The main problem is that RI coefficients and the Coulomb matrix are cast into a compressed format, but have to be expanded again for later matrix multiplications. This scheme only allows to carry out dense matrix algebra for small or medium sized batches. Thus, with the current algorithm the full capabilities of GPUs cannot be used, and speedups would be limited by communication. An overhaul of the algorithm, and GPU-specific storage and communication patterns will be needed to make it amenable for heterogeneous, GPU-accelerated architectures.

From a technological standpoint, the availability of sufficiently large memory per compute node and task will be crucial for any advanced electronic structure methods, particularly since non-local operators, which often require a careful balance between data locality and communication across nodes and tasks, are typically evaluated. The closer integration of accelerators within HPC nodes, as newly introduced with the Nvidia Grace-Hopper and AMD MI300A processors, shows great promise. However, communication bottlenecks between GPU and CPU tend to vary significantly across different platforms. In general, library APIs for solving mathematical and physical problems are often either vendor-specific or not optimized for performance across various platforms. Addressing these issues by creating vendor-agnostic, high-level APIs that perform efficiently on different architectures would extend the applicability of scientific code, reduce the need for code duplication, and significantly lower research costs. Notably, the ELPA eigensolver has such a vendor-agnostic infrastructure API already implemented. Nevertheless, optimizing communication pat-

*Roadmap on Advancements of the FHI-aims Software Package**3.2. HYBRID DENSITY FUNCTIONALS FOR LARGE SCALE SIMULATIONS*

terns between CPUs and GPUs for specific architectures remains a complex task. Additionally, new workload distribution models may be required to fully utilize all available resources, such as enabling simultaneous computation on both GPUs and CPUs, as current practices often leave CPUs idle while GPUs perform most of the work.

Acknowledgements

The authors would like to acknowledge the contribution of Rainer Johanni, who implemented important infrastructure for the hybrid DFT code part in FHI-aims, but passed away much too early. We also acknowledge Arvid Ihrig for numerous contributions to the RI-LVL implementation.

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

3.3 Density-Functional Based Methods for Dispersion Interactions

Alberto Ambrosetti¹, Kyle R. Bryenton², Robert A. DiStasio Jr.³, Simiam Ghan⁴, James A. Green⁵, Jan Hermann^{6,7a}, Johannes Hoja⁸, *Erin R. Johnson^{2,9,10}, Reinhard J. Maurer^{11,12,13}, Alberto Otero-de-la-Roza¹⁴, Alastair J. A. Price^{15,16}, Victor Ruiz¹⁷, Matthias Scheffler⁶, and *Alexandre Tkatchenko^{7,18}

¹Università degli Studi di Padova

²Department of Physics and Atmospheric Science, Dalhousie University, 6310 Coburg Road, Halifax, Nova Scotia, B3H 4R2, Canada

³Department of Chemistry and Chemical Biology, Cornell University, Ithaca, New York 14853, USA

⁴Department of Physics, Technical University of Denmark, Fysikvej, 313, 524, 2800 Kgs. Lyngby, Denmark

⁵Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁶The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Department of Physics and Materials Science, University of Luxembourg, L-1511 Luxembourg City, Luxembourg

⁸Department of Chemistry, University of Graz, Heinrichstraße 28, 8010 Graz, Austria

⁹Department of Chemistry, Dalhousie University, 6243 Alumni Crescent, Halifax, Nova Scotia, B3H 4R2, Canada

¹⁰Yusuf Hamied Department of Chemistry, University of Cambridge, Lensfield Road, Cambridge, CB2 1EW, United Kingdom

¹¹Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

¹²Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

¹³Faculty of Physics, University of Vienna, Vienna A-1090, Austria

¹⁴Departamento de Química Física y Analítica and MALTA-Consolider Team, Facultad de Química, Universidad de Oviedo, 33006 Oviedo, Spain

¹⁵Department of Chemistry, University of Toronto, St. George Campus, Toronto, ON, Canada

¹⁶Acceleration Consortium, University of Toronto, 80 St. George Street, Toronto, ON M5S 3H6

¹⁷Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹⁸Institute for Advanced Studies, University of Luxembourg, L-1511 Luxembourg, Luxembourg

*Coordinator of this contribution.

^aCurrent Address: Microsoft Research AI for Science, Karl-Liebknecht-Str 32, 10178 Berlin, Germany

Summary

Density-functional theory (DFT) is widely recognized as the standard computational technique for modeling the electronic structure of molecules and materials with up to 10,000 atoms. Despite its proven capabilities, achieving reliability and high accuracy in DFT simulations of intermolecular interactions necessitates the inclusion of London dispersion or van der Waals (vdW) forces. While they are formally a part of E_{xc} in Eq. (3.1), such forces result from highly non-local electron correlations that are not captured by local,

3.3. DENSITY-FUNCTIONAL BASED METHODS FOR DISPERSION INTERACTIONS

semi-local, or hybrid exchange-correlation functionals. Numerous post-SCF (self-consistent field) and full SCF vdW/dispersion methods have been developed to pair with popular DFT functionals, demonstrating excellent performance at low computational cost. Here, we describe the vdW/dispersion methods currently implemented in the Fritz Haber Institute *ab initio* materials simulation (FHI-aims) program [8].

Current Status of the Implementation

Currently, there are several post-SCF and full SCF vdW/dispersion methods available in FHI-aims from which to choose. These include the Tkatchenko-Scheffler (TS) model [269]; two variants of the many-body dispersion (MBD) method—the range-separated, self-consistently screened version (MBD@rsSCS) [270, 271] and the nonlocal version (MBD-NL) [272]; and the exchange-hole dipole moment (XDM) method [273–275]. TS, MBD@rsSCS, and MBD-NL are implemented within the libmbd library, while XDM is implemented in the `xdm.f90` routine. With each of these methods, the dispersion energy is computed using the self-consistent (SC) electron density from a DFT calculation and, in some cases, other density-dependent properties. It is then added to the self-consistent energy to give the total DFT energy:

$$E_{\text{DFT}} = E_{\text{SC}} + E_{\text{disp}}. \quad (3.13)$$

As a result, all of the vdW/dispersion methods listed above have 1-2 empirical parameters within their formulation that depend on the choice of density functional. Similarly, the forces and stresses are derived from the dispersion energy and can be added to the DFT forces and stresses.

Both TS and MBD@rsSCS are also implemented with full self consistency [276–278], meaning that the vdW/dispersion interactions not only affect the energy (and forces/stresses), but also the electron density, orbitals, and all the properties of interest derived from the orbitals and electron density. These effects can be important for large and/or polarizable systems [277, 278].

Because the evaluation of the pairwise dispersion energy depends on a summation over all atomic pairs, it formally scales as $\mathcal{O}(N^2)$, where N is the number of atoms. On the other hand, the MBD Hamiltonian requires diagonalization of the oscillator coupling matrix, hence MBD/MBD-NL methods scale as $\mathcal{O}(N^3)$. Typically, both pairwise (TS/XDM) and MBD dispersion methods do not increase the cost of the underlying semilocal or hybrid DFT calculation. This is because the pairwise energy expression can be evaluated in a linearly scaling fashion by employing suitable and controlled cutoffs. The MBD expression is efficiently implemented with Ewald summation and employs modern diagonalization routines, whereas the size of the MBD matrix is small compared to the Kohn-Sham matrix. Hence, the cost of non-self-consistent dispersion calculations is negligible compared to the underlying Kohn-Sham calculation. In contrast, the self-consistent TS/MBD calculations can increase the computational cost by a factor of 2-3 or higher depending on the system size. This is because the Hirshfeld partitioning and the calculation of the dispersion potential is done at every SCF step of the DFT calculation. This computational overhead is essentially independent of the specific method used for dispersion interactions.

In addition to the four dispersion methods discussed above, the non-local van der Waals density functional of Dion and coworkers [224, 225] is also present within FHI-aims, although it has not been extensively tested and it is still considered experimental. The D3 dispersion correction by Grimme et. al. [279, 280] is also implemented in FHI-aims via the `s-dftd3` library [281, 282], and is capable of computing energies, forces and stresses.

Roadmap on Advancements of the FHI-aims Software Package

3.3. DENSITY-FUNCTIONAL BASED METHODS FOR DISPERSION INTERACTIONS

Usability and Tutorials

The TS, MBD@rsSCS, MBD-NL, and XDM methods can easily be used in FHI-aims by specifying their keywords in the `control.in` file. These are:

TS	<code>vdw_ts</code>
MBD@rsSCS	<code>many_body_dispersion</code>
MBD-NL	<code>many_body_dispersion_nl</code>
XDM	<code>xm [basis] or xm [a1 a2]</code>

Note that an older implementation of the TS method (not included with `libmbd`) can also be called via the `vdw_correction_hirshfeld` keyword. While additional options can be called with each keyword, the default settings are recommended for most applications. The options for the TS, MBD@rsSCS, and MBD-NL methods, specified on the same line as the main keyword, are:

<code>self_consistent</code>	Add MBD or TS contribution to the XC potential
<code>vdw_params_kind</code>	"ts" for standard or "tssurf" for surface-adjusted vdW parameters
<code>beta</code>	Damping parameter β for MBD
<code>sr</code>	Damping parameter s_R for TS
<code>k_grid</code>	Integration k -grid for the MBD energy of periodic system
<code>zero_negative</code>	Zero out negative MBD eigenvalues
<code>do_rpa</code>	Calculate MBD energy via the imaginary-frequency integral
<code>rpa_rescale_eigs</code>	Rescale negative eigenvalues in the imaginary-frequency integral

We suggest employing "tssurf" parameter set only when calculating adsorption processes on surfaces and using TS or MBD@rsSCS methods. The damping parameter for TS, MBD@rsSCS, and MBD-NL methods depend only weakly on the basis set in FHI-aims from "light" to "really tight" settings. The fitting of the damping parameter was done using the "tight" settings.

For XDM, the damping parameters depend on the basis set. Hence, additional options must be specified on the same line as the main keyword to inform the code what Becke-Johnson damping parameters to use. As of FHI-aims release 240920, two keyword options are available for this purpose:

<code>xm [basis]</code>	If using a default basis from <code>species_defaults</code> , specify its name here
or	
<code>xm [a1 a2]</code>	Set the damping parameters, a_1 (unitless) and a_2 (in Å), manually

The exchange-correlation functional is detected from the `control.in` and, when the `[basis]` optional parameter is specified, the XDM routine automatically sets optimal damping parameters for that combination. Most common basis sets are supported by this automatic parameter setting, including `light`, `intermediate`, `tight`, and `lightdenser`; the calculation will terminate with an error if the basis-set name is not recognised. The FHI-aims manual provides a comprehensive list of basis defaults and functionals for which XDM damping parameters are available. If your functional/basis combination is not natively supported by XDM, you may manually enter `[a1 a2]` as two floating-point numbers (a_2 in Ångstrom units). Detailed instructions on fitting XDM's damping parameters may be found in the XDM tutorial [283].

Dispersion/vdW methods are not recommended for use with the local density approximation (LDA) due to its well-established systematic overbinding tendency [284]. However, damping parameters for TS, MBD@rsSCS,

MBD-NL, and XDM are available for use with the popular GGA and hybrid functionals, PBE, HSE06, and PBE0. Specifically for XDM, slightly better performance can be obtained using B86b rather than PBE exchange [285], as in the B86bPBE GGA and the B86bPBE0 hybrid functional [275], which is an analogue of PBE0.

The D3 dispersion correction can be activated via the `d3` keyword in the `control.in` file. Both the zero [279] and Becke-Johnson [280] damping options are possible, and may be selected via the `damping` option on the same line as the `d3` keyword. After detection of the exchange-correlation functional in `control.in`, automatic selection of the D3 parameters is supported for over 100 functionals, comprising both internally implemented functionals, and those from LibXC. Alternatively, user specific parameters may be entered on the same line as the `d3` keyword, with all options documented in the FHI-aims manual.

Future Plans and Challenges

An extension of MBD (MBD+C) is currently being developed [286] to explicitly account for the dynamical response of metallic electrons. With respect to MBD@rsSCS and MBD-NL, that rely on harmonically bound charges, MBD+C additionally includes interatomic electron-hopping terms, in analogy to tight-binding models. MBD+C aims to provide a unified many-body description of bound and delocalized electrons, which is necessary in order to capture correct power-law scalings and to automatically include plasmon contributions and metallic screening.

For XDM, a key challenge is the inclusion of additional force and stress terms that are currently neglected. The present implementation assumes that, when calculating the atomic forces or stress tensor, the dispersion coefficients do not change with atomic positions or unit-cell lengths. While this is a fair assumption for isolated molecules and molecular crystals, it is poorer for harder solids, where geometry optimizations can result in overly compact unit-cell volumes (by $\leq 1\%$) and higher energies (by ≤ 0.04 eV) than the true potential-energy minimum [285]. While the additional stress and force terms are almost always neglected in all dispersion corrections, their omission disproportionately affects XDM due to its highly varying dispersion coefficients near equilibrium geometries [287, 288]. Accounting for these missing force and stress terms is an area of active work, and ultimately depends on implementing the derivatives of the Hirshfeld weights that partition the total electron density into atomic pieces with respect to atomic coordinates.

Additional work seeks to improve the atomic polarizabilities used with XDM by replacing the simple scaling of free-atomic values using ratios of Hirshfeld atomic volumes. Options to address this include implementing an atomic polarizability functional such as that used by MBD-NL [272], or replacing the current simple Hirshfeld partitioning with a more sophisticated iterative Hirshfeld [289] or fractionally ionic [290] scheme. While iterative Hirshfeld partitioning is already implemented in FHI-aims, it currently only works reliably for the aperiodic case. Lastly, a manuscript is in preparation where the XDM dispersion energy is damped by a different, 1-parameter function depending on atomic numbers [291]; this method will soon be available in FHI-aims.

Finally, it is also possible to include beyond-dipole dispersion terms in the TS and MBD methods [292, 293]. To determine the multipolar parameters, it is possible to use scaling laws based on quantum Drude oscillators and/or their ratios determined by the XDM approach [273].

Acknowledgements

The authors gratefully acknowledge contributions from Nicola Ferri and Ville Havu. RAD gratefully acknowledges financial support from the National Science Foundation under Grant No. CHE-1945676 and an Alfred P. Sloan Research Fellowship. ERJ thanks the Natural Sciences and Engineering Research Council (NSERC) of Canada for financial support, and the Royal Society for a Wolfson Visiting Fellowship. AOR thanks the Spanish Ministerio de Ciencia e Innovación and the Agencia Estatal de Investigación, projects PGC2021-125518NB-I00 and RED2022-134388-T co-financed by EU FEDER funds; the Spanish MCIN/AEI/10.13039/501100011033 and European Union NextGenerationEU/PRTR for grants TED2021-129457B-I00 and CNS2023-144958. AT acknowledges the European Research Council (ERC AdG FITMOL) for funding. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

3.4 Random Phase Approximation and Beyond for Total Energies and Forces

Volker Blum^{1,2}, Francisco A. Delesma³, Dorothea Golze³, Florian Merz⁴, *Xinguo Ren^{5,10,a}, Patrick Rinke^{6,7,8,9}, Matthias Scheffler¹⁰, Muhammad N. Tahir⁵, Alexandre Tkatchenko¹¹, Igor Ying Zhang^{10,12,b}, and Tong Zhu¹³

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁴Lenovo HPC Innovation Center, Meitnerstr. 9, D-70563 Stuttgart, Germany

⁵Institute of Physics, Chinese Academy of Sciences, 3rd South Str. 8, Beijing 100190, China

⁶Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

⁷Physics Department, TUM School of Natural Sciences, Technical University of Munich, Garching, Germany

⁸Atomistic Modelling Center, Munich Data Science Institute, Technical University of Munich, Garching, Germany

⁹Munich Center for Machine Learning (MCML), Munich, Germany

¹⁰The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹¹Department of Physics and Materials Science, University of Luxembourg, L-1511, Luxembourg City, Luxembourg

¹²Collaborative Innovation Center of Chemistry for Energy Materials, Shanghai, Key Laboratory of Molecular Catalysis and Innovative Materials, MOE Key Laboratory of Computational Physical Sciences, Department of Chemistry, Fudan University, Shanghai 200433, China

¹³Department of Electrical and Computer Engineering, University of Toronto, Toronto, Ontario M5S 1A4, Canada

*Coordinator of this contribution.

^aCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

^bCurrent Address: Collaborative Innovation Center of Chemistry for Energy Materials, Shanghai, Key Laboratory of Molecular Catalysis and Innovative Materials, MOE Key Laboratory of Computational Physical Sciences, Department of Chemistry, Fudan University, Shanghai 200433, China

Summary

The random phase approximation (RPA) is classified under the fifth rung of Jacob's ladder, which organizes exchange-correlation (XC) functionals in Kohn-Sham (KS) density functional theory [209]. Over the past two decades, RPA has gained recognition in computational chemistry, physics, and materials science [294, 295] for its ability to accurately capture non-local electron correlations and effectively describe subtle energy differences, such as adsorption energies of molecules on surfaces and energy differences between crystal polymorphs. The RPA concept also led to the development of the many-body dispersion method [270], which addresses van der Waals interactions beyond atomic pairwise summations. While the standard RPA approach

Roadmap on Advancements of the FHI-aims Software Package

3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

tends to underestimate the atomization energies of molecules and the cohesive energies of solids, this issue can be mitigated by incorporating corrections from renormalized single excitations (rSE) and second-order screened exchange contributions [296–298]. Since 2009, the FHI-aims code has included an RPA implementation for molecular geometries using numeric atomic-centered orbitals [8, 62, 299]. Recently, this implementation has been extended to periodic systems and force calculations for molecules [68]. The RPA energy expression in FHI-aims is applied as a post-processing correction, i.e., it is computed following a self-consistent calculation with a different DFA and then effectively replaces the E_{xc} term in Eq. (3.1). One prominent feature of RPA-based methods is that they can describe vdW interactions seamlessly and automatically capture the non-additive and non-anisotropic vdW effects. However, compared to the dedicated vdW-inclusive approaches presented in Sec. 3.3, RPA is computationally much more demanding. Nevertheless, in complex chemical environments and when there are strong variations among the results obtained by the empirical/semi-empirical vdW-inclusive approaches, RPA can be used as a benchmark method.

The implementation of RPA in FHI-aims is based on the resolution of identity (RI) approximation, where the Kohn-Sham density response function and Coulomb operator are represented in terms of a set of auxiliary basis functions (ABFs). Different RI flavors are implemented, including a global RI based on the Coulomb-metric (RI-V) [62] and a localized RI based on a two-center approximation (LRI, also known as RI-LVL) [63]. The single-point RPA calculations for finite systems (molecular geometries) can be performed using both RI-V and RI-LVL, while the geometry relaxations and single-point RPA energy calculations for periodic systems are exclusively based on RI-LVL. Currently, RPA calculations can be done for molecules with over 100 atoms, and RPA (and RPA+rSE) geometry relaxation can be performed for systems with a few tens of atoms [300]. The current periodic RPA implementation in FHI-aims enables the computation of relatively simple crystal structures with finite \mathbf{k} -point sampling. However, the code infrastructure for RPA force calculations for periodic systems is still under development. Large-scale periodic calculations can be done by interfacing FHI-aims with LibRPA [301], a standalone package for evaluating RPA correlation energy based on low-scaling, real-space algorithm.

Current Status of the Implementation

In standard RPA calculations, the total energy is given by $E^{\text{RPA}} = E^{\text{HF}}[\{\psi_i^{\text{KS}}\}] + E_c^{\text{RPA}}$, where $E^{\text{HF}}[\{\psi_i^{\text{KS}}\}]$ is the Hartree-Fock total energy evaluated using the KS orbitals and E_c^{RPA} is the RPA correlation energy. A convenient expression to evaluate E_c^{RPA} is

$$E_c^{\text{RPA}} = \frac{1}{2\pi} \int_0^\infty d\omega \text{Tr} [\ln(1 - \chi^0(i\omega)V) + \chi^0(i\omega)V] \quad (3.14)$$

where χ^0 and V can be understood as matrices in the ABF basis. Specifically,

$$\chi_{\mu\nu}^0(i\omega) = \sum_{m,n,\sigma} \frac{(f_{m,\sigma} - f_{n,\sigma})C_{m,n,\sigma}^\mu C_{n,m,\sigma}^\nu}{\epsilon_{m,\sigma} - \epsilon_{n,\sigma} - i\omega} \quad (3.15)$$

where $C_{m,n,\sigma}^\mu$ are the RI expansion coefficients of products of KS spin-orbitals in terms of ABFs ($\{P_\mu(\mathbf{r})\}$),

$$\psi_{m,\sigma}(\mathbf{r})\psi_{n,\sigma}(\mathbf{r}) = \sum_{\mu} C_{m,n,\sigma}^\mu P_\mu(\mathbf{r}) \quad (3.16)$$

Roadmap on Advancements of the FHI-aims Software Package

3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

and V is the Coulomb matrix

$$V_{\mu\nu} = \int d\mathbf{r}d\mathbf{r}' \frac{P_{\mu}(\mathbf{r})P_{\nu}(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}. \quad (3.17)$$

For an in-depth discussion of the theoretical basis and algorithmic formulation behind these equations, the readers are referred to Refs. [294, 302]. In FHI-aims, we use an “on-the-fly” procedure to generate suitable atom-centered ABFs for a given single-particle NAO basis set [62, 63]. The $C_{m,n,\sigma}^{\mu}$ expansion coefficients in Eq. 3.16 are obtained by first computing the RI coefficients $\tilde{C}_{i,j}^{\mu}$ that expand pair products of atomic basis functions $\phi_i\phi_j$ in terms of ABFs, and then transform $\tilde{C}_{i,j}^{\mu}$ into $C_{m,n,\sigma}^{\mu}$ by multiplying with the KS eigenvectors $c_{n\sigma}^i$. The different methods for determining $\tilde{C}_{i,j}^{\mu}$ distinguish the various flavors of RI, such as RI-V and RI-LVL. The primary challenge in RPA correlation energy calculations is evaluating the χ^0 matrix as described in Eq. 3.15, which scales with the system size as $O(N^4)$.

Extending RPA energy calculations from finite systems to periodic systems with finite \mathbf{k} -point sampling presents a significant challenge because the KS orbitals carry a Bloch wavevector. Consequently, $C_{m,n,\sigma}^{\mu}$, $\chi_{\mu\nu}^0$, and $V_{\mu\nu}$ all become momentum-dependent, i.e., $C_{m,n,\sigma}^{\mu} \rightarrow C_{m,n,\sigma}^{\mu}(\mathbf{k}, \mathbf{q})$, $\chi_{\mu\nu}^0 \rightarrow \chi_{\mu\nu}^0(\mathbf{q})$, and $V_{\mu\nu} \rightarrow V_{\mu\nu}(\mathbf{q})$ and additional sums over \mathbf{q} (or \mathbf{k}) points are required for the trace or summation operations in Eq. 3.14 and 3.15. The extension to reciprocal space is straightforward except for $\mathbf{q} = 0$ where certain elements of $V_{\mu\nu}(\mathbf{q} = 0)$ diverge. In our implementation, the truncated Coulomb operator is used at $\mathbf{q} = 0$ to avoid any such divergences, while the full Coulomb operator is used elsewhere in the Brillouin zone (BZ).

As previously noted, FHI-aims also supplies gradients of the RPA total energy with respect to nuclear displacements, facilitating structure relaxations based on RPA forces. In our implementation, the total RPA force on atom I located at spatial position τ_I is decomposed into the following terms

$$\begin{aligned} \mathbf{F}_I^{\text{RPA}} &= -\frac{dE^{\text{RPA}}}{d\tau_I} \\ &= -\frac{dE^{\text{DFA}}}{d\tau_I} + \frac{dE_{xc}^{\text{DFA}}}{d\tau_I} - \frac{dE_x^{\text{EX}}}{d\tau_I} - \frac{dE_c^{\text{RPA}}}{d\tau_I} \end{aligned} \quad (3.18)$$

where $-\frac{dE^{\text{DFA}}}{d\tau_I}$ is the gradient of a local or semilocal density functional approximations (DFA) and has been long available in FHI-aims. The RPA gradient is obtained by subtracting the contribution of the local/semilocal XC functional from the DFA force, and adding the exact exchange and RPA correlation terms instead. To calculate the latter two, the derivatives of the KS eigenvectors with respect to the nuclear displacements are calculated with density functional perturbation theory (DFPT) [303]. Further details can be found in Ref. [68]. Currently, the RPA force calculation functionality in FHI-aims is fully developed for molecular geometries, while the implementation for periodic systems is still in progress. The main computational steps for the correlation part of the RPA energy and force calculations, as implemented in FHI-aims (and LibRPA), are shown in Fig. 3.4.

Usability and Tutorials

FHI-aims RPA total energy and force calculations are well-established by now. Here, we provide a brief overview of how to perform these calculations using FHI-aims.

Total Energies: All functionalities for RPA-based total energy calculations are now accessible through simple

Roadmap on Advancements of the FHI-aims Software Package

3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

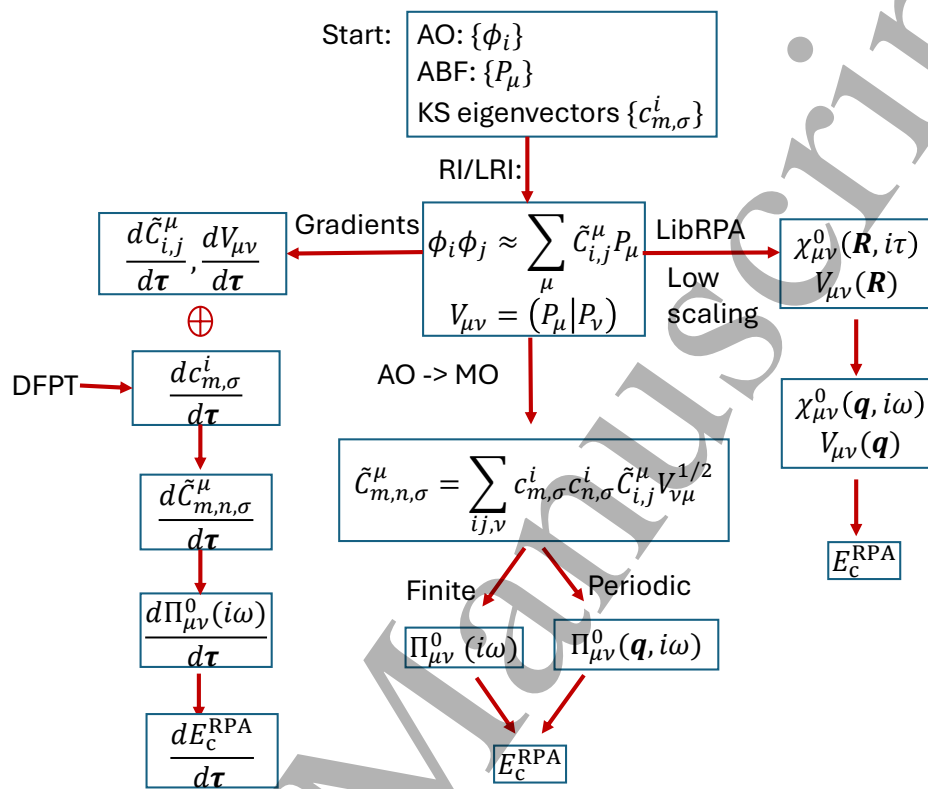


Figure 3.4: Illustration of the major computational steps in FHI-aims for evaluating the correlation part of the RPA ground-state energies and forces. Here AO denotes atomic orbitals and MO denote KS molecular orbitals. Note that $\tilde{C}_{m,n,\sigma}^{\mu}$ here is related to $C_{m,n,\sigma}^{\mu}$ in Eq. 3.16 by $\tilde{C}_{m,n,\sigma}^{\mu} = \sum_{\nu} C_{m,n,\sigma}^{\nu} V_{\nu\mu}^{1/2}$, and similarly Π^0 is related to χ^0 by $\Pi^0 = V^{1/2} \chi^0 V^{1/2}$.

keywords in the `control.in` file, as documented in the FHI-aims manual. Currently, RPA-type calculations are carried out in a post-processing manner, and to invoke RPA-based energy calculations, the keywords

```
xc dfa
total_energy_method type
```

need to be set in the `control.in` file. While `dfa` specifies the prior density functional approximation (dfa) that generates input orbitals and orbital energies for RPA-type calculations, `type` is a keyword (string) that specifies a chosen post-processing RPA or beyond-RPA method. For example, for a standard RPA@PBE calculation, `dfa` should be PBE and `type` should be RPA. In addition to RPA, other choices like `rpa+sosex` and `rpt2` are also legitimate, which yields RPA+SOSEX and the renormalized 2nd-order perturbation the-

Roadmap on Advancements of the FHI-aims Software Package

3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

ory (rPT2) total energies, respectively. Such beyond-RPA methods usually give improved results when the standard RPA shows pronounced underbinding behavior. However, RPA+SOSEX and rPT2 are substantially more expensive than RPA. On the other hand, the rSE correction is very cheap, and this term is automatically calculated whenever one performs RPA calculations. RPA+rSE typically describes van der Waals bonded systems more accurately than the standard RPA. Further details about the performance of RPA and beyond-RPA methodologies can be found in Refs. [294, 298]. Periodic RPA energy calculations using FHI-aims can be done as usual by setting the `k_grid` in `control.in` and `lattice_vector` in the `geometry.in` file. An example of such calculations for the Ar crystal was described in Ref. [304]. Periodic RPA calculations can be efficiently performed using the LibRPA library, which implements a low-scaling real-space RPA algorithm [301]. These low-scaling RPA calculations utilize the minimax imaginary time-frequency grids provided by the GreenX library [305], accessible in FHI-aims with the keyword `freq_grid_type minimax`. Recent studies have shown that minimax imaginary frequency grids converge approximately 3.5 times faster than modified Gauss-Legendre grids for integrating the RPA correlation energy, Eq. (3.14). This enhancement significantly reduces the computational prefactor in canonical RPA calculations for both solids and molecules [306]. At present, the coupling between FHI-aims and LibRPA is implemented via a file-based interface. To prepare the data, the user only needs to add the keyword “`output librpa`” to the FHI-aims input, which triggers the export of all quantities required by LibRPA in the expected format. No additional manual post-processing of the generated files is needed. With this exported dataset and a separate LibRPA input file specifying the desired calculation settings, the user can directly run the low-scaling RPA calculation using LibRPA. For more details, please refer to Ref. [307].

Forces: The force calculations can be done for molecules at the level of RPA and RPA+rSE using FHI-aims. To run such calculations, additional keywords related to DFPT and choice of the RPA force method also need to be set. For example, structure relaxations based on RPA@PBE gradients can be performed using the following set of keywords in `control.in` file

```
xc pbe
relativistic none
occupation_type gaussian 0.001
RI_method lvl
DFPT vibration_reduce_memory
relax_geometry bfgs 1e-2
sc_accuracy_forces 1e-2
least_memory_4 .true.
rpa_force freq_formula_method
frequency_points 16
my_prodbas_threshold 1.e-12
```

Both RPA energy and force implementations in FHI-aims are MPI-parallelized. Depending on the system size, parallel calculations can scale up to hundreds to thousands of CPU cores. A recent performance benchmark and applications to water clusters of the RPA force implementation can be found in Ref. [300]. Regarding the timings, the current state of the art of computational efficiency of molecular RPA and RPA+rSE force calculations using FHI-aims can be found in Ref. [308]. As for the improvement from the canonical scaling in FHI-aims to the sub-quadratic scaling in LibRPA for periodic RPA calculations, detailed timings together with the scaling behavior has been reported in Ref. [301]. The basic functionalities of RPA and beyond-RPA energy and force calculations are detailed in an online tutorial [309].

An important issue in RPA calculations is the choice of the single-particle basis set. The standard FHI-aims

Roadmap on Advancements of the FHI-aims Software Package

3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

basis sets (*tier's*) can be used, but it is mandatory to perform a counterpoise (CP) correction to the basis set superposition error (BSSE). Furthermore, it is not possible to extrapolate to the complete basis set (CBS) limit using *tiers*. A recommended choice of basis sets for RPA calculations using FHI-aims are NAO-VCC-*n*Z [20] or its localized variants. In practice, it is often a good idea to perform a CP correction and compare the CP-corrected and uncorrected results. For elements where NAO-VCC-*n*Z is not available, correlation-consistent Gaussian basis sets provide a viable alternative.

Future Plans and Challenges

RPA calculations based on NAO basis sets encounter several challenges that we aim to address in future work. Firstly, the RPA correlation energy converges slowly with respect to single-particle basis sets, often necessitating extrapolation to the CBS limit to achieve high-quality results. In FHI-aims, such “correlation consistent” NAO-VCC-*n*Z [20] that facilitate CBS extrapolation are officially available only for light elements up to Ar. Recent efforts have focused on developing NAO-VCC-*n*Z basis sets beyond Ar, but so far, success has been limited to a few selected main-group elements [310]. Additionally, BSSE is significant in NAO-based RPA calculations. While BSSE can be corrected using the counterpoise scheme for binding energies of finite systems, counterpoise corrections are not straightforward for periodic systems. Practical periodic RPA calculations depend on the relatively small BSSE of the newly developed NAO-VCC-*n*Z basis sets and the cancellation between BSSE and basis set incompleteness errors. However, the remaining uncertainty is often difficult to estimate. Therefore, a better strategy is urgently needed to address the basis set issues for RPA calculations with NAOs.

The second issue concerns the handling of the Coulomb singularity at the Γ point in periodic RPA calculations. Ideally, the RPA integrand does not exhibit diverging behavior as $q \rightarrow 0$ in the Brillouin zone (BZ), at least for insulating systems, because the $1/q^2$ divergence in the Coulomb operator is offset by the q^2 factor in the KS response function $\chi^0(\mathbf{q})$. However, in the matrix representation using auxiliary basis functions (ABFs), this cancellation is incomplete, necessitating special numerical treatment. Practically, a mixed scheme—using the truncated Coulomb operator for $\mathbf{q} = 0$ and the full Coulomb operator elsewhere in the BZ—works quite well. Nonetheless, the performance of this mixed scheme degrades for very sparse \mathbf{k} grids or Γ -point-only calculations. Developing a more consistent Γ -point treatment scheme for periodic RPA calculations remains a goal for future development in FHI-aims. A related Coulomb singularity issue in the periodic *GW* calculations and how it was addressed in FHI-aims can be found in Ref. [36].

The third issue pertains to fractional occupations in RPA calculations. These fractional occupations frequently arise in preceding KS-DFT calculations, particularly in systems with open-shell transition metal atoms, defects, or radicals. While the RPA formalism can theoretically accommodate fractional occupations in the KS response function, the practical results are highly sensitive to these occupation numbers, often leading to a decline in RPA performance. Consequently, there is an urgent need to advance beyond the standard RPA to effectively handle such cases. This challenge is also closely linked to the broader goal of extending the RPA formalism to address systems with strong multi-configurational characteristics and finite temperatures.

Acknowledgements

We acknowledge the funding support from the National Natural Science Foundation of China (Grant Nos 12374067 and 12188101) and the National Key Research and Development Program of China (Grant Nos 2022YFA1403800 and 2023YFA1507004). The authors acknowledge the European Union's Horizon 2020

*Roadmap on Advancements of the FHI-aims Software Package**3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES*

research and innovation program for financial support under the grant number 951786 (Nomad Center of Excellence) and the Horizon Europe MSCA Doctoral network grant n.101073486 (EUSpecLab). D. G. acknowledges funding by the Emmy Noether Program of the German Research Foundation (project number 453275048). We also wish to acknowledge CSC - IT Center for Science (Finland), the Jülich Supercomputer Computer Center (Germany) and Aalto Science-IT project for computational resources. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233.

*Roadmap on Advancements of the FHI-aims Software Package*3.4. RANDOM PHASE APPROXIMATION AND BEYOND FOR TOTAL ENERGIES AND FORCES

Accepted Manuscript

4 Electronic Excited States

4.1 Coupled-Cluster Theory for the Ground State and for Excitations

Andreas Grüneis¹, *Evgeny Moerman², *Matthias Scheffler², Tonghao Shen^{2,a}, and Igor Ying Zhang^{2,a}

¹Institute for Theoretical Physics, TU Wien, Wiedner Hauptstraße 8–10/136, 1040 Vienna, Austria

²The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

^aCurrent Address: Department of Chemistry, Fudan University, Shanghai 200433, People's Republic of China

Summary

In the molecular quantum chemistry community, coupled-cluster (CC) methods are well-recognized for their systematic convergence and reliability. The extension of the theory to extended systems has been comparably recent [311], so that developments and studies of periodic CC methods for both the ground-state and for excited states are still active fields of research and provide valuable benchmark data when the reliability of density functional approximations is questionable. In this contribution we describe the CC-aims interface between the FHI-aims and the Cc4s software packages. This linkage makes a variety of correlated wave function-based ground-state methods including Møller-Plesset perturbation theory (MP2), the random-phase approximation (RPA) and the *gold-standard of quantum chemistry* CCSD(T) method for both molecular and periodic applications accessible. This contribution discusses these ground-state methods for clusters and molecules, as well as for periodic systems. In particular, we discuss recent advancements and the implementation of the equation-of-motion CC method for the calculation of ionization (IP-EOM-CCSD) and electron attachment (EA-EOM-CCSD) processes. Open questions and routes to solutions are discussed as well.

Current Status of the Implementation

The Cc4s code constitutes an open-source quantum chemistry software package, which features several correlated wave function methods. As a post-SCF code, Cc4s requires single-particle eigenenergies and wave functions from a mean-field calculation, which another electronic structure package must provide. Using the CC-aims interface [312] the relevant quantities are conveniently obtained from a FHI-aims Hartree-Fock calculation. CC-aims, then, parses and converts the needed quantities (e.g the single-particle eigenenergies) to a Cc4s-compatible format and computes additional quantities (e.g the Coulomb vertex). The files generated by CC-aims are then used to launch a Cc4s calculation with any of the therein implemented wave-function methods. This workflow is illustrated in Figure 4.1.

Below, we discuss the coupled-cluster approach for the electronic ground-state and for excited states for molecules and clusters and for periodic solids.

In practice, the limiting factor of CC methods is the substantial memory requirement, which results from the size of the Coulomb integral tensor

Roadmap on Advancements of the FHI-aims Software Package

4.1. COUPLED-CLUSTER THEORY FOR THE GROUND STATE AND FOR EXCITATIONS

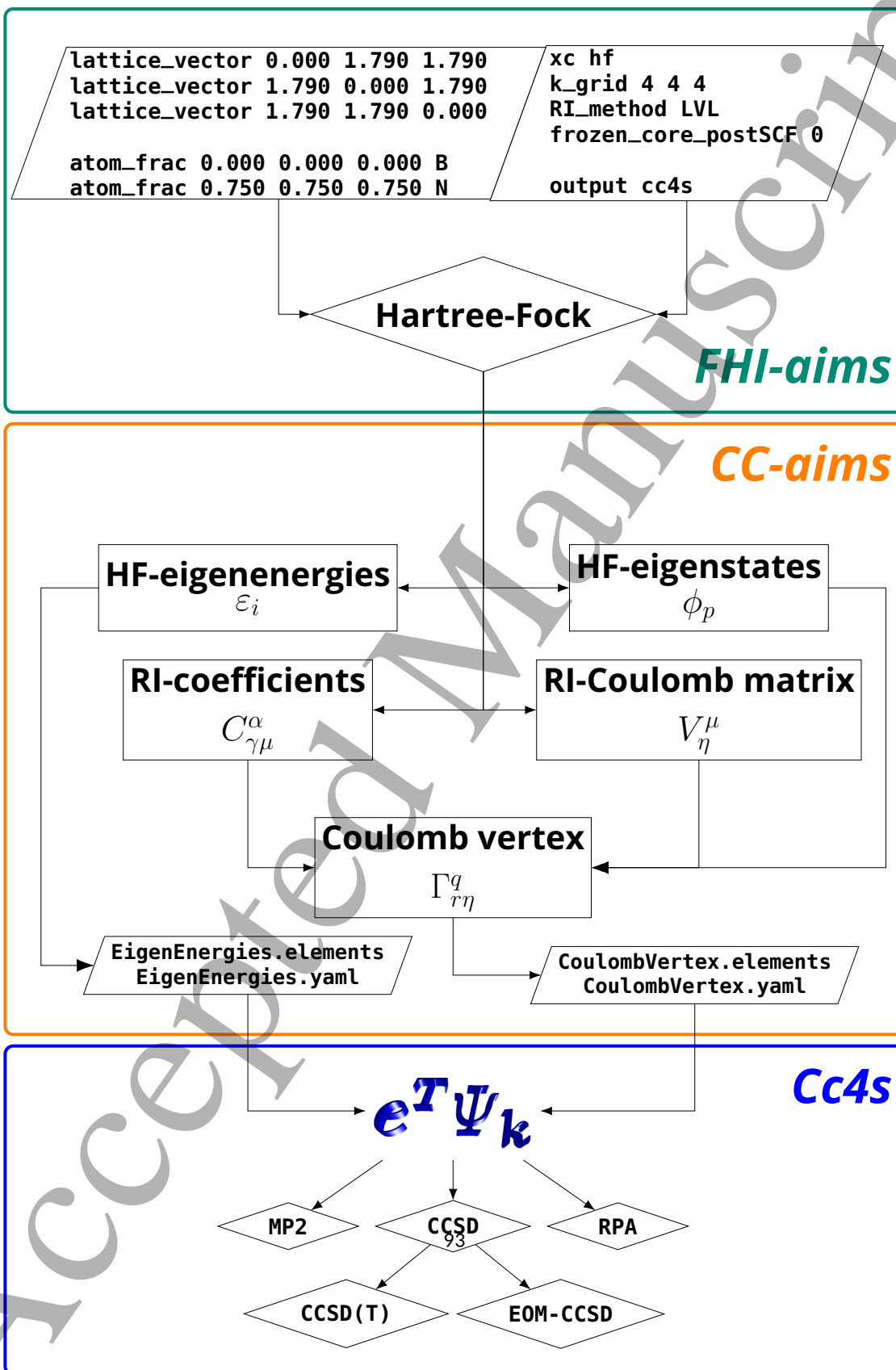


Figure 4.1: The Cc4s@FHI-aims workflow

Roadmap on Advancements of the FHI-aims Software Package

4.1. COUPLED-CLUSTER THEORY FOR THE GROUND STATE AND FOR EXCITATIONS

$$V_{rs}^{pq} = \int d\mathbf{r}\mathbf{r}' \frac{\phi_p^*(\mathbf{r})\phi_q^*(\mathbf{r}')\phi_r(\mathbf{r})\phi_s(\mathbf{r}')}{|\mathbf{r} - \mathbf{r}'|}, \quad (4.1)$$

where ϕ_p , ϕ_q , ϕ_r and ϕ_s denote four single-particle wave functions. As a consequence, the memory to store this tensor scales with the fourth power of the basis set size. By performing a low-rank decomposition of V_{rs}^{pq} , the Coulomb vertex $\Gamma_r^{p,\eta}$, a rank three tensor with a significantly smaller memory footprint [313] is defined

$$V_{rs}^{pq} \approx \sum_{\eta} \Gamma_r^{p,\eta} \Gamma_{s\eta}^{q*}, \quad (4.2)$$

where η indexes the basis functions of an auxiliary basis. As a memory-saving measure, CC-aims computes the Coulomb vertex. The auxiliary basis is that of the resolution-of-the-identity (RI) employed in FHI-aims. In combination with CC-aims, one can choose between the RI-V scheme for molecular applications and its localized approximation, the RI-LVL scheme for periodic ones. While the former is deemed very accurate, the latter is more memory-efficient but in general of insufficient accuracy [63]. However, the incompleteness of the RI-LVL auxiliary basis for Hartree-Fock, MP2 and RPA methods can be satisfactorily resolved by manually adding a few auxiliary *f*-, *g*- and *h*-type basis functions with small effective charges [63]. This approach is also applicable for CC calculations. The memory footprint of the Coulomb vertex can be further reduced by performing a principal component analysis, with which up to 70% of the Coulomb vertex can be discarded in many applications [313].

Currently, the periodic CC infrastructure of FHI-aims does not allow to perform spin-polarized calculations and the starting point needs to be canonical Hartree-Fock orbitals. For periodic applications, the most important constraint is given by the inability of Cc4s to perform an explicit *k*-point summation. Instead, Cc4s requires a super cell based treatment of a periodic system. As a consequence, Cc4s does not make use of the translational symmetry of crystalline systems, which would reduce the memory scaling by a factor of N_k and the computational scaling by a factor of N_k^2 , with N_k being the number of *k*-points.

Usability and Tutorials

A variety of FHI-aims based calculations with the CCSD, CCSD(T) and the EOM-CCSD implementations in Cc4s have already been performed for small and medium-sized molecules and for crystalline insulators and semiconductors. Tutorials for using the workflow involving FHI-aims, CC-aims and Cc4s for both molecules and crystalline materials are available on the FHI-aims platform [314]. These tutorials include a detailed step-by-step guide on the installation of the necessary software (i.e CC-aims and Cc4s) and the calculation of ground-state and excitation energies. Currently, the tutorial consists of three parts. The first one introduces the FHI-aims/CC-aims/Cc4s workflow by an exemplary calculation of the MP2 correlation energy of the paracetamol molecule. The second tutorial focuses on a periodic application and demonstrates the importance of a careful consideration of the system-size dependent error for correlated wave function methods. To that end, the necessary steps involved in the calculation of the CCSD cohesive energy of Neon are presented, the results of which are also shown in Figure 4.2c. In the third tutorial, the additional steps to compute quasi-particle energies via the IP- and EA-EOM-CCSD method are described in detail, for which the ionization potential of uracil (see Figure 4.2b) and the band gap of *trans*-polyacetylene is computed [315].

Figure 4.2 shows four typical applications of CC theory using FHI-aims for molecular and extended system

Roadmap on Advancements of the FHI-aims Software Package

4.1. COUPLED-CLUSTER THEORY FOR THE GROUND STATE AND FOR EXCITATIONS

involving both the ground state and charged excited states. Figure 4.2a and 4.2b confirm the accuracy of both ground state and excited state properties for molecules using our valence-correlation consistent NAO-VCC- nZ basis sets [20]. Via extrapolation to the complete basis set (CBS) limit, we are able to reproduce the stacking energy of uracil (Figure 4.2a) within a few meV. Going beyond the ground-state, in Figure 4.2b we show results of the IP-EOM-CCSD method to obtain the ionization potentials of the five nucleobases, which are in good agreement (within ≈ 100 meV) with previously reported values by Tripathi et al. [316].

The remaining deviation stems from the use of different finite basis sets: cc-pVTZ in Reference [316] and NAO-VCC-3Z in our study. For ground-state properties, we have found that the NAO-VCC- nZ basis set performs better than the cc-pV nZ basis set for advanced correlation methods, including MP2, RPA, and CCSD(T) [20, 317]. This discrepancy can be further minimized by extrapolating to the CBS limit.

The remaining deviation is attributable to the utilization of a 3Z basis set, for which a remaining difference of that magnitude to the CBS limit is expected.

Figure 4.2d demonstrates the applicability of the EOM-CCSD method to obtain quasi-particle band gaps. For that we studied the fundamental, indirect $K \rightarrow \Gamma$ band gap of two-dimensional hexagonal boron nitride (hBN), for which different state-of-the-art methods like generalized KS-DFT and the GW approximation, and highly accurate methods like Diffusion Monte Carlo (DMC) yielded very different results. Our analysis shows good agreement with the DMC result by Hunt et al. [318], confirming that an accurate treatment of electronic correlation is crucial in this system. Figure 4.2c shows the cohesive energy of the Neon crystal. The stability of this crystal is almost exclusively determined by van-der-Waals interactions, which CC methods are known to precisely capture. With Cc4s@FHI-aims we obtain a cohesive energy of -17.9 meV/atom at the CCSD level. The experimental value, already adjusted for zero-point fluctuations, is -27 meV/atom, with an estimated zero-point contribution of 7 meV/atom [319]. For these calculations the NAO-VCC-2Z and -3Z basis sets were employed. Performing a conventional two-point extrapolation to the CBS limit yields the results in Figure 4.2c with a remaining difference to the CBS limit of 8 meV/atom, which was added to all of the data points. A more accurate cohesive energy was obtained by accounting for triple excitations in a perturbative manner. That triple (T)-correction was found to be mostly independent of the system size, and determined to lie between -8.7 meV/atom and -12.6 meV/atom. By adding these values to the extrapolated CCSD cohesive energy, a CCSD(T) result of -29.2 ± 1.9 meV/atom was found, which is in excellent agreement with the experimental finding.

Future Plans and Challenges

As mentioned before, currently the biggest limitation in performing periodic CC calculations via Cc4s is the lack of an explicit k -point summation. The translational symmetry of a periodic system is reflected in the block-sparsity of the CC tensors. A block-sparse implementation of the tensor contraction engine in Cc4s is currently under development and is expected to severely reduce the computational and memory requirements of CC calculations. Further future plans involve the extension of the Cc4s functionalities to include spin-polarized and non-HF based calculations. The convergence of CC ground and excited state properties with respect to system size can be very slow. As has been in part demonstrated in Figure 4.2c, one can correct the corresponding finite-size errors by additionally employing another, computationally cheaper non-CC method (e.g the RPA for the ground-state or the G_0W_0 method for quasi-particles [321]), which exhibits a similar system size convergence. However, via the implementation of a block-sparse tensor treatment in Cc4s, more work has to be done to verify the accuracy and the limitations of this methodology. Development of new strategies to reduce the basis set incompleteness error is the second, very important

Roadmap on Advancements of the FHI-aims Software Package

4.1. COUPLED-CLUSTER THEORY FOR THE GROUND STATE AND FOR EXCITATIONS

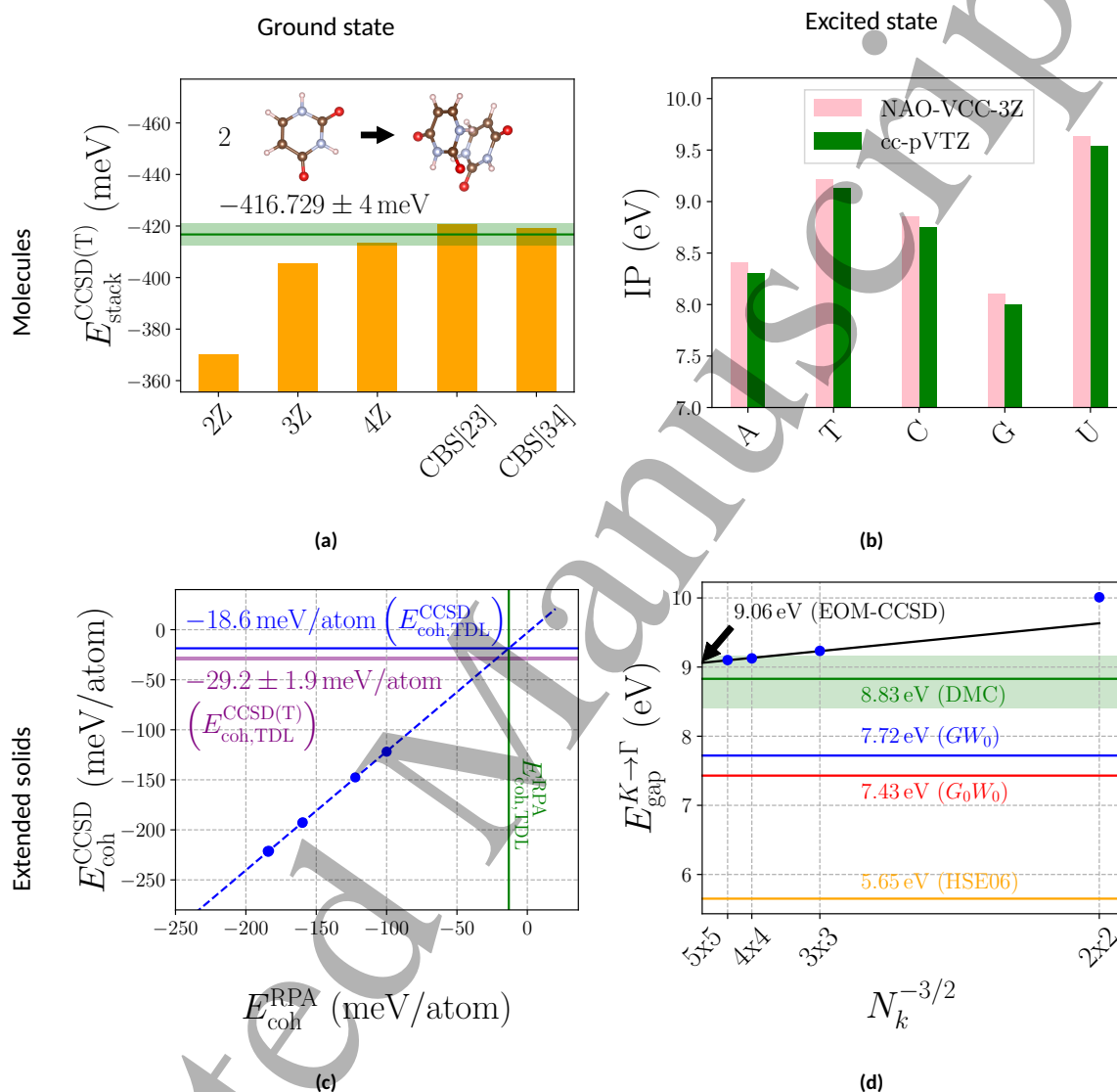


Figure 4.2: A selection of molecular and periodic CC calculations with the NAO-VCC- n Z basis sets: Figure 4.2a stacking energy of a uracil dimer on the CCSD(T) level of theory compared to Reference [320], Figure 4.2b vertical ionization potentials of adenine (A), thymine (T), cytosine (C), guanine (G) and uracil (U) via IP-EOM-CCSD compared to Reference [316], Figure 4.2c Finite-size convergence of the cohesive energy of Neon using the CCSD method plotted against the RPA results. The (T)-correction has been found to be mostly system-size-independent and has been added to the extrapolated CCSD result. Figure 4.2d Electronic band gap of two-dimensional boron nitride via IP- and EA-EOM-CCSD compared to HSE06 and higher-level correlated methods in Reference [318]. The opaque green area illustrates the uncertainty of the DMC result.

*Roadmap on Advancements of the FHI-aims Software Package***4.1. COUPLED-CLUSTER THEORY FOR THE GROUND STATE AND FOR EXCITATIONS**

future goal, to improve the precision of CC calculations. In ground-state CC applications in particular, where the quantity of interest itself is sometimes on the order of a few meV, one is forced to use very big basis sets, even if extrapolation schemes are accessible to achieve the necessary precision. One solution is to use the much more compact natural orbital basis instead of the canonical HF basis [322]. One way to construct natural orbitals for correlated wave function methods is to construct the one-electron reduced density matrix at the MP2 level [323]. Diagonalization of the density matrix yields the natural orbitals and selection of only those with a sufficiently high occupation numbers allows to significantly reduce the number of single-particle states involved in the subsequent CC calculation, thus reducing the overall computational cost. The possibility to construct natural orbitals in FHI-aims or CC-aims is therefore a very desirable feature. Alternative approaches to reduce the computational cost of CC methods for larger molecules and periodic structures include embedding methods and local CC schemes. Embedding approaches are particularly valuable when a small region requiring highly accurate CC treatment can be identified, while remaining atoms are treated with less expensive methods such as DFT [324], MP2 [325], or RPA [326]. Local CC methods, including local natural orbital (LNO) [327] and the domain based local pair natural orbital (DLPNO) [328] CC approaches directly address the steep computational scaling by localizing molecular orbitals and considering only those contributing most significantly to the correlation energy. By approximating electronic correlation as local, these methods achieve nearly linear system-size scaling. Recently, the LNO-CC framework has been expanded to periodic solids [329], paving the way to broader application of coupled-cluster methods in more complex materials. While these frameworks enable treatment of systems inaccessible to canonical CC methods [320, 330–332], errors from the local approximation can become problematic for larger systems where long-range interactions are significant. Despite these advances in approximate methods, canonical CC implementations remain essential for rigorous benchmarking and method development. The systematic hierarchy of CC approximations provides well-defined reference data that are crucial for assessing the accuracy of emerging electronic structure methods, including local and embedding approaches, as well as machine learning potentials and density functional approximations. The development and optimization of canonical CC methods for both molecules and solids, as presented in this work and now accessible to all FHI-aims users, therefore continues to play a vital role in advancing the field of computational electronic structure theory.

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

4.2 All-Electron GW for Finite Systems

Volker Blum^{1,2}, Fabio Caruso^{3,4,a}, *Dorothea Golze⁵, Jannis Kockläuner⁵, Moritz Leucke⁵, Qinglong Liu⁵, Ramón L. Panadés-Barrueta⁵, *Xinguo Ren^{6,7,b}, Patrick Rinke^{8,9,10,11}, Yanyong Wang¹², and Tong Zhu¹³

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Christian-Albrechts-Universität zu Kiel, 24118 Kiel, Germany

⁵Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁶Institute of Physics, Chinese Academy of Sciences, 100190 Beijing, China

⁷The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁸Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

⁹Physics Department, TUM School of Natural Sciences, Technical University of Munich, Garching, Germany

¹⁰Atomistic Modelling Center, Munich Data Science Institute, Technical University of Munich, Garching, Germany

¹¹Munich Center for Machine Learning (MCML), Munich, Germany

¹²CAS Key Laboratory of Quantum Information, University of Science and Technology of China, Hefei, Anhui 230026, China

¹³University of Toronto, Toronto, Ontario M5S 1A4, Canada

* Coordinator of this contribution.

^aCurrent Address: Christian-Albrechts-Universität zu Kiel, 24118 Kiel, Germany

^bCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

Summary

The GW approximation [333] to Hedin's equations is a widely used method for predicting charged excitations, as measured by direct or inverse photoemission spectroscopy. While GW was initially applied to compute the band structures of materials, its use has since expanded to finite systems [334]. For atoms, molecules or clusters, our all-electron GW implementation is suitable for the computation of deep core excitations with energies larger than 100 eV [335–337], semi-core and valence excitations as well as electron affinities and fundamental gaps, see Figure 4.3. Additionally, the total spectral function can be computed, including both quasiparticle excitations and satellite features.

FHI-aims offers different GW flavors for finite systems, including single-shot G_0W_0 with and without a Hedin shift in the Green's function, fully self-consistent, partially self-consistent, and eigenvalue self-consistent schemes. Scalar relativistic effects and spin-orbit coupling can both be taken into account. The frequency integration of the self-energy is performed with one of two full-frequency methods: imaginary frequencies and analytic continuation or contour deformation. Our implementation scales $O(N^4)$ with respect

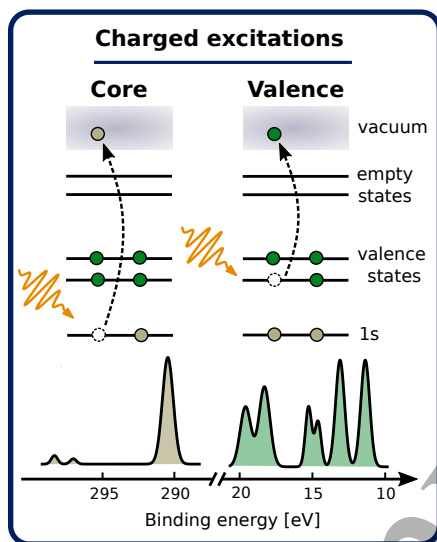


Figure 4.3: The GW implementation in FHI-aims for finite systems spans the entire energy spectrum, from deep core to valence excitations.

to system size N , enabling calculations up to 200-250 atoms on current high-performance computing (HPC) platforms. Furthermore, FHI-aims offers the following vertex corrections to go beyond GW : second-order screened exchange and full second-order self-energy corrections and cumulant expansion of the Green's function.

Current Status of the Implementation

GW Quantities: Self-Energy, Spectral Function and Quasiparticle Energies

The self-energy Σ is the key quantity in GW , encapsulating quantum mechanical effects of correlation and exchange between the excited electron or hole and the surrounding electrons. In practice, GW is often performed as a first-order perturbation (denoted G_0W_0), where the self-energy is given by

$$\Sigma(\mathbf{r}, \mathbf{r}', \omega) = \frac{i}{2\pi} \int d\omega' e^{i\omega'\eta} G_0(\mathbf{r}, \mathbf{r}', \omega + \omega') W_0(\mathbf{r}, \mathbf{r}', \omega'). \quad (4.3)$$

G_0 denotes the mean-field Green's function and W_0 is the screened Coulomb interaction in the random phase approximation (RPA), and η a small positive infinitesimal. G_0 and W_0 are computed from the mean-field single-particle orbitals $\{\psi_n\}$ and corresponding eigenvalues $\{\varepsilon_n\}$, typically obtained from Kohn-Sham density functional (KS-DFT)^a. The poles of the full propagator $G = G_0 + G_0\Sigma G$ correspond to the electron removal and addition energies measured in direct and inverse photoemission spectroscopy, respectively. For more details on the GW theory and basic equations, we refer the reader to a recent review article [334].

^aA less common starting point alternative is Hartree-Fock theory.

Roadmap on Advancements of the FHI-aims Software Package

4.2. ALL-ELECTRON GW FOR FINITE SYSTEMS

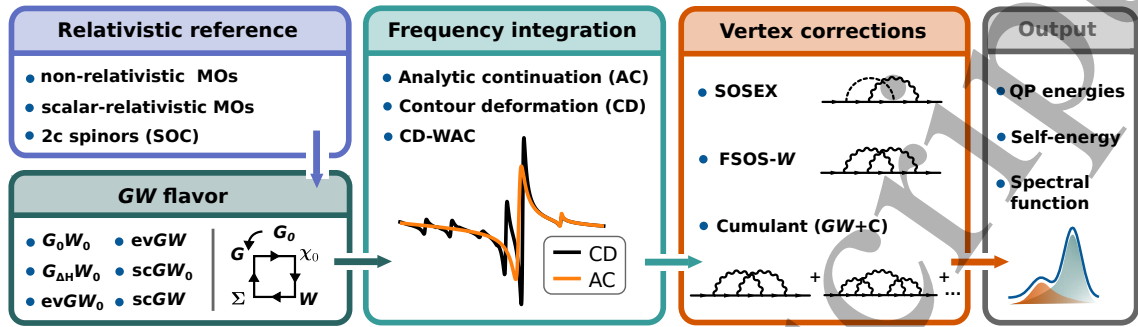


Figure 4.4: Workflow and choices available for GW calculations of finite systems with FHI-aims.

A link to the measured photocurrent can be approximately established via the spectral function $A(\omega)$

$$A(\omega) = \sum_m \frac{1}{\pi} |\text{Im} \langle \psi_m | G(\omega) | \psi_m \rangle| = \frac{1}{\pi} \sum_m \frac{|\text{Im} \Sigma_m(\omega)|}{[\omega - \varepsilon_m - (\text{Re} \Sigma_m(\omega) - v_m^{xc})]^2 + [\text{Im} \Sigma_m(\omega)]^2} \quad (4.4)$$

where m runs over all states and where $v_m^{xc} = \langle \psi_m | v^{xc} | \psi_m \rangle$ is the exchange-correlation potential from DFT and $\Sigma_m = \langle \psi_m | \Sigma(\omega) | \psi_m \rangle$ denotes the projection into the KS orbital basis. The main peaks in the spectral function are the quasiparticle (QP) energies $\varepsilon_n^{\text{QP}}$. Additionally, the spectral function includes satellite features, which arise from many-body effects such as plasmon or shake-up processes.

For QP energies, solutions to Equation (4.4) simplify to the quasiparticle equation:

$$\varepsilon_n^{\text{QP}} = \varepsilon_n + \text{Re} \Sigma_n(\varepsilon_n^{\text{QP}}) - v_n^{xc}. \quad (4.5)$$

Equation (4.5) is a non-linear equation, which is solved iteratively in FHI-aims. To avoid recalculating the self-energy in each iteration of Equation (4.5), a linearized approach, referred to as the Z -shot method, is also available [334]. Additionally, the output of the self-energy and the spectral function in a specified frequency range can be requested. The workflow and the available choices are summarized in Figure 4.4.

Relativistic Reference

Relativistic effects are introduced into the GW equations through the reference state. In FHI-aims, users can choose between non-relativistic or scalar-relativistic molecular orbitals (MOs) and 2-component (2c) spinors. The atomic zeroth-order regular approximation (ZORA) is the default scalar-relativistic method. For non-relativistic calculations and those using atomic ZORA in preceding DFT steps, relativistic corrections can be applied as a post-processing step for 1s core-level excitations, referencing fully relativistic calculations [90]. To include spin-orbit coupling (SOC), we recently implemented a 2c- GW scheme [338]. Our 2c- GW method [339] supports 2c spinors from different levels of theory, with the perturbative second-variational SOC implementation [48] currently used by default.

GW Flavors

FHI-aims offers various GW flavors beyond the G_0W_0 default. In eigenvalue self-consistency, the eigenvalues are iteratively updated in G (ev GW_0) or in both G and W (ev GW). Eigenvalue self-consistency increases the computational cost relative to G_0W_0 . To address this, we recently implemented a Hedin shift in the Green's function ($G_{\Delta H}W_0$) [337], which approximates the ev GW_0 scheme with minimal additional computational cost compared to G_0W_0 . The most conceptually rigorous and computationally demanding approach is fully self-consistent GW , denoted as sc GW [340, 341]. In sc GW , all four quantities – G , the polarizability χ_0 , W , and Σ – are iterated until self-consistency in the Green's function is achieved, as shown in the dark-green box in Figure 4.4. Additionally, a partial self-consistent scheme (sc GW_0) is available, where W is fixed at the W_0 level, and only G is determined self-consistently. The approximate self-consistent schemes reduce the dependence on the mean-field starting point, but only sc GW removes it fully [340]. In addition to the QP energies, sc GW_0 and sc GW provide also access to the total energies [340]. Further details on differences, advantages and disadvantages of the various GW flavors can be found in Ref. [334].

Frequency Integration

FHI-aims offers two techniques, both with canonical scaling with respect to system size, to solve the frequency integral in Equation (4.3): the analytic continuation (AC) of the self-energy [62] and the contour-deformation (CD) technique [335]. For a comprehensive comparison of both methods, we refer the reader to Ref. [334], restricting the description here to the key ideas. The AC scheme is the default method [62], available for all GW flavors. In this approach, the self-energy Σ is computed on the imaginary frequency axis and then analytically continued to the real-frequency axis by fitting the matrix elements $\Sigma_n(i\omega)$ to a multipole model function. Two common models are available for this purpose: a 2-pole model and a Padé approximant.

Conversely, in the CD approach the integrand in Equation (4.3) is expanded in the complex plane. The self-energy matrix elements $\Sigma_n(\omega)$ are obtained as the sum of two terms: an integral term along the imaginary frequency axis and a residue term that depends on real-frequency arguments. The calculation of the latter typically increases the computational cost compared to the AC approach. To mitigate the computational demands, we recently implemented the CD-WAC method [342] (CD with W analytic continuation), which applies an AC for the screened Coulomb interaction W in the residue term. CD and CD-WAC are only implemented for G_0W_0 , $G_{\Delta H}W_0$, ev GW_0 and ev GW and are currently only available for non- and scalar-relativistic references.

Most recently [343], a third method has been introduced: an exact frequency treatment in which the frequency convolution is performed analytically, yielding a sum-over-states expression for the correlation part of the self-energy. The RPA excitation energies and transition moments entering this expression are obtained by solving Casida-like equations. This approach is substantially more computationally demanding than AC, CD, or CD-WAC and is therefore not intended for routine calculations. Instead, it serves as a high-accuracy reference for validating more scalable frequency-integration techniques.

Algorithm, Performance and Numerical Accuracy

The AC [62], CD [335] and CD-WAC [342] implementations are based on numeric atom centered orbitals (NAOs) and on the resolution-of-the-identity (RI) approach, which reformulates the four-center two-electron

Roadmap on Advancements of the FHI-aims Software Package

4.2. ALL-ELECTRON GW FOR FINITE SYSTEMS

Coulomb integrals as product of two and three-center integrals. Our implementation employs the Coulomb fitting scheme (RI-V) [344], which is the most widely used and most accurate RI technique. The AC and CD-WAC schemes scale $O(N^4)$ with respect to system size N , while the scaling of CD depends on the energy range. Specifically, CD scales as $O(N^4)$ for valence states, but $O(N^5)$ for deep core-levels [335]. The AC, CD and CD-WAC implementations are parallelized using a standard message passing interface (MPI) and scale well up to 2000 - 3000 CPU cores. More details on the parallel performance can be found in Ref. [335]. With the CD method, core-level calculations are feasible for systems with up to 100–120 atoms on current HPC platforms. With CD-WAC and AC, both core and valence level calculations can be performed for systems with up to 200–250 atoms. We refer the reader to Refs. [335, 342] for performance benchmarks.

Among the three implementations, the CD method is the most accurate, reproducing all features of the self-energy with deviations in the sub-millielectronvolt range when compared to an exact frequency treatment, i.e., the full diagonalization of Casida-like equations at the RPA level [335]. At the G_0W_0 level, CD-WAC approximates the CD results with errors < 1 meV for valence states and < 5 meV for 1s core levels [342]. While AC is computationally most efficient and has the smallest computational prefactor, it fails to accurately describe the self-energy for deep core and semi-core levels [335]. For frontier orbital excitations, however, AC with a suitable Padé model reproduces exact frequency calculations to within 3 meV on average [41].

Beyond-GW: Vertex Corrections

FHI-aims also offers vertex corrections to the self-energy, specifically: second-order screened exchange (SOSEX) [345] and an approximate full second-order self-energy in terms of W (FSOS- W) [346]. Additionally, it includes propagator corrections, such as the cumulant expansion of the Green's function G [347]. The corresponding Feynman diagrams are displayed in Figure 4.4. Diagrammatically, SOSEX adds a second-order exchange diagram to the GW self-energy, while W is still evaluated at the RPA level. G_0W_0 + SOSEX mitigates the dependence on the mean-field reference compared to G_0W_0 and can improve the relative energy positions in molecular valence photoemission spectra [345]. FSOS- W can be considered as a dynamic extension of SOSEX, replacing the bare Coulomb interaction (dashed line in Figure 4.4) in the diagram by the frequency-dependent screened Coulomb interaction W (wiggly line). The GW +FSOS- W self-energy was also referred to as $GW\Gamma^{(1)}$ in Ref. [346] or $GW+G_3W_2$ in Ref. [348]. Currently, FHI-aims implements an approximate single-shot version of $GW\Gamma^{(1)}$ ($G_0W_0\Gamma_0^{(1)}$) and neglects all terms in the self-energy that involve three occupied or three virtual MOs [348]. $G_0W_0\Gamma_0^{(1)}$ is very similar in performance to G_0W_0 +SOSEX. Both schemes are only implemented with AC and exhibit an $O(N^5)$ scaling.

The cumulant expansion improves the prediction of satellite positions and intensities in the spectral function [349–354], which are poorly described in GW . The $GW + C$ approach is based on an exponential ansatz for the Green's function, which adds an infinite number of bosonic diagrams to the propagator G , while W remains again fixed at the RPA level. The FHI-aims $G_0W_0 + C$ implementation [343, 347] employs the spectral representation of the self-energy, as previously suggested [349, 353]. In this approach, the cumulant expansion is performed as a post-processing step after the computation of the GW self-energy, without significantly increasing the computational cost. The scaling of $G_0W_0 + C$ is determined by the GW implementation. $G_0W_0 + C$ is implemented exclusively with CD and CD-WAC because satellites are related to pole-features in the self-energy, which are inadequately captured with AC.

Usability and Tutorials

Comprehensive tutorials on how to perform GW calculations for finite systems with FHI-aims are available at <https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids>, covering general aspects [355] and core-level spectroscopy specifically [356]. The finite-system GW implementation in FHI-aims is widely used and has contributed to the development of well-established molecular benchmark sets, such as GW100 [41] and CORE65 [335]. Additionally, it has been employed to create comprehensive computational databases for machine learning applications. One example is the GW5000 dataset, which includes frontier orbital excitations of molecules with up to 100 atoms [357]. Another example is the 1s core-level database, containing more than 15,000 entries for CHO-containing molecules and clusters with up to ~ 110 atoms [358].

For QP energies, the vertex-corrected G_0W_0 +SOSEX and $G_0W_0\Gamma_0^{(1)}$ schemes outperform G_0W_0 when using conventional Perdew-Burke-Ernzerhof [27] (PBE) and PBE0 [254, 255] starting points. However, G_0W_0 with judiciously chosen starting points provides a similarly accurate, yet computationally more efficient alternative [345, 346, 348]. For satellites, we recommend $G_0W_0 + C$, though we note it yields QP energies at the $G_{\Delta H}W_0$ level. To reduce the starting point dependence, we recommend the partially self-consistent schemes $evGW_0$ or $scGW_0$ [334], as $evGW$ and $scGW$ underscreen and thus overestimate excitation energies [337, 359, 360].

Safe recommendations for finite-system GW calculations in FHI-aims, considering both core and valence levels, are summarized in Figure 4.5. Starting with the valence excitations, we suggest to perform G_0W_0 on-top of KS-DFT calculations with the PBE0 functional, which we denote as $G_0W_0@PBE0$. The expected average deviation from experiment is around 0.2 eV [334, 345, 360]. Similar or even better performance is expected when using $evGW_0@PBE$ [361], albeit at the cost of increased computational cost. Using AC with a Padé model with 16 parameters is usually sufficient for frontier orbitals [41] and deeper valence states with binding energies < 20 eV [362]. Alternatively, CD can be used, which has a slightly higher computational prefactor for valence states.

Turning to 1s core-levels, $G_{\Delta H}W_0@PBE$ is the optimal choice, yielding deviations of 0.3 eV and 0.2 eV for absolute and relative 1s core-level excitations, respectively [337]. $evGW_0@PBE$ performs equally well, but is at least an order of magnitude more expensive for core-level calculations. Alternatively, we proposed a $G_0W_0@PBEh(\alpha = 0.45)$ approach [336], using a hybrid functional with 45% of exact exchange. However, the optimal amount of exact exchange varies with atomic number [336] and was specifically optimized for second-row elements, making it unsuitable for heavier elements. We thus recommend $G_{\Delta H}W_0@PBE$ over $G_0W_0@PBEh(\alpha = 0.45)$. The CD method is mandatory for deep core states. For systems larger than 20 atoms, the CD-WAC approach will be computationally more efficient than CD.

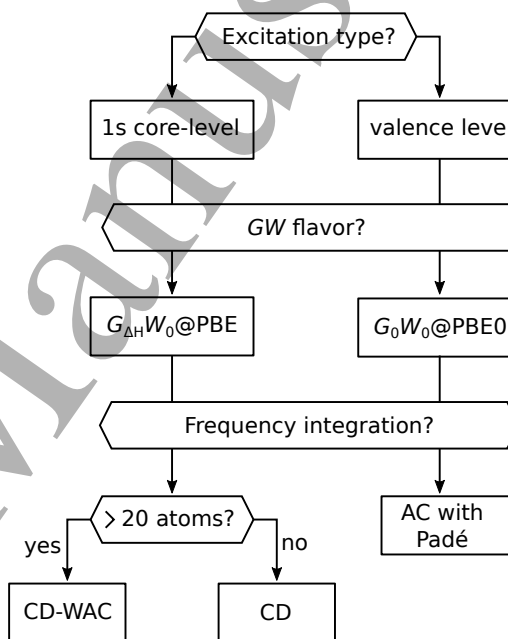


Figure 4.5: Choices regarding GW flavor and frequency integration depending on the excitation type for FHI-aims calculations.

Roadmap on Advancements of the FHI-aims Software Package

4.2. ALL-ELECTRON GW FOR FINITE SYSTEMS

Another important consideration is the choice of the basis set. As with other correlated methods, GW converges slowly with respect to basis set size. The results, even with the largest localized basis sets, are not fully converged, as demonstrated in Ref. [334]. A basis set extrapolation procedure must be employed, which is detailed in the tutorials [355, 356]. For valence states, the NAO-VCC- nZ basis set family [20] can be used. Alternatively, Gaussian basis sets, which are treated numerically in FHI-aims, such as the Dunning basis sets cc-p VnZ [32, 363] or the def2-TZPV/def2-QZVP basis sets [364] extrapolate well in our experience [41, 334]. For $1s$ states, the cc-p VnZ series works sufficiently [336], but core-rich Gaussian basis sets such as ccX- nZ [365, 366] have shown a better convergence behavior [366]. If basis set extrapolation cannot be performed due to computational cost, or if the focus is on the spectral function rather than QP energies, we recommend using NAO-VCC-5Z for valence states and tier2+STO3 for deep core states. The latter combines NAOs with Slater-type orbitals (STOs), as described in the Supporting Information of Ref. [35]. These basis sets typically reproduce extrapolated results within 0.1 eV.

Future Plans and Challenges

Future developments will improve the computational efficiency and add new features. On the feature side, core-level calculations are currently restricted to $1s$ excitations, and we plan to extend the implementation to SO-coupled p , d and f excitations in the near future. This will require adding CD and CD-WAC to the 2c- GW implementation, with an experimental 2c-CD version already available. Additionally, we plan to enable the usage of 2c- GW in combination with the quasi-four-component (Q4C) [26] scheme, which is also available in FHI-aims. The Q4C spinors should provide a more accurate relativistic reference compared to the second-variational SOC approach. Furthermore, we plan to extend the Sternheimer RPA implementation [367] to GW for benchmarking purposes.

On the performance side, work on low-scaling GW algorithms is on-going. Low-scaling RPA is already functional and the extension to GW is in progress. Our low-scaling RPA and GW algorithms rely on the separable RI scheme, which was recently implemented [368]. The separable RI coefficients are currently obtained as fit to the RI-V coefficients. To reduce the computational prefactor, we plan to replace the Coulomb metric in RI-V with the similarly accurate truncated Coulomb metric. Furthermore, the GPU acceleration for all steps in the low-scaling algorithms is currently ongoing.

Acknowledgements

D. Golze acknowledges financial support by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) for funding via the Emmy Noether Programme (project number 453275048) and for funding within the CRC 1415 (Chemistry of Synthetic Two-Dimensional Materials, project number 417590517) as well as within the RTG 2861 (Research Training Group, project number 491865171). D.G. also acknowledges support by the Innovation Study Exa4GW. The Innovation Study Exa4GW has received funding through the Inno4scale project, which is funded by the European High-Performance Computing Joint Undertaking (JU) under Grant Agreement No 101118139. The JU receives support from the European Union's Horizon Europe Programme. P. Rinke acknowledges the European Union's Horizon 2020 Research and Innovation Program for financial support under Grant No. 951786 (Nomad Center of Excellence). We further acknowledge CSC-IT Center for Science (Finland), the Jülich Supercomputer Center (Germany), the Paderborn Center For Parallel Computing (PC2, Germany) and the Aalto Science-IT project for computational resources. X. Ren acknowledges the funding support by the National Natural Science Foundation of China (Grant Nos 12374067 and

Roadmap on Advancements of the FHI-aims Software Package

4.2. ALL-ELECTRON GW FOR FINITE SYSTEMS

12188101) and the National Key Research and Development Program of China (Grant Nos 2022YFA1403800 and 2023YFA1507004).

Accepted Manuscript

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

4.3 GW Calculations for Periodic Systems

Volker Blum^{1,2}, Francisco A. Delesma³, Uthpala Herath¹, Hermann Lederer⁴, Florian Merz⁵, *Xinguo Ren^{6,11,a}, Patrick Rinke^{7,8,9,10}, Matthias Scheffler¹¹, Yi Yao^{11,12}, and Min-Ye Zhang^{6,11}

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁴Max Planck Computing and Data Facility, Giessenbachstrasse 2, D-85748 Garching, Germany

⁵Lenovo HPC Innovation Center, Meitnerstr. 9, D-70563 Stuttgart, Germany

⁶Institute of Physics, Chinese Academy of Sciences, 3rd South Str. 8, Beijing 100190, China

⁷Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

⁸Physics Department, TUM School of Natural Sciences, Technical University of Munich, Garching, Germany

⁹Atomistic Modelling Center, Munich Data Science Institute, Technical University of Munich, Garching, Germany

¹⁰Munich Center for Machine Learning (MCML), Munich, Germany

¹¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹²Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

*Coordinator of this contribution.

^aCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

Summary

The *GW* approximation [333] is a state-of-the-art approach for computing single-electron excitation energies of real materials from first principles [369]. It can be applied to a large variety of chemical systems across different characteristics and dimensionalities (for a review of the underlying theory, see Refs.[334, 370]). Compared to the (generalized) Kohn-Sham density functional theory (KS-DFT), *GW* has a much more rigorous foundation as an approach for determining single-electron excitation energies and usually yields much better electronic band structures, particularly the band gaps. However, the improvement of *GW* over KS-DFT comes with a significantly higher computational cost, which stimulates continuing efforts to improve the algorithms for *GW* calculations, aiming at more efficient (and sufficiently accurate) implementations. Since its first application to real materials nearly four decades ago [369], various *GW* codes have been developed based on different numerical frameworks [123, 371–375]. These implementations largely fall into those based on (augmented) plane waves and those based on local atomic orbitals. Typically, the plane-wave based implementations are more suitable for simulating periodic systems whereas atomic-orbital based implementations are preferred for molecular systems. The *GW* method was implemented in FHI-aims [8] at the early stage of its development [62]. The initial implementation was for finite systems [62], as described in the previous Contribution 4.2. Here, we focus on the extension of the *GW* implementation to periodic systems [36]. The periodic implementation of *GW* in the FHI-aims is based on the localized res-

Roadmap on Advancements of the FHI-aims Software Package

4.3. GW CALCULATIONS FOR PERIODIC SYSTEMS

olution of identity (termed RI-LVL) [63]. This significantly reduces the amount of integrals to evaluate and store, and makes the periodic GW implementation with NAO basis sets feasible. However, there is a higher demand for the size and quality of the auxiliary basis sets to ensure the adequate accuracy of RI-LVL for periodic calculations. Furthermore, special care must be taken to deal with the singularity of the screened Coulomb interaction at the Γ point. Currently, the periodic GW implementation in FHI-aims allows us to compute the quasiparticle energies for extended systems with small or medium-sized unit cells with finite \mathbf{k} points, and plot the band structures across the Brillouin zone (BZ).

Current Status of the Implementation

The periodic GW method in FHI-aims employs the canonical algorithm operating solely in the reciprocal space and (imaginary) frequency domain. For KS spin state $\psi_{n\sigma}^{\mathbf{k}}$ with orbital energy $\epsilon_{n\sigma}^{\mathbf{k}}$, the correlation self-energy from one-shot G_0W_0 is expressed as [36]

$$\Sigma_n^c(\mathbf{k}\sigma, i\omega) = -\frac{1}{2\pi} \sum_{m,\mathbf{q}} \sum_{\mu,\nu} \int_{-\infty}^{\infty} d\omega' \frac{C_{n,m,\sigma}^{\mu}(\mathbf{k}, \mathbf{k}-\mathbf{q}) W_{0,\mu\nu}^c(\mathbf{q}, i\omega') C_{m,n,\sigma}^{\nu}(\mathbf{k}-\mathbf{q}, \mathbf{k})}{i\omega - i\omega' + E_F - \epsilon_{m,\sigma}^{\mathbf{k}-\mathbf{q}}} \quad (4.6)$$

where $C_{n,m,\sigma}^{\mu}$ are the RI decomposition coefficients in KS space and $W_{0,\mu\nu}^c = W_{0,\mu\nu} - V_{\mu\nu}$ is the correlation part of screened Coulomb interaction expanded by auxiliary basis functions (ABFs). The summation runs over auxiliary basis indices μ, ν , band index m and wave-vector \mathbf{q} sampled in the first BZ. The frequency integral is evaluated numerically using standard quadrature grids. The self-energy at real frequency is then obtained by analytically continuing the data on the imaginary axis and used to compute the quasi-particle (QP) energy ϵ^{QP} of interest by solving the QP equation.

In the current implementation, RI-LVL plays an important role in reducing the time and memory cost for the RI triple coefficients in KS space, which are obtained by transforming those in atomic orbital (AO) basis with wave function expansion coefficients $\{c_{n\sigma}^i(\mathbf{k})\}$

$$C_{n,m,\sigma}^{\mu}(\mathbf{k}, \mathbf{k}-\mathbf{q}) = \sum_{ij} c_{n\sigma}^{i*}(\mathbf{k}) c_{m\sigma}^j(\mathbf{k}-\mathbf{q}) \tilde{C}_{i,j}^{\mu}(\mathbf{k}, \mathbf{k}-\mathbf{q}). \quad (4.7)$$

The AO coefficients $\tilde{C}_{i,j}^{\mu}(\mathbf{k}, \mathbf{k}-\mathbf{q})$ are computed by Fourier transforming their real-space counterparts. In global RI, indices i, j have to run over all the atomic basis with non-negligible Coulomb interaction with specific μ , and the Fourier transformations with respect to \mathbf{k} and $\mathbf{k}-\mathbf{q}$ are coupled. The local approximation forces either i or j basis to be located at the same atom and unit cell as the auxiliary function μ in the real space expansion. As a result, the transformation involves only two nonzero sectors, each dependent on a single wave vector. This makes the storage of real-space triple coefficients affordable and the entire transformation efficient, hence allowing on-the-fly evaluation of $C_{n,m,\sigma}^{\mu}$ to avoid its expensive storage.

To handle the finite size effect and achieve fast convergence of \mathbf{q} -point sampling in periodic GW , the singular nature of the Coulomb interaction near the Γ point has to be carefully addressed. This singularity is involved in both calculations of the screened Coulomb interaction and the self-energy. In FHI-aims, the first case is approached by computing the dielectric matrix at the Γ point in Coulomb eigenvector representation using relevant terms from $\mathbf{k} \cdot \mathbf{p}$ theory. For the self-energy calculation, the divergence of screened Coulomb is circumvented by synthesizing it with a truncated Coulomb interaction as used in the Hartree-Fock and hybrid functional calculations.

The overall cost for the self-energy evaluation of Eq.(4.6) scales as $\mathcal{O}(N^4 N_\omega N_{\mathbf{k}} N_{\mathbf{q}})$, with N being the system size of the unit cell, and has a large prefactor due to the use of all unoccupied states. This high computational load necessitates an efficient and scalable parallelization strategy to fully leverage modern massively parallel computers. In FHI-aims, we have implemented a two-level parallelization scheme based on message passing interface (MPI) and its workflow is sketched in Figure 4.6. Upon a converged DFT calculation and RI initialization, \mathbf{k} - and \mathbf{q} -grids required for GW are determined. The \mathbf{q} -grids are further grouped and distributed to a batch of process grids, each consisting of MPI tasks mapped to a square array. Within each grid, row ABF indices are distributed across process rows, while column ABFs and second AO indices of RI coefficients are distributed across process columns. The calculation of W and contributions to Σ is then performed in an embarrassingly parallel manner among process grids. Finally, the self-energy operator is reduced from all tasks for analytic continuation and solution of QP equation. We note that the MPI tasks are scheduled to favor more batches of process grids while minimizing the load imbalance. This scheme allows FHI-aims to effectively handle both small systems using dense \mathbf{q} sampling and large ones with only a few \mathbf{q} points.

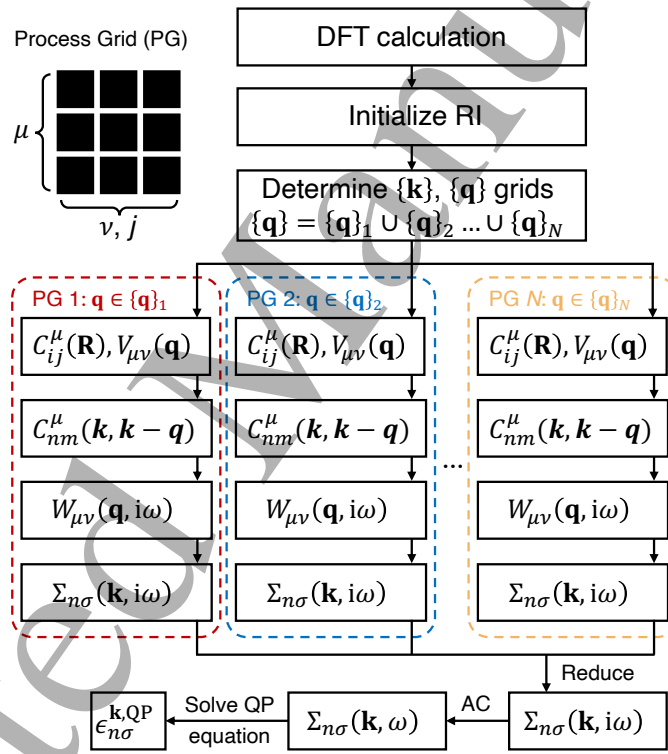


Figure 4.6: Workflow of canonical periodic G_0W_0 algorithm in FHI-aims

Usability and Tutorials

To ensure periodic GW is a reliable feature in FHI-aims, its accuracy and efficiency requires careful investigation. The accuracy of our periodic GW implementation has been benchmarked in a previous publication [36] against the all-electron LAPW-based GW code FHI-gap [374, 376]. It has also been compared recently

Roadmap on Advancements of the FHI-aims Software Package

4.3. GW CALCULATIONS FOR PERIODIC SYSTEMS

with implementations in ABINIT, GPAW, and exciting for several representative systems with diverse characteristics [377]. Among many aspects, the efficacy of RI-LVL, the BZ sampling convergence and the impact of one-electron basis are demonstrated. It is found that additional basis functions to the one-electron basis set for on-the-fly construction of ABFs are required to mitigate the error due to the local approximation to RI and obtain accurate band gaps. On top of the tier2 species defaults, FHI-aims is shown to deliver G_0W_0 band gaps for semiconductors, in similar accuracy (~ 0.2 eV) to those with local-orbital-enhanced LAPW basis from FHI-gap, except for challenging systems like ZnO and LiF. The agreement can be further improved by including Slater-type orbitals in the one-electron basis. We also note that the k -points on the regular k -grid of a self-consistent calculation and band paths are treated on equal footing through an on-the-fly Fourier transform for necessary RI coefficients, so that the band structure calculation is free of error arising from self-energy interpolation.

In order to obtain accurate G_0W_0 band gap results and band structures, it is essential to minimize the RI error due to the local approximation in RI-LVL. Although more basis functions in the ABFs construction can lead to better accuracy, it will considerably slow down the calculation. Therefore, it is of great interest to identify the optimal additional functions that lead to adequate accuracy gain, but with as few extra ABFs as possible. For this purpose, we have benchmarked the impact on G_0W_0 band structure due to different combinations of additional hydrogen-like functions in a few systems, from which promising candidates are selected and tested on a larger set of materials. It results in improved species defaults for periodic GW calculations with optimal additional functions on top of the DFT defaults. They are available in the FHI-aims species collection under “_gw”-suffixed folders.

In terms of performance, the aforementioned MPI parallelization scheme is designed to run FHI-aims periodic GW efficiently on massive cores for both small- and large-size systems. We have conducted a strong scaling test for the implementation using different sizes of ZrO_2 cell, as shown in Figure 4.7. For the 3-atom unit cell with tight-tier1 basis (77 AOs, 2220 ABFs) and 105 band k -points, we observe nearly perfect scaling up to 1440 MPI tasks (20 nodes). Calculation of an 81-atom super cell with tight basis (3429 AOs, 31617 ABFs) and only Γ point scales reasonably well up to 20736 MPI tasks (288 nodes).

To facilitate long calculation times for systems with many q -points, for example for when there is time-limit restrictions on HPC resources, a restart functionality has been implemented for periodic GW calculations, controlled by the `restart_periodic_gw` keyword. When activated, each MPI task generates a separate checkpoint file, which records the number of q points processed locally by the task (see Figure 4.6), along with the accumulated contribution to the correlation self-energy. If a job is interrupted, restarting the calculation with the same number of MPI tasks allows the checkpoint files to be loaded, enabling the calculation to resume from the next unprocessed q -point.

An informative tutorial for periodic GW is available on the official site as part of the tutorial series “GW and BSE for molecules and solids” [309]. It covers a comprehensive list of technical factors mentioned above for converging G_0W_0 calculations, and walks through the procedure in great detail. In particular, we include a worked example for monolayer MoS_2 , a prototypical two-dimensional material, to illustrate how the GW approach can be applied in the context of low-dimensional functional materials.

Future Plans and Challenges

Several directions can be explored for the further development of the periodic GW functionalities in FHI-aims, which will be briefly discussed below.

4.3. GW CALCULATIONS FOR PERIODIC SYSTEMS

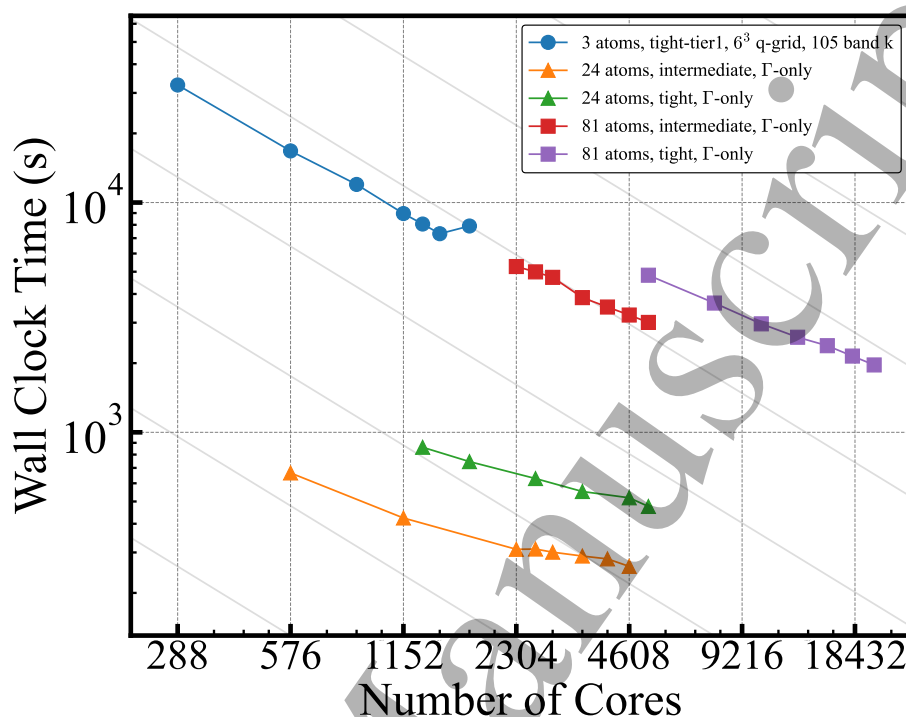


Figure 4.7: Strong scaling test of FHI-aims periodic G_0W_0 calculations for ZrO_2 unit cell (3 atoms) and different super cells. The test is conducted on the Raven system (MPCDF, Germany). Each computer node is equipped with 72 cores (Intel Platinum 8360Y processors) and 256 GB memory.

First, the current periodic GW implementation is based on the k -space formalism, and its $O(N^4)$ scaling limits its application to relatively small systems. To treat real-world materials problems such as defects and interfaces, where large supercells are typically required, efficient low-scaling implementations are highly desirable, analogous to the low-scaling strategies developed for hybrid functionals (see Contrib. 3.2). In principle, interfacing FHI-aims with LibRPA [301], where the low-scaling real-space GW algorithm has been implemented, enables one to treat large systems. However, in this case, the high memory cost for storing the RI coefficients and Coulomb matrix often becomes the computational bottleneck. Thus, for further improvement, it is necessary to develop a more efficient strategy for memory usage and distributions for large-scale periodic GW calculations.

Second, for periodic GW , currently only the one-shot G_0W_0 scheme with (semi-)local density functional approximation is implemented in FHI-aims. It would be highly desirable to implement a certain self-consistent GW scheme, particularly those that can deliver wavefunctions. Such quasiparticle wavefunctions are necessary to describe, e.g., topological materials at a theoretical level that goes beyond KS-DFT.

Third, the current periodic implementation of G_0W_0 in FHI-aims can only be used to reliably calculate the band structures of the insulating materials. To deal with metallic systems reliably, additional implementation work is in order, including the incorporation of intraband transitions in dielectric function calculations and accurate treatment of the Fermi surface in the BZ integration. Such an extension is a prerequisite for

*Roadmap on Advancements of the FHI-aims Software Package***4.3. GW CALCULATIONS FOR PERIODIC SYSTEMS**

applying FHI-aims *GW* to correlated metallic systems, such as transition metals and doped cuprates.

Fourth, there has been considerable interest in combining electronic self-energy and vibrational self-energy from electron-phonon coupling in a unified framework [378]. The quasiparticle energies determined from such a combined self-energy will naturally capture the temperature effect on the electronic band structures. Additionally, such a combined self-energy can also be used to calculate the transition temperature of superconductors via the Eliashberg theory [379]. It is also of interest to extend the combined self-energy beyond the phonon picture by integrating G_0W_0 with molecular dynamics and the band unfolding technique (see Contrib. 6.5).

Lastly, when it comes to studying properties of materials consisting of heavy elements, relativistic effects such as spin-orbit coupling (SOC) need to be taken into account. FHI-aims provides spin-orbit coupled energy corrections through a second-variational method [48] for (semi-)local and hybrid DFT calculations. It would be of great interest to incorporate this into periodic GW calculations by calculating the correlation self-energy shown in Eq. 4.6 using the SOC corrected wave-functions leading to a QP equation with relativistic corrections.

Acknowledgements

We acknowledge the funding support from the National Natural Science Foundation of China (Grant Nos 12374067 and 12188101) and the National Key Research and Development Program of China (Grant Nos 2022YFA1403800 and 2023YFA1507004). MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

 4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

4.4 Neutral Excited States: Casida Equation for Linear-Response Time-Dependent Density Functional Theory and Bethe-Salpeter Equation

*Yosuke Kanai^{1,2}, Matthias Kick³, Ruiyi Zhou¹, Chi Liu⁴, Jan Kloppenburg⁵, Yi Yao^{1,6,7}, Dorothea Golze⁸, Marc Dvorak⁹, Patrick Rinke^{9,10,11,12}, Xinguo Ren^{13,14,a}, and Volker Blum^{4,6}

¹Department of Chemistry, University of North Carolina at Chapel Hill

²Department of Physics and Astronomy, University of North Carolina at Chapel Hill

³Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Department of Chemistry, Duke University, Durham, NC 27708, USA

⁵Department of Chemistry and Materials Science, Aalto University, Finland

⁶Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

⁷Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁸Faculty of Chemistry and Food Chemistry, Technische Universität Dresden, 01062 Dresden, Germany

⁹Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

¹⁰Physics Department, TUM School of Natural Sciences, Technical University of Munich, Garching, Germany

¹¹Atomistic Modelling Center, Munich Data Science Institute, Technical University of Munich, Garching, Germany

¹²Munich Center for Machine Learning (MCML), Munich, Germany

¹³Institute of Physics, Chinese Academy of Sciences, 100190 Beijing, China

¹⁴The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

(*) Coordinator of this contribution

^aCurrent Address: Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

Summary

By transforming to the frequency domain, the time-dependent electronic structure theory of a system can be straightforwardly cast within the linear response theory. First-order responses to external stimuli such as electromagnetic fields are largely responsible for experimentally measurable quantities like optical or X-ray absorption spectra. These response properties can be written in the Lehmann representation, and computing resonance frequencies from corresponding eigenstates using this approach is of great practical interest. In the quasi-particle picture, neutral excitations, especially bound states ("excitons"), include the screened interaction between the excited electron and hole as schematically depicted in Figure 4.8. In FHI-aims, the Bethe-Salpeter equation (BSE) based on the GW method (Section 4.2), i.e., $BSE@GW$, and Casida's equation of linear-response time-dependent density functional theory (LR-TDDFT) are implemented for calculating electronic excited states using ground-state DFT as the starting point. While these two methods derive from very different theoretical formalisms (i.e. BSE from the Green's function theory and the Casida's equation for TDDFT), the practical computational algorithms share close similarities and are thus convenient for

4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

software development. In the FHI-aims code, the BSE@ GW and LR-TDDFT approaches are implemented for molecular systems, based on scalar-relativistic density functional approximations, spanning both valence excitations as well as core-level excitations. The BSE@ GW approach to compute absorption spectra is also available for extended periodic solids.

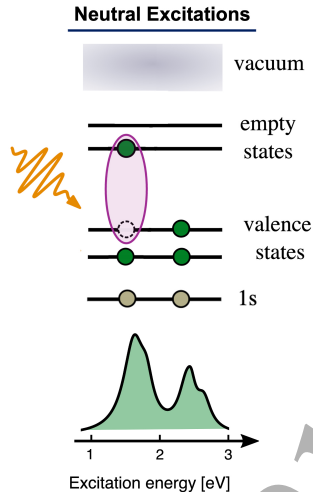


Figure 4.8: A schematic representation of a neutral excitation (i.e. exciton) modeled using BSE@ GW or LR-TDDFT via Casida equation. In this simplified picture, a valence electron is promoted to an empty state by optical absorption but retains an interaction (pink ellipse) with the resulting hole via a screened Coulomb interaction, which incorporates an effective response from all other particles in the system. Experimentally measurable properties such as the optical absorption spectrum can be computed as a function of the excitation energy using BSE@ GW or LR-TDDFT.

Current Status of the Implementation

Bethe-Salpeter Equation with GW method

Based on the quasi-particle description obtained using the GW method as discussed in Section 4.2, the BSE takes advantage of the two-particle Green's function to model the screened two-particle interaction between the hole and excited electron. Therefore, the approximations made at the GW calculation carries over to the BSE calculation as well. In particular, the GW approximation to the self-energy is the basis of the BSE implementation in the FHI-aims code [33, 380]. Additionally, as done in most cases, the frequency dependence of the screened interaction, W , is neglected at the BSE stage so that the BSE can be solved efficiently as a linear problem. Within linear response theory, the BSE is formulated as solving the generalized eigenvalue equation [381, 382]

$$\begin{bmatrix} \mathbf{A} & \mathbf{B} \\ \mathbf{B}^* & \mathbf{A}^* \end{bmatrix} \begin{bmatrix} \mathbf{X} \\ \mathbf{Y} \end{bmatrix} = \Omega \begin{bmatrix} \mathbf{I} & \mathbf{0} \\ \mathbf{0} & -\mathbf{I} \end{bmatrix} \begin{bmatrix} \mathbf{X} \\ \mathbf{Y} \end{bmatrix} \quad (4.8)$$

where Ω is the excitation energy, \mathbf{X} and \mathbf{Y} are the transition amplitudes in a product basis of occupied single-quasiparticle orbitals i and unoccupied single-quasiparticle orbitals a . More specifically, in Eq.4.8,

4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

the \mathbf{A} , \mathbf{B} matrices are defined as

$$A_{ia,jb} = \delta_{ij}\delta_{ab}(\epsilon_a^{\text{QP}} - \epsilon_i^{\text{QP}}) + v_{ia,jb} - W_{ij,ab}(\omega = 0) \quad (4.9)$$

$$B_{ia,jb} = v_{ia,bj} - W_{ib,aj}(\omega = 0) \quad (4.10)$$

where $\epsilon_{i/a}^{\text{QP}}$ is the quasi-particle energy which is typically obtained from a separate GW calculation as discussed in Section 4.2. v is the bare Coulomb interaction, and $W(\omega = 0)$ denotes that the frequency-dependence of the screened interaction is typically neglected as also done in the FHI-aims implementation. Refs. [33, 380] focused on optical excitations, i.e., from valence bands to low-lying conduction bands. However, as an all-electron implementation, FHI-aims also allows one to compute core-electron excitations as relevant, e.g., for X-ray absorption spectroscopy [35]. High numerical precision for the $1s$ excitation of a simple molecule (H_2O) with available experimental reference data was demonstrated in Ref. [35], including an extension of the applicable basis sets by tight Slater-type orbitals to capture the core orbitals' response to the excitation. In particular, the efficacy of the BSE@ GW method was compared to the equation-of-motion coupled-cluster theory methods and experiments.

Building on the BSE implementation for isolated systems [33] and GW method for extended systems [36], our all-electron NAO-based implementation of the BSE method was recently generalized also for extended periodic systems with the Brillouin zone sampling by Zhou *et al.*[380]. Currently, only the Tamm-Dancoff approximation (TDA), which amounts to neglecting \mathbf{B} in Eq. 4.8, is implemented for the periodic version.

Building on the non-periodic BSE@ GW implementations, FHI-aims includes a quantum embedding theory called dynamical configuration interaction (DCI) for finite systems. The foundations of the theory are covered in Refs. [383, 384]. DCI is an active-space method, in which a fixed active space of strongly correlated orbitals is treated with configuration interaction (CI). This active space is then embedded in a high-energy, dynamically correlated BSE@ GW -like bath. By combining CI and BSE@ GW , DCI aims to leverage the strengths of both theories, providing a balanced treatment of static and dynamic correlation. DCI offers access to total energies of both ground- and excited-state potential energy surfaces. Neutral excitation energies are obtained as the difference between these total energies.

Casida Equation: Linear-Response Formulation of Time-Dependent DFT

The Casida equation (i.e., LR-TDDFT) is derived by formulating the polarizability as a sum over states of the many-body system. In 1995, Casida showed that the square of excitation energies can be obtained as an eigenvalue problem in the matrix form:

$$\mathbf{C}\mathbf{F}_s = \Omega^2\mathbf{F}_s. \quad (4.11)$$

Here, the \mathbf{C} is the so-called Casida matrix, which has the same dimension of \mathbf{A} and \mathbf{B} in the BSE Eq.4.8. Ω are the neutral many-body excitation energies. \mathbf{F}_s are the associated eigenvectors and can be related to the oscillator strengths via the dipole operator[385]. In Eq. (4.11), the TDA was adapted, which is widely used for LR-TDDFT. The \mathbf{C} matrix can be written in the basis of product of the (g)KS orbitals

$$C_{ia,jb}(\omega) = \delta_{i,j}\delta_{a,b}(\epsilon_a - \epsilon_i)^2 + 2\sqrt{(\epsilon_a - \epsilon_i)K_{ia,jb}(\omega)}\sqrt{(\epsilon_b - \epsilon_j)}, \quad (4.12)$$

4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

where ϵ_a, ϵ_i are KS eigenvalues. The kernel $K_{ia,jb}$ is defined as

$$K_{ia,jb}(\omega) = \iint \psi_i^\dagger(\mathbf{r}) \psi_a(\mathbf{r}) \left[\frac{1}{|\mathbf{r} - \mathbf{r}'|} + f_{xc}[n_0](\mathbf{r}, \mathbf{r}', \omega) \right] \psi_j(\mathbf{r}') \psi_b^\dagger(\mathbf{r}') d\mathbf{r} d\mathbf{r}' \quad (4.13)$$

where n_0 is the (g)KS ground state electron density and $f_{xc}[n_0]$ is the exchange-correlation (XC) kernel, which is a functional derivative that describes how the exchange-correlation potential changes with respect to the electron density [386]. FHI-aims currently supports the LDA XC kernel, implemented within the adiabatic approximation (i.e. $f_{xc}(\mathbf{r}, \mathbf{r}', \omega) = f_{xc}(\mathbf{r}, \mathbf{r}')$) for isolated (non-periodic) systems.

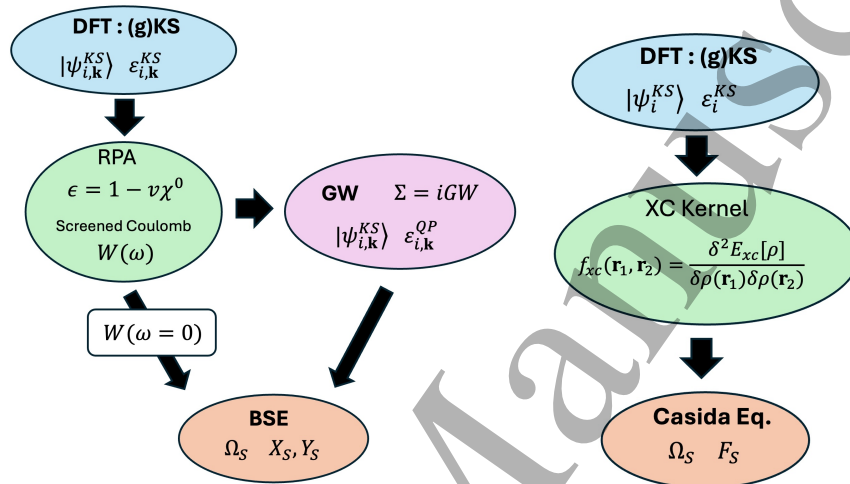


Figure 4.9: A schematic representation of the flow of the BSE and Casida Equations in the FHI-aims code. Both the BSE and the Casida equation start from performing a ground-state DFT calculation, with periodic boundary conditions supported for the BSE. The BSE additionally relies on the GW method for calculation of the quasi-particles energies. The screened Coulomb interaction is computed within the random phase approximation (RPA). Casida's equation, on the other hand, does not involve an extra step for the single-particle energies and orbitals. However, the exchange-correlation (XC) kernel needs to be approximated. Currently the adiabatic LDA kernel is implemented and the implementation works only for isolated (non-periodic) systems.

Usability and Tutorials

The BSE tutorial for FHI-aims,^b offers a concise introduction to the BSE@ GW method. It includes several straightforward examples for running BSE calculations in molecular systems and extended periodic systems using the FHI-aims code. This tutorial covers the fundamentals of performing neutral excitation calculations with the BSE@ GW method. Importantly, it also delves into numerical topics such as basis set convergence and various levels of approximation and algorithms used in the GW method, which computes the quasi-particle energies needed as inputs for BSE calculations.

The BSE@ GW implementation in FHI-aims is suitable for both molecular valence excitations, as studied in UV-vis spectroscopy [33], and molecular K -edge core-electron excitations, as measured in X-ray absorption spectroscopy [35]. For molecular valence excitations, reasonable accuracy is often achieved by

^b<https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids/>

4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

using $BSE@G_0W_0$ on top of DFT calculations using a hybrid density functional with 20-30% of exact exchange [387], such as PBE0 [254, 255]. In the preceding G_0W_0 step, employing an analytic continuation of the self-energy with a Padé model containing 16 parameters is sufficient [33]. For fast basis set convergence of low-lying excitation energies in small molecules, augmenting standard NAO basis sets with additional extended, diffuse functions is desirable, in line with the broader experience of quantum chemistry. Benchmarks by Liu *et al.* [33], show that a basis set “tier2+aug2”, where aug2 represents two augmentation functions from the aug-cc-pV n Z Gaussian basis sets [34, 388], facilitates excellent numerical convergence of low-lying excitation energies. The tier2+aug2 basis sets agree with results from the much larger, GTO-based aug-cc-pV5Z basis sets within 0.1 eV [33].

For molecular K -edge transitions in light elements, Ref. [35] demonstrated good performance when executing $BSE@G_0W_0$ on top of the $PBEh(\alpha)$ hybrid DFT functional with at least 45% exact exchange (α). The value $\alpha = 0.45$ is optimal for second-row elements, but should be gradually increased for heavier elements. We consider the $G_0W_0@PBEh$ scheme superior to partially self-consistent schemes, which we recommend for GW -based charged excitations, as the calculation of W in BSE requires high-quality underlying KS orbitals. At the DFT level, K -edge calculations should be performed using the atomic zeroth-order regular approximation (ZORA), with the scalar-relativistic correction from Ref. [90] applied to the GW quasiparticle energies. The contour deformation technique should be used to compute the GW self-energy. For K -edge calculations, we recommend using the tier2+aug2+STO3 basis set, where STO3 refers to the set of Slater-type orbitals provided in the Supporting Information of Ref. [35]. With this setup, mean absolute errors of 0.63 eV relative to experiment were achieved [35].

The option for periodic systems is a new addition for $BSE@G_0W_0$ calculations in the FHI-aims code at the time of writing. Benchmark calculations for Si and MgO are reported in Ref. [380]. Here, the localized RI framework of Ref. [63] makes the expansion of the Coulomb operator tractable for periodic systems with demonstrated high precision. In the preceding G_0W_0 calculation, one may employ analytic continuation of the self-energy either with a two-pole model or Padé model [36]. Based on the work in Ref. [380], we recommend the standard FHI-aims tier2 basis sets, which converge to 0.2 eV or better. For the Brillouin Zone (BZ) sampling, only the Γ -centered sampling is currently implemented in the FHI-aims code. For the absorption spectrum of some semiconductor materials, a large number of k points is required and a systematic convergence test for BZ sampling is necessary [380].

The tutorial on LR-TDDFT for molecules, using the Casida equation^c provides a concise introduction of how to use FHI-aims code to compute excited-state energies for simple molecules, using the LDA XC kernel. For molecular valence excitations, we recommended the same tier2+aug2 basis set as for the $BSE@G_0W_0$ calculations, which reproduces aug-cc-pV5Z results within 0.1 eV [33].

DCI generally targets multireference problems that $BSE@GW$ alone struggles to address. The study of double excitations and complicated spin multiplets lies squarely in DCI's domain. The method is well-suited for systems in which a small number of strongly interacting electrons (on the order of 2-10) is surrounded by a bath of weakly interacting electrons. Frontier orbitals are a natural choice for the active space, making defect and impurity systems prime candidates for DCI. Active spaces of up to eight levels and a bath of approximately 300 orbitals are recommended. A comprehensive documentation can be found in the FHI-aims manual. The largest system calculated with DCI so far was gas-phase free-base and Mg-porphyrin [389].

^c<https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids/Part-1/LRTDDFT/>

4.4. NEUTRAL EXCITED STATES: CASIDA EQUATION FOR LINEAR-RESPONSE TIME-DEPENDENT DENSITY FUNCTIONAL THEORY AND BETHE-SALPETER EQUATION

Future Plans and Challenges

The so-called “starting-point dependence” of the BSE as well as GW methods (i.e., the choice of the density functional on which the initial self-consistent ground state DFT calculations of orbitals and polarizabilities is based) continues to be an active area of research for practical application of Green's function theory in electronic structure calculations[334]. Using the FHI-aims code, some progress has been made for molecular systems with the renormalized singles method[390]. Practical applications of the periodic BSE method are still limited by the necessarily dense Brillouin Zone (BZ) sampling. Γ -centered sampling is currently implemented in the FHI-aims code. Optical absorption spectra of extended systems are known to converge slowly with respect to BZ sampling. However, explicitly calculating the screened Coulomb interaction matrix, W , on a dense grid is often computationally intractable due to severe memory and CPU constraints. To address this, future implementations will incorporate interpolation strategies, such as the double-grid technique (calculating W on a coarse grid and interpolating to a fine grid) [391, 392] or Fourier interpolation methods [393]. As we make further progress with the Brillouin Zone sampling, the transitions such as those mediated by phonons with a finite momentum can be computed. This will permit further usage of this many-body method for computing the exciton-phonon coupling in various applications.

In principle, the implementation of Casida's equation in FHI-aims can be straightforwardly extended for periodic systems by utilizing the periodic BSE implementation. At the same time, the proper description of XC kernel for extended systems is necessary. In particular, XC kernel of the adiabatic LDA fails to behave correctly in the long-wavelength limit[394, 395]. Several remedies have been proposed in the literature, including the use of screened exact exchange[396, 397] and corrections motivated by many-body approaches like BSE[398]. We intend to examine and incorporate some of these recent advancements in the future. On the technical side of software development, a matrix-free version of the Casida equations is also planned, employing iterative eigenvalue solver techniques. This approach will enable simulations of larger systems by avoiding the storage of the full Casida matrix, while also supporting hybrid XC functionals.

Acknowledgments

D. Golze acknowledges financial support by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) for funding via the Emmy Noether Programme (project number 453275048). We are grateful to Dr. Heiko Appel for his insights and support with the LR-TDDFT implementation in FHI-aims.

4.5 Real-Time Time-Dependent Density Functional Theory

Hannah Bertschi¹, Joscha Hekele², *Yosuke Kanai^{3,4}, *Peter Kratzer², Christopher Shepard³, Jianhang Xu³, and Yi Yao^{5,6,7}

¹Max Planck Institute for the Structure and Dynamics of Matter, D-22761 Hamburg, Germany

²Faculty of Physics, University of Duisburg-Essen, D-47057 Duisburg, Germany

³Department of Chemistry, University of North Carolina at Chapel Hill

⁴Department of Physics and Astronomy, University of North Carolina at Chapel Hill

⁵The NOMAD Laboratory at the FHI of the Max Planck Society, D-14195 Berlin, Germany

⁶Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, North Carolina 27708, USA

⁷Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

*Coordinator of this contribution.

Summary

The explicit real-time propagation approach for time-dependent density functional theory (RT-TDDFT) has become an increasingly popular first-principles computational method for modeling various time-dependent electronic properties of complex matter. Readers are referred to the comprehensive review by Li *et al.* [399] on the development of the real-time propagation approaches for electronic structure theory methods in general. The theoretical formalism of TDDFT is based on the Runge-Gross theorem [400], and TDDFT is widely used within the linear response theory approach as has been discussed in Section 4.4. The real-time propagation approach allows us to further expand the scope of the applicability beyond the linear response or the traditional perturbative regimes, and RT-TDDFT simulations are increasingly employed to help answer various outstanding questions, especially pertaining to non-equilibrium electron dynamic phenomena, including interfacial charge transfer, atom-cluster collisions, topological quantum matter, etc. [401–403]. In advantage over linear response theory based on the Casida equation [385], RT-TDDFT solely requires the application of the system's Hamiltonian to some starting wavefunction; hence its computational efforts scales linearly with the system size. Excited-state energies, including many-body interactions, can be obtained via Fourier transformation of time series. However, as becomes apparent from the Fourier sampling theorem, determining highly resolved excitation spectra will require long propagation times, and hence efficient propagation methods. Furthermore, Ehrenfest dynamics [404] can be performed on top of RT-TDDFT so that non-adiabatic effects of electrons can be taken into account in first-principles molecular dynamics simulations. Here, FHI-aims [8] with its atom-centered basis functions has the advantage that basis functions are 'carried around' with the atoms as they move, resulting in an accurate description already with modest computational effort. In contrast to plane-wave basis sets, Pulay forces due to the continuously occurring basis set transformations need to be taken into account.

Current Status of the Implementation

RT-TDDFT The expansion coefficients $c_{in}(t) \in \mathbf{C}$ of the KS orbitals here contain the time-dependence of the electronic system as a matrix $\mathbf{C} \in \mathbf{C}^{N_{\text{basis}} \times N_{\text{occ}}}$. Only the N_{occ} initially occupied orbitals are evolved in time. The time-dependent Kohn-Sham (TDKS) equation (also used for the NAO implementation in the FHI-aims code, see below) is

$$\frac{d}{dt}\mathbf{C}(t) = -i\mathbf{S}^{-1}\mathbf{H}(t)\mathbf{C}(t) \quad (4.14)$$

with the overlap matrix $S_{ij} = \langle \phi_i | \phi_j \rangle$ and the Hamiltonian matrix $H_{ij} = \langle \phi_i | \mathcal{H}^{\text{KS}} | \phi_j \rangle$ is then solved to describe electron dynamics. Note the implicit time-dependence via the electron density (i.e. $\mathbf{H}(t) \equiv \mathbf{H}[n(t), t]$). The efficient and accurate solution of this equation is the key functionality in every RT-TDDFT code. Starting with release 210716, the FHI-aims code has several options for integrating the TDKS equations. These include the Crank-Nicolson scheme [405] to variable order, the exponential midpoint rule (including its variant with enforced time-reversal symmetry), but also more complicated schemes such as the commutator-free Magnus expansion of order four, and many more. The latter schemes require the computation of an exponential of the Hamiltonian matrix; for this task the user can choose between a straightforward eigen decomposition or a 'scaling and squaring' method that is often computationally more affordable since only matrix products are required. Maintaining self-consistency between the charge density and the Hamiltonian is crucial in any integration of the TDKS equations and often limits the time step δt that can be used for integrating Eq. 4.14. This problem can be mitigated by using one of the implemented predictor-corrector schemes [406] for the propagation, see Fig. 4.10. All options are documented in the FHI-aims manual.

The fundamental approximation in most RT-TDDFT simulations is the so-called adiabatic approximation to the XC potential that ignores the (in principle mandated) time-non-local character of the XC potential ('history dependence'). Likewise, our present implementation does not consider the dependence of the functional on the current as required in TD-current-DFT.

In RT-TDDFT, we are generally interested in the time-dependent response of the system to some external perturbation. Interaction with the light is often modeled using the classical electromagnetic field. Within the dipole (or long-wavelength) approximation for the electromagnetic field (i.e. neglecting any spatial dependence), the external potential is given in the velocity gauge as $\mathcal{V}_{\text{ext}}(t) = -i\mathbf{A}(t) \cdot \nabla + \mathbf{A}^2$ or in the length gauge as $\mathcal{V}_{\text{ext}}(t) = \hat{\mathbf{r}} \cdot \mathbf{E}(t)$ as connected to each other by a gauge transformation. $\mathbf{A}(t)$ is the vector potential and $\mathbf{E}(t) = -\partial_t \mathbf{A}(t)$ is the electric field.

Ehrenfest Dynamics When the external perturbation is slowly varying, the electrons remain in the equilibrium ground state for a given set of atomic nuclear positions. In this adiabatic limit, the system's response can be studied by solving for the instantaneous eigenstate of the time-dependent Hamiltonian. First-principles molecular dynamics (FPMD) has found great success in various areas of chemistry and condensed matter physics. When the external perturbation is fast-varying, however, the time evolution of the quantum state of electrons needs to be explicitly modeled instead of assuming them to remain in the ground state. In the Ehrenfest dynamics approach, the force on classical atomic nuclei can be obtained from the electrostatic interaction with the time-dependent electron density, which can be described using RT-TDDFT.

Roadmap on Advancements of the FHI-aims Software Package

4.5. REAL-TIME TIME-DEPENDENT DENSITY FUNCTIONAL THEORY

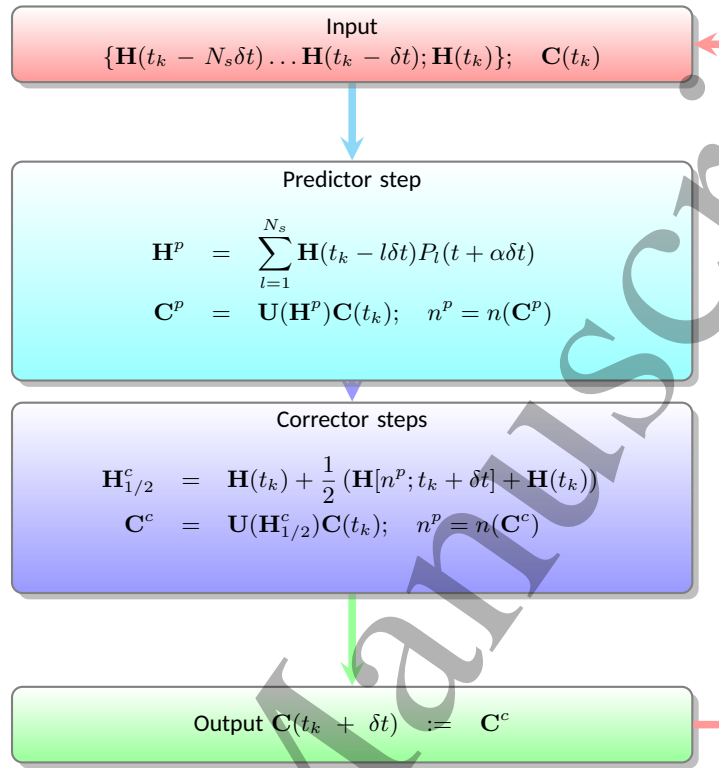


Figure 4.10: Flowchart of a predictor-corrector scheme with time step δt and a unitary propagator \mathbf{U} applied to expansion coefficients \mathbf{C} in Eq. 4.14, Charge density n^p is predicted by polynomial expansion with polynomials P_l from N_s previous Hamiltonians \mathbf{H} . The corrector step works with a mid-point Hamiltonian $\mathbf{H}_{1/2}^c$ and may require iteration to converge n^p .

Electrons are described using the TDKS equations with a non-adiabatic term \mathbf{G} :

$$\frac{d}{dt}\mathbf{C}(t) = -i\mathbf{S}^{-1}(\mathbf{H} + \mathbf{G})\mathbf{C}(t),$$

$$G_{ij} = -i \sum_I \dot{\vec{R}}_I \cdot \vec{\mathbf{B}}_I, \quad (4.15)$$

where \vec{R}_I is the coordinate of nucleus I , and $\vec{\mathbf{B}}_I = \langle \phi_i | \nabla_I | \phi_j \rangle$. The nuclei are propagated using the

conventional energy gradient with non-adiabatic contributions,

$$\begin{aligned}\vec{F}_I &= \vec{F}_I^{\text{HF}} + \vec{F}_I^{\text{MP}} + \vec{F}_I^{\text{XC}} + \vec{F}_I^{\text{DBC}} + \vec{F}_I^{\text{NC}}, \\ \vec{F}_I^{\text{DBC}} &= - \sum_n^{N_{\text{occ}}} \sum_{ij}^{N_{\text{basis}}} f_n c_{in}^* c_{jn} \left[\langle \nabla_I \phi_i | \hat{H} | \phi_j \rangle + \langle \phi_i | \hat{H} | \nabla_I \phi_j \rangle \right], \\ \vec{F}_I^{\text{NC}} &= \sum_n^{N_{\text{occ}}} f_n \mathbf{c}_n^\dagger \left[\mathbf{H} \mathbf{S}^{-1} \vec{\mathbf{B}}_I + \vec{\mathbf{B}}_I^\dagger \mathbf{S}^{-1} \mathbf{H} + i \left(\vec{\mathbf{W}}_I^\dagger - \vec{\mathbf{W}}_I + \mathbf{D}_I^\dagger \mathbf{S}^{-1} \vec{\mathbf{B}}_I - \vec{\mathbf{B}}_I^\dagger \mathbf{S}^{-1} \mathbf{D}_I \right) \right].\end{aligned}\tag{4.16}$$

The first three terms are existing forces in FHI-aims, \vec{F}_I^{HF} are the standard Hellmann-Feynman forces, \vec{F}_I^{MP} are multipole force contributions and \vec{F}_I^{XC} are XC related forces. \vec{F}_I^{DBC} is the dynamical basis correction term that can be seen as the time-dependent analogue to Pulay forces. \vec{F}_I^{NC} is the non-adiabatic coupling force with $\mathbf{W}_I = \dot{\mathbf{R}}_I \langle \nabla_I \phi_i | \nabla_I \phi_j \rangle$ and $\mathbf{D}_I = \dot{\mathbf{R}}_I \cdot \vec{\mathbf{B}}_I$. For finite systems, D3 [279] and Tkatchenko-Scheffler [269] dispersion corrections on the forces are available. Ehrenfest dynamics is a mean-field theory and thus disregards electron-nuclear dynamical correlations. It can be viewed as a controlled short-time semiclassical expansion of the exact Ehrenfest theorem. It can perform well for ultrafast, nearly coherent processes where the nuclear wave-packet remains sufficiently localised. It fails when electronic states have widely different forces and different possible pathways for the nuclear wave-packet would have to be considered.

In some applications, e.g. in the context of non-adiabatic friction or optical breakthrough in dielectrics, the deviation of the time-propagated wavefunctions from their stationary counterparts can be interpreted as promoting electrons into energetically higher-lying Kohn-Sham states. For instance, the energy lost by a projectile due to electronic friction (i.e. electronic stopping) is spent for the creation of electrons and holes in the target material. To obtain the probability $n_{\text{ex}}(t)$ of creating such electrons and holes, the time-propagated TDKS wavefunctions $\psi_{n\mathbf{k}}(t)$ are projected onto a complete set of electronic ground-state wavefunctions $\psi_{n\mathbf{k},t}(0)$ for the particular atomic geometry at time t , thus yielding

$$n_{\text{ex}}(t) = \sum_{nn'\mathbf{k}} (\delta_{nn'} - |\langle \psi_{n'\mathbf{k},t}(0) | \psi_{n\mathbf{k}}(t) \rangle|^2)$$

This option is available as well in FHI-aims.

RT-TDDFT can be performed also with the Nuclear Electronic Orbital (NEO) method[407], referred to as “RT-NEO-TDDFT”, such that coupled quantum dynamics of atomic nuclei (usually protons) and electrons can be simulated on an equal footing[408, 409]. The NEO approach is discussed in Section 6.3.

Finally, we make some comments with respect to other RT-TDDFT implementations. For materials simulations, the plane-wave-pseudopotential formalism remains the most popular implementation of electronic structure methods, and this is the case also for RT-TDDFT, as implemented in codes like Qbox[410] and Qb@II[411]. Our NAO implementation in the FHI-aims code, due to its all-electron description, has some unique advantages. For instance, it allows users to model core-electron excitation dynamics in materials. At the same time, studying phenomena such as ionization would require careful consideration of the NAO basis set flexibility.

Usability and Tutorials

A tutorial that describes how to set up RT-TDDFT simulations using FHI-aims is available under [this link](#)^d. The tutorial provides a brief overview of the theory along with several simple examples. The first section focuses on molecular systems, outlining all the necessary parameters for RT-TDDFT in FHI-aims and explaining how to calculate the absorption spectrum of a single water molecule. The second section addresses the calculation of the optical absorption spectra for periodic systems, using crystalline silicon as the example. Periodic simulations introduce specific challenges that require adjustments to certain parameters, which are thoroughly explained in the tutorial. With both absorption spectrum examples, two python scripts are provided to assist with data evaluation. The first one, called `eval_tddft.py`, allows the user to extract observable quantities, such as the externally applied field, the time-dependent dipole moments and current densities, from the output, and to perform Fourier analysis to obtain the spectra. For atomic and molecular systems, determining the precise spectral location of electronic transitions as well as their oscillator strengths is a major concern. The script `ifit_abs.py` offers graphical support for this task: by clicking on a peak in the spectrum, an algorithm based on Padé approximants [412] outputs the precise peak position. For more refined techniques of extracting excited energy levels, we refer to the recent literature [413].

Finally, the last section of the tutorial demonstrates how to perform Ehrenfest dynamics with RT-TDDFT in the FHI-aims code through a simple example of H_2 bond dissociation. While the tutorial covers only a few small applications of RT-TDDFT, the implementation allows for many others, such as high-harmonic generation simulations or ion bombardment simulations based on non-adiabatically coupled electron-ion dynamics (Ehrenfest dynamics)[414]. Additional information and example inputs for the optical absorption spectrum of benzene, high-harmonic generation on silicon, and Ehrenfest dynamics of $CH_2=NH_2^+$ can be found in the appendix of Ph.D. thesis of Ref. [414]. Moreover, we point out that electronic transport properties in exotic phase of matter, such as a warm, dense hydrogen plasma, have been addressed recently by means of RT-TDDFT simulations [415].

Future Plans and Challenges

Hybrid XC functionals have become increasingly popular in recent years. Currently, hybrid XC functionals in RT-TDDFT are implemented only for molecular (non-periodic) systems. Since the Hamiltonian needs to be evaluated in each timestep, efficient methods for calculating exchange integrals are key to the efficiency of hybrid functional RT-TDDFT. In contrast to molecular systems, the methods such as the resolution of the identity are crucial for efficiently evaluating exchange integrals for periodic systems, and they are currently being developed in the context of RT-TDDFT.

The applications of RT-TDDFT within the FHI-aims code so far have been dealing with systems of light atoms, e.g. first-row elements or silicon (cf. Ref. [416]). FHI-aims, being an all-electron method, includes (scalar) relativistic treatment of the core shells of atoms. While this treatment (called the zeroth-order approximation ZORA of the Koelling-Harmon scheme) is active and usable also in the present RT-TDDFT implementation, it could lead to stability problems if medium to heavy elements (from Fe onward) shall be treated via RT-TDDFT. The highly negative eigen energies of core levels may lead to inaccuracies in the propagation of the phase of the KS wavefunctions in exponential schemes. Similarly, the large spectral range of the all-electron Hamiltonian is a challenge for solving the linear system of equations associated with the implicit Crank-Nicolson propagation scheme. As a future development, one may implement a method that uses different

^d<https://gitlab.com/FHI-aims-club/tutorials/real-time-time-dependent-dft-in-fhi-aims>

*Roadmap on Advancements of the FHI-aims Software Package***4.5. REAL-TIME TIME-DEPENDENT DENSITY FUNCTIONAL THEORY**

schemes to propagate the core and valence states of condensed-matter systems containing heavy elements. Moreover, the fully relativistic treatment using four-component spinor wavefunctions (see Section 2.3) yet awaits its implementation in the RT-TDDFT module.

For applications that involve excitation of electrons to continuum or near-continuum (e.g. Rydberg) states, the atom-centered basis sets used in FHI-aims obviously poses some limitations. Here, a key point is the accurate description of the density difference between the ground state and the excited state. This could be achieved by introducing some additional unbiased basis functions, and ultimately by an additional real-space grid if required. Such a major code extension could be a long-term project for the future.

Acknowledgements

This work was partly (Y.K., J.X., C.S.) funded by the National Science Foundation (United States of America) under Award Nos. CHE-1954894 and partly (J.H.,P.K.) by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) Project No. 278162697-CRC1242 and H.B. was funded by the European Union through the Marie Skłodowska-Curie Grant Agreement no. 101118915 (TIMES).

4.6 All-Electron Δ -Self-Consistent-Field Calculations in Molecular and Periodic Systems

Samuel J. Hall¹, *J. Matthias Kahk², Benedikt P. Klein³, *Reinhard J. Maurer^{4,5,6}, Georg S. Michelitsch⁷, Dylan Morgan⁴, and Karsten Reuter⁷

¹Helmholtz-Zentrum Berlin für Materialien und Energie GmbH, Berlin, DE 10409, Germany

²Institute of Physics, University of Tartu, W. Ostwaldi 1, 50411 Tartu, Estonia

³Research Center for Materials Analysis, Korea Basic Science Institute, Daejeon 34133, Republic of Korea

⁴Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

⁵Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

⁶Faculty of Physics, University of Vienna, Vienna A-1090, Austria

⁷Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

Summary

In a conventional density functional theory (DFT) calculation, the Kohn-Sham orbitals are populated according to the Aufbau principle. However, it is also possible to perform calculations in which the self-consistent field (SCF) is allowed to converge for an excited electronic configuration [417]. DFT calculations with non-Aufbau-principle occupation numbers are most commonly used to model core-excited states of materials and molecules, e.g. in theoretical simulations of X-ray Photoelectron Spectroscopy (XPS) and X-ray Absorption Spectroscopy (XAS) [418–427].

As an example, consider the removal of a C 1s core electron from a methane molecule. The electron's binding energy is defined as:

$$E_B = E_{N-1,\text{ch}} - E_{N,\text{ground}}, \quad (4.17)$$

where E_B is the binding energy, $E_{N,\text{ground}}$ is the total energy of the N -electron ground state, and $E_{N-1,\text{ch}}$ is the total energy of the $N-1$ electron final state with a core hole. $E_{N,\text{ground}}$ is straightforward to calculate using ground state DFT. For $E_{N-1,\text{ch}}$, one can perform a DFT calculation in which the occupation number of the lowest energy eigenstate in one of the spin channels is fixed at zero throughout the SCF cycle. When both total energies are known, the core electron binding energy is directly obtained as the difference between the two – this is known as the Δ SCF method. As DFT is a ground-state theory, the formal justification for Δ SCF calculations has been an issue of intense debate. Several works explore the formal theoretical underpinnings of the Δ SCF method [428–434].

FHI-aims allows the user to perform all-electron Δ SCF calculations for both periodic and aperiodic systems, and it contains tools for creating, tracking, and visualizing a localized core hole. Simulations of both charged and neutral excitations from core orbitals, as well as valence excitations are possible – see for example references [421, 424, 435–441].

Roadmap on Advancements of the FHI-aims Software Package

4.6. ALL-ELECTRON Δ -SCF CALCULATIONS IN MOLECULAR AND PERIODIC SYSTEMS**Table 4.1:** A summary of the currently supported types of Δ SCF calculations in FHI-aims.

Geometry	Kohn-Sham eigensolver	deltascf_projector	deltascf_basis
Aperiodic	Serial	✓	✓
	Parallel	✓	✓
Periodic (Γ -point only)	Serial	✓	✗
	Parallel	✓	✗
Periodic (multiple k-points)	Serial	✓	✗
	Parallel	✗	✗

Current Status of the Implementation

In FHI-aims, there are currently two ways to set a fixed occupation number for a given Kohn-Sham eigenstate.

In the first method, invoked by the keyword `deltascf_projector`, the eigenstate whose occupation number is constrained is identified at the start of the SCF cycle by its index. Then, during subsequent SCF iterations, the maximum overlap method (MOM) [442, 443] is used to keep track of that state, even if the energy ordering of the eigenstates were to change.

In the second method, invoked by the keyword `deltascf_basis`, the occupation number constraint is applied to the eigenstate that has the largest contribution from a particular basis function specified by the user. This is best used in cases when an eigenstate is clearly dominated by contributions from a single basis function as is often the case for core states.

The keywords `deltascf_projector` and `deltascf_basis` were introduced into FHI-aims as part of a major overhaul of the occupation-constrained DFT functionality [444]. These new routines replace an older implementation, internally referenced by the keywords `force_occupation_projector` and `force_occupation_basis`, that was not compatible with the use of parallel eigensolvers. In the new implementation, the application of occupation number constraints is performed with the (pre-)exascale electronic structure interface ELSI [7, 103]. The corresponding routine that creates non-Aufbau occupations can also be used within other codes. As a result of using ELSI, subject to the current limitations listed in Table 4.1, Δ SCF calculations now run at approximately the same speed as regular DFT calculations in FHI-aims.

An illustration of the computational speed-up that the new `deltascf` routines offer over the previous, `force_occupation` routines is provided in Figure 4.11. The legacy keywords `force_occupation_projector` and `force_occupation_basis` are now considered obsolete and will be removed from FHI-aims in a future release.

Usability and Tutorials

In the following, the workflow for modelling core-excited states in FHI-aims is outlined for each of the `deltascf` methods described above. A more detailed tutorial on performing Δ SCF calculations in aims is available at <https://fhi-aims-club.gitlab.io/tutorials/core-level-with-delta-scf>.

The general workflow consists of three steps: (i) initializing a localized core hole, (ii) running an SCF calculation in which the remaining electrons are allowed to relax in the presence of the core hole, and (iii) verifying

Roadmap on Advancements of the FHI-aims Software Package

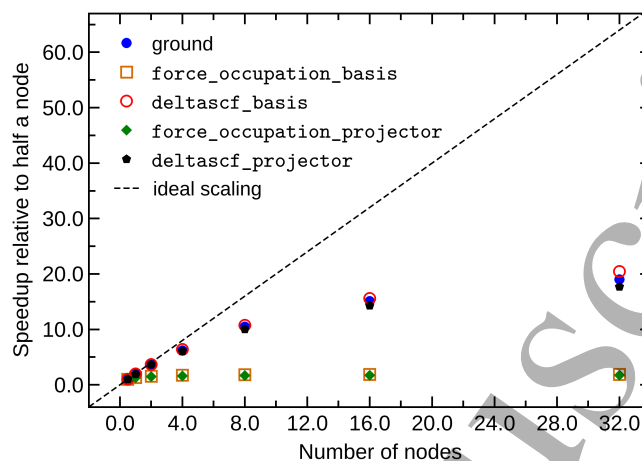
4.6. ALL-ELECTRON Δ -SCF CALCULATIONS IN MOLECULAR AND PERIODIC SYSTEMS

Figure 4.11: The scalability of the new `deltascf` routines in FHI-aims, in comparison to regular ground state DFT calculations, and the legacy `force_occupation` routines. The data, originally presented in [444], shows how the average time per SCF step depends on the number of nodes used to run the calculation. The new `deltascf` routines show similar scalability to regular ground state calculations, whereas with the old implementation, there was almost no improvement in performance when using more than 5 nodes. The test system for the shown scalability calculations was a peroxide-terminated diamond cluster containing 205 atoms - the calculations were run using the PBE functional and the FHI-aims default 'intermediate' level basis sets [8, 27]. The tests were performed on a machine with 128 CPU cores per node, and speedup was calculated relative to the calculation run on half a node (64 cores). The dashed line indicates an ideal 1:1 speedup with respect to the number of processes.

the correct localization of the core hole at the end of the SCF calculation.

The procedure for creating a core hole depends on which `deltascf` method is being used. For `deltascf_projector`, the calculation must be started from an orbital-based restart file (or files). The restart files are typically obtained from a ground state calculation of the same system. To create a core hole on a particular atom, the user must identify the Kohn-Sham eigenstate that best represents the relevant atomic core orbital. In simple cases, this can often be done by examination of the orbital eigenvalues, but tools such as Mulliken analysis [160] or visualization using "cube" files can also be used for this purpose. When the structure contains multiple equivalent atoms, an additional symmetry-breaking step may be required to create a properly localized core hole. Often, simply using a different basis set on the atom whose core electron is to be removed is sufficient for localizing the core orbitals. Alternatively, for aperiodic systems, Boys localization can be applied to the core eigenstates at the end of the ground state calculation [445–448]. Finally, as a last resort, localization of the core orbitals can be achieved by obtaining the restart files from a calculation in which the nuclear charge on the chosen atom where the core hole is meant to be introduced is artificially increased by a small amount, e.g. 0.1 e. For `deltascf_basis`, the use of restart files is not mandatory, although the user must still ensure that the core hole correctly localizes in systems with equivalent atoms.

Performing a DFT calculation with non-Aufbau-principle occupation numbers is in general no different from a ground state DFT calculation in FHI-aims. When a calculation of a periodic system with a charged unit cell is requested, FHI-aims automatically introduces a compensating uniform background charge. When analyzing the results of periodic calculations, the effect of the spurious interactions between periodic images of the core hole must be taken into account. Due to the highly localized nature of core holes, in homogeneous bulk

*Roadmap on Advancements of the FHI-aims Software Package*4.6. ALL-ELECTRON Δ -SCF CALCULATIONS IN MOLECULAR AND PERIODIC SYSTEMS

materials the finite size effect is often well approximated by the first term of the Makov-Payne correction [438, 449]. In other cases, systematic convergence testing with respect to unit cell sizes is required [436].

Finally, for verifying the correct localization of the core hole, again, Mulliken analysis or direct visualization of the Kohn-Sham orbitals can be used. In systems with a closed shell ground state, Mulliken spin analysis is a particularly convenient tool, as the spin density of a core hole will normally be almost entirely localized onto a single atom.

Future Plans and Challenges

As noted in Table 4.1, the current implementation of occupation-constrained DFT in FHI-aims has some limitations in calculations of periodic systems. Specifically, at present, `deltascf_basis` type constraints cannot be applied in periodic calculations at all, and periodic calculations with more than one k-point that use `deltascf_projector` type constraints can presently only be performed using the serial eigensolver. Removing these limitations is the most important short-term goal of the ongoing development work, as calculations with complex periodic structures are often desirable for the interpretation of practical XPS or XAS spectra. Another aim of ongoing work is to remove the limitation that the `deltascf` keywords cannot be currently used in calculations with a fixed overall spin moment.

In the medium term, our aim is to develop localized numerical basis sets that have been optimized for calculations of atoms with a core hole. To facilitate this, the keyword `core_shell_occ` for the species definition section of the `control.in` file has been recently introduced. `core_shell_occ` allows the minimal basis functions, that form a part of the FHI-aims default basis sets to be generated for electronic configurations in which the atomic core orbitals are not fully occupied. However, for complete and usable basis sets, additional sets of augmentation functions (analogous to the Tier basis functions used in FHI-aims default basis sets) optimized for atoms with a core hole must also be developed.

In the longer term, there are plans to also integrate the current Δ SCF functionality with the implementation of fully relativistic DFT in FHI-aims [26], to permit calculations of systems with a core hole in a spin-orbit split p -, d -, or f -shell.

Acknowledgements

This project has received funding from the European Union's Horizon Europe research and innovation programme under grant agreement no. 101131173. Refactorization of the Δ SCF code in FHI-aims was funded under the embedded CSE programme of the ARCHER2 UK National Supercomputing Service (<http://www.archer2.ac.uk>). RJM acknowledges support through the UKRI Future Leaders Fellowship programme (MR/S016023/1 and MR/X023109/1), and a UKRI frontier research grant (EP/X014088/1). JMK acknowledges support from the European Union's Horizon Europe project EXANST (contract 101159716), from the Estonian Ministry of Education and Research grant number TK210 and from the Estonian Research Council grant number PSG1037.

*Roadmap on Advancements of the FHI-aims Software Package*4.6. ALL-ELECTRON Δ -SCF CALCULATIONS IN MOLECULAR AND PERIODIC SYSTEMS

Accepted Manuscript

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

5 Linear Response Methods

Accepted Manuscript

5.1 Density-Functional Perturbation Theory with Numeric Atom-Centered Orbitals

Connor L. Box¹, Reinhard J. Maurer^{1,2,3}, Honghui Shang^{4,5,a}, Nathaniel Raimbault⁴, Danilo Simoes Brambila⁴, Matthias Scheffler⁶, Volker Blum^{7,8}, Christian Carbogno^{6,9,b}, and *Mariana Rossi¹⁰

¹Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

²Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

³Faculty of Physics, University of Vienna, Vienna A-1090, Austria

⁴Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵Key Laboratory of Precision and Intelligent Chemistry, University of Science and Technology of China, Hefei, China

⁶The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁷Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

⁸Department of Chemistry, Duke University, Durham, NC 27708, USA

⁹Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

¹⁰Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

* Coordinator of this contribution.

^aCurrent Address: Key Laboratory of Precision and Intelligent Chemistry, University of Science and Technology of China, Hefei, China

^bCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

Density-functional perturbation theory (DFPT) extends the framework of DFT to the calculation of response properties of the electronic density. DFPT can give access to a large variety of properties, depending on the type of external stimulus that generates the perturbation [450]. An incomplete list of these properties includes force constant matrices and phonons [451], molecular polarizabilities [452] and dielectric properties of solids [450, 453], vibrational scattering cross sections [454, 455], electron-phonon coupling and non-adiabatic matrix elements [456, 457], NMR shifts and J-couplings [31].

The implementation of these techniques within a numeric atom-centered orbital (NAO) framework and real-space context, aligned with the FHI-aims code architecture, necessitates specific choices that we outline in the next section. The original implementation, as detailed in papers by H. Shang and coworkers [458, 459], employed independent routines for different perturbation scenarios. Recently, this approach has been streamlined by centralizing these routines, resulting in a more robust implementation and improved support for the code. [460]

Like other functionalities of FHI-aims, both periodic and non-periodic evaluations are possible within the same all-electron, NAO infrastructure. For periodic phonons, only limited support (LDA functional, and systems that do not require large memory) is provided in the current implementation. A more comprehensive and very efficient handling of the calculation of phonons with different functionals and larger system sizes is provided through the FHI-aims connection to the FHI-vibes infrastructure [461], detailed in another contribution of this Roadmap.

Current Status of the Implementation

In the following, we summarize the main aspects of the implementation regarding atomic displacement and electric field perturbations. A dedicated section on the implementation for magnetic field perturbations is given in contribution 5.2 of this Roadmap.

In FHI-aims, the Kohn-Sham orbitals ψ are expanded in a local NAO basis (cf. (2.1)):

$$\psi_l^{(0)} = \sum_i c_{il}^{(0)} \varphi_i, \quad (5.1)$$

where in the above notation l is the orbital index, i is an index running over the basis functions, the superscript (0) indicates that this is a ground-state quantity, and φ are atom-centered basis functions in finite systems and Bloch-like superpositions of atom-centered basis functions in the periodic case. For the latter case, φ also carries a complex phase (see Eqs. 23 and 24 in Ref. [459]) and the orbitals would depend on k , which we are not showing to simplify notation. Denoting first-order response quantities with the superscript (1), standard first-order perturbation theory yields the Sternheimer equation [462] (for each k point)

$$(\hat{h}_{KS}^{(0)} - \epsilon_l^{(0)})|\psi_l^{(1)}\rangle = -(\hat{h}_{KS}^{(1)} - \epsilon_l^{(1)})|\psi_l^{(0)}\rangle. \quad (5.2)$$

In order to solve it, we can expand the response of the wavefunction analogously to Eq. 5.1,

$$\psi_l^{(1)} = \sum_i [c_{il}^{(1)*} \varphi_i^{(0)} + e_{il}^{(0)*} \varphi_i^{(1)}]. \quad (5.3)$$

For nuclear displacements, the atom-centered basis sets φ indeed change upon displacement and thus yield a non-zero $\varphi_i^{(1)}$, whilst in the case of an electric-field perturbation this term is zero and Eq. 5.3 simplifies to only the first term. While DFPT formalisms concentrate on a self-consistent procedure to determine directly the coefficients $c^{(1)}$, the procedure commonly named coupled perturbed self-consistent field (CPSCF) writes $c_{il}^{(1)} = \sum_{l'} U_{ll'} c_{il'}^{(0)}$ as a linear expansion on the unperturbed coefficients. The new coefficients $U_{ll'}$, where l denotes an occupied orbital and l'

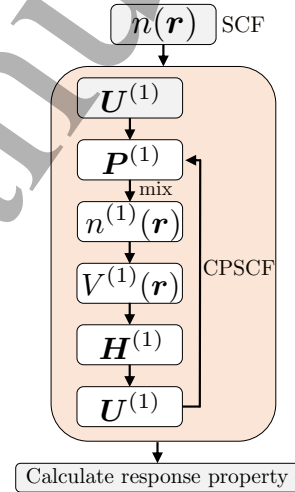


Figure 5.1: Self-consistent workflow for response properties in FHI-aims. The ground-state density $n(\mathbf{r})$ is used for a first guess of the expansion coefficients $U^{(1)}$, which enters the response density-matrix $P^{(1)}$, which then defines the response density $n^{(1)}$. $n^{(1)}$ enters the perturbation potential $V^{(1)}$, the Hamiltonian matrix $H^{(1)}$ is defined and new expansion coefficients $U^{(1)}$ are built. The procedure is evolved self-consistently until the change in $P^{(1)}$ falls below a certain threshold. The desired properties are subsequently calculated.

Roadmap on Advancements of the FHI-aims Software Package

5.1. DENSITY-FUNCTIONAL PERTURBATION THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

an unoccupied one, take the following form

$$U_{ll'} = \begin{cases} \frac{\sum_{ij} (c_{il'}^{(0)})^* H_{ij}^{(1)} c_{jl}^{(0)}}{\epsilon_l^{(0)} - \epsilon_{l'}^{(0)}}, & \text{el. field} \\ \frac{\sum_{ij} (c_{il'}^{(0)})^* H_{ij}^{(1)} c_{jl}^{(0)} - \sum_{ij} (c_{il'}^{(0)})^* S_{ij}^{(1)} (c^{(0)} \mathbf{E}^{(0)})_{jl}}{\epsilon_l^{(0)} - \epsilon_{l'}^{(0)}}, & \text{nuclear displ.} \end{cases} \quad (5.4)$$

and a self-consistent procedure is also necessary to determine them (see Fig. 5.1). Note that the response of the overlap matrix ($S^{(1)}$) is only necessary for nuclear displacement perturbations. The CPSCF formulation, while completely equivalent to DFPT, turns out to be advantageous in FHI-aims because it allows us to make use of existing algorithms for the massively parallel evaluation of matrix elements in this representation. This expansion also allows computing the density response

$$n^{(1)} = \sum_l f(\epsilon_l) [\psi^{(1)*} \psi^{(0)} + \psi^{(0)*} \psi^{(1)}] = \quad (5.5)$$

$$= \begin{cases} \sum_{ij} P_{ij}^{(1)} \varphi_i^{(0)} \varphi_j^{(0)}, & \text{el. field} \\ \sum_{ij} [P_{ij}^{(1)} \varphi_i^{(0)} \varphi_j^{(0)} + P_{ij}^{(0)} (\varphi_i^{(1)} \varphi_j^{(0)} + \varphi_i^{(0)} \varphi_j^{(1)})] & \text{nuclear displ.} \end{cases} \quad (5.6)$$

within a response density-matrix $P^{(1)}$ formalism, analogous to the ground-state density evaluation in the code. The full self-consistent procedure is summarized in Fig. 5.1.

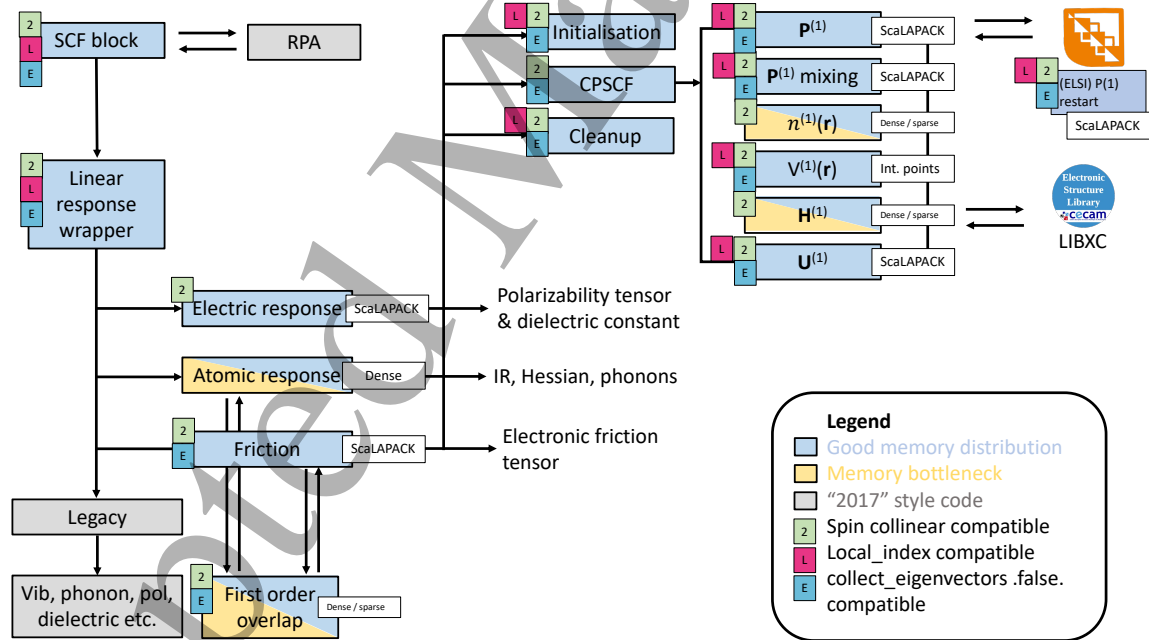


Figure 5.2: Schematic representation of the current code structure in FHI-aims. “Dense / sparse” refers to when the full global dense (aperiodic) or sparse (periodic) response matrices are needed, “Int. points” refers to distribution over integration grid points and “ScaLAPACK” refers to ScaLAPACK support for distribution of matrices. Mixed coloured regions currently represent a memory bottleneck for aperiodic systems due to the use of full dense matrices.

Roadmap on Advancements of the FHI-aims Software Package

5.1. DENSITY-FUNCTIONAL PERTURBATION THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

Table 5.1: DFPT Functionality matrix. Columns represent aperiodic (AP) and periodic (P) implementations across electric field response, electron-phonon coupling, and phonon drivers; “X” marks unavailable functionality. Solid-black cells (■) denote combinations that are not meaningful. The symbol q represents the phonon wavevector.

	Electric		EPC		Phonon	
	AP	P	AP	P	AP	P
LDA						
GGA						X
Hybrid		X	X	X	X	X
MetaGGA	X	X	X	X	X	X
Frozen core						
Spin Collinear					X	X
ScaLAPACK						X
<code>collect_eigenvectors .false.</code>	X	X	X	X	X	X
<code>local_index .true.</code>	X	X	X	X	X	X
Real Eigenvectors						X
Complex Eigenvectors	■		■		■	
$q = 0$						
$q > 0$	■	■	■	X	■	X
Fractional occupations				X	X	X
Dipole correction (surface)	■		■		■	

An overview of the currently supported functionality is summarized in Table 5.1, whilst the code layout and support of features for each specific routine is given in Fig. 5.2. Focusing on Fig. 5.2, the DFPT implementation in FHI-aims is structured around a central *linear-response wrapper* that is invoked following self-consistent ground-state convergence. This wrapper calls the different response drivers currently available, which include: (i) *electric response* (e.g., polarizabilities and dielectric constants), (ii) *atomic response* (phonon spectra and molecular vibrations), and (iii) *electronic friction* (electron-phonon coupling). These response drivers use the same common DFPT/CPSCF module. The modular design facilitates straightforward extension of the framework: each functionality module adheres to a shared and well-documented interface template, enabling future developments that build upon the same DFPT infrastructure. A natural forthcoming addition is the random-phase approximation (RPA) framework, whose force evaluations depend on response quantities that are provided by this same infrastructure.

Recent developments extend the *electric response* and *electronic friction* modules to fractionally occupied systems, including metals, which require specialized handling of the denominator in Eq. 5.4. Further avenues for optimizations are clear: for example, enabling the `use_local_index` flag reduces memory footprint and can improve parallel efficiency, particularly for large-scale systems, routines that support this are labelled in Fig. 5.2. Table 5.1 additionally highlights current capabilities such as spin-collinear support (two spin channels) and the use of `collect_eigenvectors .false.`, which – when paired with ScaLAPACK-based matrix operations – offers significant memory savings for large matrix problems. We note that the cells marked with “X” in Table 5.1 represent improvements that should be achieved in the near future. In

Roadmap on Advancements of the FHI-aims Software Package

5.1. DENSITY-FUNCTIONAL PERTURBATION THEORY WITH NUMERIC ATOM-CENTERED ORBITALS

particular, the robust support of hybrid and meta-GGA functionals for the responses is an ongoing effort.

The central CPSCF module now features interfaces to external open-source libraries such as ELSI [7] and LibXC [463]. ELSI enables a restart of the CPSCF routine by parallelised I/O of the first order density matrix and LibXC enables the calculation of the first order Hamiltonian for arbitrary LDA, GGA, and (when supported) hybrid exchange-correlation functionals

Routines that represent significant bottlenecks in memory distribution and parallelisation are shaded yellow in Fig. 5.2, whilst routines with optimised memory distribution are shaded blue. The half-shading refers to having full dense matrices allocated in the aperiodic case which is memory inefficient when compared to cases that support ScaLAPACK-type distribution, whilst real-space sparse matrices are allocated in the periodic case, which is generally a memory efficient approach. For the latter case, `use_local_index` is desirable for large scale systems.

Usability and Tutorials

All functionality is now accessible through simple keywords in the `control.in` file, documented in the FHI-aims manual since release 240507. Currently, there are keywords that control the central CPSCF shared by all driver routines, these all start with the prefix `dfpt_` and control accuracy thresholds, mixing and restart behaviour. The actual DFPT calculation is triggered by one of the driver-related keywords, currently these are `electric_field_response DFPT`, `atomic_pert_response DFPT`, or `calculate_friction DFPT`.

We have recently released online tutorials, which will be continuously updated and augmented, in order to make the functionality more accessible to new users and showcase what the DFPT functionality can achieve. One tutorial employs the electric-field response functionality to calculate non-resonant vibrational Raman spectra of an isolated molecule and a molecular crystal. Notably, this tutorial also showcases the new implementation of the Berry-phase polarization through the use of Wannier orbitals (showcased in another article of this Roadmap) in order to calculate IR spectra of periodic systems. Another tutorial makes use of the electronic friction driver to calculate electron-phonon couplings and vibrational lifetimes including non-adiabatic effects through electronic friction for a molecular overlayer adsorbed on a metallic surface. These effects are discussed in depth in another contribution of this Roadmap. These tutorials can be accessed through <https://fhi-aims-club.gitlab.io/tutorials/tutorials-overview/>.

Future Plans and Challenges

The newly refactored infrastructure [460] is significantly more compact and easier to maintain than the original. The total number of code lines has been reduced by just over 60% for the CPSCF code, with the newer interface also including more features. Several existing versions of multiple routines have been consolidated, reducing code duplication and increasing clarity for future developers. Specific run-mode cases are now only dealt with at the lowest level and high-level developments at the level of the CPSCF cycle are decoupled. In the near future, we plan to extend the support of local dense matrix computations and the distribution of eigenvectors (currently, a copy of the full eigenvector is created on every MPI task for the k -point(s) it is working on). These changes will improve memory usage and computational efficiency of not just the CPSCF calculations but allow these features to be utilised in the SCF part of the calculation as well, where they are already supported. A scalable implementation of DFPT based on an all-electron NAO basis that performs across various HPC systems has been reported [464]. These implementations can address

*Roadmap on Advancements of the FHI-aims Software Package**5.1. DENSITY-FUNCTIONAL PERTURBATION THEORY WITH NUMERIC ATOM-CENTERED ORBITALS*

systems up to 200,000 atoms. The code scalability using the ELSI library support has also been reported for systems of more than 3000 atoms in Ref. [465]. These developments are compatible with FHI-aims.

It is also important to stress the role that machine-learning (ML) plays in the calculation of electronic response properties. While this field is still less advanced than other topics of ML for materials science, FHI-aims is already able to produce the density-response data that is needed to train models that learn the electronic density response directly, or its derived quantities. The success of these frameworks based on FHI-aims data in predicting Raman spectra, dielectric properties of materials, and electronic friction tensors has already been established [466–469]. This machinery can massively speed up computations and is expected to have an even larger impact on the calculation of electronic response properties in the coming years.

Acknowledgements

We acknowledge fruitful discussions with Andrew Logsdail, Ville Havu, and Xinguo Ren. We acknowledge financial support through the UKRI Future Leaders Fellowship programme (MR/S016023/1 and MR/X023109/1), and a UKRI frontier research grant (EP/X014088/1). MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233.

5.2 Nuclear Magnetic Resonance with Numeric Atom-Centered Basis Sets

*Volker Blum^{1,2}, Xingtao He¹, William P. Huhn¹, Raul Laasner¹, Iuliia Mandzhieva³, Jack Morgenstein¹, *Thomas Theis³, and Franziska Theiss³

¹Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

²Department of Chemistry, Duke University, Durham, NC 27708, USA

³Department of Chemistry, North Carolina State University, Raleigh, North Carolina 27606, United States

(*) Coordinator of this contribution

Summary

Magnetic resonance (MR) spectroscopies are disproportionately important in chemistry, physics, materials science and medicine as atomic-scale probes of the structure and dynamics of matter, as well as in the form of non-destructive diagnostic tools, by exerting quantum control over ensembles of spins [470]. Key atomic-scale parameters of relevance for MR include nuclear shieldings (i.e., the degree to which the response of an individual nucleus is altered by the presence of a surrounding system), nuclear spin-spin couplings or J -couplings (i.e., the effective parameters that govern the quantum-mechanical interaction of two nuclear spins in their surrounding medium), but also molecular magnetizabilities (i.e., the diamagnetic response of the electronic system to an external magnetic field).

In FHI-aims, shieldings and J -couplings for nuclear magnetic resonance (NMR), as well as magnetizabilities are accessible for molecules at the scalar-relativistic level of theory and have been demonstrated for large systems [31]. NAO basis sets capture magnetizabilities and shieldings efficiently. For J -couplings, the response at the nucleus is captured by a benchmarked group of specialized NAO basis sets ("NAO-J-n"), which add highly localized Gaussian-type orbitals (GTOs) near the nucleus, following Jensen's GTO-based pc-J-n basis sets [38, 39]. NMR spectra can be interpreted from these observables, either by employing empirical principles or by interfacing with spin dynamics computations. FHI-aims supports the standardized SpinXML output format [471], which allows it to communicate calculated magnetic resonance parameters directly to the SPINACH spin-dynamics package [472].

Current Status of the Implementation

FHI-aims follows the density-functional perturbation theory (DFPT) formalism [473] to compute magnetic resonance observables. The general DFPT formalism is briefly summarized in contribution 5.1 of this Roadmap. For detailed expressions pertaining to MR as implemented in FHI-aims and for further references, see Ref. [31].

The magnetizability $\overleftrightarrow{\xi}$ of a closed-shell molecule, shielding $\overleftrightarrow{\sigma}_A$ of a nucleus of A , and J -coupling $\overleftrightarrow{J}_{AB}$

Roadmap on Advancements of the FHI-aims Software Package

5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS

of two nuclei of atoms A and B are defined as the following derivatives:

$$\overleftrightarrow{\xi} = - \left. \frac{\partial^2 E}{\partial \mathbf{B}^2} \right|_{\mathbf{B}=0}, \quad (5.7)$$

$$\overleftrightarrow{\sigma}_A = \left. \frac{\partial^2 E}{\partial \mathbf{B} \partial \boldsymbol{\mu}_A} \right|_{\mathbf{B}=0; \boldsymbol{\mu}_A=0}, \quad (5.8)$$

$$\overleftrightarrow{J}_{AB} = \left. \frac{1}{h} \frac{\partial^2 E}{\partial \mathbf{I}_A \partial \mathbf{I}_B} \right|_{\mathbf{I}_A=\mathbf{I}_B=0} = h \frac{\gamma_A}{2\pi} \frac{\gamma_B}{2\pi} \left. \frac{\partial^2 E}{\partial \boldsymbol{\mu}_A \partial \boldsymbol{\mu}_B} \right|_{\boldsymbol{\mu}_A=\boldsymbol{\mu}_B=0}. \quad (5.9)$$

The \leftrightarrow notation indicates that each quantity is a 3×3 tensor and depends on molecular orientation, though the trace is frequently used as an average if the molecular orientation is not known. E is the total energy, \mathbf{B} is the externally applied magnetic field, $\boldsymbol{\mu}_A$ and $\boldsymbol{\mu}_B$ are nuclear magnetic moments, γ_A and γ_B their gyromagnetic ratios, and \mathbf{I}_A and \mathbf{I}_B are the nuclear spins. h is Planck's constant. It is worth noting that $\overleftrightarrow{J}_{AB}$ is not the total spin-spin tensor but rather only the contribution due to the electronic response. The classical dipole-dipole tensor between the nuclei can be tabulated separately (it does not depend on the electronic system) and must be added to $\overleftrightarrow{J}_{AB}$ to obtain the total coupling tensor.

FHI-aims implements all three tensor types by DFPT for non-periodic system geometries, tested for over 1,000 atoms [31]. The local-density approximation and generalized-gradient (GGA) density functionals are fully supported. A full self-consistent field cycle without fields or nuclear spins is performed first, followed by a post-processing computation of the desired tensor for each degree of freedom – a single tensor for $\overleftrightarrow{\xi}$, N tensors $\overleftrightarrow{\sigma}_A$ for N nuclei considered, and $N(N-1)/2$ tensors for $\overleftrightarrow{J}_{AB}$. The current implementation implements non-relativistic and scalar-relativistic theory, limiting the applicability to light elements, where this level of theory is standard in quantum chemistry. For heavy elements, spin-orbit coupling effects will be necessary but are not yet considered.

In quantum mechanics, a magnetic field dependence is represented through the vector potential \mathbf{A} that generates the field. Since $\mathbf{A} = \mathbf{B} \times \mathbf{r}$, the resulting operator depends on the chosen origin and the perturbed wave function acquires a position-dependent phase factor. The standard solution in the computation of shieldings and magnetizabilities with localized basis functions is to include the gauge as a phase factor in the basis functions (the GIAO approach [474]). Thus, in FHI-aims, gauge-including NAOs $\varphi_j^{\text{GIAO}} = e^{-\frac{i}{2}(\mathbf{R}_{A(j)} \times \mathbf{r}) \cdot \mathbf{B}} \varphi_j(\mathbf{r})$ are employed. $A(j)$ denotes the atom at which basis function j is centered. Due to the nature of the perturbation, the first-order perturbation of the orbitals is imaginary-valued and the first-order density perturbation associated with shieldings and magnetizabilities is zero.

In contrast, J -couplings do not require the inclusion of a gauge-dependent phase factor. However, the overall perturbation is associated with a non-zero first-order perturbed density. This response of the electronic system to the presence of a nucleus must be represented by an adequate basis set close to the nucleus. By construction, standard NAO basis sets intended for ground-state calculations do not provide this flexibility since the near-nuclear ground state wave function is already well represented by the minimal basis. Therefore, J -couplings cannot be computed with any degree of reliability using unmodified basis sets originally constructed for ground-state calculations. Instead, numerically precise J -couplings require both additional basis functions very close to the nucleus for stability, as well as dense integration grids very close to the nucleus in order to integrate the near-nuclear perturbation correctly. Therefore, for J -couplings, specifically designed basis sets and integration grids must be employed and are available in FHI-aims [31].

Ref. [31] reports extensive precision benchmarks for molecular magnetizabilities, shieldings, and J -couplings

Roadmap on Advancements of the FHI-aims Software Package

5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS

computed with FHI-aims, covering light elements in the range $Z=1-18$ and conducted at the DFT-PBE [27] level of theory. Importantly, Ref. [31] focused on numerical precision of the implementation, not the physical accuracy of the DFT-PBE functional, while separate and broader benchmark studies [475-477] have shown reasonable qualitative accuracy of DFT-PBE for all quantities considered. The central findings of Ref. [31] are:

- Magnetizabilities are captured with excellent precision using FHI-aims' standard "tier" basis sets for ground-state DFT [8]. For example, FHI-aims' "tier 2" basis sets achieve a precision of $\sim 10^{-30}$ J/T² for a set of 25 benchmark molecules, at around 2/3 the size of comparable general-purpose GTO basis sets tested.
- For shieldings, either FHI-aims' standard basis sets [8] or the NAO-VCC-nZ basis sets, originally designed for basis set convergence of correlated methods [20], can be employed. The average precision of shieldings for 25 benchmark molecules at the "tier 2" level is around 3 ppm, similar to the quality of Jensen's dedicated GTO-based pcS-n basis sets [478] and also similar to the NAO-VCC-3 basis set level. More precise shielding calculations require significantly larger, more expensive basis sets; e.g., the NAO-VCC-4 basis sets approach an average precision of ~ 1 ppm.
- For J -couplings involving light elements ($Z=1-18$), FHI-aims provides a set of dedicated basis sets "NAO-J-n" [31], constructed by amending the standard NAO-VCC-nZ basis sets by primitive tight GTO functions using exponents from Jensen's specialized pc-J basis sets [38, 39], used together with dense integration grids. J -couplings involving elements H, B, C, N, O, Al, Si, S, and Cl are captured with an average precision level of ~ 0.3 Hz already at the moderate NAO-J-3 basis set. The elements F and P show somewhat larger absolute deviations, limited to an average precision of ~ 3 Hz (around 2% of the overall J -couplings targeted for these elements, the magnitudes of which can be quite large).
- For calculations of shieldings and J -couplings in a single run, the NAO-J-n basis sets can be employed.
- Given the larger numerical demands of basis sets and grids for J -coupling calculations, it is possible to use the respective NAO-J-n species defaults only on the specific atoms of interest in a structure (e.g., a molecule of interest surrounded by a solvent), while computing the electronic structure of unrelated atoms with standard ground-state basis sets.
- Finally, Jensen's GTO-based, dedicated pcS and pcJ basis sets for shieldings and J -couplings, respectively, extend to larger size than FHI-aims' basis sets and can also be used for benchmark precision studies in FHI-aims.

Overall, computing magnetizabilities, shieldings and J -couplings for light-element molecules and nanostructures is thus well supported in FHI-aims at the DFT-GGA level of theory (particularly DFT-PBE) - both using moderately sized NAO basis sets with good precision or larger, dedicated GTO basis sets (pcS or pcJ) for verification.

Usability and Tutorials

The DFPT formalism for magnetic resonance spectroscopies in FHI-aims is implemented for local-density and generalized-gradient approximation functionals. Non-relativistic and atomic ZORA scalar-relativistic calculations [8] are supported. The standard FHI-aims "tier" basis sets as well as the NAO-VCC-nZ basis sets

Roadmap on Advancements of the FHI-aims Software Package

5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS

are tabulated as species defaults and can be used for magnetizabilities and shieldings. For the NAO-J-n basis sets, dedicated species defaults with stringent numerical settings exist, using FHI-aims' really_tight numerical defaults as a foundation and increasing the radial density of integration grids by a factor of four compared to the standard really_tight settings. In FHI-aims' terminology, this choice corresponds to the radial_multiplier keyword set to a value of 8 [20]. Jensen's polarization consistent GTO basis sets (pcS for shieldings [478], pcJ for J -couplings [38, 39] and larger, augmented version of the basis sets) are also tabulated as species defaults, with similarly tight settings for grids and for the Hartree potential. It is important to note that the NAO-J-n basis sets and the pcS and pcJ species defaults are thus numerically costly compared to FHI-aims' standard DFT basis sets, for which much lighter numerical settings have been benchmarked.

The actual selection of MR observables to be calculated is made in FHI-aims' input files control.in and geometry.in. At a minimum, the magnetic_response keyword in control.in must be set to specify which type(s) of response observable(s) are requested. Additionally, the atoms to be included in shielding and J -coupling calculations must be identified in geometry.in – specifically, by placing a magnetic_response keyword after any selected atom. This is the minimum information needed by the code to instigate a response calculation. The details of the computation and of the DFPT cycle are extensively customizable by further keywords, all of which are documented in a dedicated chapter in FHI-aims' manual. A dedicated tutorial [479] explains the necessary steps to compute magnetizabilities, shieldings and J -couplings for individual molecules. Furthermore, comments on efficiency and on comparisons to experimentally measured NMR spectra are included.

On their own, computed shielding values and J -couplings from static molecular structures can be used to interpret experimental NMR spectra, but directly plotting calculated shielding and J -coupling values will usually result in poor agreement with experiment. For rigid molecules in which all ^1H atoms are equivalent, such as tetramethylsilane (TMS), calculated shielding values from a single point calculation may serve as a reference that is comparable to experiment. However, molecular motion, solvent environments, hydrogen exchange processes and other phenomena mean that the shape of NMR spectra is rarely a direct mapping of calculated shieldings and J -couplings. For example, the interaction timeframe of a radio frequency pulse with a nuclear spin is of the order of a microsecond, much longer than timescales associated with nuclear motion and certain exchanges.

We exemplify the process of interpreting an experimentally obtained ^1H NMR spectrum of ethanol, using experimental data obtained from ChemicalBook [480]. First, the experimental spectrum of ethanol itself is far from unique, but instead depends heavily on factors such as solvent used and concentration. Among the five very different spectra included in ChemicalBook at the time of writing, we focus on a spectrum obtained for 0.04 ml ethanol in 0.5 ml CDCl_3 obtained at 89.56 MHz. The spectrum exhibits a set of well defined peaks, schematically redrawn in Figure 5.3(a). In this instance, the spectrum shows clearly resolved groups of peaks that may be attributable to the CH_2 group (red), CH_3 group (green) and OH group (blue). The CH_2 and CH_3 peaks show splittings that can be attributed to interaction between both groups via J -couplings.

A first computation of all ^1H chemical shifts (with respect to TMS, i.e., subtracting the chemical shift obtained in a separate static computation of TMS) associated with each H atom in a static, isolated ethanol molecule (inset) is shown in Figure 5.3(b) (DFT-PBE, atomic ZORA scalar relativity, NAO-VCC-5Z basis sets [20] and numerical settings). In comparison to Figure 5.3(a), although no J -couplings are yet considered in the computed spectrum, there is more than one peak per CH_2 and CH_3 group, respectively, and the peak associated with the OH group is far off. One obvious measure to achieve physically more appropriate re-

Roadmap on Advancements of the FHI-aims Software Package

5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS

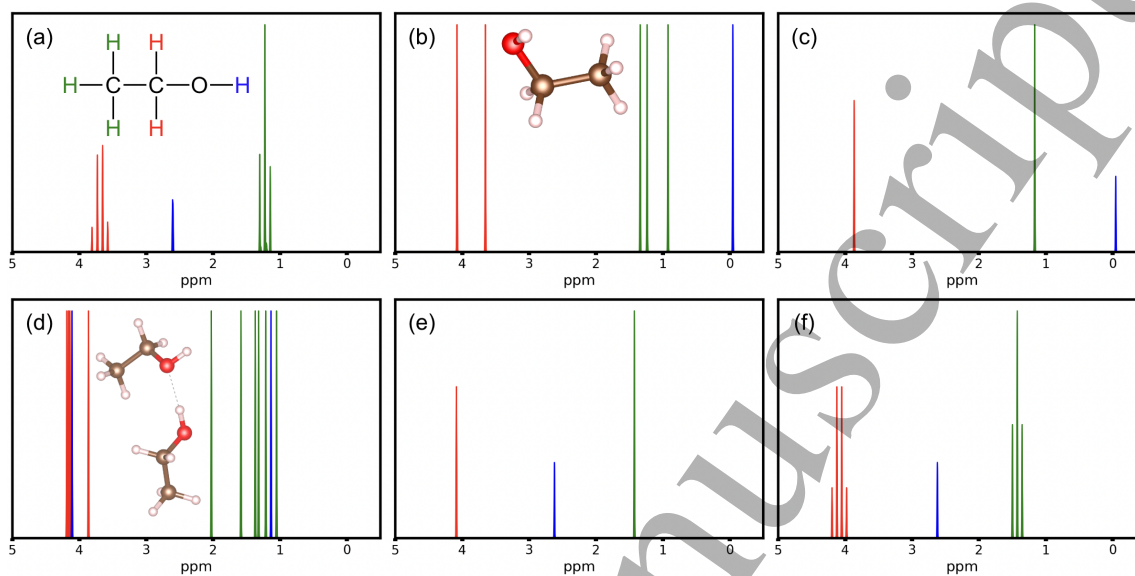


Figure 5.3: (a) Peaks in the experimental Ethanol ^1H NMR spectrum at 89.56 MHz for 0.04 ml ethanol in 0.5 ml CDCl_3 , data taken from [480]. Red, blue, and green peaks are tentatively attributed to signals from the CH_2 group, OH group, and CH_3 group, respectively. (b) DFT-PBE single point calculation of ^1H chemical shifts with respect to TMS, for a single ethanol molecule (inset). (c) Same data as (b) with signals averaged across the CH_2 and CH_3 groups in ethanol. (d) Data pertaining to a DFT simulation of the molecular dynamics (see text for details) of a hydrogen-bonded ethanol dimer. Structures were sampled from this ensemble and the NMR response for each atom was averaged across these samples and visualized. (e) Same data as (d) with signals averaged across each of CH_2 , CH_3 and OH. (f) Same data as (e) but including the peak splitting due to J -coupling between H atoms in the CH_2 and CH_3 groups.

sults is to account for averages between the H atoms within the CH_2 and CH_3 groups, respectively, leading to the spectrum shown in Figure 5.3(c). In this figure, the peak intensity is scaled to the number of atoms represented by a given peak. The CH_2 and CH_3 peaks are now reassuringly close to the related peaks in the experimental spectrum, though still without accounting for J -couplings. However, the peak associated with OH is far off. Similar observations can be made for spectra in different solvents and under conditions in [480] (not shown here), in which the CH_2 and CH_3 peaks remain roughly in the same locations but the position of the OH peak varies widely. Typically, the OH peak is expected to be impacted by hydrogen bonding and H exchange, depending on the solvent environment.

For the specific conditions in Figure 5.3(a), a better interpretation of the experimental spectrum can be arrived at by testing specific hypotheses for the ethanol coordination and environment in direct computations. First, each ethanol molecule undergoes molecular motion; second, even in dilute conditions, aggregation of two or more ethanol molecules to form hydrogen bonds could occur. Figure 5.3(d) tests these hypotheses by averaging over shifts for each H atom individually, obtained from an *ab initio* molecular dynamics (AIMD) trajectory of a hydrogen-bonded ethanol dimer (FHI-aims tight settings, 10 ps, 1 fs AIMD step, stochastic velocity rescaling thermostat [481] at $T=300$ K with 0.5 ps collision time settings) – note that the signal from the hydrogen-bonded OH group (blue, ~ 4 ppm) is drastically shifted compared to the non-hydrogen bonded OH (blue, ~ 1 ppm). Further averaging over chemical ^1H associated with CH_3 , CH_2 and OH groups in both molecules, respectively, leads to the spectrum shown in Figure 5.3(e), with peaks in good agreement

*Roadmap on Advancements of the FHI-aims Software Package***5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS**

with Figure 5.3(a). Finally, the average J -coupling between the protons in the CH_2 group and those in the CH_3 group is computed to be 6.49 Hz for the DFT-PBE functional and NAO-J-5 basis set. Converted to ppm for the experimental frequency used, this amounts to 0.0725 ppm. This J -coupling can be applied to the spectrum using multiplicity rules for the interaction between CH_2 and CH_3 , leading to fourfold splitting of the former and threefold splitting of the latter peak in Figure 5.3(f). This exercise illustrates how mechanistic hypotheses for the behavior of a substance can be tested by comparing to its experimental spectrum, though clearly, experience and significant familiarity with NMR beyond simple static DFT calculations is required. Furthermore, more general solvation effects are not yet included in the NMR calculations (e.g., by implicit solvation or QM/MM) and represent an important future direction for this work.

Finally, we briefly mention that a more automated, less heuristic derivation of NMR spectra from first principles is possible using spin dynamics simulation packages such as SPINACH [472] and the standardized SpinXML format [471] to communicate FHI-aims' computed shieldings and J -couplings between the packages. Importantly, SPINACH supports the computation of non-standard NMR spectra such as low-field spectra which do not require expensive superconducting high-field magnets for their collection. In such spectra, shieldings (the separation of which is scaled by the applied magnetic field) collapse into single peaks and J -couplings (which reflect absolute interaction strengths, i.e., not relative to an external field) shape the spectra. Some of us have recently published a catalogue of low-field spectra (zero-field and spin-locking induced spectra) for a set of over 200 small molecules [482], calculated using FHI-aims and SPINACH and showing that sensitivity to specific molecules is retained in these spectra, offering potential future avenues for low-cost chemical analyses by magnetic resonance.

Future Plans and Challenges

Molecular magnetic resonance parameters in FHI-aims are reliable and well benchmarked for light-element molecules using DFT at the GGA level of theory, enabling detailed interpretation of NMR spectra, with a direct connection to spin dynamics calculations that allow for quantitative simulations, e.g. of low-field spectra.

Beyond the current state of the implementation, significant future improvements are possible. A simple but effective modification would be an approach that contains the need for denser integration grids for J -coupling to the near-nuclear region only, a task that is not simple due to the need for consistently spaced grid shells around each atom for other numerical operations in FHI-aims. Obviously, further benchmarks for potentially improved density functionals, such as hybrid DFT, are highly desirable. Indeed, support for magnetic resonance observables from hybrid DFT is partially implemented (for J -couplings) but not yet fully tested [31].

Beyond these technical points, there are multiple desirable extensions of the physical reach of magnetic resonance computations in FHI-aims. Support for periodic systems should be straightforward for J -couplings, whereas shieldings and diamagnetic magnetizabilities will require an extension of the GIAO approach to periodic systems, e.g., using a modulated magnetic field [483, 484]. For heavier elements, NMR signals will be affected both by relativity (especially spin-orbit coupling [485, 486]) and by the actual, finite shape of a nucleus [487].

Given high-precision, accurate first-principles data, a host of opportunities exist to advance our understanding and interpretation of NMR-based observables, important in fields as diverse as chemical analysis, geological probes, and medical imaging. Continued development of this functionality in FHI-aims will, over time, enable accurate, tractable "ground truth" data across the periodic table.

*Roadmap on Advancements of the FHI-aims Software Package**5.2. NUCLEAR MAGNETIC RESONANCE WITH NUMERIC ATOM-CENTERED BASIS SETS*

Acknowledgements

Acknowledgment is made to the Donors of the American Chemical Society Petroleum Research Fund for partial support of this research.

5.3 Polarization, Born Effective Charges, and Topological Invariants via a Berry-Phase Approach

***Christian Carbogno**^{1,2,a}, **Nikita Rybin**^{1,b}, **Sara Panahian Jand**¹, **Alaa Akkoush**^{1,3}, **Carlos Mera Acosta**^{1,4}, **Zhenkun Yuan**^{1,c}, and **Mariana Rossi**³

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

³Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

⁴Center for Natural and Human Sciences, Federal University of ABC, Santo André, SP, Brazil

*Coordinator of this contribution.

^a*Current Address:* Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

^b*Current Address:* Skolkovo Institute of Science and Technology, Bolshoi bulvar 30, build.1, 121205, Moscow, Russia

^c*Current Address:* Thayer School of Engineering, Dartmouth College, Hanover, NH 03755, USA

Summary

The Berry connection $\mathbf{A}_{mn}(\mathbf{k})$, the Berry curvature $\nabla \times \mathbf{A}_{mn}(\mathbf{k})$, and the Berry phase $\phi = \oint \mathbf{A}_{mn}(\mathbf{k}) \cdot d\mathbf{k}$ are key properties describing the reciprocal-space topology, here the connection between two electronic states labeled m and n . They provide a profound link between the phase of a quantum wave function and macroscopic observables as well as material properties. Most prominently, these quantities are central to our understanding of topological materials and provide a route to classify phases in terms of topology [488]. For instance, all the aforementioned quantities enter the definition of the topological-invariant \mathbb{Z}_2 as given by Fu and Kane [489]:

$$\mathbb{Z}_2 = \frac{1}{2\pi} \sum_n^{\text{occ}} \left\{ \oint_{\partial B} \mathbf{A}_{nn}(\mathbf{k}) \cdot d\mathbf{k} - \int_B [\nabla \times \mathbf{A}_{nn}(\mathbf{k})] \cdot d^2\mathbf{k} \right\} \text{ mod } 2 \quad (5.10)$$

Here, B is half the Brillouin zone (BZ) and ∂B its boundary, the sum runs over all occupied states, and the gauge of $\mathbf{A}_{nm}(\mathbf{k})$ is constrained to respect time-reversal symmetry.

Furthermore, these quantities play a fundamental role for computing the polarization in periodic systems [490, 491], or, more precisely, for calculating the polarization density \mathbf{P} and its derivatives, the Born effective

5.3. POLARIZATION, BORN EFFECTIVE CHARGES, AND TOPOLOGICAL INVARIANTS

charges \mathbf{Z}_I^* , via

$$\mathbf{P} = \frac{1}{V} \left[\sum_I Z_I \mathbf{R}_I - \frac{eV}{(2\pi)^3} \sum_n^{\text{occ}} \oint_{\text{BZ}} \mathbf{A}_{nn}(\mathbf{k}) d^3\mathbf{k} \right] \text{ mod } \mathbf{P}_0 \quad \text{and} \quad Z_{\alpha,I\beta}^* = \frac{V}{e} \frac{\partial P_\alpha}{\partial R_{I\beta}}. \quad (5.11)$$

Here, V denotes the volume of the unit cell, $\mathbf{P}_0 = \frac{e}{V} (|\mathbf{a}_1|, |\mathbf{a}_2|, |\mathbf{a}_3|)$ the polarization quanta along the lattice vectors \mathbf{a}_α , Z_I the nuclear charge, \mathbf{R}_I the nuclear positions, and e the elementary charge. The first term describes the trivial contribution of the bare nuclei, whereas the second term covers the contributions stemming from the electronic states $\Psi_n(\mathbf{k})$. Besides providing the theoretical foundation for understanding the quantization of adiabatic charge transport [492], the polarization is a key property for describing the electrodynamics in solids, e.g., for modeling light-matter interactions and for studying ferroelectric and piezoelectric effects.

In addition, these quantities play a central role in the assessment of currents, fluxes, magnetization, and, last but not least, in the transformation of delocalized electronic wave functions into a localized Wannier basis [493–495]. We refer the interested reader to Ref. [496] for a more thorough discussion of all these effects.

Current Status of the Implementation

To compute all the aforementioned material properties, the fundamental quantity that needs to be calculated in a first-principles code is the Berry connection [493]:

$$A_{mn}(\mathbf{k}) = i \langle u_m(\mathbf{k}) | \partial u_n(\mathbf{k}) / \partial \mathbf{k} \rangle. \quad (5.12)$$

Here, $u_l(\mathbf{k}) = \exp(-i\mathbf{k}\mathbf{r})\psi_l(\mathbf{k}, \mathbf{r})$ is the lattice-periodic part of the electronic wave function $\psi_l(\mathbf{k}, \mathbf{r})$ for state l with wave vector \mathbf{k} and $\langle \cdot | \cdot \rangle$ denotes the scalar product in Hilbert space. Before evaluating this definition, let us recall that FHI-aims uses a Bloch-like representation of the wave functions

$$\psi_l(\mathbf{k}, \mathbf{r}) = \sum_i c_{i,l}(\mathbf{k}) \chi_i(\mathbf{k}, \mathbf{r}) \quad \text{with} \quad \chi_i(\mathbf{k}, \mathbf{r}) = \sum_N \exp(i\mathbf{k}\mathbf{L}_N) \varphi_{i,N}(\mathbf{r} - \mathbf{R}_i - \mathbf{L}_N), \quad (5.13)$$

as detailed in Contrib. 2. Here, $c_{i,l}(\mathbf{k})$ are the Kohn-Sham expansion coefficients and $\varphi_{i,N}(\mathbf{r} - \mathbf{R}_i - \mathbf{L}_N)$ are the numeric atomic orbitals (NAOs) associated with the basis function with index i for the periodic image in the cell \mathbf{L}_N of the atom located at \mathbf{R}_i . With that, the Berry connection can be expressed as

$$A_{mn}(\mathbf{k}) = \underbrace{i \sum_{i,j} c_{j,m}^*(\mathbf{k}) \frac{\partial c_{i,n}(\mathbf{k})}{\partial \mathbf{k}} S_{ij}(\mathbf{k})}_{A_{mn}^{(1)}(\mathbf{k})} - \underbrace{\sum_{i,j} c_{j,m}^*(\mathbf{k}) c_{i,n}(\mathbf{k}) [D_{ij}(\mathbf{k}) - \mathbf{R}_i S_{ij}(\mathbf{k})]}_{A_{mn}^{(2)}(\mathbf{k})}. \quad (5.14)$$

Roadmap on Advancements of the FHI-aims Software Package

5.3. POLARIZATION, BORN EFFECTIVE CHARGES, AND TOPOLOGICAL INVARIANTS

The first term in Eq. (5.14), i.e., $\mathbf{A}_{mn}^{(1)}(\mathbf{k})$, denotes the gauge-dependent Berry-connection term, which here includes the overlap matrix $S_{ij}(\mathbf{k}) = \sum_N e^{i\mathbf{k}\cdot\mathbf{L}_N} \langle \varphi_{j,0} | \varphi_{i,N} \rangle$ due to the non-orthogonality of the employed NAO basis set. For evaluating the associated contribution to the Berry phase, i.e., the closed-path integrals required for Eq. (5.10) and Eq. (5.11), the reciprocal-space path is discretized on $K+1$ points $(\mathbf{k}_0, \mathbf{k}_1, \dots, \mathbf{k}_K)$, whereby the initial and final point are equivalent with respect to the BZ's periodicity $\mathbf{k}_0 = \mathbf{k}_K \bmod \frac{2\pi}{V}$. By expressing the \mathbf{k} -derivatives as two-point finite-differences, one obtains [494]

$$\sum_n^{\text{occ}} \oint_{\text{BZ}} \mathbf{A}_{nn}^{(1)}(\mathbf{k}) \cdot d\mathbf{k} = -\text{Im} \ln [\det (\underline{\mathbf{M}}_{0,1} \cdot \underline{\mathbf{M}}_{1,2} \cdots \underline{\mathbf{M}}_{K-2,K-1} \cdot \underline{\mathbf{M}}_{K-1,0})] . \quad (5.15)$$

Note that the Kohn-Sham coefficients $\underline{c}^{\text{occ}}(\mathbf{k})$ used for computing the matrices

$$\underline{\mathbf{M}}_{a,b} = \underline{c}^{\text{occ}\dagger}(\mathbf{k}_a) \underline{\mathbf{S}}(\mathbf{k}_a) \underline{c}^{\text{occ}}(\mathbf{k}_b)$$

entering the above expression only cover the subspace of occupied states.

The second term entering Eq. (5.14), i.e., $\mathbf{A}_{mn}^{(2)}(\mathbf{k})$, is gauge-invariant and features the matrix

$$D_{ij}(\mathbf{k}) = - \sum_N e^{i\mathbf{k}\cdot\mathbf{L}_N} \langle \varphi_{j,0} | [\mathbf{r} - \mathbf{R}_i - \mathbf{L}_N] | \varphi_{i,N} \rangle , \quad (5.16)$$

which captures the contributions of the NAO basis functions. Since this form exhibits the exact same periodicity as the overlap matrix $\underline{\mathbf{S}}(\mathbf{k})$, it can be integrated using the real-space routines already present in FHI-aims [19, 54]. Similarly, the associated contribution to the Berry-phase required for Eq. (5.10) and Eq. (5.11) can be computed straightforwardly by performing the trace over occupied states and by numerically integrating $\mathbf{A}_{mn}^{(2)}(\mathbf{k})$ along the exact same path used in Eq. (5.15).

In the current implementation, the polarization is calculated in the basis of the reciprocal-lattice vectors. To perform the \mathbf{k} -derivative along the reciprocal-lattice vector of interest, the closed-loop path is chosen parallel to it. The remaining integrations $\int d^2\mathbf{k}$ perpendicular to this path are performed by splining the Berry phases. As an example for such a calculation, Fig. 5.4 shows the polarization of PbTiO_3 , here for smoothly interpolated geometries and lattices between the tetragonal, symmetry-degenerate $P4mm$ equilibrium structures and the cubic, centrosymmetric $Pm\bar{3}m$ structure. While the latter structure must have a vanishing polarization due to symmetry, the tetragonal configurations do not. This highlights that the absolute values of the polarization are meaningless, only relative differences of the polarization matter. To evaluate such differences, “branch-

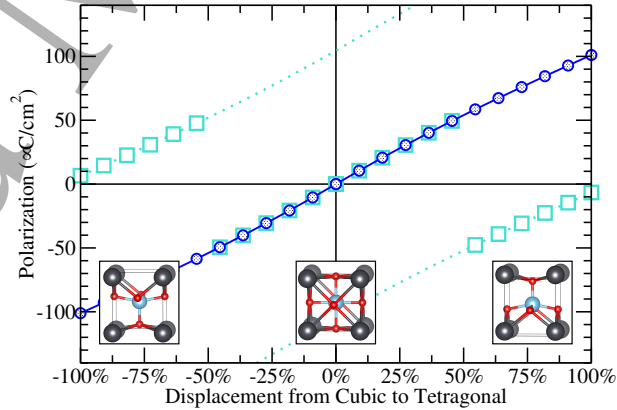


Figure 5.4: Polarization ($24 \times 8 \times 8$ \mathbf{k} -points for the Berry-phase) of PbTiO_3 (HSE06, 8^3 \mathbf{k} -points) along the minimum-energy path connecting the tetragonal, symmetry-degenerate $P4mm$ and the cubic, centrosymmetric $Pm\bar{3}m$ structure. Cyan squares denote the bare output, blue circles the “branch-matched” polarization, for which the discontinuities associated to the mod \mathbf{P}_0 -operation are resolved.

Roadmap on Advancements of the FHI-aims Software Package

5.3. POLARIZATION, BORN EFFECTIVE CHARGES, AND TOPOLOGICAL INVARIANTS

matching” [497], i.e., ensuring that the actual polarization values lie on the branch associated with the same multiples of \mathbf{P}_0 value, is crucial^a.

For the evaluation of the \mathbb{Z}_2 invariant, the current implementation follows the formalism proposed in Refs. [498, 499], which is equivalent to the definition given in Eq. (5.10), but does not require gauge-fixing. In practice, it requires tracking the evolution of the individual Wannier centers

$$\mathbf{X}_n(\mathbf{k}_2) = \oint_{-\pi}^{\pi} A_{nn}(\mathbf{k}_1) \cdot d\mathbf{k}_1 \quad \text{mod } 2\pi \quad (5.17)$$

across a path in the BZ described by \mathbf{k}_2 . In practice, one evaluates the line-path integral in Eq. (5.17) for varying values of \mathbf{k}_2 . Each of these integrals is solved for all occupied states as discussed above, i.e., by using $A_{mn}^{(1)}(\mathbf{k})$ as given by Eq. (5.15) and $A_{mn}^{(2)}(\mathbf{k})$ along a discretized path $\mathbf{k}_1 \perp \mathbf{k}_2$. The determinant viz. trace is, however, not evaluated. Rather, the obtained matrix is diagonalized and the complex phase of the resulting eigenvalues is then tracked, as the example in Fig. 5.5 shows. If an arbitrary continuous line across the whole \mathbf{k}_2 -axis crosses the evolution of the Wannier centers an even number of times, \mathbb{Z}_2 is 0 and otherwise 1. Similarly, this can be judged by tracking the largest gap between the individual Wannier centers [498].

Usability and Tutorials

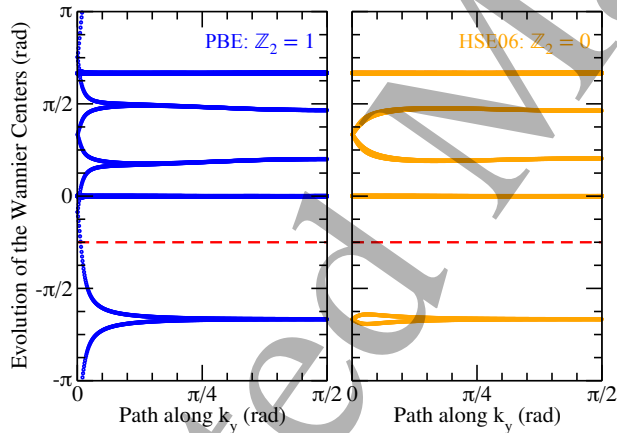


Figure 5.5: Evolution of the Wannier Centers of Charge for the functionalized 2D-honeycomb structure of GeF_2 , as computed with PBE (left) and HSE06 (right) along $\mathbf{k}_2 = k_y$. Both calculations use spin-orbit coupling [48]. As indicated by the red, dashed line, a continuous path along k_y must cross the Wannier Centers an odd (PBE) viz. even (HSE06) number of times, resulting in $\mathbb{Z}_2 = 1$ and $\mathbb{Z}_2 = 0$, respectively. This showcases the influence of the exchange-correlation functional on topological invariants [500, 501].

^aNote that an internal branch matching in FHI-aims is already performed for splining and integrating over the perpendicular \mathbf{k} -directions.

For the evaluation of the polarization, it is sufficient to add one keyword to the control.in file:

```
output polarization alpha n1 n2 n3
```

For instance, the line `output polarization 2 5 10 5` will compute the polarization along the reciprocal-lattice vector number 2 using a grid of $5 \times 10 \times 5$ \mathbf{k} -points in the BZ. As the example highlights, the discretization used along the reciprocal-lattice vector of interest, i.e., the one along which the closed-path integral in Eq. (5.15) is performed, usually requires denser grids. For convenience, multiple `output polarization` statements can be combined and, if all three directions are requested, the code will also report the polarization in Cartesian coordinates. For the evaluation of Born effective charges \underline{Z}_I , a Python script `BEC.py` is provided in the utilities folder of the FHI-aims distribution to perform the

5.3. POLARIZATION, BORN EFFECTIVE CHARGES, AND TOPOLOGICAL INVARIANTS

required derivatives, cf. (5.11) via finite differences. Although these functionalities are rather self-explanatory, a tutorial is provided at <https://fhi-aims-club.gitlab.io/tutorials/phonons-with-fhi-vibes/>, which also shows how the computed Born effective charges can be used to account for long-range dipole interactions in the calculations of phonon spectra in polar crystals, see [450] and references therein.

Similarly, the evaluation of \mathbb{Z}_2 viz. of the evolution of the Wannier centers of charge only requires to add one keyword to the `control.in` file:

```
output Z2_invariant  $\gamma$   $n_{\parallel}$   $n_{\perp}$ 
```

Here, γ is an index that encodes which Cartesian planes shall actually be targeted, e.g., $\gamma = 1$ implies that Eq. (5.17) is evaluated along the first reciprocal-lattice vector using a \mathbf{k} -discretization of n_{\parallel} points and that this procedure is repeated for n_{\perp} closed-paths that have equidistant $k_2 \in [0, \pi]$ and $k_3 = 0$. The latter scan over k_2 can, for instance, be used to discern strong from weak topological insulators [502].

Eventually, let us note that the implementation supports all exchange-correlation functionals, i.e., all semi-local and hybrid functionals, as well as spin-orbit coupling as described in Ref. [48]. Also, given that this functionality targets unit cells and requires rather dense \mathbf{k} -grids, the parallelization occurs over \mathbf{k} -points using LAPACK for the compute-intense linear-algebra operations.

Future Plans and Challenges

So far, the existing implementation has proven useful, accurate, and performant for targeting relatively simple materials with few (< 100) atoms in the unit cell. However, there is increased scientific and technological interest in targeting materials with structural or compositional disorder, e.g., for alloyed topological insulators featuring thousands of atoms in the unit cell [503]. For such kind of systems, the current \mathbf{k} -point-based parallelization strategy is not efficient and limited by memory usage, given that each thread stores a local copy of the required matrices in memory. To reduce the memory footprint per thread and retain computational efficiency, support for distributed linear algebra (ScaLAPACK) is needed and is currently being pursued. This also requires the implementation of functionality that goes beyond the ScaLAPACK memory and communicator layout currently used in FHI-aims, e.g., for concurrent access to quantities at distinct \mathbf{k} -points, as required for evaluating the matrix elements in the closed-loop integral in Eq. (5.15).

Furthermore, the described Berry-phase approach does not only give access to polarization, Born effective charges, and topological invariants, but to a multitude of other material properties, as described in the introduction. In this context, a systematic interface between the methodologies described in this contributions and the density-functional perturbation theory implementation described in Contrib. 5.1 is desirable for the accelerate assessment of response properties, e.g., piezoelectric tensors, but also Born effective charges, or other properties pivotal for electrodynamics viz. light-matter interactions. Another route that is being exploited is machine-learning the polarization using systems with a reduced number of atoms to then gain access to much larger unit cells, e.g., for simulating nuclear dynamics with light-matter coupling [504, 505]. Learning a topological quantity requires special care and several strategies have been proposed in literature for this purpose, e.g., learning the Wannier center positions [506], learning Born effective charges [507, 508], or learning multi-valued polarization surfaces [505].

Finally, let us emphasize that the implemented infrastructure also makes a transformation to Wannier func-

Roadmap on Advancements of the FHI-aims Software Package

5.3. POLARIZATION, BORN EFFECTIVE CHARGES, AND TOPOLOGICAL INVARIANTS

tions straightforward. While this functionality would not be particularly useful with FHI-aims itself, given that the NAOs already provide a localized representation, it would be beneficial for interfacing to other community codes based on a Wannier representation. For instance, this would give access to all the functionality provided by, e.g., Wannier90 [509], EPW [510], and Perturbo [511], and, in turn, enable more systematic, community-wide benchmarks and collaborations across “code-boundaries”.

Acknowledgements

The authors gratefully acknowledge the support of Matthias Scheffler and funding provided by his TEC1p Advanced Grant under the European Research Council (ERC) Horizon 2020 research and innovation programme (Grant Agreement No. 740233).

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

6 Vibrations and Dynamics

Accepted Manuscript

6.1 Phonons, Anharmonicity Quantification, and Strongly Anharmonic Vibrations in Solids

***Thomas A. R. Purcell**^{1,2,a}, **Florian Knoop**^{1,3}, **Shuo Zhao**¹, **Marcel F. Langer**^{1,4,5}, **Marcel Hülsberg**⁸, **Daniel T. Speckhard**^{1,6}, **Maja O. Lenz-Himmer**¹, **Jörg Meyer**^{7,8,b}, ***Matthias Scheffler**¹, and ***Christian Carbogno**^{1,9,c}

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

³Department of Physics, Chemistry and Biology (IFM), Linköping University, SE-581 83, Linköping, Sweden

⁴Machine Learning Group, Technische Universität Berlin, 10587 Berlin, Germany

⁵Laboratory of Computational Science and Modeling, Institut des Matériaux, École Polytechnique Fédérale de Lausanne, 1015 Lausanne, Switzerland

⁶Physics Department and CSMB, Humboldt-Universität zu Berlin, Zum Großen Windkanal 2, 12489 Berlin, Germany

⁷Leiden Institute of Chemistry, Gorlaeus Laboratories, Leiden University, P.O. Box 9502, 2300 RA Leiden, The Netherlands

⁸Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁹Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

^a*Current Address:* Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

^b*Current Address:* Leiden Institute of Chemistry, Gorlaeus Laboratories, Leiden University, P.O. Box 9502, 2300 RA Leiden, The Netherlands

^c*Current Address:* Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

In condensed-matter physics and materials science, the vibrational modes in a material are the primary drivers of materials properties [512]. At the lowest level of theory, it is common to describe these modes using the *harmonic* approximation as a set of uncoupled harmonic oscillators, i.e., phonons [512]. Because all phonon modes are uncoupled, both the classical equations of motion and the quantum Schrödinger equation can be solved analytically and thermodynamic quantities such as the harmonic free-energy and heat capacity can be readily computed. However, this approximation comes with severe deficiencies and inaccuracies, especially for describing processes at elevated temperatures or featuring corrugated potential-energy surfaces. For instance, lattice expansion and vibrational thermal transport cannot be assessed solely within the harmonic approximation, but necessitate the inclusion of additional scattering terms, most prominently those stemming from anharmonic effects [513–516]. Several techniques [517–521] exist that account for such

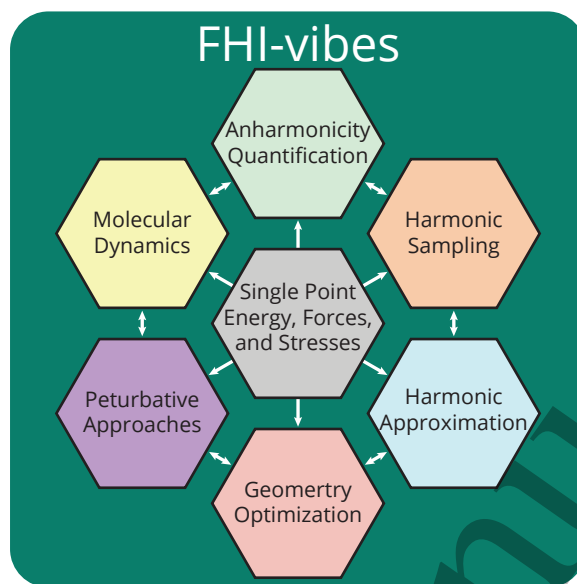


Figure 6.1: Schematic of how FHI-vibes can calculate vibrational properties of materials across all orders of anharmonicity.

effects at various degrees of approximations. Accounting for all orders of anharmonic effects in an unbiased manner, however, typically requires *ab initio* molecular dynamics (*aiMD*) simulations.

FHI-vibes is a Python package that aims to facilitate the investigation of harmonic and anharmonic effects with the electronic-structure theory code FHI-aims. It seamlessly bridges between harmonic phonons, approximate treatments of anharmonicity, and a fully anharmonic description of the vibrational dynamics with *aiMD* [461], as illustrated in Figure 6.1. To achieve this, it builds on the Atomistic Simulation Environment (ASE) [43] to internally represent materials and to interface to first- and second-principles calculators for obtaining energy and forces. From here it links to external codes such as spglib [66], phonopy [522], phonopy [523], TDEP [524], and hiphive [518] to generate harmonic or anharmonic force-constants. Both a command line and a Python API is provided to perform these calculations as well as useful tools for postprocessing the results. In particular, this includes an interface to calculate a material's anharmonicity called σ^A [525], which highlights the advantages of being able to seamlessly process both harmonic and anharmonic dynamics on the same footing. Finally, let us note that these workflows can be run in a high-throughput fashion via FireWorks [526].

Current Status of the Implementation

The central quantity of interest is the potential-energy surface (PES) $U(\{\mathbf{R}\})$, on which the dynamics of the N nuclei with coordinates $\{\mathbf{R}\} = \{\mathbf{R}_1, \dots, \mathbf{R}_N\}$ evolve. In the harmonic approximation, this PES is

Roadmap on Advancements of the FHI-aims Software Package

6.1. PHONONS, ANHARMONICITY QUANTIFICATION, AND STRONGLY ANHARMONIC VIBRATIONS

approximated via the truncated Taylor expansion

$$U(\{\mathbf{R}\}) \approx U^{\text{ha}}(\{\mathbf{R}\}) = U(\{\mathbf{R}^0\}) + \frac{1}{2} \sum_{I,J,\alpha,\beta} \underbrace{\frac{\partial^2 U(\{\mathbf{R}\})}{\partial R_{I\alpha} \partial R_{J\beta}} \Big|_{\{\mathbf{R}\}=\{\mathbf{R}^0\}}}_{\Phi_{IJ}^{\alpha\beta}} (R_{I\alpha} - R_{I\alpha}^0)(R_{J\beta} - R_{J\beta}^0) \quad (6.1)$$

around a (local) minimum-energy configuration $\{\mathbf{R}^0\}$. This already highlights that the dynamics is restricted to oscillations around this specific configuration in the harmonic approximation. The force constants $\Phi_{IJ}^{\alpha\beta}$ characterize the Hessian matrix at this configuration, whereby Greek indices denote Cartesian axes x, y, z . In solid-state physics, it is common to not directly inspect $\Phi_{IJ}^{\alpha\beta}$, but its mass(M)-weighted Fourier-transform with respect to lattice vectors, i.e., the dynamical matrix $\underline{\mathbf{D}}(\mathbf{q})$, which is a function of the reciprocal-space vector \mathbf{q} [512]. Its eigenvalues $\omega_s^2(\mathbf{q})$ and eigenvectors characterize the harmonic phonon modes of a material.

The term *anharmonicity* subsumes all effects of the atomic dynamics not captured by the harmonic approximation, i.e., the differences between $U(\{\mathbf{R}\})$ and $U^{\text{ha}}(\{\mathbf{R}\})$. When the phonon picture is valid, i.e., in the case of weak anharmonicity as discussed in Contribution 7.1, these effects can be captured by extending the Taylor expansion in Eq. (6.1) to higher orders, e.g., third- or fourth order [518, 523], and/or by effectively incorporating anharmonicity in $\Phi_{IJ}^{\alpha\beta}$ [527]. For instance, this can be achieved by fitting $\Phi_{IJ}^{\alpha\beta}$ to *aiMD* trajectories, as done in *hiphive* [518]. For strongly anharmonic materials, the phonon picture breaks down and *aiMD* is needed for an accurate description of the dynamics, also see Contribution 7.1. Comparing fully anharmonic *aiMD* with the harmonic approximation is for instance useful to gauge the degree of anharmonicity present in a material at specific thermodynamic conditions.

In the following, a standard workflow for FHI-vibes is described. It is broken down into ordered subsections to highlight the individual components.

Geometry Optimization

The goal of the geometry optimization step is to find the minimum-energy structure $\{\mathbf{R}^0\}$ around which the Taylor expansion in Eq. (6.1) is performed. Accordingly, $\{\mathbf{R}^0\}$ is the reference geometry around which the vibrational modes are forced to oscillate in the harmonic approximation. FHI-vibes supports geometry optimizations via the algorithms natively implemented in FHI-aims and via those implemented in ASE. This dual approach allows for a larger flexibility, especially when it comes to enforcing constraints on the relaxation [70]. This is particularly useful when addressing different crystal polymorphs for a given composition.

Phonons

Once a minimum-energy structure $\{\mathbf{R}^0\}$ is identified, the harmonic force constants $\Phi_{IJ}^{\alpha\beta}$ can be computed using the finite-difference method [528] as implemented in *phonopy*. *Phonopy* first generates a list of symmetry-reduced displacements from the reference geometry and then FHI-vibes calculates the forces acting on each atom using FHI-aims. All relevant data is then stored in a trajectory file and passed back to *phonopy* to compute $\Phi_{IJ}^{\alpha\beta}$ and all derived properties, i.e., the dynamical matrix, the phonon frequencies and the associated displacement eigenvectors. Furthermore, FHI-vibes also provides an interface to *phonopy*'s postprocessing tools to calculate and plot properties such as phonon bandstructures, densities-

Roadmap on Advancements of the FHI-aims Software Package

6.1. PHONONS, ANHARMONICITY QUANTIFICATION, AND STRONGLY ANHARMONIC VIBRATIONS

of-states (DOS), and thermodynamic properties such as the vibrational free energy and heat capacity. In this phonon picture, it is assumed that the harmonic approximation is a valid approximation for the PES.

We note in passing that performing such phonon calculations at different cell volumes enables an approximate treatment of anharmonicity within the *quasi-harmonic* approximation [529–531]. For instance, this allows to model lattice expansion, as explained in the FHI-vibes tutorials. Further capabilities of FHI-vibes for the treatment of anharmonicity at different levels of approximation are covered in detail in Chapter 7.1 on heat transport.

Ab Initio Molecular Dynamics

For performing *aiMD* calculations, FHI-vibes supports the integrators, thermo- and barostats natively implemented in FHI-aims and those in ASE. As in the case of geometry optimizations, this allows for more flexibility and a wider variety of thermodynamic ensembles to be sampled. For the fast initialization and thermalization of *aiMD* trajectories, information from the harmonic approximation can be used to prepare a system close to the right thermodynamic equilibrium via harmonic sampling [532]. The actual *aiMD* trajectories including position and velocities are then stored as concatenated JSON strings to include metadata for each calculation. This format also facilitates post-processing of the computed *aiMD* data and several routines are provided to the user for this purpose. For instance, (harmonic) force constants $\Phi_{IJ}^{\alpha\beta}$ can be incorporated in the trajectory file and then subsequently be used to map the trajectory onto harmonic phonon modes.

Anharmonicity Quantification

One key advancement enabled by the seamless integration of harmonic models and fully anharmonic *aiMD* present in FHI-vibes is the development of the anharmonicity measure σ^A [525]. This measure defines the anharmonicity of a material as

$$\sigma^A(T) = \sqrt{\frac{\sum_{i,\alpha} \langle (F_{I,\alpha}^{(2)} - F_{I,\alpha})^2 \rangle}{\sum_{i,\alpha} \langle (F_{I,\alpha})^2 \rangle}}, \quad (6.2)$$

where $\langle \cdot \rangle$ denotes a thermodynamic ensemble average at temperature T , $F_{I,\alpha}^{(2)}$ is the force predicted by the harmonic model and $F_{I,\alpha}$ is the fully anharmonic force on atom I and component α . In practice, this metric can be used to quantify the relevance of anharmonic effects under different thermodynamic conditions, e.g., for phase-transition detection [525], for estimating thermal conductivities [533, 534], for analyzing trends in electronic mobilities [535], and even for improving the accuracy and validity of machine-learning potentials for vibrations [536]. Essentially, it normalizes the standard error of the harmonic model to the standard deviation of the DFT forces, and can act as a measure of what fraction of the effects are anharmonic. A per-sample definition of $\sigma^A(T, t)$ can be defined by replacing the thermodynamic ensemble average with the average force on all atoms for each sample. This metric has been used to successfully describe the dynamics of thermal transport [537] and to capture, in an adapted formulation, elastic and inelastic thermal transport across a barrier [538]. This measure can also be used to detect when rare events occur in *aiMD* trajectories, as illustrated in Figure 6.2 for defect formation in CuI [533]. As mentioned above, FHI-vibes allows to decompose σ^A into atomic contributions and to map it onto the individual phonon modes. This decomposition allows for determining the role of strongly anharmonic effects for heat transport (see Contribution 7.1 and

Roadmap on Advancements of the FHI-aims Software Package

6.1. PHONONS, ANHARMONICITY QUANTIFICATION, AND STRONGLY ANHARMONIC VIBRATIONS

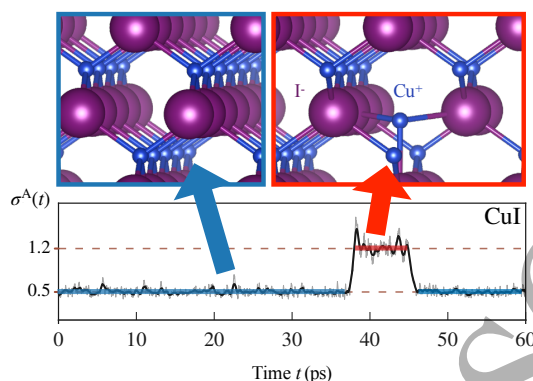


Figure 6.2: The anharmonicity score per time step for an *aiMD* trajectory of CuI. The outset structures represent the average positions of the atoms in the CuI supercell over the blue (left) and red (right) highlighted regions of the trajectory. This is reproduced from [533]

Ref. [533]), clarifying the origin of anharmonicity in β -zeolites [539], and to design machine-learning based descriptors for heat transport [534].

Usability and Tutorials

In this section we will provide a brief tutorial on how to use FHI-vibes, for a complete set of instructions see our documentation here: <https://vibes-developers.gitlab.io/vibes/> and the FHI-aims-specific tutorials provided here: <https://fhi-aims-club.gitlab.io/tutorials/phonons-with-fhi-vibes/>.

Setting up a Single Point Calculation

One of the most important steps in using FHI-vibes is specifying the parameters used for a single-point calculation that is done using a single geometry without optimizing the structure. No matter what task is being completed FHI-vibes needs to know how to calculate the energy, forces, and stresses for a given material. Because the interface is built on top of the ASE interface, all supported ASE calculators can be used with FHI-vibes. An example of what must be defined in this section is shown below.

```
[files]
geometries:    geometry.in.???
```

```
[calculator]
name:          aims
socketio:     true
```

```
[calculator.parameters]
xc:           pw-lda
compute_forces: true
```

*Roadmap on Advancements of the FHI-aims Software Package***6.1. PHONONS, ANHARMONICITY QUANTIFICATION, AND STRONGLY ANHARMONIC VIBRATIONS**

```
[calculator.kpoints]
density:      3

[calculator.basissets]
Si:          light
```

All input structures that we want to perform the calculation on are listed in the `files` section, under either the `geometry` keyword for a single file or the `geometries` one for multiple files matching a particular pattern. In the example the `???` signifies the pattern to be matched, any three character string. The `calculator` section and corresponding subsections are used to define which ASE calculator object to create, the `name` keyword mapped to a calculator class, `socketio` keyword or subsection outlining which socket to use for the SocketIO calculators, the `parameters` subsection listing all parameters passed to the ASE calculator, and `basissets` listing all basis set information for the materials. Because FHI-vibes is designed to work on one or many input structures, the `k_grid` information can be passed both as a parameter, or via a separate section that specifies the desired density of k-points in each crystal direction for consistent calculations. If no other section is defined in the input file, FHI-vibes will assume that a single-point calculation is requested.

Running Individual Tasks

Once the calculation settings are created each individual task can be then be run by creating a new section titled with the task. The current supported tasks are

- `relaxation`: ASE relaxation
- `phonopy`: 2nd order force constant generator/phonopy calculation
- `phono3py`: 3rd order force constant generator/phono3py calculation
- `md`: Molecular dynamics

Additional types of calculations such as harmonic sampling and anharmonicity quantification can be done through utilities and setting up a series of single point calculations. Each task should have its own input file with tasks chained together through a command line interface. For a comparison of the predicted and experimentally measured vibrational properties using these methods we point the reader to Refs. [522, 523] and the FHI-vibes tutorials.

Using FireWorks for High-Throughput Calculations

Finally FHI-vibes supports high-throughput calculations using fireworks. We provide a set of instructions on how to setup a FireWorks job server (LaunchPad) to run these calculations, but importantly a single input file can be used to run a multi-step job. For these jobs each type of calculation requested should be listed in the input file, with FHI-vibes knowing what order to place each run, e.g. a relaxation should occur before calculating the phonons of a material. It also adds a set of new sections, e.g. `statistical_sampling`, to replace steps that would normally be done by hand. Additionally a `use_aims_relax` keyword was added to the relaxation step to ensure that if a local FHI-aims relaxation is requested it would not affect the phonon calculations.

Future Plans and Challenges

From a physical point of view, all fundamental properties to compute key harmonic and anharmonic properties of a solid are implemented in FHI-vibes already, making it a useful and important tool within the FHI-aims software-verse. However, this does not mean that its development has reached an end. Beside maintenance to adapt to updates in the API's of the employed libraries, the main focus is on improving the usability for both common and less common tasks. For example, the inclusion of long-range dipole-dipole interactions [540] is currently being supported and explained in the tutorials, but requires manual calls to phonopy. In this regard, incorporating native support via FHI-vibes is planned for the near future. Along these lines, also the computation of the properties needed for computing this long-range effects, namely the Born-effective charges (see Contribution 5.3) and the dielectric tensor (see Contribution 5.1) will be incorporated to allow for seamless workflows within FHI-vibes. Similarly, further improvements will be application-driven and revolve around easier access to FHI-vibes' internal routines. For example, our experience in the evaluation of σ^A and of heat transport has shown that the mapping of the anharmonic motion onto harmonic phonon modes can be pivotal to obtain qualitative understanding of the dynamics. Accordingly, exposing the relevant routines so that this kind of analysis can be easily used also for other kind of applications is an important task for our users. All of these improvements will focus on expanding the types of problems accessible using FHI-vibes.

Although FHI-vibes was originally developed as an interface for FHI-aims, its MIT licence and the fact that it builds on top of ASE, allows it to be used with all first- and second-principle codes supported by ASE calculators, including those for machine-learned interatomic potentials (MLIPs, see Contribution 8.2), which would extend the systems accessible by FHI-vibes to those with up to 10,000 atoms. This was demonstrated for heat transport calculations in Ref. [541]. Such MLIP MD simulations are less computationally limited in time- and length-scales compared to *aiMD*, but still benefit from the features of FHI-vibes when it comes to analyzing and understanding the simulations. Naturally, this calls for incorporating training-data creation, training, and testing of MLIPs into FHI-vibes, to allow for a seamless user experience. In this regard, those results suggests that these plans might not only be beneficial for the usability, but also important for improving the reliability and accuracy of MLIPs.

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

6.2 Performing Advanced Ab Initio Molecular Dynamics

Yuanyuan Zhou^{1,a}, Herzain I. Rivera-Arrieta¹, Luca M. Ghiringhelli^{2,3}, Christian Carbogno^{1,4,b}, Matthias Scheffler¹, and *Mariana Rossi⁵

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Department of Materials Science and Engineering, Friedrich-Alexander Universität, Erlangen-Nürnberg, Germany

³Erlangen National High Performance Computing Center, Martensstraße 1, Erlangen, 91058, Bayern, Germany

⁴Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

*Coordinator of this contribution.

^aCurrent Address: State Key Laboratory of Quantum Functional Materials, Department of Physics, and Guangdong Basic Research Center of Excellence for Quantum Science, Southern University of Science and Technology, Shenzhen 518055, China

^bCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

Since its inception [542], *ab initio* molecular dynamics (*aiMD*) has established itself as one of the most successful numerical techniques in chemistry, solid-state physics, and materials science. It offers an appealing connection between two prominent fields, namely electronic-structure theory and statistical mechanics. It allows to compute, analyze, and understand thermodynamic processes and properties without relying on potentially erroneous assumptions for the interatomic interactions. In addition, the electronic-structure that is computed along the whole trajectory enables to assess thermodynamic expectation values of observables that cannot be computed from the nuclear motion alone.

Since its first release [8], FHI-aims has supported a native implementation of *aiMD*, as evidenced by early applications that were based on this functionality [543, 544]. Over time, FHI-aims has also been interfaced with codes that are able to perform advanced flavors of *aiMD*.

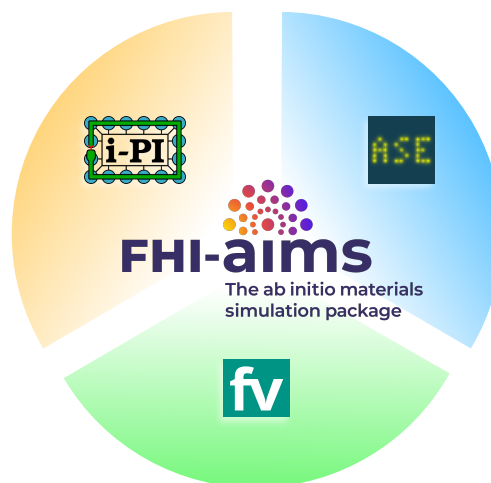


Figure 6.3: The software packages i-PI [504], ASE (Atomistic Simulation Environment) [43], and FHI-vibes [461] provide additionally *aiMD*-related functionalities beyond what natively supported in FHI-aims.

Prominent interfaces that we can mention are i-PI [504], ASE (Atomistic Simulation Environment) [43], and FHI-vibes [461]. Several successful applications have built on this feature, as discussed for instance the Roadmap contributions 6.1, 6.5, 7.2, 7.1, 6.3, and 8.2, as well as a vast array of publications that are too numerous to be singled out.

In this contribution, we summarize the status of the implementation of *aiMD* in FHI-aims, the usability of interfaces, and future plans. As a highlight, we discuss the use of *aiMD* in the context of grand-canonical replica exchange molecular dynamics (REGC) simulations, a method that was first applied with a bespoke interface to FHI-aims [545].

Current Status of the Implementation

The Born-Oppenheimer approximation [546] (BOA), i.e., the separation of the dynamics of electronic $\{\mathbf{r}\} = \{\mathbf{r}_1, \mathbf{r}_2, \dots\}$ and nuclear degrees of freedom $\{\mathbf{R}\} = \{\mathbf{R}_1, \mathbf{R}_2, \dots\}$, is at the very heart of *aiMD*. The assumption of classical nuclear motion and instantaneous interaction with the ground-state electronic state yields the following equations,

$$\hat{H}_{\{\mathbf{R}\}}(\{\mathbf{r}\})\Psi_{\{\mathbf{R}\}}(\{\mathbf{r}\}) = E_{\{\mathbf{R}\}}\Psi_{\{\mathbf{R}\}}(\{\mathbf{r}\}) \quad \text{and} \quad M_I\ddot{\mathbf{R}}(t) = \nabla_I E_{\{\mathbf{R}\}} \quad (6.3)$$

Here, the first equation is the electronic-structure theory problem, that depends *parametrically* on the instantaneous nuclear coordinates. Its ground-state solution $E_{\{\mathbf{R}\}}$ describes the potential-energy surface, on which the nuclei move. Accordingly, it defines the forces acting on the nuclei with masses M_I in the second equation, which is a classical Newtonian equation of motion. Related molecular dynamics methods, where the BOA is relaxed, as for example in Ehrenfest dynamics, or where the assumption of classical nuclei is relaxed, or even where both assumptions are relaxed are discussed in contributions 6.3 and 4.5.

The nuclear equations of motion from Eq. (6.3) can be step-wise integrated [547] to simulate the dynamics and obtain time-discretized trajectories $\{\mathbf{R}(t_l)\}$. At each time step t_l , it is however necessary to solve the associated electronic-structure problem for the Hamiltonian $H_{\{\mathbf{R}(t_l)\}}(\{\mathbf{r}\})$ to obtain the potential and the forces acting on the nuclei, as sketched in Fig. 6.4. While the straight solution to these equations of motion can only lead to sampling of the microcanonical ensemble (NVE), it has been long recognized [548] that coupling the dynamics to thermostats and barostats can lead to sampling the canonical (NVT), isothermal-isobaric (NPT) and the isothermal-isostress (NsT) ensemble. In these ensembles, it is possible to control temperature, pressure and anisotropic stress. Last but not least, *ab initio* replica-exchange molecular dynamics can enable the calculation of the μ VT ensemble with variable particle number, but constant chemical potential, as shown below.

FHI-aims natively supports basic MD functionality, which can sometimes be advantageous on HPC installa-

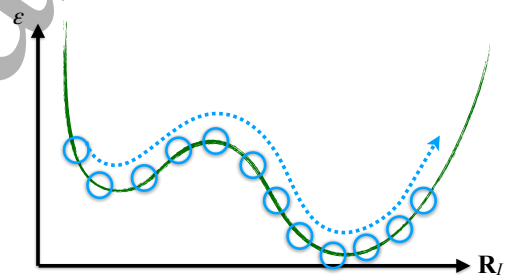


Figure 6.4: Sketch of an *aiMD* calculation on a potential-energy surface ε (green). By step-wise integrating the equations of motion, one can simulate the dynamics. This requires to solve the electronic-structure theory problem in each time step, as indicated by the blue circles.

Roadmap on Advancements of the FHI-aims Software Package

6.2. PERFORMING ADVANCED AB INITIO MOLECULAR DYNAMICS

tions with, e.g., limited or restricted access to (potentially dated) libraries. For more advanced molecular dynamics simulations, FHI-aims is linked to external packages. Under the BOA, the interface comes naturally, as it exploits the separation of electronic and nuclear degrees of freedom that underlies *aiMD*. The external codes take care of solving the nuclear equation of motion, while FHI-aims tackles the electronic-structure problem. Communication between the codes can happen either via input and output file parsing (e.g., ASE, FHI-PANDA^a) or, more efficiently, via a UNIX or internet socket communication (e.g., ASE, i-PI, FHI-vibes).

In more detail, the following functionality is accessible through different implementations:

1. The native MD functionality of FHI-aims supports NVE calculations with a Velocity-Verlet integrator [547], as well as NVT calculations using the Andersen [549], Nosé-Hoover [550], stochastic velocity rescaling [481], and a colored-noise thermostat based on a generalized Langevin ansatz [551]. All necessary infrastructure is provided within the FHI-aims code, so that no additional software package is needed. Related native infrastructure for Ehrenfest dynamics, regarding the coupled motion of electrons and nuclei beyond the BOA are also implemented and discussed in Section 4.5.
2. ASE, the Atomistic Simulation Environment [43], provides a rapid and flexible route to interact with first- and second principles codes via Python. It provides NVE-MD functionality, but also includes support for thermo- and barostats. Besides, it offers a variety of other algorithms to manipulate nuclear coordinates, see contribution 8.1. Accordingly, it is particularly useful for rapid prototyping during development and for linking MD calculations to other approaches, c.f. see FHI-vibes below.
3. FHI-vibes [461] is a Python package to compute vibrational properties of solids from first principles. It can also perform *aiMD* calculations by using the respective functionality built into ASE. By linking such *aiMD* calculations to harmonic and quasi-harmonic descriptions of the dynamics it offers a rich set of analysis tools, e.g., to map and compare with lattice-dynamics calculations for quantifying anharmonicity and for computing heat transport coefficients, see contribution 6.1 and 7.1 for more details.
4. i-PI [504] is a versatile code that is interfaced with a variety of electronic-structure packages, machine-learning potential packages, and empirical-potential packages. When interfaced with electronic structure codes, it can perform several flavors of nuclear motion, ranging from standard ones such as geometry optimizations, nudged-elastic-band, and harmonic phonons, to advanced ones, such as molecular dynamics with higher-order integrators, NVE, NVT, NPT and NsT dynamics, replica-exchange molecular dynamics, and other enhanced sampling techniques. In particular, its interface with FHI-aims allows transferring not only energies, forces, and virial stresses, but also electronic-structure quantities that can be used in non-adiabatic or light-driven dynamics. For the particular use of i-PI for the simulation of nuclear quantum effects, one of its main strengths, see contribution 6.3.

Naïvely employing *aiMD* as a phase-space sampling tool is inefficient in general. At times, this inefficiency is so pronounced (systems with multiple minima and large free-energy barriers, rare reactive events, etc.) that a “vanilla” *aiMD* sampling is effectively impossible. To address this limitation, many enhanced-sampling methods were introduced. These methods can be classified based on their conceptual approach to overcoming barriers into two main categories [548]: (i) incorporating bias terms into the Hamiltonian of the original system, e.g., metadynamics, umbrella sampling, and accelerated molecular dynamics; (ii) generating a generalized ensemble of the original system. Generalized-ensemble simulations involve a broader

^a<https://gitlab.com/zhouyuanyuan/fhi-panda>

Roadmap on Advancements of the FHI-aims Software Package

6.2. PERFORMING ADVANCED AB INITIO MOLECULAR DYNAMICS

sampling of the potential energy landscape. Techniques such as simulated tempering, multicanonical sampling, and parallel tempering (also known as replica exchange) fall into this second category. These methods are particularly effective in overcoming kinetic barriers.

During the last several decades, *ab initio* atomistic thermodynamics [552] has been widely and successfully applied to study the thermodynamic stability of surfaces under realistic conditions. This technique requires a prior knowledge of relevant surface phases. Furthermore, in most cases, the vibrational contributions are neglected or at most treated by harmonic approximation. To free from these limitations, an unbiased sampling of the surface phase space is required. In this context, based on the grand-canonical extension [553, 554] of the original parallel-tempering MD scheme [555, 556], Zhou, Scheffler and Ghiringhelli [545, 557] have recently developed the grand-canonical replica-exchange (REGC) method coupled to *aiMD* and implemented it as a wrapper for FHI-aims. Open ensembles (i.e., exchanging both energy and matter with a reservoir) are described at equilibrium by the grand-canonical-ensemble formalism and provide an effective mean to overcome slow-diffusion problems: Thermodynamically possible defect states can be efficiently generated and sampled by means of the atoms' insertion and removal as well as the parallel coupling between different temperatures and different chemical potentials. In REGC, an extended ensemble including a set of $S = L \times M$ replicas of the studied system is created. The replicas span L values of temperature and M values of the chemical potential of the selected chemical species. Presently, only one species can be exchanged with a reservoir at given chemical potential, but the extension to multiple species is under development. The partition function of this extended ensemble is the product of the partition functions of the individual $\mu_m \nu T_l$ ensembles, where $l = 1, 2, \dots, L$ and $m = 1, 2, \dots, M$,

$$Q_{\text{extended}} = \prod_{l=1}^L \prod_{m=1}^M \frac{e^{\beta_l \mu_m N_{l,m}} \Lambda^{N_{l,m}}}{\Lambda_l^{N_{l,m}} N_{l,m}!} \int d\mathbf{R} e^{-\beta_l E_{\{\mathbf{R}\}}(N_{l,m})}. \quad (6.4)$$

In the equation above, \mathbf{R} denotes the nuclear coordinates that define the configuration of the system, $\beta = 1/k_B T$, Λ is the thermal wavelength, and $E_{\{\mathbf{R}\}}$ is the potential energy of a configuration \mathbf{R} of the N -particle system. During the REGC simulation, short *aiMD* trajectories are run for each replica in parallel, after which a Metropolis Monte Carlo (MMC) move is attempted. In this MMC move, either a replica exchange or a particle insertion/removal is performed. This decision is based on a random criterion. In the replica-exchange move, a swap of configurations between pairs of replicas at different temperatures and/or chemical potential is tried. This means that an MMC criterion based on the difference in β and μ between the pair of replicas and the difference in energy between the corresponding pair of configurations, accepts or rejects the swap of configurations. The detailed-balance conditions imposes a specific form for the probability of swapping configurations, which is given by Eq. (6) in Ref. [545]. In the particle insertion/removal move, the MMC acceptance criterion is based on the difference between the change in energy upon insertion or removal of a randomly positioned particle and the imposed chemical potential, given by Eqs. (2) and (3) in Ref. [545].

The REGC algorithm as described above and sketched in Fig. 6.5, samples the multi-canonical ensemble at each of the L temperatures and M chemical potentials. In REGC, the interface with FHI-aims is based on output parsing. The configuration at a certain thermodynamic condition (T_l, μ_m) for each replica after every particle exchange or replica exchange provides the initial geometry file and the target temperature for FHI-aims to perform *aiMD* in the canonical ensemble. At the end of one iteration of the algorithm, after all the *aiMD* trajectory have run for the predefined number of steps, the code FHI-PANDA reads all the configurations and the energies for the next iteration. In order to estimate ensemble averages of observables at any temperature and chemical potential within (and slightly outside) the sampled intervals, one has to

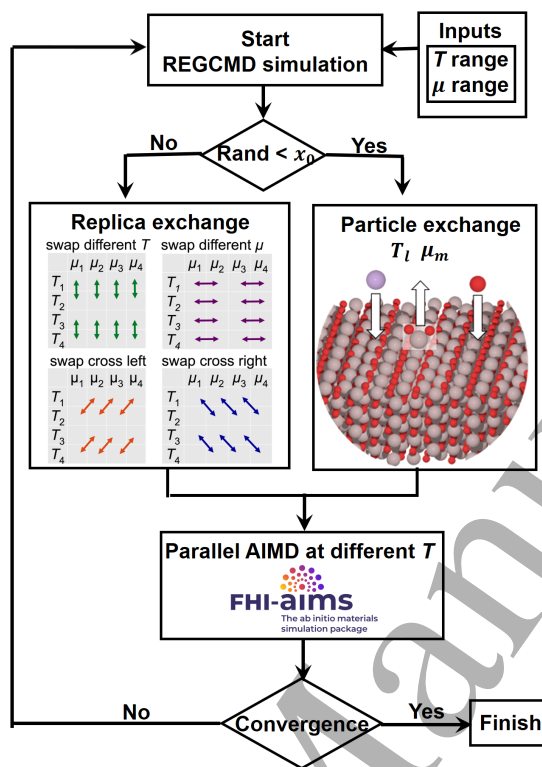


Figure 6.5: Schematic workflow for the replica-exchange grand-canonical (REGC) *ab initio* molecular dynamics. AIMD: *ab initio* molecular dynamics, which is performed within the FHI-aims code; T : temperature; μ : chemical potential. DFT: density-functional theory. x_0 is the probability to perform the RE attempt. ($0 \leq x_0 \leq 1$).

apply a re-weighting algorithm, such as the multistate Bennett acceptance ratio (MBAR) [558]. In this way, one can construct phase diagrams for given observables [545]. In particular, if the chosen observable is the heat capacity of the system, one can infer phase boundaries and construct a complete configurational phase diagram without any prior knowledge of the expected phases [557]. We note that the REGC approach as implemented for FHI-aims is particularly suited to study adsorption on surfaces from the gas phase. In case of bulk systems, the particle insertion MMC move has very low acceptance, leading to poor statistics, unless dedicated biasing mechanisms are implemented.

Usability and Tutorials

The native implementation of *aiMD* in FHI-aims is extensively described in section 3.12 of the code manual (year 2024). The central control of the run is accessed via the `MD_run` flag. A number of flags defining the time-step, velocity initialization, wave-function extrapolation scheme, and restarting options complete the implementation. Since 2021, the code supports a non-self consistent (NSC) Hartree-potential correction to

Roadmap on Advancements of the FHI-aims Software Package

6.2. PERFORMING ADVANCED AB INITIO MOLECULAR DYNAMICS

the Hellmann-Feynman forces of the form [559]

$$\mathbf{F}_I^{(\nu),\text{NSC}} = \int d\mathbf{r} \left[\left(n^{(\nu-1)}(\mathbf{r}) - n_{\text{KS}}^{(\nu)}(\mathbf{r}) \right) \frac{\partial v_H^{(\nu-1)}(\mathbf{r}; \mathbf{R})}{\partial \mathbf{R}_I} \right] \quad (6.5)$$

where ν is the current self-consistent step, $\nu - 1$ is the previous step, n_{KS} is the unmixed Kohn-Sham (KS) density, n is the mixed density, and v_H is the Hartree potential. The correction is computed at each SCF step and converges to zero as n approaches the self-consistent solution. In practice, its explicit inclusion allows obtaining very accurate forces with a less-converged density. This implementation, realized by Rivera-Arrieta and Rossi, required a rearrangement of the SCF loop in FHI-aims, in which the density update and mixing step was moved after the solution of the KS equations, instead of being performed at the beginning of the SCF loop. For *aiMD* simulations, this correction term represented a major improvement in efficiency. In our tests, this correction allows us to obtain energy drifts with a density convergence threshold of $10^{-3} e/a_0^3$ that is comparable to the drift obtained with a $10^{-5} e/a_0^3$ threshold without the correction. In practice, for example for the Zundel cation, this represents a 40% reduction in the number of SCF cycles in a simulation of 10000 steps. This correction is now automatically included in any force calculation of the code, as it does not incur any appreciable computational overhead.

In regards of the functionality of the code, one important addition compared to the original release is the support of variable-cell *aiMD* under periodic boundary conditions. This is enabled by the implementation of the stress tensor, which describes the changes of the PES with respect to symmetric strain deformations of the unit cell [54]. Besides enabling the usage of barostats to sample the isothermal-isobaric (NPT) and the isothermal-isostress (NsT) ensemble, this also allows to monitor the stress during *aiMD* runs, .e.g. to compute thermal-expansion coefficients [78]. Furthermore, this gives access to the virials, i.e., the atom-resolved stress contributions, which allow to compute the virial contributions to the heat flux, see contribution 7.1.

The massively parallel REGC algorithm requires no prior knowledge of the phase diagram and takes only the potential energy function together with the desired μ and T ranges as inputs. As shown in Figure 6.5, during a REGC simulation, each replica is assigned a random number which is compared to a probability x_0 ($0 \leq x_0 \leq 1$) that determines whether to attempt exchanging a particle with the reservoir or to attempt a replica-exchange move. After the particle/replica-exchange attempt, parallel *AIMD* runs follow to diffuse the system in the canonical ensemble, i.e., at temperature T_i , with fixed number of particles N and volume V of the system (NVT ensemble). The procedure is then iterated until the convergence of defined quantities is achieved. A 2D (one dimension is T and the other dimension is μ) replica-exchange scheme is performed for the system in contact with a single gas reservoir. Each replica has 3-8 neighbors, compared to 1-2 neighbors in conventional 1D schemes, which improves exchange efficiency. The tutorial for studying a Si_2 cluster in contact with gas-phase H_2 , where the whole workflow of the REGC method is demonstrated, is available [under this link](https://fhi-aims-club.gitlab.io/tutorials/introduction-of-ab-initio-thermodynamics-and-regc).^b

Future Plans and Challenges

With the development of fast and reliable machine-learning interatomic potentials (MLIPs), *aiMD* simulations have become quickly less popular. The reason is that molecular dynamics with a very similar accuracy

^b<https://fhi-aims-club.gitlab.io/tutorials/introduction-of-ab-initio-thermodynamics-and-regc>

Roadmap on Advancements of the FHI-aims Software Package

6.2. PERFORMING ADVANCED AB INITIO MOLECULAR DYNAMICS

as *aiMD* can be performed at a computational cost that is 4 to 8 orders of magnitude cheaper [504] than the underlying electronic-structure method on which the MLIP was trained. Therefore, the system sizes that can be treated with MLIPs are much larger and statistical convergence of thermodynamical observables can be routinely obtained. However *aiMD* is still useful (and at times still essential), and will likely continue to be for years to come, as MLIP require *ab initio* data to be trained on. In addition, there are systems for which the physics that need to be captured is still challenging for current MLIPs, such as systems with complex charged defects and where electronic spin is particularly relevant. Regarding FHI-aims, a bespoke interface for MLIP active learning based on *aiMD* will soon be released, which will offer several advantages in terms of data generation and rare-event sampling. Wrappers and workflows around FHI-aims that perform active learning already exist and are described in contribution 8.2. It is only natural that the REGC method is also integrated with MLIPs.

Regarding the REGC algorithm, current and near-future development is dedicated to i) a smart handling of the load unbalance generated by the parallel *aiMD* run with different number of particles and ii) the extension to multiple species with independent chemical potentials for multi-component phase diagrams, where the challenge is to create an adaptive grid for the sampled temperatures and chemical potentials.

On a more technical note, there are algorithmic improvements related to *aiMD* that would benefit any situation where the method is used, including data generation. For example, regarding the force-correction term that was discussed in this contribution, one could also compute similar terms for the Hartree-Fock exchange and for the computation of the stress tensor. This implementation would make the convergence of forces from hybrid-functionals and stresses from any functional even faster, making data generation more efficient.

Acknowledgements

We acknowledge Felix Hanke and Jürgen Wieferink, who have contributed at fundamental points in the implementation of molecular dynamics in FHI-aims, and Konstantin Lion, who has more recently worked on wave-function propagation schemes. MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233.

6.3 Electronic Structure Calculations with Quantum Nuclei

*Yosuke Kanai^{1,2}, Yair Litman^{3,4,a}, Songrui Liu¹, *Mariana Rossi⁵, Elia Stocco⁵, Jianhang Xu¹, Xingchen Zhang¹, and Ruiyi Zhou¹

¹Department of Chemistry, University of North Carolina at Chapel Hill, USA

²Department of Physics and Astronomy, University of North Carolina at Chapel Hill, USA

³Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Yusuf Hamied Department of Chemistry, Cambridge University, UK

⁵Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

*Coordinator of this contribution.

^aCurrent Address: Max Planck Institute for Polymer Research, Mainz, Germany

Summary

Electronic structure simulations are most commonly performed within the Born-Oppenheimer approximation (BOA) and under the assumption that nuclei are fixed point charges. However, accounting for quantum nuclear motion often goes beyond a simple correction to such simulations. Quantum nuclei delocalize, interfere, and tunnel through barriers. These effects strongly impact the thermodynamics and reactive properties of several materials. FHI-aims currently allows atomic nuclei to be treated as quantum-mechanical particles within the BOA, via path-integral methods, or going beyond the BOA through nuclear-electronic orbital (NEO) methods.

Path-integral molecular dynamics (PIMD) has emerged as a practical and accurate methodology to treat nuclear quantum effects, when the BOA holds. Proposed by Rahman and Parrinello [560], it relies on the molecular dynamics sampling of quantum-mechanical thermodynamical observables within the imaginary-time path integral (PI) representation of quantum mechanics. Approximations to time-dependent thermodynamic observables can also be obtained from this formalism. A tight integration of FHI-aims with the i-PI code [504, 561] allows advanced *ab initio* PI simulations to be performed.

The nuclear-electronic orbital method by Hammes-Schiffer and co-workers [407, 562-564] enables specified atomic nuclei (usually protons) to be treated on an equal footing as electrons in electronic structure calculations. NEO can be combined with various electronic structure methods including DFT [565, 566]. In FHI-aims, the NEO method is employed as a practical approach for implementing multicomponent DFT [567], which allows the atomic nuclei to be modeled as quantum-mechanical particles without the usual BOA.

Current Status of the Implementation

Exploiting the fact that the quantum time propagator is equal to the quantum density matrix $\exp(-\beta\hat{H})$, where $\beta = 1/(k_B T)$, at imaginary time $i\beta\hbar$, one can cast quantum thermodynamics in the Feynman path

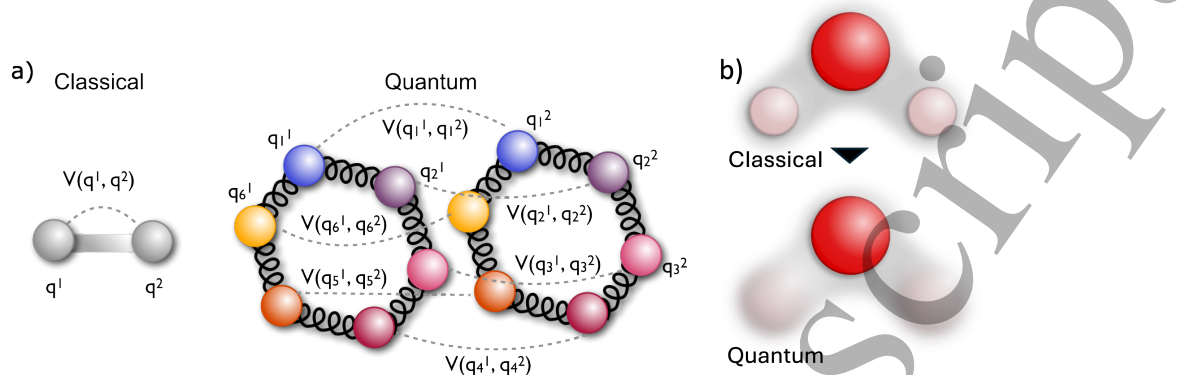


Figure 6.6: a) A depiction of a classical system of two particles interacting through a potential V and the corresponding quantum path-integral representation of the same system, considering that the particles are distinguishable. Each particle becomes a ring polymer. Each bead in the first particle interacts with the corresponding bead in the second particle through the same potential V . Nearest neighbor beads within a ring-polymer are connected by a harmonic spring. b) A water molecule described using the conventional *ab initio* approach and the corresponding nuclear-electronic orbital approach is shown. In the classical picture, both oxygen (red sphere) and hydrogen (pink spheres) atoms are treated as classical nuclei, while electrons (grey cloud) are quantum particles. In the NEO approach, protons (pink cloud) are also treated as quantum particles, at the same level of theory as electrons.

integral formalism. The canonical partition function $Z = \text{Tr}[\exp^{-\beta\hat{H}}]$ can be calculated by performing a Trotter factorization of the trace in the position representation, producing,

$$Z = \lim_{P \rightarrow \infty} \left(\frac{1}{2\pi\hbar} \sqrt{\frac{m}{m'}} \right)^P \int d\mathbf{R} d\mathbf{p} \exp \{-\beta_P H_P(\mathbf{p}, \mathbf{R})\} \quad (6.6)$$

where the m is the physical mass of the system, m' are fictitious masses for sampling, $\beta_P = \beta/P$ and P corresponds to the number of identities introduced to factorize the trace. The non-commuting nature of the position operator \hat{R} and momentum operators \hat{p} gives rise to a classical ring-polymer Hamiltonian H_P , consisting of P "beads" (system replicas) connected by harmonic springs. It is given by

$$H_P = \sum_{j=1}^P \left[\frac{p_j^2}{2m'} + \frac{m\omega_P^2}{2} (R_{j+1} - R_j)^2 + V[n, R_j] \right], \quad (6.7)$$

where, in *ab initio* PIMD based on DFT, $V[n; R_j]$ is the ground-state Born-Oppenheimer potential calculated at the (fixed) nuclear positions R_j of replica j . The spring frequency $\omega_P = P/(\beta\hbar)$ and $R_1 = R_{P+1}$. A pictorial representation of the classical and quantum system represented by ring polymers is given in Fig. 6.6a. The definition of the Hamiltonian in Eq. 6.7 allows one to evolve the nuclear positions following Newton's equations of motion and use molecular dynamics techniques (see Section 6.2) in order to sample the phase space.

The most advanced PIMD algorithms can reduce the cost of PI simulations by many orders of magnitude, making them almost as cheap as an *ab initio* simulation where nuclei are treated classically. These algorithms involve combinations of higher-order integrators, multiple-time-stepping algorithms, ring-polymer

Roadmap on Advancements of the FHI-aims Software Package

6.3. ELECTRONIC STRUCTURE CALCULATIONS WITH QUANTUM NUCLEI

contraction algorithms, and much more [561, 568–571]. Because PIMD and related PI methods largely rely on the BOA, it has been deemed most efficient to interface FHI-aims with a code in which a team of developers implements all the most modern algorithmic alternatives: the i-PI code [504, 561]. This interface is automatically compiled with FHI-aims and using it requires setting only a few keywords in the input file of FHI-aims, as detailed in the next section. FHI-aims provides the BO potential calculated from Kohn-Sham density-functional theory, together with the corresponding forces and stress, and i-PI evolves the equations of motion for the (quantum) nuclei with these quantities. Advantages of PIMD are that it can be applied to large and complex systems, it naturally captures quantum statistics from all nuclei, and it is exact for distinguishable particles, ensuring the accuracy of the results as long as the BOA holds.

Approximations to time-correlation functions can be obtained as approximations from this formalism and give access to system response properties such as vibrational spectra, thermal rate constants, heat transport, etc [572, 573]. The semiclassical instanton rate theory can also be cast in the same formalism [574, 575], usually referred to as a ring-polymer instanton rate theory and constitutes a method to calculate nuclear tunnelling splittings and tunnelling rates [576].

NEO: The NEO method is widely used to treat protons quantum-mechanically in electronic structure calculations while any other atomic nuclei can be made quantum in principle. It allows us to formulate a practical method for multicomponent DFT. Kohn-Sham (KS) Hamiltonians for electrons and protons are coupled via the classical electrostatic and also the electron-proton correlation functional as in

$$\hat{H}^e = -\frac{1}{2}\nabla_e^2 - \hat{V}_{\text{es}}^c(\mathbf{r}^e) - \hat{V}_{\text{es}}^p(\mathbf{r}^e) - \hat{V}_H^e(\mathbf{r}^e) + \frac{\delta E_{XC}^e[n^e]}{\delta n^e} + \frac{\delta E_{epc}[n^e, n^p]}{\delta n^e} \quad (6.8)$$

$$\hat{H}^p = -\frac{1}{2M^p}\nabla_p^2 + \hat{V}_{\text{es}}^c(\mathbf{r}^p) + \hat{V}_H^p(\mathbf{r}^p) + \hat{V}_{\text{es}}^e(\mathbf{r}^p) + \frac{\delta E_{XC}^p[n^p]}{\delta n^p} + \frac{\delta E_{epc}[n^e, n^p]}{\delta n^p} \quad (6.9)$$

where \hat{V}_{es}^c and $\hat{V}_{\text{es}}^{e/p}$ are the electrostatic potential from classical atomic nuclei and electron/protons, respectively. \hat{V}_H is the Hartree potential. E_{XC}^e , E_{XC}^p , and E_{epc} represent the exchange-correlation (XC) energy of electrons, the XC energy of protons, and the correlation energy between electron and proton, respectively. \mathbf{r}^e and \mathbf{r}^p represent the electron and proton coordinates. The KS eigenvalue equations thus must be solved together in a self-consistent fashion, and therefore the electron density is indirectly modified by the proton quantization, as discussed in Ref. [577]. In practice, we approximate E_{XC}^p with the Hartree-Fock exchange. It is motivated by the fact that the correlation among protons is negligible in most cases and using the exact exchange makes the protons to be free of the infamous self-interaction error, which would otherwise cause artificial delocalization of the quantum protons. Additionally, we implemented only Γ -point sampling of the Brillouin zone for the proton KS orbitals. For the electron-proton correlation functional, E_{epc} , several approximations have been proposed in literature. Currently, the EPC-17 and EPC-17-2 (LDA equivalent) [578, 579] are implemented in the FHI-aims code. For quantum protons, special Gaussian-type orbital basis sets have been developed [580] and FHI-aims code also employs them but using the algorithms described in Ref. [577].

In addition to treating protons quantum-mechanically in equilibrium, NEO method also allows us to study coupled quantum dynamics of protons and electrons via real-time time-dependent density functional theory (RT-TDDFT) and via Ehrenfest method, which are discussed in Section 4.5. The underlying dynamics of the mixed classical-quantum system is given by the Lagrangian

6.3. ELECTRONIC STRUCTURE CALCULATIONS WITH QUANTUM NUCLEI

$$\begin{aligned}
L^{\text{NEO}}(t) = & \int d\mathbf{k} \sum_n \langle \psi_{n,\mathbf{k}}^e(t) | i \frac{\partial}{\partial t} + \frac{1}{2} \nabla_e^2 | \psi_{n,\mathbf{k}}^e(t) \rangle + \sum_n \langle \psi_n^p(t) | i \frac{\partial}{\partial t} + \frac{1}{2M^p} \nabla_p^2 | \psi_n^p(t) \rangle \\
& - E_H[n^e] - E_H[n^p] - E_{XC}^e[n^e] - E_{XC}^p[n^p] \\
& + \int d\mathbf{r}^e d\mathbf{r}^p \frac{1}{|\mathbf{r}^e - \mathbf{r}^p|} n^e(\mathbf{r}^e, t) n^p(\mathbf{r}^p, t) - E_{epc}[n^e, n^p] \\
& + \sum_I \frac{1}{2} M_I \dot{\mathbf{R}}_I^2(t) - \sum_{I < J} \frac{Z_I Z_J}{|\mathbf{R}_I(t) - \mathbf{R}_J(t)|} \\
& + \int d\mathbf{r}^e n^e(\mathbf{r}^e, t) \sum_I \frac{Z_I}{|\mathbf{r}^e - \mathbf{R}_I(t)|} - \int d\mathbf{r}^p n^p(\mathbf{r}^p, t) \sum_I \frac{Z_I}{|\mathbf{r}^p - \mathbf{R}_I(t)|},
\end{aligned} \tag{6.10}$$

where $\mathbf{R}_I(t)$ and Z_I are the position coordinates and the charge, respectively, of classical nucleus I . The upper dot denotes the time derivative, and N^c is the total number of classical atomic nuclei. By applying the variational principle to this NEO action, Euler-Lagrange equation allows us to derive necessary equations of motion (EOM) for quantum (electrons and protons) and classical (non-proton atomic nuclei) degrees of freedom, allowing us to formulate RT-TDDFT and Ehrenfest Dynamics in the context of multicomponent DFT formalism [408, 581]. The approach is particularly convenient for studying large complex condensed matter because one can select which parts of the system are treated with quantum dynamics and which parts are governed by the classical equations of motion (see Section 6.2).

Additionally, the constrained NEO (cNEO) method [582] has been also recently implemented in FHI-aims [583]. cNEO constrains the expectation value of the quantum nuclear position operator at specific points in space. This approach effectively constrains the centers of the quantum nuclear probability densities to designated positions in real space, allowing NEO electronic structure calculations with controlled nuclear localization and providing a straightforward framework to defining nuclear equations of motion for the constrained position and momenta. [584] Its connection to the quantum-corrected effective potential, as usually formulated using quantum-mechanical path integrals [585], is discussed in Ref. [583].

Usability and Tutorials

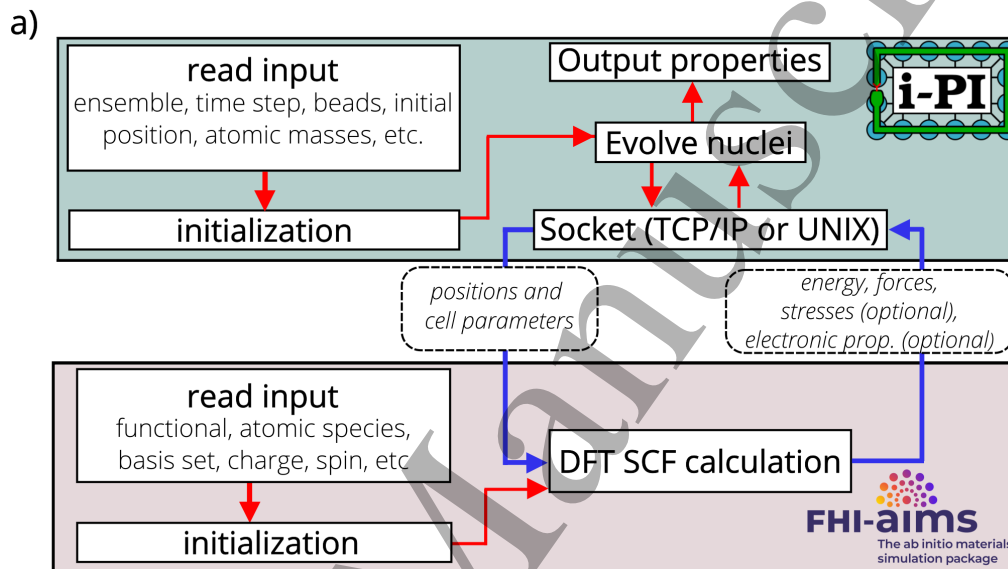
We begin by considering the inclusion of NQEs within the BOA. For this purpose, a server-client paradigm is adopted, where i-PI takes the role of server and FHI-aims serves as a client (see Fig. 6.7a). The communication between both codes is established through UNIX or TCP/IP sockets. The latter allows for communication across different nodes as it might be required when running in high-performance computing (HPC) facilities. The framework enables trivial parallelization over beads by running as many instances of FHI-aims as there are beads in the simulation. As shown in Fig. 6.7b, the FHI-aims input file requires minimal additions to specify the type and port address of the socket, as well as the communication of properties beyond energy, forces and stresses. At the moment it is possible to communicate polarizabilities, Hirshfeld charges, atomic stresses, dipole moments and electronic friction, some of which can be used by i-PI to determine nuclear dynamics.

A tutorial that describes how to set up an *ab initio* molecular dynamics simulation using FHI-aims and i-PI

Roadmap on Advancements of the FHI-aims Software Package

6.3. ELECTRONIC STRUCTURE CALCULATIONS WITH QUANTUM NUCLEI

is available [under this link^c](#). The tutorial covers the basics of performing these simulations, such as describing how the codes communicate, setting up the simulation, determining the time-step and assessing the thermostat. It also covers advanced topics like anharmonic free energy calculations and constant pressure simulations. The combination of FHI-aims with i-PI has enabled studies on a wide range of systems and topics including metallic interfaces [570, 586–588], molecular crystals [589, 590], nuclear quantum tunnelling in gas-phase molecules [591–593], quantum dissipative dynamics in metals [594, 595], biomolecules [596, 597] and low-dimensional materials [571, 598].



b)

in `control.in`:

```

use_pimd_wrapper <host> <port>
communicate_pimd_wrapper dipole
communicate_pimd_wrapper polarizability
communicate_pimd_wrapper friction <energy>
communicate_pimd_wrapper atomic_stress

```

Figure 6.7: a) A schematic representation of the communication between FHI-aims and i-PI. Once a connection is established, the simulation loop proceeds as follows: i) i-PI sends the nuclear positions and cell parameters to FHI-aims; ii) FHI-aims computes the necessary properties, such as energies and forces, and communicates them back; and iii) i-PI updates the atomic positions using the received data, thus restarting the loop. The red and blue arrows show intra-code and inter-code flow of data, respectively. b) A snippet of the FHI-aims input file showing the keywords that manage communication with i-PI. In this example, FHI-aims communicates the dipole moment, polarizability, electronic friction tensor, and atomic stress to i-PI.

The NEO tutorial, which can be accessed at [this link^d](#), provides a brief overview of the NEO theory along with several simple examples for running NEO-DFT, RT-NEO-TDDFT simulations in FHI-aims code. The first part of

^c<https://fhi-aims-club.gitlab.io/tutorials/molecular-dynamics-with-i-pi/>

^d<https://fhi-aims-club.gitlab.io/tutorials/NEO/>

Roadmap on Advancements of the FHI-aims Software Package

6.3. ELECTRONIC STRUCTURE CALCULATIONS WITH QUANTUM NUCLEI

the tutorial focuses on standard NEO-DFT calculation, outlining all the necessary parameters and explaining how to calculate the ground state with quantized protons for both isolated systems and extended periodic systems. It also has an example for performing NEO 'geometry optimizations' in which the centers of proton basis sets are also optimized. For protons, a number of Gaussian-type basis functions have been specifically designed as detailed in Ref. [580]. The second part of the tutorial is focused on performing RT-NEO-TDDFT simulation for studying coupled quantum dynamics of electrons and protons. The case of an excited state proton transfer in a small organic molecule is used as an example.

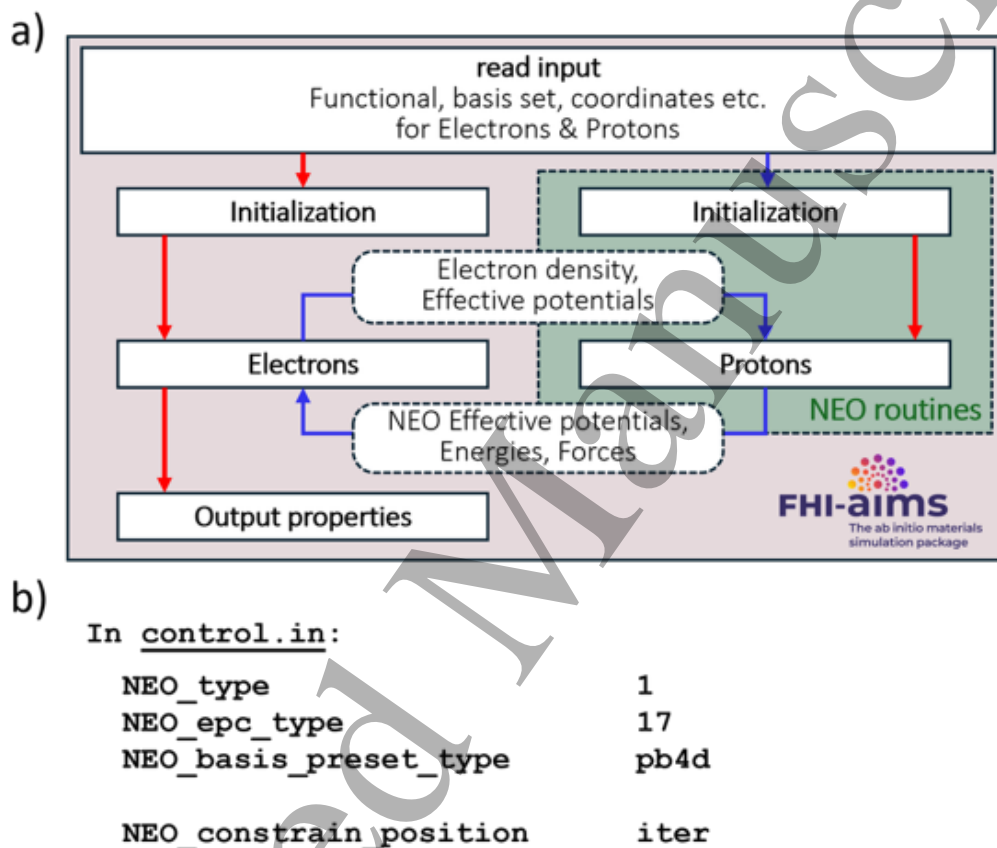


Figure 6.8: a) A schematic representation of the implementation of the NEO method in FHI-aims. NEO create a quantum subsystem for protons. The initialization and SCF cycles are done independently. Only necessary information is passed between the electron system and the proton system. The red and blue arrows show intra-system and inter-system flow of data, respectively. b) A snippet of the FHI-aims input file showing the keywords for running NEO simulations. The keyword in the last line is for cNEO calculations.

Future Plans and Challenges

Ab initio PIMD has been extensively and successfully employed in material simulations for decades. However, at best, these simulations carry the same cost as *ab initio* molecular dynamics (AIMD) and often they are at least one order of magnitude more expensive. With the development of the last-generation machine-

6.3. ELECTRONIC STRUCTURE CALCULATIONS WITH QUANTUM NUCLEI

learning interatomic potentials (MLIPs), the usage of Born-Oppenheimer AIMD and *ab initio* PIMD is becoming restricted as data-generation tool to train or refine these potentials. MLIPs currently offer 5-7 orders of magnitude gain in the speed of force evaluations when compared to DFT, with minimal accuracy loss.

Nevertheless, it has been repeatedly shown that it is necessary to include data from PIMD simulations to train MLIPs when one is targeting an accurate description of nuclear quantum effects. Therefore, we expect that the necessity of *ab initio* PIMD simulations will continue to exist, and a better integration of i-PI with FHI-aims through MPI communicators is desirable. That would remove the need for socket communication and would make the infrastructure more robust and easier to use in high-performance architectures. Going beyond the BOA with path-integral based approximations is a topic of forefront research [599]. We expect that future methods that follow this approach will directly profit from these implementations.

The NEO method has become an exciting avenue for treating atomic nuclei quantum-mechanically on an equal footing as electrons in electronic structure calculations over the last several years. The method is actively developed in the community, and new functionalities will be added also to the FHI-aims code in the near future. Because quantum nuclei are represented by Gaussian functions centered on a particular point in real space, one needs to ensure that enough proton basis functions are present such that the quantum dynamics of protons can be modeled accurately. In order to overcome this inconvenience, the idea of Ehrenfest RT-NEO-TDDFT has been introduced[581], and it is under active development. Another theoretical challenge is the development of a correlation functional for protons. While its effect is insignificant in most molecular/materials systems, such an effect might induce interesting behavior when protons are situated very closely such as under extremely high pressure.

Acknowledgements

M.R. acknowledges Eszter S. Pócs and Karen Fidanyan for essential contributions to the tutorial regarding the use of FHI-aims with i-PI.

6.4 Electron-Phonon Coupling and Electronic Friction

Connor L. Box¹, George Trenins², Honghui Shang³, Yair Litman^{2,4,a}, Mariana Rossi², and *Reinhard J. Maurer^{1,5,6}

¹Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

²Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

³Key Laboratory of Precision and Intelligent Chemistry, University of Science and Technology of China, Hefei, China

⁴Yusuf Hamied Department of Chemistry, Cambridge University, UK

⁵Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

⁶Faculty of Physics, University of Vienna, Vienna A-1090, Austria

*Coordinator of this contribution.

^aCurrent Address: Max Planck Institute for Polymer Research, Mainz, Germany

Summary

The electron-phonon interaction manifests in many physical properties of materials, such as their thermodynamic, transport and superconducting properties. Particularly for metals, phonons effectively couple to low-energy electron-hole pair excitations. In these situations, standard molecular dynamics techniques neglect the energy transfer mechanism between lattice vibrations and electronic excitations, and can provide qualitatively incorrect dynamical properties. [600] When nuclear motion can be considered classical, this energy transfer can be accounted for through molecular dynamics with electronic friction [601], which is evaluated in a Langevin framework:

$$M\ddot{R}_{a\kappa} = -\frac{\partial V(\mathbf{R})}{\partial R_{a\kappa}} - \sum_{a'\kappa'} \Lambda_{a\kappa,a'\kappa'} \dot{R}_{a'\kappa'} + \mathcal{R}_{a\kappa}(t), \quad (6.11)$$

where an atom with index a , mass M and position \mathbf{R} at time t experiences forces from the adiabatic potential energy V , a friction force from the electronic friction tensor Λ and a random force \mathcal{R} . The index κ indicates the 3 Cartesian directions x , y , and z . The Markovian friction tensor, Λ , describes the energy dissipation that arises due to electron-phonon coupling in the quasi-static (zero-frequency) limit. The random force establishes detailed balance between the electronic bath and the atoms. FHI-aims provides an infrastructure to evaluate the electronic friction tensor, Λ , and more generally, the electron-phonon-induced vibrational lifetime (or relaxation rates) based on time-dependent perturbation theory for both aperiodic and periodic systems. The friction module was written by Reinhard J. Maurer, and released in 2016. [602] This was re-written by Connor L. Box in 2021, improving parallelism, allowing direct output of electron-phonon coupling matrix elements and completing Honghui Shang's previous efforts in interfacing to the DFPT infrastructure in FHI-aims. [457, 460] Additional improvements were made by George Trenins in 2024 to improve numerical stability.

This module has previously supported research efforts to investigate nonadiabatic energy loss of atoms and

Roadmap on Advancements of the FHI-aims Software Package

6.4. ELECTRON-PHONON COUPLING AND ELECTRONIC FRICTION

molecules scattering from metal surfaces [603, 604], the interplay of nonadiabatic effects and quantum effects in hydrogen diffusion in bulk metals [594], and the phonon lifetimes of molecular overlayers on metal surfaces. [457, 602] Machine learning representations of the FHI-aims-predicted Markovian electronic friction calculated have also been proposed and successfully employed [467, 605].

Current Status of the Implementation

For the theory of electron-phonon coupling calculations, the interested reader is referred to Ref [378], for the specifics of the implementation within FHI-aims, we suggest Ref [457]. Considering a phonon of frequency $\omega_{q\nu}$ and mass M_ν , characterized by two quantum numbers, namely the band index, ν , and the momentum vector, \mathbf{q} , and electronic states, characterized by band indices m, n , and momentum \mathbf{k} , the EPC matrix element is given by:

$$g_{mn\nu}(\mathbf{k}, \mathbf{q}) = \left(\frac{\hbar}{2M_\nu\omega_{q\nu}} \right)^{1/2} \tilde{g}_{mn\nu}(\mathbf{k}, \mathbf{q}), \quad (6.12)$$

with

$$\tilde{g}_{mn\nu}(\mathbf{k}, \mathbf{q}) = \langle m\mathbf{k} + \mathbf{q} | \partial_{\mathbf{q}\nu} V | n\mathbf{k} \rangle. \quad (6.13)$$

The EPC matrix elements describe the excitation of an electron from a state $n\mathbf{k}$ to a state $m\mathbf{k} + \mathbf{q}$ by absorption of a phonon $\mathbf{q}\nu$. V in Eq. 6.13 is the self-consistent ("screened") effective potential from a Kohn-Sham DFT calculation. These screened EPC elements can be output using ELSI infrastructure for post-processing. They are evaluated in NAO basis and Cartesian framework as:

$$\tilde{g}_{mn}^{a\kappa}(\mathbf{k}, \mathbf{q}) = \sum_{ij} \left(C_{m\mathbf{k}+\mathbf{q}}^j \right)^* C_{n\mathbf{k}}^i \left(H_{ij}^{a\kappa, (1)}(\mathbf{k}, \mathbf{q}) - \epsilon_{n\mathbf{k}} S_{ij}^{a\kappa, L}(\mathbf{k}, \mathbf{q}) - \epsilon_{m\mathbf{k}+\mathbf{q}} S_{ij}^{a\kappa, R}(\mathbf{k}, \mathbf{q}) \right), \quad (6.14)$$

furthermore, an approximation, proposed by Head-Gordan and Tully, [606] can be applied for the term in response matrices:

$$C_{ij}^{a\kappa, \text{HGT}}(\mathbf{k}, \mathbf{q}) = \left(H_{ij}^{a\kappa, (1)}(\mathbf{k}, \mathbf{q}) - \epsilon_F S_{ij}^{a\kappa, (1)}(\mathbf{k}, \mathbf{q}) \right), \quad (6.15)$$

where $H^{(1)}$ ($S^{(1)}$) is the first order Hamiltonian (overlap) response to the atomic coordinate perturbation, and L(R) superscripts refer to the one-sided left (right) derivatives of the overlap matrix such that $S^{(1)} = S^L + S^R$. In FHI-aims, this approximation is applied by default but can be switched off by the user. Its effect on the EPC matrix elements has previously been assessed. [457, 602] FHI-aims supports several different variations of the electronic friction tensor, by default, the following expression is used:

$$\Lambda_{a\kappa, a'\kappa'}^{\mathbf{q}, \text{Ib}}(\hbar\omega) = \pi\hbar \sum_{\sigma mn} \int \frac{d\mathbf{k}}{\Omega_{BZ}} \tilde{g}_{mn}^{a\kappa}(\mathbf{k}, \mathbf{q}) (\tilde{g}_{mn}^{a'\kappa'})^*(\mathbf{k}, \mathbf{q}) (f_{n\mathbf{k}} - f_{m\mathbf{k}+\mathbf{q}}) \frac{\delta(\epsilon_{m\mathbf{k}+\mathbf{q}} - \epsilon_{n\mathbf{k}} - \hbar\omega)}{\epsilon_{m\mathbf{k}+\mathbf{q}} - \epsilon_{n\mathbf{k}}}. \quad (6.16)$$

The friction tensor can be used in MDEF dynamics directly through on-the-fly evaluation [603] or via machine-learning models, [467, 604] or projected along normal modes (e.g calculated by ASE [43]) to calculate phonon lifetimes or linewidth broadenings. [457, 602]

The current implementation supports ScaLAPACK-type distribution of the first-order response and EPC matrices. The response matrices can be evaluated using finite-difference or DFPT (see Sec. 5.1). Collinear spin is also supported. A restart mechanism exists that enables the reading and writing of first-order response

matrices to restart electronic friction calculations or to calculate the response atom by atom before evaluating the full friction tensor. The implementation is currently limited to phonon momenta of zero ($\mathbf{q} = 0$), consequently, $\mathbf{q} > 0$ is available only through post-processing with supercells. However, the code is highly scalable and efficient, enabling the evaluation of EPC for unit cells containing > 250 atoms and several k -points. For example, Box et al., employed the code to evaluate the Brillouin-zone averaged vibrational linewidth of the internal stretch vibration of a carbon monoxide layer on a Cu(100) surface. [457]

The current code has been benchmarked against plane-wave approaches in Ref [457], the interested reader is directed there for a more detailed benchmark and scaling performance.

Usability and Tutorials

The aforementioned features have been comprehensively documented in the FHI-aims manual since release 240507. All keywords related to the friction calculations are prefixed with `friction_`, with the exception of `calculate_friction`. This keyword accepts the options `numerical_friction` (for finite-difference evaluation of the electron-phonon coupling matrix elements) or `DFPT` (for density functional perturbation theory evaluation of the electron-phonon coupling matrix elements) and initiates the friction calculation.

An online tutorial utilizing the friction module is available. This tutorial employs the electronic friction driver to calculate electron-phonon couplings and vibrational lifetimes, including the effect of non-adiabatic effects via electronic friction for a molecule adsorbed on a metallic surface. Figure 1 presents the landing page and a brief description of this tutorial. The tutorials can be accessed at <https://fhi-aims-club.gitlab.io/tutorials/tutorials-overview/>.

Electron-Phonon Coupling and Electronic Friction

Prepared by Connor L. Box, Mariana Rossi and Reinhard J. Maurer

Summary of the tutorial

The tutorial is organised as follows:

Part 1 introduces theory of electron-phonon coupling and electronic friction

Part 2 calculates the vibrational lifetime of CO adsorbed on a Cu(100) surface

Figure 6.9: Landing page for electronic friction/electron-phonon coupling tutorial.

Future Plans and Challenges

Future work focused on expanding the capabilities of the code, will be aimed at expanding the properties that can be calculated, either within FHI-aims or by interfacing to other electron-phonon coupling codes

*Roadmap on Advancements of the FHI-aims Software Package***6.4. ELECTRON-PHONON COUPLING AND ELECTRONIC FRICTION**

such as Perturbo. [511] For example, the calculation of the Hessian/dynamical matrix to evaluate linewidths requires post processing currently, this can be circumvented by interfacing to existing parts of FHI-aims that calculates these properties. Another example is the evaluation of the Eliashberg function, which is not yet implemented within the friction module. Support for $q > 0$ calculations is desirable, and some existing DFPT routines are available for both reciprocal-space and supercell approaches. Finally, use `local_index` is currently not supported, changing this would improve efficiency and memory usage for systems with hundreds of atoms, comparable to the effect for ground-state SCF.

The combination of the electronic friction formalism with quantum nuclear dynamics is a topic of current research. [594, 607] A connection of this formalism with instanton theory to describe thermal rates of nuclear tunneling reactions has been derived and implemented in FHI-aims and i-PI. [608] In the future, we plan extensions of this interface to include MDEF and new algorithms joining path-integral molecular-dynamics approximations to thermal rates and electronic friction.

Acknowledgements

We acknowledge financial support through the UKRI Future Leaders Fellowship programme (MR/S016023/1 and MR/X023109/1), and a UKRI frontier research grant (EP/X014088/1). We acknowledge the contribution of Eszter S. Pócs for her work in the implementation of an interface for the friction tensor output to the i-PI code.

 6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

6.5 Temperature-Dependent Electronic Spectral Functions from Band-Structure Unfolding

***Jingkai Quan**^{1,2}, **Min-Ye Zhang**^{1,3}, **Nikita Rybin**^{1,a}, **Marios Zacharias**^{4,5}, **Xinguo Ren**^{3,1,b}, **Hong Jiang**⁶, **Matthias Scheffler**¹, and ***Christian Carbogno**^{1,7,c}

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Max-Planck Institute for the Structure and Dynamics of Matter, Luruper Chausse 149, 22761, Hamburg, Germany

³Institute of Physics, Chinese Academy of Sciences, Beijing 100190, China

⁴Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵Computation-Based Science and Technology Research Center, The Cyprus Institute, Aglantzia, 2121, Cyprus

⁶Beijing National Laboratory for Molecular Sciences, College of Chemistry and Molecular Engineering, Peking University, 100871 Beijing, China

⁷Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

^a*Current Address:* Skolkovo Institute of Science and Technology, Bolshoi bulvar 30, build.1, 121205, Moscow, Russia

^b*Current Address:* Institute of Physics, Chinese Academy of Sciences, Beijing, 100190, China

^c*Current Address:* Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

The electronic band structure $\varepsilon_n(\mathbf{k})$, which describes the periodic dependence of the electronic quantum states n on the lattice momentum \mathbf{k} in reciprocal space, is one of the fundamental concepts in solid-state physics. It is key to our understanding of solid-state devices, since it allows rationalizing the electronic properties of periodic materials, e.g., to discern semiconductors with a direct band gap from those with an indirect one. In spite of that, the electronic band structure $\varepsilon_n(\mathbf{k})$ is actually only well-defined for static nuclei, i.e., immobile nuclei fixed at their crystallographic positions. This constitutes a severe approximation that does not even hold in the limit of zero Kelvin due to the quantum-nuclear zero-point motion. To account for these thermodynamics effects, the band-structure concept can be generalized by introducing a temperature-dependent (T) spectral-function $\langle A(\mathbf{k}, E) \rangle_T$, as shown in Fig. 6.10. In this case, the electronic quantum states at each reciprocal-vector \mathbf{k} are no longer sharp values $\varepsilon_n(\mathbf{k})$, but are characterized by a finite-width distribution. Several fundamental physical properties and mechanisms can only be understood and computed when the coupling between nuclear and electronic degrees of freedom is accounted for. For instance, this includes the temperature-dependence of key electronic properties such as the band gap [609–612], optical absorption spectra [613, 614], and electronic transport coefficients [354, 615, 616],

6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

also see Contrib. 7.2.

One possible route to compute spectral functions and the associated observables is many-body perturbation theory, a formally elegant and computationally efficient framework for treating electron-phonon coupling that is applicable within and beyond the Born-Oppenheimer approximation [378]. However, both the dynamics of the nuclei and that of the electrons are treated approximately in these methods by using the harmonic approximation for the nuclear degrees of freedom, also see Contribs. 6.1 and 7.1, and expressing the electronic response (including some second-order effects [378]) in terms of linear-order electron-phonon coupling elements. These approximations may well fail at elevated temperatures and/or for mobile atoms. For instance, it has been demonstrated that the complete concept of phonons can break down in systems exhibiting spontaneous defect formation, even if these defects are short-lived [533]. To avoid potential inaccuracies from the aforementioned approximations, the electronic spectral function can be equally obtained in a non-perturbative fashion within the Born-Oppenheimer approximation, hence capturing higher-order couplings between electronic and vibrational degrees of freedom. To this end, *ab initio* molecular dynamics (*aiMD*) can be run to accurately sample the full anharmonic potential-energy surface. The spectral-function is then obtained as the thermodynamic average $\langle A(\mathbf{k}, E) \rangle_T$ of the instantaneous electronic-band structures $\varepsilon_n(\mathbf{k})$ observed during this dynamics [617–620].

From an electronic-structure point of view, one major hurdle in the implementation of the non-perturbative method arises from the fact that accurately sampling the dynamics in solids requires extended supercells, see Contrib. 6.1. However, the electronic band-structures' topology is fundamentally connected to the translational symmetries of a crystal, which results in band-structure folding in supercell electronic-structure theory calculations. In this case, electronic states associated to different \mathbf{k} -vectors in the first Brillouin zone are mapped onto the same \mathbf{K} -vector in the reduced Brillouin zone when extended supercells are used in real space, see Fig. 6.11. Accordingly, the uttermost topological information is lost if the spectral function is computed as simple average over folded supercell band structures. Rather, it is necessary to first recover the representation in the first Brillouin zone, a process often referred to as *unfolding* [621–625] and visualized in Fig. 6.11. In this contribution, we shortly describe the implementation of the band-structure unfolding technique in the electronic-structure theory pack-

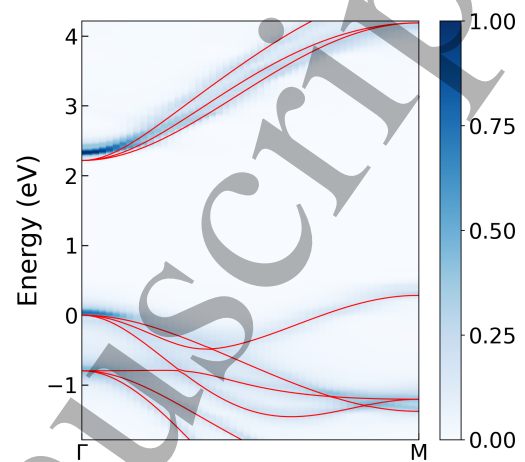


Figure 6.10: Electronic spectral function at 300K and the static band structure (red lines) of SrTiO₃.

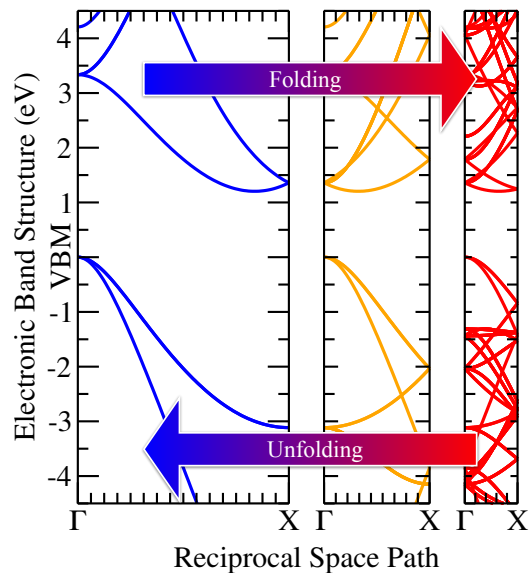


Figure 6.11: Folding viz. unfolding visualized for the HSE06 band structure of silicon along the high-symmetry $\Gamma \rightarrow X$ path for the primitive unit cell (left), an eight-atom cubic conventional cell (middle), and a 64-atom cubic supercell (right).

6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

age FHI-aims and the updates made since its original development [617].

Current Status of the Implementation

The main difficulty in the implementation of band-unfolding techniques in FHI-aims stems from the usage of a non-orthogonal, atomic-centered basis. In this numeric atomic-orbitals (NAO) basis, atom displacements also affect the basis functions, so that changes in the basis set and in the overlap matrix need to be incorporated in the unfolding weight derivation [617]. We discuss the unfolding technique by establishing a relationship between the electronic structure obtained for a primitive cell (PC) and the one obtained for a supercell (SC). Notationwise, properties associated to the PC or SC are denoted by using lower- and upper-case letters, respectively, and/or PC and SC subscripts when needed for clarity. Accordingly, the PC is characterized by the set of lattice vectors $\mathbf{a} = (\mathbf{a}_1, \mathbf{a}_2, \mathbf{a}_3)$ and the supercell (SC) by the lattice vectors $\mathbf{A} = \mathbf{a} \cdot \mathbf{M}$. Here, the lattice vectors \mathbf{a} and \mathbf{A} are column vectors, as by FHI-aims convention, and the \mathbf{M} is a non-singular matrix with integer entries, implying that the volume of the SC is $m = \|\det(\mathbf{M})\|$ times larger than that of the PC. Similarly, the lattice vectors \mathbf{b} of the first Brillouin zone (BZ) associated with the PC and the ones of the reduced BZ associated with the SC are related via $\mathbf{B} = \mathbf{M}^{-1} \mathbf{b}$. In turn, the volume of the reduced BZ is m times smaller than that of the first BZ, so that m different \mathbf{k} -vectors in the first BZ zone are mapped onto one and the same \mathbf{K} -vector in the reduced BZ. The unfolding technique reverses this mapping and re-establishes a representation in the first BZ.

To perform the mapping, the projection operator

$$\mathbf{P}_{\mathbf{k}} = |\mathbf{k}\rangle \langle \mathbf{k}| \quad (6.17)$$

is used. Here, $|\mathbf{k}\rangle$ are the eigenvectors with eigenvalue $\exp(i\mathbf{k} \cdot \mathbf{a})$ that solve the eigenvalue problem^e

$$\mathbf{t} |\mathbf{k}\rangle = \exp(i\mathbf{k} \cdot \mathbf{a}) |\mathbf{k}\rangle \quad (6.18)$$

for the translational operators \mathbf{t} associated to the lattice vectors \mathbf{a} . With that, it is possible to obtain the weights

$$W_{\mathbf{KN}}^{\mathbf{k}} = \langle \Psi_{\mathbf{KN}} | \mathbf{P}_{\mathbf{k}} | \Psi_{\mathbf{KN}} \rangle = |\langle \mathbf{k} | \Psi_{\mathbf{KN}} \rangle|^2, \quad (6.19)$$

i.e., the contribution stemming from the subspace spanned by $|\mathbf{k}\rangle$ associated with the translational symmetry of the PC to the SC wave function $\Psi_{\mathbf{KN}}$. For a “perfect” SC, i.e., one consisting of identical PC replicas, these weights are either 0 or 1, and one can hence reconstruct the exact band structure in the first BZ from a SC calculation. For a disturbed SC, the weights $W_{\mathbf{KN}}^{\mathbf{k}}$ become fractional, since the SC system is not invariant under PC-translations \mathbf{t} . Accordingly, the desired remapping can be obtained by summing over all SC wave functions $\Psi_{\mathbf{KN}}$ and computing the electronic spectral function

$$A(\mathbf{k}, E) = \sum_{\mathbf{KN}} W_{\mathbf{KN}}^{\mathbf{k}} \delta(E - E_{\mathbf{KN}}). \quad (6.20)$$

In practice, the unfolding implementation in FHI-aims start from the representation of the wave func-

^eFor the sake of clarity, the derivation here assumes non-degenerate eigenvalues. A discussion of the more general case including degeneracy can be found in Ref. [620].

6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

tion $\Psi_{\mathbf{K}N} = \sum_i C_{Ni}(\mathbf{K})\chi_i(\mathbf{K})$ in the SC, where $C_{Ni}(\mathbf{K})$ denotes a Kohn-Sham expansion coefficient and $\chi_i(\mathbf{K})$ is a Bloch-like basis function build from numeric atomic orbitals. In this SC Bloch-like basis, the algebraic representation of the PC translational operator is then constructed via $t_{ij} = \langle \chi_i(\mathbf{K}) | \mathbf{t} | \chi_j(\mathbf{K}) \rangle$. Let us emphasize that these functions are not orthogonal in real space, i.e., the overlap matrix $S_{ij}(\mathbf{K}) = \langle \chi_i(\mathbf{K}) | \chi_j(\mathbf{K}) \rangle \neq \delta_{ij}$ is not diagonal. Accordingly, Eq. (6.18) becomes a generalized eigenvector problem in this representation, also see Contrib. 2.4. Its solution yields the eigenvalues $\exp(i\mathbf{k} \cdot \mathbf{a})$, i.e., those set of \mathbf{k} -points in the first BZ that this \mathbf{K} -point in the reduced BZ can be mapped to, and the set of eigenvectors $\mathbf{k} = \sum_i F_i(\mathbf{k})\chi_i(\mathbf{K})$ that are needed for constructing the projection operator $\underline{\mathbf{P}}_{\mathbf{k}}$ defined in Eq. (6.17). With that one obtains the following formula for the weight

$$W_{\mathbf{K}N}^{\mathbf{k}} = |\mathbf{F}^\dagger(\mathbf{k})\underline{\mathbf{S}}_{\mathbf{K}}\mathbf{C}_N(\mathbf{K})|^2 = |\mathbf{F}'^\dagger(\mathbf{k})\mathbf{C}'_N(\mathbf{K})|^2. \quad (6.21)$$

In the last step, we have introduced the orthogonal representation $\mathbf{C}'_N(\mathbf{K}) = \underline{\mathbf{S}}_{\mathbf{K}}^{1/2}\mathbf{C}_N(\mathbf{K})$ and $\mathbf{F}'(\mathbf{k}) = \underline{\mathbf{S}}_{\mathbf{K}}^{1/2}\mathbf{F}(\mathbf{k})$ via a Löwdin transformation. Although the square root calculation $\underline{\mathbf{S}}_{\mathbf{K}}^{1/2}$ constitutes a computational overhead, the latter orthogonal representation is advantageous, since this allows to find analytical expressions of the eigenvector expansion coefficients $\mathbf{F}'(\mathbf{k})$ at each \mathbf{K} -point in the reduced BZ [620]. With that the numerical solution of the eigenvalue problem in Eq. (6.18) is circumvented, leading to an overall speed-up of the procedure. Eventually, let us note that translations in three-dimensional systems form an Abelian group, so that the translations \mathbf{t}_α associated to different lattice vectors \mathbf{a}_α can be tackled consecutively.

Usability and Tutorials

The overall workflow for computing temperature-dependent spectral functions $\langle A(\mathbf{k}, E) \rangle_T$ is sketched in Fig. 6.12. First, one or multiple aiMD runs are performed in a SC at temperature T to sample the thermodynamic phase-space. From these trajectories, L single, uncorrelated “samples”, i.e., atomic configurations $\mathbf{R}_I^{\text{SC}}(t_l)$ at sufficiently distant times t_l , are extracted and their electronic band-structure $\varepsilon_N(\mathbf{K}, t_l)$ is computed at a sufficiently dense \mathbf{K} -grid. In this step, the unfolding routines are used to map back the band-structure $\varepsilon_N(\mathbf{K}, t_l)$ onto the first BZ, i.e., the weights $W_{\mathbf{K}N}^{\mathbf{k}}(t_l)$ are computed and stored on file together with $\varepsilon_N(\mathbf{K}, t_l)$. In the last steps, the outputs are post-processed by computing the spectral functions $A(\mathbf{k}, E, t_l)$ of the individual samples, see Eq. (6.20), and by eventually computing the thermodynamic average

$$\langle A(\mathbf{k}, E) \rangle_T = \frac{1}{L} \sum_{l=1}^L A(\mathbf{k}, E, t_l). \quad (6.22)$$

A more detailed description including scripts to pre- and post-process the calculations and to analyze the data are

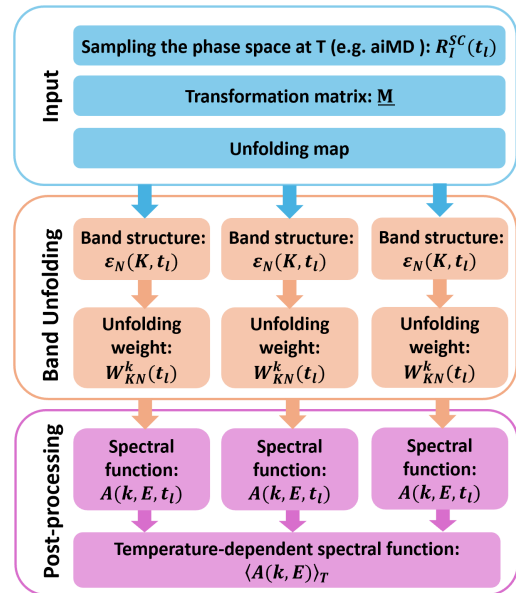


Figure 6.12: Schematics of a typical workflow used to obtain temperature-dependent spectral functions $\langle A(\mathbf{k}, E) \rangle_T$.

6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

given in the tutorial at <https://fhi-aims-club.github.io/tutorials/band-unfolding>. In this context, let us note that the method is not strictly restricted to *ai*MD, but any kind of sampling method can be used to explore the relevant phase-space. For instance, path-integral MD, see Contrib. 6.3 can be used instead, if it is necessary and desirable to account for quantum-nuclear effects. Along this lines, also more approximative methods, e.g., harmonic sampling as implemented in FHI-vibes [461] and described in Contrib. 6.1 or MD on machine-learned interatomic potentials as described in Contrib. 8.2, can in principle be used. In such cases, however, correct spectral functions can only be obtained if the employed approximations hold, as discussed in the respective contributions.

In more detail, the unfolding procedure in FHI-aims is part of the native band-structure post-processing tool. Accordingly it is invoked by using the keyword `output band` to define the reciprocal-space path for the SC, i.e., the \mathbf{K} -points that shall be targeted and by setting the keyword `bs.unfolding` in the `control.in` file. Additionally, two more input files are required: `transformation_matrix.dat`, where one defines the transformation matrix $\underline{\mathbf{M}}$ between PC lattice vector $\underline{\mathbf{a}}$ and SC lattice vector $\underline{\mathbf{A}}$, and `unfolding_map.dat`, describing the mapping between atoms in the PC and the SC that is later used for constructing the translational operator t . The latter is an index map $I \rightarrow j$ that relates the atomic coordinates in an unperturbed supercell $\mathbf{R}_I^{\text{SC}} = \mathbf{R}_j^{\text{PC}} + \sum_{\alpha} n_{\alpha} \mathbf{a}_{\alpha}$ $n_{\alpha} \in \mathbb{Z}$ to the ones of the atoms \mathbf{R}_j^{PC} in a primitive cell. Upon completion, the unfolding weights for each SC \mathbf{K} -point are written into separate files named `unfold_k_###.out` using the same `NXY` format used in the standard band structure output.

Currently, the band-unfolding implementation in FHI-aims supports all Bravais lattice, all exchange-correlation functionals, and all type of transformation matrices $\underline{\mathbf{M}}$, including non-diagonal ones typically needed to map primitive fcc and bcc structures to conventional cubic supercells. Both support for LAPACK and ScaLAPACK is implemented, allowing for a trivially parallel parallelization over \mathbf{K} -points in the case of small systems requiring many \mathbf{K} -points (LAPACK) and for a block cyclic distribution and distributed linear algebra in the case of large systems with few \mathbf{K} -points (ScaLAPACK). With that, the band unfolding implementation in FHI-aims can routinely handle large supercells with small computational overhead. For instance, the unfolding only requires approx. 20% of the total runtime when computing a 4,096-atom Silicon with one \mathbf{k} -point, > 100,000 basis functions and > 40,000 states at the semi-local level of exchange-correlation.

Eventually, let us note that also *electron-electron* coupling, i.e., electronic many-body effects, can play an important role for spectral functions [626, 627]. To this end, the current implementation also supports the unfolding of G_0W_0 calculations. In this case, the complex-valued self-energy is treated as a correction to the orbital energy, while the wave function is assumed to remain as in the Kohn-Sham scheme. The unfolded spectral function can then be computed in close analogy to Eq. (6.20). The required weights are obtained at the Kohn-Sham level, while the delta-function $\delta(E - E_{\mathbf{KN}})$ is replaced by the spectral function obtained at the G_0W_0 level. By this means,

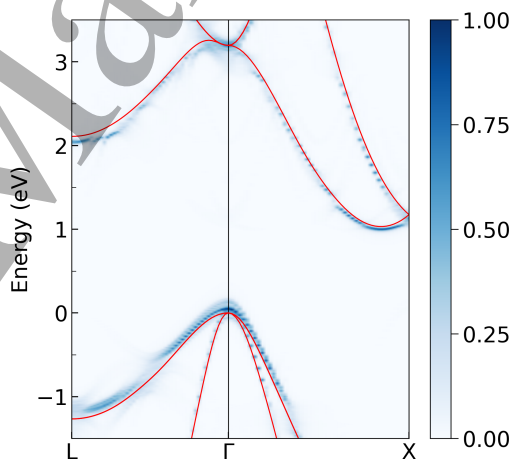


Figure 6.13: G_0W_0 spectral function (blue) at 300K for 64-atom silicon supercell unfolded onto the first Brillouin zone. The G_0W_0 band structure obtained in a PC is shown as comparison in red.

6.5. TEMPERATURE-DEPENDENT ELECTRONIC SPECTRAL FUNCTIONS FROM BAND-STRUCTURE UNFOLDING

electron correlation is explicitly accounted for. As an example, Fig. 6.13 shows the unfolded G_0W_0 band structure for silicon from 64-atom supercell configurations.

Future Plans and Challenges

As described in this contribution, the band-structure unfolding procedures implemented in FHI-aims enable a routine assessment of temperature-dependent spectral functions both at the density-functional theory and at the G_0W_0 level. From a workflow perspective, the presented methodology is independent on the actual methodologies used (i) for accurately sampling the thermodynamic phase-space and (ii) for obtaining the electronic band-structures $\varepsilon_N(\mathbf{K})$ in the SC, as detailed above. With that, the availability of (sufficiently accurate) machine-learned interatomic potentials [628], also see Contrib. 8.2, and of (sufficiently accurate) machine-learned electronic-structure theory [83, 337, 629], can give access to much larger system sizes, i.e., up to 10,000 atoms already to date, c.f. Contrib. 8.3. Performance-wise, already the current implementation can handle such cases. However, access to such larger system sizes also enables the study of new physical questions, e.g., the influence of point defects, of grain boundaries, and of solid-solid interfaces. While the overall theory holds also in such cases, the construction of the translational t operator will require case-specific adaptations to reflect the different types of breaking in translational symmetry.

Furthermore, the above-mentioned machine-learning methods might even help to solve a more fundamental open question with respect to non-adiabatic effects [630], which can play a fundamental role for spectral functions, as evidence from many-body perturbation theory shows [631]. So far, however, such kind of effects could not be studied with the herein presented non-perturbative approach, since it would require prohibitively expensive time-dependent electronic-structure theory calculations. With the advent of such machine-learning methods, this hurdle may fall and hence allow to study the connection between anharmonic and non-adiabatic effects.

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

7 Transport

Accepted Manuscript

7.1 Vibrational Heat Transport in Weakly, Moderately, and Strongly Anharmonic Solids

***Christian Carbogno**^{1,2,a}, **Florian Knoop**^{1,3}, ***Thomas A.R. Purcell**^{1,4,b}, **Shuo Zhao**¹, **Marcel F. Langer**^{1,5,6,c}, **Marcel Hülsberg**⁷, and ***Matthias Scheffler**¹

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

³Department of Physics, Chemistry and Biology (IFM), Linköping University, SE-581 83, Linköping, Sweden

⁴Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

⁵Machine Learning Group, Technische Universität Berlin, 10587 Berlin, Germany

⁶Laboratory of Computational Science and Modeling, Institut des Matériaux, École Polytechnique Fédérale de Lausanne, 1015 Lausanne, Switzerland

⁷Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

^a*Current Address:* Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

^b*Current Address:* Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

^c*Current Address:* Laboratory of Computational Science and Modeling, Institut des Matériaux, École Polytechnique Fédérale de Lausanne, 1015 Lausanne, Switzerland

Summary

Macroscopic heat transport in materials is characterized by the pressure- and temperature-dependent thermal conductivity tensor $\kappa(T, p)$, which relates heat flux \mathbf{J} and temperature gradient ∇T in Fourier's law $\mathbf{J} = -\kappa(T, p)\nabla T$. For insulators and semiconductors, the dominant contribution to κ stems from the vibrational motion of the atoms [632]. However, the harmonic phonon approximation that is commonly used for describing the motion in solids, see Contrib. 6.1, yields an infinite thermal conductivity by construction and is hence insufficient to capture heat transport, as first shown by Peierls [633]. To correctly model and compute thermal conductivities, it is thus quintessential to take into account anharmonic effects, i.e., deviations of the potential-energy surface (PES) from the approximative parabolic PES introduced in Contrib. 6.1.

From a bird's eye view, two different approaches to compute vibrational thermal conductivities exist and both are supported in FHI-aims. While *perturbative* methods inherently assume the validity of a harmonic quasi-particle picture, *non-perturbative* methods overcome this approximation and hence account for strongly anharmonic effects, e.g., defect formation and the exploration of multiple minima on the PES [533]. In *perturbative* methods, the PES is approximated by a truncated Taylor expansion. The terms up to second order, i.e., the *harmonic* approximation, are used to analytically describe the nuclear motion in terms of

7.1. VIBRATIONAL HEAT TRANSPORT IN WEAKLY, MODERATELY, AND STRONGLY ANHARMONIC SOLIDS

phonons with frequencies $\omega_s(\mathbf{q})$, in which s is the mode index and \mathbf{q} the reciprocal-space wave vector, see Contrib. 6.1. Higher-order terms describing anharmonicity are accounted for via perturbation theory under the assumption that they only make up for a minor correction of the PES. Conversely, *non-perturbative* methods do not make any approximations about the PES and hence require an explicit simulation of the nuclear dynamics, e.g., via *ab initio* molecular dynamics (*aiMD*), cf. Contrib. 6.2.

The validity of the quasi-particle picture and, with that, of the applicability of a perturbative approach can be rationalized by comparing phonon frequencies $\omega_s(\mathbf{q})$ and lifetimes $\tau_s(\mathbf{q})$ with respect to the Ioffe-Regel limit [635], as exemplified in Fig. 7.1. Perturbative approaches are no longer applicable in the overdamped, strongly anharmonic regime, in which the lifetimes become comparable to or even shorter than a single oscillation period $\tau_s(\mathbf{q}) \ll \omega_s^{-1}(\mathbf{q})$, because phonons are no longer well defined quasi-particles. Conversely, the perturbative phonon picture holds when the lifetimes span several oscillation periods $\tau_s(\mathbf{q}) \gg \omega_s^{-1}(\mathbf{q})$. Although the Ioffe-Regel limit does not constitute a strict quantitative rule, it is qualitatively helpful in identifying different transport regimes. Furthermore, similar considerations allow to identify particle-like (Peierls) and wave-like (Wigner) transport regimes within the quasi-particle regime [636].

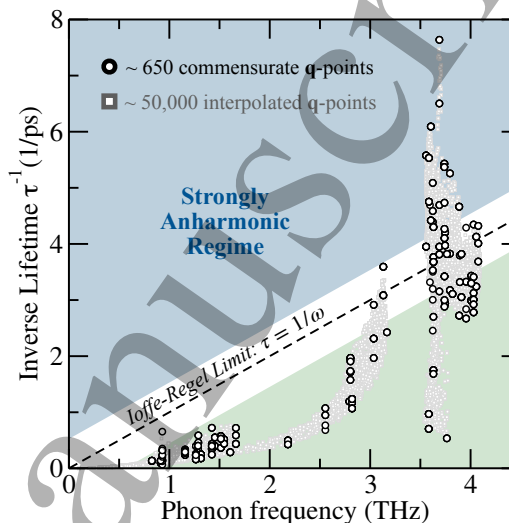


Figure 7.1: Phonon lifetimes obtained from fully anharmonic *ab initio*MD, adapted from Ref. [634]. The dashed line represents the Ioffe-Regel limit that qualitatively separates the quasi-particle regime (light green) from the strongly anharmonic regime. The quasi-particle picture becomes increasingly questionable when the lifetimes are getting close to ω^{-1} and breaks down in the strongly anharmonic regime, for which the Ioffe-Regel limit is violated.

Current Status of the Implementation

Both *perturbative* approaches and *non-perturbative* approaches are supported in FHI-aims for the calculation of vibrational thermal conductivities, whereby FHI-vibes [461] typically serves as the tool to setup, run, and evaluate the calculations.

Perturbative approaches start from the harmonic approximation, a second-order Taylor expansion of the PES, see Contrib. 6.1. By analytically treating the dynamics in this harmonic potential, quantum-nuclear effects can be accounted for, so that perturbative approaches are applicable also in the low-temperature regime. Anharmonic deviations from the harmonic PES are treated as a minor perturbation. They are accounted for by determining at least the third-order [637], sometimes also the fourth-order Taylor expansion of the PES [638, 639]. In turn, this allows one to determine scattering-cross-sections viz. phonon-lifetimes $\tau_s(\mathbf{q})$ that are the final ingredient for obtaining κ by solving the Boltzmann-Peierls transport equation. Let us note that this formalism also allows one to account for isotopic disorder [640] as well as glass-like Wigner-transport beyond Peierls' particle-like transport [636]. Similarly, higher-order anharmonic effects can be approximatively accounted for by extracting temperature-dependent force-constants from finite-temperature *aiMD* [527]; detailed discussions on practical implementations can be found in Refs. [517, 518, 520, 521, 641].

Non-perturbative approaches rely on explicit simulations of the nuclear motions via classical *ab initio* molecular dynamics (*aiMD*), cf. Contrib. 6.2. Then, fully anharmonic thermal conductivities can be obtained from

Roadmap on Advancements of the FHI-aims Software Package

7.1. VIBRATIONAL HEAT TRANSPORT IN WEAKLY, MODERATELY, AND STRONGLY ANHARMONIC SOLIDS

non-equilibrium *aiMD* trajectories, e.g., by explicitly simulating a thermal gradient [642], by imposing a heat flux [643], by monitoring equilibration from non-equilibrium [644] or by imposing close-to-equilibrium temperature profiles [645]. However, such non-equilibrium approaches are typically plagued by poor convergence with respect to simulation time and cell size. For this reason, it is generally advantageous to compute thermal conductivities from equilibrium *aiMD* trajectories by leveraging the fluctuation-dissipation theorem, e.g., by monitoring energy-density fluctuations [646, 647] or energy-current fluctuations via the Green-Kubo formalism [648].

In the following, we shortly summarize the implementations and workflows available for the perturbative and non-perturbative ansatz.

Perturbative Approaches:

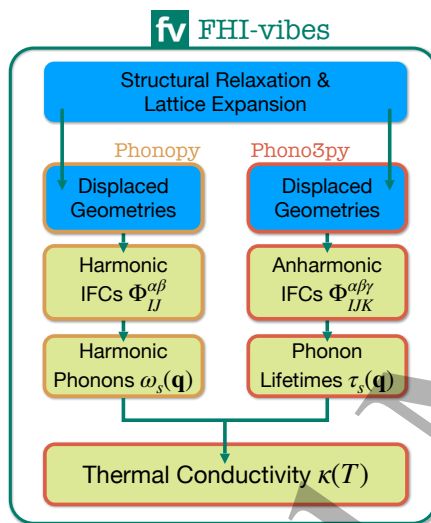


Figure 7.2: Workflow for computing vibrational thermal conductivities κ using a perturbative approach, for which interatomic force constants (IFCs) are obtained from finite differences. Blue cells highlight the steps in which FHI-aims is used for electronic-structure theory calculations to obtain, e.g. forces.

obtained via linear regression from thermodynamic snapshots generated, e.g., by *aiMD*. To this end, FHI-vibes offers an interface (`vibes utils trajectory 2tdep`) to the temperature-dependent effective potentials method TDEP described in Refs. [524, 641, 649].

Non-Perturbative Approaches:

For computing thermal conductivities in a *non-perturbative* fashion with FHI-aims, the method of choice is the *ab initio* Green-Kubo method [650]. In this formalism, it is necessary to run *aiMD* simulations and to

A typical FHI-aims workflow for the calculation of thermal conductivities in a perturbative approaches starts with a geometry relaxation to obtain the equilibrium geometry $\{\mathbf{R}^0\}$ and a subsequent *harmonic* phonon calculation to obtain the properties of the phonons with frequencies $\omega_s(\mathbf{q})$ with the Python package FHI-vibes, see Contrib. 6.1 for details. Subsequently, third-order derivatives of the PES, the so-called third-order force constants $\Phi_{IJK}^{\alpha\beta\gamma}$ need to be computed. They are defined as

$$\Phi_{IJK}^{\alpha\beta\gamma} = \left. \frac{\partial^3 U(\{\mathbf{R}\})}{\partial R_{I\alpha} \partial R_{J\beta} \partial R_{K\gamma}} \right|_{\{\mathbf{R}\}=\{\mathbf{R}^0\}}, \quad (7.1)$$

i.e., by extending the Taylor expansion introduced for the definition of the harmonic force constants $\Phi_{IJ}^{\alpha\beta}$ given in Contrib. 6.1. As shown in Fig. 7.2, the computation of $\Phi_{IJK}^{\alpha\beta\gamma}$ via finite differences can be orchestrated via FHI-vibes, which (i) generates geometries with symmetry-reduced displacements using Phono3py [523], (ii) evaluates the forces for these structures via FHI-aims, and (iii) postprocesses these calculations with Phono3py to determine the $\Phi_{IJK}^{\alpha\beta\gamma}$. Eventually, the thermal conductivity is obtained via FHI-vibes viz. Phono3py by computing the phonon lifetimes $\tau_s(\mathbf{q})$ using the $\Phi_{IJK}^{\alpha\beta\gamma}$, so to evaluate the Boltzmann transport equation. Besides using $\Phi_{IJ}^{\alpha\beta}$ and $\Phi_{IJK}^{\alpha\beta\gamma}$, FHI-vibes also supports the usage of temperature-renormalized force constants, which are obtained via linear regression from thermodynamic snapshots generated, e.g., by *aiMD*. To this end, FHI-vibes offers an interface (`vibes utils trajectory 2tdep`) to the temperature-dependent effective potentials method TDEP described in Refs. [524, 641, 649].

7.1. VIBRATIONAL HEAT TRANSPORT IN WEAKLY, MODERATELY, AND STRONGLY ANHARMONIC SOLIDS

compute the heat flux $\mathbf{J}(t)$ along the trajectory. The heat-flux implementation in FHI-aims [650] supports the calculation of *virial* heat fluxes, i.e., the dominant contribution in solids in which mass transport is negligible [651]. To this end, the contributions of each atom I to the virial $\underline{\sigma}_I(t)$ needs to be computed, details regarding the evaluation of these terms can be found in Refs. [54, 650]. In practice, such Green-Kubo calculations are best orchestrated with FHI-vibes, which takes care of running *aiMD* with FHI-aims and storing all relevant information including positions $\mathbf{R}_I(t)$, velocities $\dot{\mathbf{R}}_I(t)$, and virials $\underline{\sigma}_I(t)$ along the trajectory, as shown in Figure 7.3.

During postprocessing, FHI-vibes then also takes care to assemble the virial heat flux $\mathbf{J}_v(t) = V^{-1} \sum_I \underline{\sigma}_I(t) \cdot \dot{\mathbf{R}}_I(t)$, to remove non-contributing terms [652], to reduce noise [634], and to eventually evaluate the thermal conductivity κ by integrating the heat-flux autocorrelation function [648]. In the last step, a finite-size correction can be efficiently applied by mapping the dynamics onto a harmonic model $\omega_s(\mathbf{q})$, extracting the associated fully-anharmonic phonon lifetimes $\tau_s(\mathbf{q})$ from *aiMD*, and then extrapolating to the bulk viz. interpolating to the dense \mathbf{q} limit in a BTE-type model. This last step requires force constants $\Phi_{IJ}^{\alpha\beta}$ for the harmonic model, which can be computed at a harmonic level or also extracted from the *aiMD* via FHI-vibes.

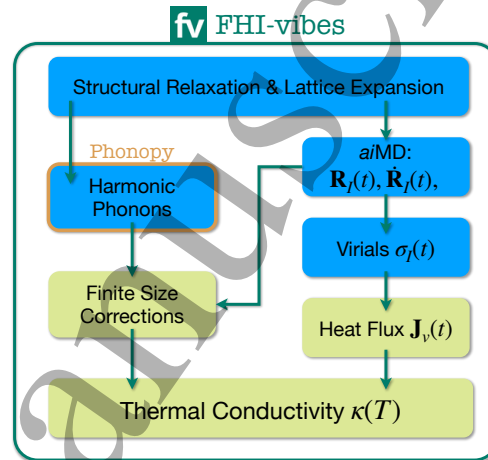


Figure 7.3: Workflow for computing vibrational thermal conductivities κ via the non-perturbative Green-Kubo approach. Blue cells highlight the steps in which FHI-aims is used for electronic-structure theory calculations to obtain, e.g. forces.

Usability and Tutorials

From an electronic-structure point of view, *perturbative* approaches based on finite differences only require the evaluation of forces, cf. Fig. 7.2. Accordingly, they can be used with all levels of theory for which FHI-aims is able to compute forces, notable all semi-local exchange-correlation functionals and hybrids, which include a fraction of Fock exchange, see Contrib. 3.2. Similarly, the Green-Kubo approach requires the evaluation of forces for performing the *aiMD* simulations and of atomic stresses for evaluating the virial heat flux. For the latter, hybrid functionals are not yet supported, but all necessary ingredients to evaluate the virials are already implemented also for these cases [54].

From a practical point of view, the computation of thermal conductivities with perturbative and non-perturbative approaches does, however, require quite complex workflows and the (system-dependent) convergence of several physical and numerical parameters beyond those important at the electronic-structure theory level. To just name a few, this includes computing accurately relaxed equilibrium structures within the symmetry group of interest, determining lattice expansion at finite temperatures, and converging the supercell size used for capturing the interactions relevant for the nuclear dynamics, i.e., for obtaining harmonic and anharmonic force constants $\Phi_{IJ}^{\alpha\beta}$ and $\Phi_{IJK}^{\alpha\beta\gamma}$ in perturbative approaches and for running *aiMD* simulations in non-perturbative approaches. To guide a novel user through this process, a set of tutorials for the computation of thermal conductivities with perturbative and non-perturbative methods are provided at <https://gitlab.com/FHI-aims-club/tutorials/thermal-transport-with-fhi-vibes>. The

*Roadmap on Advancements of the FHI-aims Software Package***7.1. VIBRATIONAL HEAT TRANSPORT IN WEAKLY, MODERATELY, AND STRONGLY ANHARMONIC SOLIDS**

tutorials build on the ones provided for the analysis of harmonic and anharmonic vibrations with FHI-vibes, cf. Contrib. 6.1, and showcase how to compute thermal conductivities using CuI, a strongly anharmonic material [533], as an example. Besides introducing the necessary physical background knowledge, the tutorials provide useful scripts and teach how to use features and functions of FHI-vibes to analyze, inspect, and evaluate this kind of calculations.

Future Plans and Challenges

As described in this contribution, substantial functionality for computing vibrational thermal conductivities $\kappa(T)$ from first principles is provided within FHI-aims and FHI-vibes. This enables to accurately calculate $\kappa(T)$ both for weakly anharmonic materials, but also for moderately and strongly anharmonic cases [533], for which approaches based on the phonon picture are uncertain or even not justified [636]. Clearly, further accelerating such kind of calculations to enable a rapid screening of material space is an important future step. Here, replacing the computationally involved *aiMD* with simulations performed with MD run with machine-learned interatomic potentials (MLIPs) [628], also see Contrib. 8.2, can be extremely beneficial. First steps in this direction have already been taken [536, 541, 653]. To provide accurate $\kappa(T)$ values, it is quintessential that the trained MLIP accurately captures the actuating (strongly) anharmonic effects. This is not guaranteed by standard training approaches, but can be facilitated by active-learning schemes [536]. Let us note that the infrastructure provided within FHI-vibes for thermal-conductivity calculations and for gauging the degree of anharmonicity [525] has also been proven useful in this MLIP context [536, 541, 653].

Eventually, it is also important to stress that the thermal conductivity of single crystals is certainly a fundamental quantity, but not the only one of interest in materials science. Here, investigating the role of disorder at different scales, be it mass disorder [640], structural disorder as in glasses [636], or interfacial resistance [654], is a very active field of research not yet covered by the implementations so far.

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233.. FK acknowledges support from the Swedish Research Council (VR) program 2020-04630, and the Swedish e-Science Research Centre (SeRC).

7.2 Electrical Transport for Anharmonic Materials

***Jingkai Quan**^{1,2}, **Florian Fiebig**¹, **Karsten Rasim**^{1,a}, **Bjoern Bieniek**³, **Yi Yao**^{1,4}, **Roman Kempt**⁵, **Zhenkun Yuan**^{1,b},
***Matthias Scheffler**¹, and ***Christian Carbogno**^{1,6,c}

¹The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

²Max-Planck Institute for the Structure and Dynamics of Matter, Luruper Chausse 149, 22761, Hamburg, Germany

³Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁵Technische Universität Dresden, Theoretische Chemie, 01069 Dresden, Germany

⁶Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

*Coordinator of this contribution.

^aCurrent Address: Miele & Cie. KG, Gütersloh, Germany

^bCurrent Address: Thayer School of Engineering, Dartmouth College, Hanover, NH 03755, USA

^cCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

Summary

Electrical transport plays a pivotal role for a multitude of scientific and industrial applications. It critically depends on the chemical and structural details of the materials, its doping, as well as temperature and pressure. From the conductivity, one can derive crucial thermoelectric quantities for heat transport and energy generation. Similarly, the coupling of mechanical motion and electronic transport gives access to electromechanical properties. For instance, band-type conductivity is at the very heart of semiconductor technology. While conductivities can be estimated [655] from electronic band-structures and the associated group velocities, an accurate assessment requires accounting for the coupling of electronic and nuclear degrees of freedom. Commonly, electronic-structure theory and many-body perturbation theory [378] are used to describe electron-phonon coupling and to compute band-type mobilities limited by phonon scattering from first principles [615, 616, 656]. In this approach, the nuclear dynamics is approximated by an (effective [657]) harmonic potential, i.e., phonons, and their first-order coupling to the electronic degrees of freedom. In turn, this allows for the calculation of scattering cross-sections that can be fed to transport equations, e.g., the Boltzmann [616] and/or the Wigner formalism [658]. The perturbative approach works well for harmonic materials [656], but the aforementioned approximations may fail in anharmonic materials and/or at elevated temperatures. For instance, it has been shown that the temperature-dependence of the mobility in the strongly anharmonic perovskites SrTiO₃ and BaTiO₃ is driven by higher-order anharmonic and electron-vibrational couplings [535]. Similarly, thermoelectric materials are known to exhibit strong anharmonicity [525, 533] that cannot be described within harmonic models and that are even challeng-

ing to describe with machine-learned potentials [536]. To capture these effects and to achieve a general, non-perturbative evaluation of mobilities, the Kubo-Greenwood formalism [659] has been implemented in FHI-aims.

Current Status of the Implementation

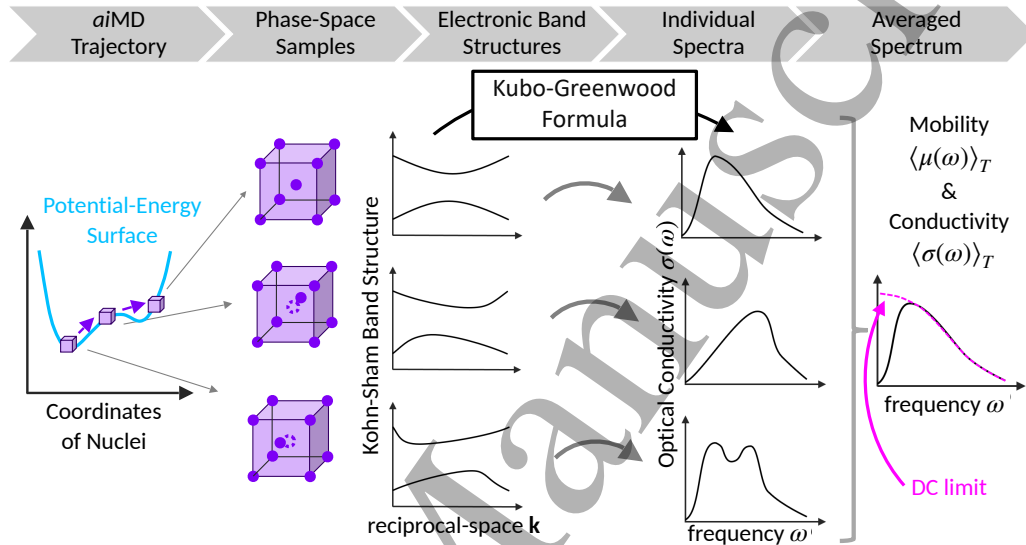


Figure 7.4: Schematic workflow of a Kubo-Greenwood calculation. For illustration, three snapshots of a bcc crystal structure, in which only the central atom moves, are shown as example. Similarly, the effects on the band structure and on the optical conductivity are exaggerated for the sake of clarity.

A typical workflow for the evaluation of mobilities with the Kubo-Greenwood formalism is shown in Fig. 7.4. In a first step, an *ab initio* molecular-dynamics (*aiMD*) simulation is run on the self-consistent potential-energy surface to account for all orders of anharmonic effects, c.f. Contrib. 6.2. In a second step, uncorrelated geometry samples are chosen from this trajectory to cover the phase-space at the thermodynamic conditions of interest. In a third step, the ground state electronic structure is determined self-consistently for each of these samples, which accounts for all orders of adiabatic electron-vibrational coupling. In a fourth step, the obtained electronic-structures are used to compute the frequency-dependent conductivity, typically referred to as optical conductivity $\sigma(\omega)$ for the individual samples. In the final step, the spectra $\sigma(\omega)$ are post-processed to determine the mobilities $\mu(\omega)$.

By using linear response theory in the Kubo formalism [648] and expressing the band-like current operator in terms of effective, single-particle eigenstates $|\mathbf{k}m\rangle$ one obtains the following Kubo-Greenwood (KG) formula for the real, diagonal part of the frequency-dependent (ω) optical conductivity for one sample of phase space [659, 660]:

$$\text{Re}(\sigma_{\alpha\alpha}(\omega)) = \frac{2\pi q^2 \hbar^2}{m^2 V \omega} \sum_{\mathbf{k}mn} (\langle \mathbf{k}n | \nabla_{\alpha} | \mathbf{k}m \rangle \langle \mathbf{k}m | \nabla_{\alpha} | \mathbf{k}n \rangle) (f_{\mathbf{k}n} - f_{\mathbf{k}m}) \delta(\epsilon_{\mathbf{k}m} - \epsilon_{\mathbf{k}n} - \hbar\omega). \quad (7.2)$$

Here, α denotes a Cartesian axis, $f_{\mathbf{k}m}$ and $\epsilon_{\mathbf{k}m}$ are the occupation number and the energy eigenvalue of eigenstate $|\mathbf{k}m\rangle$, $\langle \mathbf{k}m | \nabla_{\alpha} | \mathbf{k}n \rangle$ are the gradient matrix elements in this basis, and V is the cell volume.

Roadmap on Advancements of the FHI-aims Software Package

7.2. ELECTRICAL TRANSPORT FOR ANHARMONIC MATERIALS

The electron e and hole h conductivities $\sigma_{e/h}(\omega)$ can be obtained by limiting the summation over m to states in the conduction and valence band, respectively. In the FHI-aims implementation, the matrix elements $\langle \mathbf{k}m | \nabla_\alpha | \mathbf{k}n \rangle$ in Kohn-Sham space are obtained via

$$\langle \mathbf{k}m | \nabla_\alpha | \mathbf{k}n \rangle = \sum_{ij} [C_m^i(\mathbf{k})]^* C_n^j(\mathbf{k}) \sum_{\mathbf{N}} e^{i\mathbf{k} \cdot \mathbf{T}(\mathbf{N})} \langle \phi_{i\mathbf{0}}(r) | \nabla_\alpha | \phi_{j\mathbf{N}}(r) \rangle, \quad (7.3)$$

where $C_n^j(\mathbf{k})$ denotes a Kohn-Sham expansion coefficient, $\mathbf{T}(\mathbf{N})$ a translation by a combination of lattice vectors, and \mathbf{k} a reciprocal-space vector. More details on the used notation can be found in Contrib. 2. The real-space gradient matrix elements $\langle \phi_{i\mathbf{0}}(r) | \nabla | \phi_{j\mathbf{N}}(r) \rangle$ in the basis of the numeric atom-centered orbitals $\phi_{j\mathbf{N}}(r)$ are computed once using the integration scheme discussed in Ref. [19] and then stored as sparse matrix.

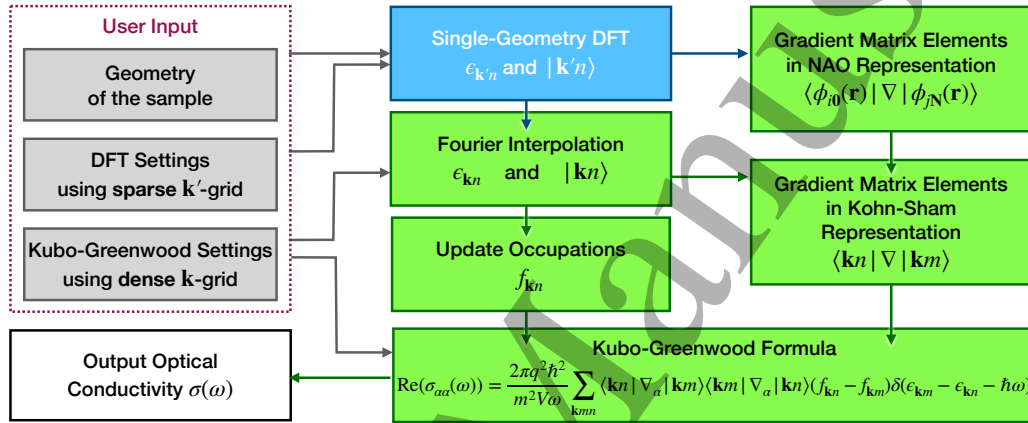


Figure 7.5: Flowchart for the calculation of the optical conductivity $\sigma(\omega)$ for *one* sample using the Kubo-Greenwood implementation in FHI-aims. Grey boxes highlight user input, the blue box a “standard” FHI-aims calculations, green boxes the steps needed for evaluating Eq. (7.2), and the white box the output.

Achieving numerical convergence of the KG formula is challenging in crystalline materials especially in the DC limit $\omega \rightarrow 0$, since only a small portion of \mathbf{k} -space contributes to Eq. (7.2) due to the fact that both $f_{\mathbf{k}n} - f_{\mathbf{k}m} \neq 0$ and $\epsilon_{\mathbf{k}m} - \epsilon_{\mathbf{k}n} = \hbar\omega$ need to be fulfilled. For the latter condition, the delta function is numerically resolved by either a Gaussian or Lorentzian broadening function with a finite, but minute width η that appropriately represent the limit of vanishing broadness.

In practice, \mathbf{k} -grids in the order of 10^3 - 10^5 \mathbf{k} -points are needed to achieve convergence for sufficiently small values of η , even in extended supercells [535, 661]. Such dense \mathbf{k} -grids are typically unnecessary for achieving convergence in the self-consistent field cycle (SCF), since the electronic density and, in turn, the associated real-space Hamiltonian, can typically be converged within a relatively sparse \mathbf{k}' -grid during the SCF cycle. Due to the locality of the real-space matrix elements, the converged real-space Hamiltonian $\langle \phi_{i\mathbf{0}} | \underline{\mathbf{H}} | \phi_{j\mathbf{N}} \rangle$ and overlap matrix $\langle \phi_{i\mathbf{0}} | \underline{\mathbf{S}} | \phi_{j\mathbf{N}} \rangle$ can then be Fourier-interpolated to arbitrarily dense \mathbf{k} -grids via:

$$\langle \phi_{i\mathbf{k}} | \underline{\mathbf{H}} | \phi_{j\mathbf{k}} \rangle = \sum_{\mathbf{N}} e^{i\mathbf{k} \cdot \mathbf{T}(\mathbf{N})} \langle \phi_{i\mathbf{0}} | \underline{\mathbf{H}} | \phi_{j\mathbf{N}} \rangle \quad (7.4)$$

$$\langle \phi_{i\mathbf{k}} | \underline{\mathbf{S}} | \phi_{j\mathbf{k}} \rangle = \sum_{\mathbf{N}} e^{i\mathbf{k} \cdot \mathbf{T}(\mathbf{N})} \langle \phi_{i\mathbf{0}} | \underline{\mathbf{S}} | \phi_{j\mathbf{N}} \rangle. \quad (7.5)$$

As sketched in Fig. 7.5, this procedure is used in the KG calculation after SCF convergence has been achieved to evaluate Eq. (7.2). By this means, the dense \mathbf{k} -grids required for converging Eq. (7.2) can be targeted without incurring into prohibitive numerical costs for the SCF that prevent achieving convergence [662]. Note that also the Fermi-level ϵ_F and the electron and hole charge-carrier densities $n_{e/h}$ can change when switching to denser \mathbf{k} -grids. Accordingly, also these quantities are updated in this step, so that the optical conductivities $\sigma_{e/h}(\omega)$ for one sample are obtained within FHI-aims.

Beyond the Kubo-Greenwood method, we have also implemented several semi-empirical methods to quickly estimate properties related to charge transport. This includes the carrier-lattice distance d_{c-1} [663] that aims at capturing electron-phonon coupling effects. To compute

$$d_{c-1}(\text{CBM/VBM}) = \int_{\text{uc}} d\mathbf{r} |\psi_{\text{CBM/VBM}}(\mathbf{r})|^2 \min_{\alpha} \{|\mathbf{r} - \mathbf{R}_{\alpha}|\}, \quad (7.6)$$

we extract the eigenstate density from FHI-aims at the conduction band minimum (or valence band maximum) and integrate the expression over the whole unit cell (uc). In a similar spirit, conductivities can be estimated by using the interface with the BoltzTraP2 [664] or with the the AICON2 code [665]. Here, electronic band-structures are used to determine group velocities; in the AICON2 case, deformation-potential theory is additionally used to approximate relaxation times.

Usability and Tutorials

The evaluation of optical conductivities $\sigma(\omega)$ for single samples is directly controlled via keywords in the input file `control.in`, most importantly `compute_kubo_greenwood` for invoking the evaluation of the KG formula and setting numerical parameters such as the broadening η . The Fourier interpolation is controlled via the keyword `dos_kgrid_factors`, since the density-of-states routines are used internally for this scope. Given that dense, interpolated \mathbf{k} -grids are necessary for converging Eq. (7.2), the parallelization in this routine occurs over \mathbf{k} -points with extensive use of LAPACK for linear algebra operations. This enables good parallel scaling with number of nodes, so that system sizes up to 1,000 atoms can be routinely treated; for even larger systems, extending the implementation to also support distributed ScaLAPACK arrays and `use_local_index` in the KG-formula evaluation would be desirable. Let us emphasize that the actual SCF cycle predating the KG-formula evaluation can already be run using distributed arrays in ScaLAPACK mode and that all exchange-correlation (xc) functionals, including hybrids, c.f. Contrib. 3.2, are supported.

To eventually obtain thermodynamically relevant conductivities at temperature T , an ensemble average $\langle \sigma(\omega) \rangle_T$ over all samples is performed. This step is independent of FHI-aims viz. the electronic-structure theory code and is performed as a post-processing procedure. At this stage, also the mobility $\langle \mu \rangle_T = \langle \sigma(\omega) \rangle_T / \langle qn \rangle_T$ is calculated, where q is the carrier's charge and n is the carrier density. The DC mobility $\langle \mu(0) \rangle_T$ in the thermodynamic bulk limit $\omega \rightarrow 0$ is determined by fitting the low-frequency part of $\langle \mu(\omega) \rangle_T$ using a Drude function with lifetime τ

$$\langle \mu(\omega) \rangle_T \approx \frac{\mu(0)}{(\omega\tau)^2 + 1}, \quad (7.7)$$

as sketched in the last step in Fig. 7.4. Note that this extrapolation is necessary since reaching the DC limit would require infinitely large supercells [535], for which the whole electronic-band structure is folded onto the Γ -point, see Contrib. 6.5.

The conductivity strongly depends on the band-gap through the carrier density n . Accordingly, semi-local xc-functionals can overestimate conductivities by orders of magnitude due the notorious band-gap problem. Conversely, mobilities are an intrinsic property that is unaffected by the carrier density in the low-doping regime and are thus reasonably predicted already with semi-local xc-functionals [616]. In this context, constraining the charge-carrier density to a fixed value can be numerically beneficial, as detailed in Ref. [535].

To guide novel user through the workflow of a KG calculation, a tutorial is provided at [the following link](#)^a. Using Al and SrTiO₃ as examples, it showcases a typical workflow to calculate the conductivity and mobility with KG formalism, provides scripts for pre- and postprocessing, and explains how the results can be analysed and understood.

Future Plans and Challenges

The main advantage of the implemented Kubo-Greenwood formalism lies in its ability to account for anharmonic and higher-order coupling effects in the evaluation of mobilities. With that, it can be used to effectively assess strongly doped materials, which are for instance common in thermoelectric applications [666]. In this regard, it is also advantageous that the implemented Kubo-Greenwood formalism can directly deal with high charge-carrier densities and defect scattering within the supercell approach and does not require additional *ad hoc* models to account for these effects. Besides the electrical conductivity, the Kubo-Greenwood formalism allows to determine electronic contributions to the thermal conductivity κ_{el} and the Seebeck coefficient S as well. Thus the exact same code base can be used to calculate charge, heat, and thermoelectrical transport properties of a material. The implementation of the respective equations, see Ref. [660], is essentially finished, but requires thorough testing and validation. Also in this regard, the implemented formalism can help validating and improving approximations taken in perturbative approaches. In turn, a systematic comparison with perturbative approaches can clarify the role of approximations taken in the Kubo-Greenwood formalism, notably using a supercell-approach and the Born-Oppenheimer approximation [631], which inhibits a systematic inclusion of long-range Fröhlich effects. Combining such studies with the evaluation of temperature-dependent spectral functions, can help including such effects more accurately and more efficiently, see Contrib. 6.5.

From a numerical point of view, accelerating KG calculations would be desirable. For instance, this would allow to rapidly address weakly or moderately anharmonic materials, for which the KG approach equally holds, but often becomes numerically impractical. To this end, interfacing to machine-learning based approaches is a promising route. Machine-learning approaches for predicting electronic properties such as densities and matrix elements [83, 337, 629] can potentially accelerate the evaluation of the KG formula, especially if one can circumvent the computationally dominant eigenvalue problem, see Contrib. 2.4. Similarly, using machine-learned interatomic potentials [628] instead of *aiMD* for the sampling of the thermodynamic phase space can reduce the computational cost in the sampling step, see Contrib. 8.2. Here, particular care has to be taken to ensure that strongly anharmonic effects, e.g., those triggered by spontaneous defect formation [533] are appropriately described by the trained potentials [536].

Eventually, let us emphasize that the developed approach covers band-type conductivity, but not ionic and polaronic conductivities. Those do not only require different flux operators, but typically also necessitate to simulate even longer time- and length scales [667]. In this regard, the development of methodologies that cover all these kind of conductivities constitutes a significant challenge for first-principles based modeling [668].

^a<https://fhi-aims-club.gitlab.io/tutorials/kubo-greenwood-formula>

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233. CC and MS acknowledge helpful discussions with Martin French and Ronald Redmer.

7.3 Molecular Transport

**María Camarasa-Gómez*^{1,2}, **Daniel Hernangómez-Pérez*³, *Jan Wilhelm*⁴, *Alexej Bagrets*⁵, and **Ferdinand Evers*⁴

¹Departamento de Polímeros y Materiales Avanzados: Física, Química y Tecnología, Facultad de Química, UPV/EHU, Apartado 1072, 20018 Donostia-San Sebastián, Spain

²Centro de Física de Materiales (CFM-MPC) CSIC-UPV/EHU, 20018 Donostia-San Sebastián, Spain

³CIC nanoGUNE BRTA, Tolosa Hiribidea 76, 20018 San Sebastián, Spain

⁴Institute of Theoretical Physics and Regensburg Center for Ultrafast Nanoscopy (RUN), University of Regensburg, 93040 Regensburg, Germany

⁵Institute of Nanotechnology, Karlsruhe Institute of Technology, 76344 Eggenstein-Leopoldshafen, Germany, and Steinbuch Centre for Computing, Karlsruhe Institute of Technology, 76344 Eggenstein-Leopoldshafen, Germany (currently at ITK Engineering GmbH.)

*Coordinator of this contribution.

Summary

Single-molecule junctions [669–672] — nanoscale systems where a molecule is connected to metallic electrodes - offer a unique platform for studying charge, spin and energy transport in non-equilibrium many-body quantum systems, with few parallels in other areas of condensed matter physics. Over the past decades, these systems have revealed a wide range of remarkable quantum phenomena, including quantum interference [669, 673], non-equilibrium spin-crossover [674], diode-like behavior [675], or chiral-induced spin selectivity [676], among many others [670, 671]. To develop a detailed understanding, it turned out essential to have available *ab initio*-based tools for accurately describing quantum transport in such systems [670, 671]. They need to be capable of capturing the intricate electronic structure of molecules, sometimes in the presence of electron-electron or electron-phonon interactions, in out-of-equilibrium environments. Such tools are also indispensable for explaining experimentally observed phenomena using parametrized tight-binding models in the context of quantum transport.

While FHI-aims also offers specialized transport routines [677–679], e.g. for chemically functionalized nanotubes or nanotube networks, our focus in this section is on the AITRANSS package [680–683] designed for simulations of single-molecule transport.

AITRANSS is an independent post-processing tool that, when combined with FHI-aims, enables the calculation of electronic transport properties, as well as atom-projected density of states, spin properties and the simulation of scanning tunneling microscope images in molecular junctions. Pilot versions of the code extend some of these capabilities to non-linear transport in the applied bias, with plans to include these features in future releases of the package.

Current Status of the Implementation

The AITRANSS code implements the non-equilibrium Green's function formalism (NEGF) [671, 672, 684, 685]; AITRANSS can handle closed-shell and spin-polarized contacts [682] and also spin-orbit coupling [681,

7.3. MOLECULAR TRANSPORT

686].

A central element in the calculation of electronic transport is the (ballistic) transmission function, which can be obtained using the trace formula [687]

$$T(E) = \text{Tr} \left[\hat{\Gamma}_L \hat{G}(E) \Gamma_R \hat{G}^\dagger(E) \right], \quad (7.8)$$

where $\hat{G}(E)$ denotes the Green's function of the extended molecule in the presence of the leads, *i.e.* source and drain. The extended molecule includes a part of the leads as metal clusters, which are in contact with the molecule; the construction of AITRANSS offers various advantages, *e.g.*, complete flexibility in the relative orientation of the electrodes. As described elsewhere [671], the Green's function can be obtained through partitioning,

$$\hat{G} = (E\mathbb{1} - \hat{H}^{\text{KS}} - \hat{\Sigma})^{-1}. \quad (7.9)$$

Here, $\mathbb{1}$ represents the identity operator, \hat{H}^{KS} the Kohn-Sham Hamiltonian and $\hat{\Sigma} = \hat{\Sigma}_L + \hat{\Sigma}_R$ the self-energy operator of the left (L) and right (R) leads. Finally, $\hat{\Gamma}_{L,R}$ denotes the anti-Hermitian part of the self-energy operators $\Sigma_{L,R}$. A distinctive feature of AITRANSS is its computationally efficient implementation of absorbing boundary condition using a model self-energy approach [680, 682, 688]. In this scheme, the self-energy is given by

$$\hat{\Sigma}_\alpha = \sum_{\tilde{\mu}, \tilde{\nu} \in \mathcal{S}_\alpha} |\tilde{\mu}\rangle [\delta\epsilon - i\eta] \delta_{\tilde{\mu}\tilde{\nu}} \langle \tilde{\nu}|, \quad (7.10)$$

where $\delta\epsilon$ is the real energy shift, and η characterizes the imaginary part, which corresponds to a local, energy-independent, material-specific leakage rate out of the scattering region. Note that the self-energy is applied only within a subspace \mathcal{S}_α corresponding to the lead atoms of the extended molecule that are farthest away from the molecule.

Calculations with post-processing only. The simplest operation mode of AITRANSS is non-self-consistent, see Fig. 7.6, *i.e.* without feedback loop. The process begins with a standard DFT calculation using the optimized geometry of the molecular junction obtained from FHI-aims. It delivers the set of Kohn-Sham energies $\{\epsilon_l\}$ and orbitals

$$\psi_l(\mathbf{r}) = \sum_{i=1}^{N_B} c_{il} \varphi_i(\mathbf{r}), \quad (7.11)$$

where N_B is the number of orbitals, c_{il} are the molecular orbital coefficients and φ_i denotes the FHI-aims basis set (cf. Eq. (2.2)). These orbitals are used by AITRANSS to reconstruct the Hamiltonian. Because the basis is non-orthogonal, AITRANSS employs the Löwdin orthogonalization procedure [689] using the overlap matrix, S ,

$$\varphi_i(\mathbf{r}) = \sum_{j=1}^{N_B} S_{j,i}^{-1/2} \varphi_j(\mathbf{r}), \quad (7.12)$$

to orthogonalize the states and reconstruct the Kohn-Sham Hamiltonian in an orthogonal basis, $\hat{H}^{\text{KS}} = S^{1/2} C \epsilon C^\dagger S^{1/2}$. The resulting Hamiltonian is then used for evaluating Eqs. (7.8)-(7.9).

Self-consistent calculation procedure. In the self-consistent calculation mode, FHI-aims and AITRANSS perform a feedback loop, see Fig. 7.7. As in the post-processing-only variant, the starting point is a standard DFT-calculation for the extended molecule. The role of feedback is to introduce self-consistency in the sense that the DFT-calculation in FHI-aims and the Green's function calculations in AITRANSS refer to the same

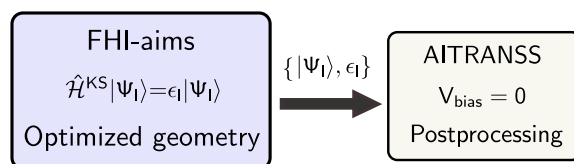


Figure 7.6: Workflow of the post-processing-only cycle combining FHI-aims and the AITRANSS package.

(non-equilibrium) density matrix ρ : during each iteration step the Kohn-Sham Hamiltonian is updated with $\hat{\rho}$, which is constructed using the NEGF [681, 682, 686]. The feedback-loop establishes, in particular, the Fermi energy, E_F , at fixed particle number, N , and the real part of the self-energy, $\delta\epsilon$. Physically, the cycle ensures correct charge redistribution in the junction, accounting for the macroscopic nature of the contacts and maintaining charge neutrality while screening the excess charge accumulated at the outermost boundaries of the finite-sized clusters [680–682, 686, 690, 691]. During the feedback loop, the real part of the self-energy, $\Sigma(\delta\epsilon^*)$, is gradually deformed so as to enforce charge-homogeneity on the far sites of the metal cluster, also under finite bias voltage, V_{bias} . Physical observables are calculated in the final post-processing step: transmission function, current-voltage characteristics, spin-orbit torques, and more.

Usability and Tutorials

AITRANSS is a project under continuous development since 2002, and is currently centered at Universität Regensburg. The current main developers are María Camarasa-Gómez, Daniel Hernangómez-Pérez and Ferdinand Evers. The source code, along with examples and a basic electrode library, is distributed with the FHI-aims package. It can be found in the subdirectory `external/aitranss`. This directory contains the source code, the script `tcontrol.aims.x` used to prepare the mandatory `tcontrol` input file for AITRANSS, a representative electrode library, and a folder with documented examples, including input and output files of FHI-aims and AITRANSS.

Compilation instructions, explanations of the code, and a list of available keywords are provided in the FHI-aims manual. As an independent package, AITRANSS has its own build system, which has also been integrated into the FHI-aims cmake-based build scheme, as documented in the FHI-aims manual release 240507. The current release of AITRANSS is parallelized using OpenMP directives, enabling the creation of a multithreaded version of the executable.

Without spin-orbit coupling, using the mandatory keyword `output aitranss` in the `control.in` file generates three ASCII files after a successful run of FHI-aims: `basis.out`, which contains information about the basis functions, `omat.aims`, which contains the overlap integrals and `mos.aims` which contains the Kohn-Sham orbitals and energies for the extended molecule. If an open-shell calculation is performed, `mos.aims` is substituted by two files called `alpha.aims` and `beta.aims`. AITRANSS is then run in the same directory where the output files of FHI-aims are located. To include spin-orbit interaction, the keywords

```

include_spin_orbit
output soc_eigenvectors 1
output soc_aitranss
  
```

must also be used, as documented in Ref. [686]. The basis and molecular orbital files are replaced by `basis-indices.soc.out` and `omat.aims.soc`. The current implementation of the self-consistent non-

Roadmap on Advancements of the FHI-aims Software Package

7.3. MOLECULAR TRANSPORT

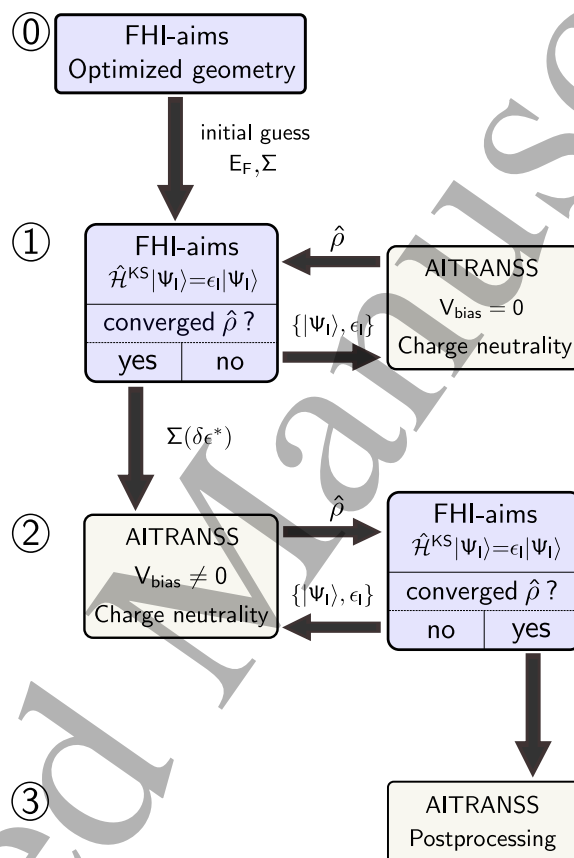


Figure 7.7: Workflow of the self-consistent cycle combining FHI-aims and the AITRANSS package. The calculation proceeds in four steps: ①, initial preparatory calculation with FHI-aims for the geometry optimization; ②, parametrization of the self-energy and the Fermi energy; ③, self-consistent loop at finite bias; and ④, postprocessing step in which observables based on the non-equilibrium density matrix can be computed. Fig. adapted from Fig. 2 in Ref. [681] (CC BY-NC 4.0).

equilibrium cycle (Fig. 7.7) is managed by an external shell script and maintains its performance in terms of memory and computational requirements for small molecular junctions. Additional details can be found as well in Ref. [686].

Finally, as a project in continuous expansion, online tutorials and updated information can be accessed from the AITRANSS webpage at <https://aitranss.ur.de/>.

Future Plans and Challenges

The AITRANSS package is an ongoing project under continuous development, with new features and capabilities to be added in the near future. From a computational perspective, we will focus on maintaining the package efficiently. The code is currently being refactored to enhance readability, making it more compact and easier to manage. Additional examples and tutorials will be added as well. In addition, we plan to extend the parallelization strategies beyond OpenMP (multithreading) by incorporating MPI parallelization, which will also function independently from the former. Further enhancements include replacing outdated ASCII files with modern formats such as HDF5.

From the functionality perspective, a significant issue, resulting from the limitations of Kohn-Sham transport calculations using semilocal functionals, is the deviation of Kohn-Sham spectral properties from the exact values. We aim to address this issue by employing scissor-like operation techniques based on hybrid functionals and image-charge correction. Furthermore, we will be expanding the study of transport and dynamical properties in the presence of spin-orbit interactions, for example incorporating spin-orbit effects in scanning tunneling microscope imaging simulations. We also plan to continue developing the implementations for current-induced forces (both mechanical and spin) and to incorporate interaction with light in the terahertz regime, local currents [692–694] as well as phonon effects. Finally, we plan to work on implementing multiterminal calculations in AITRANSS, particularly in the context of electrochemistry, where electrostatic effects are crucial.

Acknowledgments

We appreciate pleasant and fruitful discussions with many colleagues over the years: L. Venkataraman, M. S. Inkpen, R. Korytár, M. Kamenetska, M. Walz, G. Solomon, J. van Ruitenbeek. Contributions of P. Havu and V. Havu to the development of additional transport routines interfaced with FHI-aims are acknowledged. M.C.-G. acknowledges support from the Diputación Foral de Gipuzkoa through Grant 2024-FELL-000007-01 and from the Gobierno Vasco-UPV/EHU Project No. IT1569-22. D. H.-P. is grateful for funding from the Diputación Foral de Gipuzkoa through Grants 2023-FELL-000002-01, 2024-FELL-000009-01, and from the Spanish MICIU/AEI/10.13039/501100011033 and FEDER, UE through Project No. PID2023-147324NA-I00. J. W. acknowledges funding by the German Research Foundation (DFG) via the Emmy Noether Programme (Project No. 503985532). F. E. is grateful for support from DFG through the Collaborative Research Center (SFB) 1277 - Project ID 314695032 (subproject A03), from GRK 2905 (project-ID 502572516) and from the State Major Instrumentation Program, INST 89/560-1 (project number 464531296). F. E. and J. W. acknowledge the Gauss Centre for Supercomputing e.V. for providing computational resources on SuperMUC-NG at the Leibniz Supercomputing Centre under the Project ID pn36zo and the computing time provided to them on the high performance computer Noctua 2 at the NHR Center PC2. These are funded by the Federal Ministry of Education and Research and the state governments participating on the basis of the resolutions of the GWK for the national high-performance computing at universities (www.nhr-verein.de/unsere-partner).

7.4 Computing Diabatic Couplings

Simiam Ghan¹, *Harald Oberhofer², Karsten Reuter³, Christoph Schober⁴,

¹Catalysis Theory Center, Department of Physics, Technical University of Denmark

²Chair for Theoretical Physics VII and Bavarian Center for Battery Technologies, University of Bayreuth, Germany

³Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Department of Computer Science, Deggendorf Institute of Technology, Deggendorf, Germany

*Coordinator of this contribution.

Summary

The electronic coupling parameter – also known as the charge transfer integral – is defined as the Hamiltonian matrix element between two localized *diabatic* states:

$$H_{ab} = \langle \psi_a | \mathcal{H} | \psi_b \rangle. \quad (7.13)$$

In molecular systems, the electronic coupling is invoked in the theories of charge transfer[695], while on surfaces it is additionally used in theories of e.g. chemisorption and catalysis[696–698], scanning-probe microscopy[699], and quantum impurities[700]. In particular, the electronic coupling is one of the key features determining the efficiency of charge transport in molecular crystals[701].

Contrary to adiabatic states, which are the delocalized (Kohn-Sham) eigenstates of a system's Hamiltonian, localized diabatic states are nonunique and must be approximated. A large number of diabaticization methods have emerged to determine suitable diabatic states and the couplings between them[695, 702]. For FHI-aims, two efficient approaches are currently available: Fragment Orbital DFT (FO-DFT) and Projection-Operator Diabatization (POD2+). Our recent works have demonstrated the use of these methods for both molecular and surface systems.

Current Status of the Implementation

Both diabaticization methods described below have been shown to work with FHI-aims' native NAO basis as well as the popular Pople[703] and Dunning[32] basis sets. At close distances up to 5Å no difference in accuracy could be detected between a Tier 1 NAO basis set and a triple zeta Gaussian basis, with the former being more efficient due to its smaller size.

Coupling between Molecules

Fragment-Orbital DFT (FO-DFT)

In FO-DFT the initial and final diabatic states of a charge transfer reaction are constructed from non-interacting fragment densities (see fig. 7.8). As a consequence, any interactions between the fragments in the real system are neglected, making FO-DFT a method best suited for weakly interacting donor-acceptor pairs such as found in organic crystals^[704, 705] or physisorbed adsorbates on surfaces.

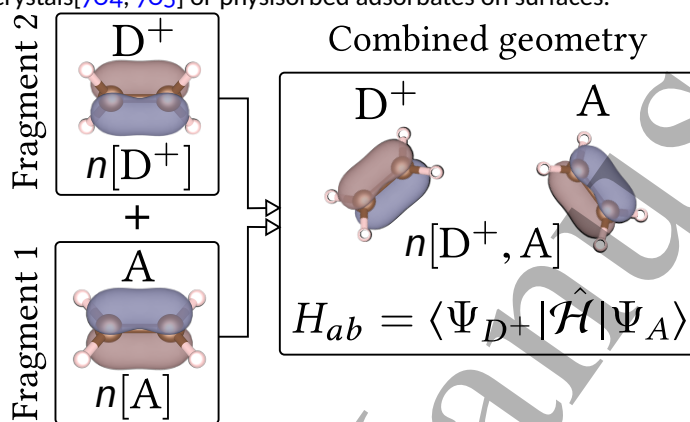


Figure 7.8: Schematic illustration of the fragment orbital DFT scheme for hole transfer in ethylene, using charged fragments.

Due to the approximations involved in constructing the fragment and combined system, FO-DFT can be described as a family of methods depending on how the fragment densities and combined Hamiltonian are defined.^[706] FHI-aims currently implements three FO-DFT flavors:

- $\mathcal{H}^{2n}@DA$ The original flavor implemented by Senthilkumar et. al^[707] with two neutral fragments and $2n$ electrons in the combined Hamiltonian (MRSE HAB11/HAB7:^[706] -24.6% / -22.4%).
- $\mathcal{H}^{2n-1}@DA/\mathcal{H}^{2n+1}@D^-A^-$ This flavor^[708] improves on the original formulation by using the correct number of electrons in the combined Hamiltonian (MRSE HAB11/HAB7:^[706] -37.7% / -27.1%).
- $\mathcal{H}^{2n-1}@D^+A/\mathcal{H}^{2n+1}@D^-A$ The most accurate flavor of FO-DFT employs charged fragment calculations, most closely mimicking the correct diabatic states in the system (MRSE HAB11/HAB7:^[706] -22.4% / -22.9%). Requires an additional full DFT calculation compared to the previous two schemes, making it computationally more expensive.

Projection-Operator Diabatization

The central idea of Projection-Operator Diabatization (POD) ^[709–711] lies in a partitioning of the system's Kohn-Sham Hamiltonian into blocks based on the (NAO) basis functions associated with either of the two fragments, labelled a (e.g. surface) and b (e.g. adsorbate). Each block is then diagonalized separately as illustrated in Figure (7.9).

Thereby, diabatic states are approximated as the block eigenstates with energies $\epsilon_{a,i}$ and $\epsilon_{b,j}$. First approximations to the coupling between two specific orbitals ψ_a and ψ_b are then simply encoded in the off-diagonal

Roadmap on Advancements of the FHI-aims Software Package

7.4. COMPUTING DIABATIC COUPLINGS

Usability and Tutorials

FO-DFT The functionality is fully integrated and documented within FHI-aims. It is available via the `fo_dft` tag in the `control.in` file. A full FO-DFT calculation requires at least 2 invocations of FHI-aims (fragment and combined system). A Python-based package automating the different calculation steps and allowing the re-use of fragment calculations for different geometries is available at <https://gitlab.com/schobler-ch/aimsutils>.

POD2+ A Python software package for POD2+ methods on surfaces, including tutorial, is available online at https://gitlab.com/simiamghan/bd_utils. This is an external post-processing routine which requires as inputs the overlap and converged Hamiltonian matrices of FHI-aims, which can be printed using the keywords `output_hamiltonian_matrix` and `output_overlap_matrix`. Methods are documented in references [711, 712].

Future Plans and Challenges

FO-DFT The calculation of heterogeneous couplings where $H_{ab} \neq H_{ba}$ is not implemented in an automated way and requires explicit fragment geometries. The current implementation also relies on the legacy restart infrastructure in FHI-aims. To further improve the efficiency of the method, future versions will use the more efficient ELSI-based infrastructure.

POD2+ The suitability of the POD2+ diabaticization methods for calculating coupling in covalently bound or chemisorbed systems is currently being studied. The POD2-based routines are currently implemented as external post-processing routines with significant storage requirements (FHI-aims must print all matrices to file for parsing). An internal FHI-aims implementation is needed.

Acknowledgements

HO acknowledges support from the German Science Foundation (DFG) under grant numbers OB 425/9-1 and OB 425/3-2.

*Roadmap on Advancements of the FHI-aims Software Package**7.4. COMPUTING DIABATIC COUPLINGS*

Accepted Manuscript

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

8 Workflows and AI

Accepted Manuscript

8.1 Code-Agnostic Materials Science Libraries, Workflow Managers, and Visualization Tools

Jonas Björk¹, Volker Blum^{2,3}, Felix Hanke^{4,a}, Florian Knoop^{5,1}, Alexander Knoll⁶, *Sebastian Kokott^{5,7}, Ask Hjorth Larsen⁸, Akhil S. Nair⁵, Thomas A. R. Purcell^{5,9,b}, Simon P. Rittmeyer¹⁰, Matthias Scheffler⁵, and *Andrei Sobolev^{5,7}

¹Department of Physics, Chemistry and Biology (IFM), Linköping University, SE-581 83, Linköping, Sweden

²Thomas Lord Department of Mechanical Engineering and Materials Science, Duke University, Durham, NC 27708, USA

³Department of Chemistry, Duke University, Durham, NC 27708, USA

⁴Theory Department (since 1/1/2020: The NOMAD Laboratory), Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁶Research Center Chemical Sciences and Sustainability, Lehrstuhl für Theoretische Chemie II, Ruhr-Universität Bochum, Gesundheitscampus Süd 25, 44801 Bochum

⁷Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

⁸CAMD, Department of Physics, Technical University of Denmark, 2800 Kgs. Lyngby, Denmark

⁹Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

¹⁰Chair for Theoretical Chemistry and Catalysis Research Center, Technische Universität München, Lichtenbergstr. 4, 85747 Garching, Germany

* Coordinator of this contribution.

^aCurrent Address: CuspAI Ltd, Cambridge, CB1 2GE, United Kingdom

^bCurrent Address: Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

Summary

With routine access to peta-scale resources and the rise of exascale computing, data-centric computational materials science faces significant challenges, such as creating and managing large datasets, visualizing and analyzing diverse data, and assessing the quality of simulation results. To address these issues, the computational materials science community has focused on developing abstractions across all layers of materials simulations — ranging from representing geometries of a material and computing physical observables to developing a unified language for workflows and providing graphical user interfaces for data visualization. Over the past decade, these concepts have been realized through cross-institutional collaborations that support most existing DFT codes. In this section, we highlight the contributions of the FHI-aims community to these developments or examples that demonstrate the use of code-agnostic frameworks. To avoid any misunderstanding, the use of any of the packages mentioned below should be acknowledged by citing the original publications given by the references in this article.

FHI-aims integrates with the Atomic Simulation Environment (ASE) [43] and the Python Materials Genomics

*Roadmap on Advancements of the FHI-aims Software Package***8.1. CODE-AGNOSTIC MATERIALS SCIENCE LIBRARIES,
WORKFLOW MANAGERS, AND VISUALIZATION TOOLS**

(pymatgen) library [713], two widely adopted Python frameworks that offer code-agnostic objects for atomic structures and controlling simulations. To concatenate many calculation steps and automate their execution, several materials science workflow frameworks have been developed, including atomate2 [714], TaskBlaster [715], and AiiDA [716], providing support for FHI-aims. Visualization tools such as Vesta [717], XCrySDen [718], VMD [719], jmol [720], the ASE GUI, and the Graphical Interface for Materials Science (GIMS) [721] enable rendering three-dimensional geometries from the FHI-aims input format and analyzing structural properties. Additionally, Vesta, jmol, and XCrySDen can visualize volumetric data (such as cube files). GIMS, a browser-based toolbox, also facilitates the generation of input files for basic workflows, along with the analysis and visualization of results derived from output files.

Current Status of the Implementation

FHI-aims is driven and controlled mainly by the input file interface. To increase efficiency by automation and to standardize the input file generation, tools written in a high-level programming language are usually used. Python has been established as the language of choice to implement such tools. Python is easy to learn due to its straightforward syntax, making scripting and programming accessible even to scientists without programming experience. Further, the vast amount of third-party scientific Python libraries provide tools to solve numerical problems quickly and efficiently. Promoted by the two initiatives, the **Materials Project** and the **NOMAD Project**, two Python software packages have been established in the materials science community: The Python Materials Genomics (pymatgen) library [713] and the Atomic Simulation Environment (ASE) [43]. These libraries provide code-agnostic classes for representing structures, providing code-specific file I/O for input and output files, and computing physical observables from elemental tasks such as ground state calculations, structure relaxation, or molecular dynamics (MD). Both libraries come with functions that allow creating the two FHI-aims input files (`control.in` and `geometry.in`) and extracting relevant physical quantities, e.g., the total energy, forces, stress, chemical potential, and structure trajectories from output files. In addition, ASE is able to run FHI-aims both externally, through the Aims calculator interface, and interactively via the socket interface (the `SocketIOCalculator` class) adapting infrastructure from the i-PI implementation [561]. Through the socket interface, the atomic structure, its forces and, for periodic structures, stress is reported to the calling ASE function after a full self-consistent field cycle (SCF) is completed. In turn, ASE can process the new inputs and return a new structure to FHI-aims; after that, another SCF cycle is evaluated until a user-defined numerical convergence criterion is fulfilled. A full list of features would exceed the scope of the contribution, but can be found in online documentation of **ASE** and **pymatgen**.

The computational analysis of a material consists of a set of standard workflows. The atomate2 [714] project, based on the pymatgen package, has collected many of these workflows in one framework. The actual execution of the workflows is facilitated by `custodian` [713], a job management framework; `jobflow` [722], an abstract language to compose and connect workflows; and `Fireworks` [526], a framework for executing calculations on high-performance computing (HPC) systems and providing advanced database infrastructure. FHI-aims has now also added support for many of the atomate2 workflows, such as structure optimization, electron band structure calculation with DFT, GW, and other excitation methods, harmonic phonon calculation, computing elastic constants and the equation of state, anharmonicity quantification, and molecular dynamics. Many more features will be added in the near future. `Taskblaster` [715] provides a lightweight Python framework for designing and managing workflows and can be extended to support ASE classes and functionality. By using the ASE interface, FHI-aims can be employed to carry out multiple tasks (e.g. structure relaxation, band structure calculation etc.) over a large number of systems. For one of the most advanced

*Roadmap on Advancements of the FHI-aims Software Package*8.1. CODE-AGNOSTIC MATERIALS SCIENCE LIBRARIES,
WORKFLOW MANAGERS, AND VISUALIZATION TOOLS

workflow managers, AiiDA [716], support for FHI-aims was also added recently. AiiDA's core ecosystem is built around high-throughput workflow construction, data provenance, and efficient facilitation of HPC resources. It helps researchers organize, share, and reproduce complex scientific workflows efficiently. All the mentioned workflow managers provide a user-friendly command-line interface for workflow inspection and support for the major job scheduling systems, e.g., SLURM.

Visualization is an integral part of understanding and analyzing data — especially the rendering of 3D structure models is an important human consistency check and source of new ideas. Many visualization tools have added support for the FHI-aims input file `geometry.in`. Among them are Vesta [717], XCrySDen [718], VMD [719], jmol [720], the ASE GUI, and the Graphical Interface for Materials Science (GIMS) [721]. These visualisation tools allow measuring interatomic distances, the angle of an atom triple, and the torsion angle of two atom pairs, obtaining crystal symmetry information, and many more features.

Web-based applications require no setup and are built around user convenience and data visualization. Examples include the Graphical Interface for Materials Simulations (GIMS) [721]. GIMS supports FHI-aims, but is easily extendable to any other DFT code, as demonstrated for the *exciting* [46] code in the current version. GIMS was designed for easy manipulation and visualization of atomic structures, as well as generating corresponding input files. A mechanism of input keyword constraints and consistency checks is implemented to ensure that the results of a calculation with the generated input files are meaningful. GIMS is also able to parse, analyze and plot important FHI-aims output files and provide downloadable figures, such as convergence graphs, band structures, densities of electronic states, and dielectric functions.

Usability and Tutorials

All of the described Python packages are available on Python Package Index and can be installed with standard Python tools, like `pip`. The majority requires Python 3.10, and the basic functionality is available to the user right after the installation. Nevertheless, further configuration is required following the installation process in order to allow for the integration of the code. The configuration for FHI-aims along with the possible usage scenarios is described in the tutorials on high-throughput calculations [723–726].

A tutorial specifically written for the use of ASE with FHI-aims is available as part of our tutorial collection [723]. The tutorial introduces the `Atoms` object and the ASE calculators, which are used for the setup of a calculation, demonstrates nudged-elastic band workflow implemented in ASE, as well as showcases the use of the socket interface for a structure optimization. A pragmatic use for the ASE calculator when experimenting with DFT calculations is to compile the complete input files using the `write_input` method. The use of `pymatgen` is documented as part of a tutorial about `atomate2`. The `Structure` object is the `pymatgen` analog to the ASE `Atoms` object. The `pymatgen` tutorial demonstrates how to create the FHI-aims input files and explains the concept of the `pymatgen` input sets, which form the basis of the workflows. Eventually, it is explained how the FHI-aims output files can be parsed with the `AimsOutput` class. It is important to note that the use cases of ASE and `pymatgen` are by far not limited to the applications in the tutorials and this article: the concepts are widely adopted in several Python packages enabling writing of computational software-agnostic Python code. Such a code greatly reduces the cost of research and is thus of great benefit to the entire community.

`Atomate2`, being made of several distinct parts, also requires an advanced setup configuration. The workflows are written in the `jobflow` language and can be executed either locally or on the HPC resources using `Fireworks` or `jobflow-remote` workflow managers. Both workflow managers also use `MongoDB` to store information about calculation inputs and outputs as well as HPC queue status. `jobflow-remote` does not need

Roadmap on Advancements of the FHI-aims Software Package

8.1. CODE-AGNOSTIC MATERIALS SCIENCE LIBRARIES,
WORKFLOW MANAGERS, AND VISUALIZATION TOOLS

Name	Content
FHI-aims with ASE	- Installation - ASE Atoms and AimsCalculator - Running calculations - Using socket interface
High-throughput workflows with FHI-aims and atomate2	- Installation and configuration - Pymatgen support for FHI-aims - Executing workflows locally and remotely
Working in FHI-aims with Taskblaster	- Installation - Preparing input files - Running workflows
Working with FHI-aims and AiiDA	- Installing AiiDA and FHI-aims plugin - Running calculations - Accessing results
Managing Research Data with AiiDA and FHI-aims	- Using <i>aiida-ase</i> plugin to work with FHI-aims - Running single point calculation - Running structure optimization

Table 8.1: The list of available tutorials for the Python infrastructure of FHI-aims

access to MongoDB from the HPC. Both workflow managers require that the remote HPC center must allow non-interactive passwordless login. The steps needed to be done to configure and use atomate2 for a simple calculation are outlined in the tutorial [724].

Taskblaster is a workflow tool similar to jobflow, integrated into the ASE ecosystem. Through plugins it allows the execution of multiple tasks (e.g. structural relaxation, bandstructure calculation, postprocessing analysis etc.) via command-line interface on HPC resources. The installation and configuration, as well as the integration of FHI-aims into Taskblaster is demonstrated by the tutorial "FHI-aims with Taskblaster" [725].

AiiDA, in its turn, requires PostgreSQL database and RabbitMQ message broker to be installed in a production environment; however, limited functionality is available even without them. AiiDA has a extensive plugin ecosystem — FHI-aims can be used alongside AiiDA with either ASE or dedicated FHI-aims plugin. The dedicated FHI-aims plugin for AiiDA supports running generic calculations; more specialized workflows will be available in the near future. Several tutorials on working with AiiDA and FHI-aims are available online [726, 727].

All Python package tutorials and their content are summarized in Table 8.1.

GIMS, as noted before, does not require any installation. It is a Web application that consists of a collection of apps, such as manipulating atomic structure (Structure Builder), generating input files (Control Generator) and parsing the outputs (Output Analyzer). These apps are called elemental apps. The elemental apps can be chained together to form workflows (Calculation Apps). There are several such apps available for different calculation kinds, including single point, MD, band structure, and GW. Different Calculation Apps have different constraints and consistency checks implemented to ensure the user get the control file with parameters most suitable for their specific calculation. We host a stable release version of GIMS [728], as well as a development version [729] in the Internet. However, also standalone versions of GIMS are released for all major operating systems. It is possible to install GIMS on local machines and laptops. Detailed installation instructions and tutorials for GIMS can be found in the manual [730].

8.1. CODE-AGNOSTIC MATERIALS SCIENCE LIBRARIES, WORKFLOW MANAGERS, AND VISUALIZATION TOOLS

Future Plans and Challenges

Maintenance, continuous support, and adding new features form the corners of successful open source projects. While new packages and features are often part of new research, the FHI-aims community will stay dedicated to continuously support and help to maintain open source projects via platforms like the nonprofit organization Molecular Simulations from First Principles e.V. As already indicated above, both major Python computational materials science ecosystems, ASE and pymatgen, require Python functions to generate input files and parse output files. At the moment, each package has its own infrastructure, which complicates their maintenance. A centralized Python package for FHI-aims specific file I/O is currently under development under the name `pyfhiaims`. The package will unify the file I/O functions and provide an API that allows the seamless integration into the ecosystems as a dependency. The big advantage of the centralized `pyfhiaims` package is that changes in the in- and output files of FHI-aims can be faster integrated. Subsequently, more parsers for the various output files, like the band structure and DOS files, will be added. Also a sophisticated API for error handling will be implemented. The error handling is an important part in any workflows to determine a next step in case of a FHI-aims calculation fails. Support for pymatgen and atomate2 was only added recently. More features and integration of FHI-aims in the atomate2 workflows will be added, such as adding anharmonic phonons, magnetic ordering, and defect workflows in atomate2.

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

FK acknowledges support from the Swedish Research Council (VR) program 2020-04630, and the Swedish e-Science Research Centre (SeRC).

8.2 Workflows for Artificial Intelligence

Jörg Behler^{1,2}, Christian Carbogno^{3,4,a}, Gábor Csányi⁵, *Lucas Foppa³, Kisung Kang^{3,b}, Roman Kempt⁶, Marcel F. Langer⁷, Johannes T. Margraf⁸, *Akhil S. Nair³, Thomas A. R. Purcell^{3,9,c}, Patrick Rinke^{10,11,12,13}, Matthias Scheffler³, Alexandre Tkatchenko^{14,15}, Milica Todorović¹⁶, Oliver T. Unke¹⁷, and Yi Yao¹⁸

¹Lehrstuhl für Theoretische Chemie II, Ruhr-Universität Bochum, D-44780 Bochum, Germany

²Research Center Chemical Sciences and Sustainability, Research Alliance Ruhr, D-44780 Bochum, Germany

³The NOMAD Laboratory at the Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁴Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

⁵Department of Engineering, University of Cambridge, Cambridge CB2 1PZ, U.K.

⁶Technische Universität Dresden, Theoretische Chemie, 01069 Dresden, Germany

⁷Laboratory of Computational Science and Modeling, Institut des Matériaux, École Polytechnique Fédérale de Lausanne, 1015 Lausanne, Switzerland

⁸Bavarian Center for Battery Technology (BayBatt), University of Bayreuth, Weiherstraße 26, 95448, Bayreuth, Germany

⁹Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

¹⁰Department of Applied Physics, Aalto University, P.O. Box 11000, FI-00076 Aalto, Finland

¹¹Physics Department, TUM School of Natural Sciences, Technical University of Munich, Garching, Germany

¹²Atomistic Modelling Center, Munich Data Science Institute, Technical University of Munich, Garching, Germany

¹³Munich Center for Machine Learning (MCML), Munich, Germany

¹⁴Department of Physics and Materials Science, University of Luxembourg, L-1511 Luxembourg, Luxembourg

¹⁵Institute for Advanced Studies, University of Luxembourg, L-1511 Luxembourg, Luxembourg

¹⁶Department of Mechanical and Materials Engineering, University of Turku, 20014 Turku, Finland

¹⁷Google DeepMind, Berlin, Germany

¹⁸Molecular Simulations from First Principles e.V., Akazienstr. 3A, 10823, Berlin, Germany

*Coordinator of this contribution.

^aCurrent Address: Theory Department, Fritz Haber Institute of the Max Planck Society, Faradayweg 4-6, D-14195 Berlin, Germany

^bCurrent Address: Department of Materials Science & Engineering, Yonsei University, 50 Yonsei-Ro, Seodaemun-Gu, Seoul 03722, Republic of Korea

^cCurrent Address: Department of Chemistry and Biochemistry, The University of Arizona, Tucson, AZ 85721, USA

Summary

The efficiency and reliability of artificial-intelligence (AI) models used for physics, chemistry, biophysics, materials science, and engineering depend on the acquisition of sufficient high-quality data. Due to its all-

Roadmap on Advancements of the FHI-aims Software Package

8.2. WORKFLOWS FOR ARTIFICIAL INTELLIGENCE

electron, full potential treatment and its scalability to larger systems without precision limitations, FHI-aims provides accurate *ab initio* data from a wide range of computer simulations, such as electronic-structure calculations and molecular dynamics. To leverage the capabilities of AI models, workflows that seamlessly integrate AI tools with FHI-aims are essential. These workflows automate the data acquisition for AI and avoid the calculation of inessential data. Thus, they facilitate the iterative data exchange between AI models and simulations, allowing FHI-aims to be used as a powerful AI-guided calculation engine which puts its emphasis on information richness and not on mere data size. Also, interpretable AI models aid in analyzing the generated data. Finally, AI complements *ab initio* studies as it enables to perform simulations at larger time and length scales. In turn, the AI models must incorporate the physics required for an accurate representation of the *ab initio* data. This contribution highlights workflows developed to integrate FHI-aims with AI and future challenges.

Current Status of the Implementation

FHI-aims [8] provides various approaches to the provision and communication of data, which can then be used in workflows for the development of AI models such as machine-learning interatomic potentials (MLIPs). The interface with the Atomic Simulation Environment (ASE) [731] library provides a framework for the generation or loading of input files, calculation of properties, and storage of calculation outcomes. Interfaces with high-throughput-calculation tools such as atomate2 [714] and Aiida [726] also allow rapid acquisition of data for training AI models (see contribution 8.1). These interfaces are particularly crucial in workflows where the AI model is retrained iteratively with more data. These active learning (AL) workflows rely on data acquisition strategies informed by the AI model, which ensures that the new data to be collected is relevant. For instance, new data might be acquired when the prediction of the AI model has a high uncertainty or the interesting materials are found *out of distribution* with respect to the training data distribution [732]. In the following, we highlight examples of workflows involving FHI-aims and AI.

AL workflows using MLIPs such as high-dimensional neural network potentials (HDNNP) [733], Gaussian Approximation Potential (GAP) [734] and, more recently, equivariant message passing neural networks like MACE [735], have been designed based on training data generated by FHI-aims. Examples of this include workflows for crystal structure prediction [736], battery materials [737] or catalysis [738, 739]. In these applications, it proved essential to have large flexibility with respect to simulation types (including global optimization, transition-state searches, molecular dynamics and enhanced sampling methods), and training set selection (e.g., via uncertainty estimation or similarity based search). These requirements have led to the development of the wfl package, a Python toolkit for interatomic potential creation and atomistic simulation workflows that emphasizes modularity and parallelisation over sets of atomic configurations [740, 741].

Being based on local atomic orbitals, FHI-aims is well suited to build large-scale material models containing defect structures, grain boundaries and stacking faults, which are often challenging for other plane-wave based DFT codes. The potential energy surface of these materials can be sampled with high accuracy in FHI-aims, yielding suitable energies, forces and stresses to train MLIPs. Any ASE-compatible framework to train interatomic potentials can be used, such as DeePMD-kit, which implements the smooth version of the Deep Potential model [742, 743]. This approach has been employed to train a Deep Potential model for PtSe₂ nanoflakes up to 8.6 nm [744].

We note that it is important that the chosen MLIP model is able to describe all the relevant physics, since increasing the amount of data alone does not guarantee a model representing the system correctly. Examples are long-range electrostatic interactions beyond the local cutoff radii often employed in the construction of

MLIPs for condensed systems, dispersion interactions, and non-local charge transfer. The latter is crucial in many types of chemical reactions, e.g., if the charge of a molecule is altered by (de)protonation or an atomic oxidation state changes due to electron transfer. For these cases, often non-local approaches like fourth-generation MLIPs may be needed, which take the global structure of the system into account for describing electrostatics [745]. An alternative solution for small systems is to employ explicit global machine-learning force fields like GDML/BIGMDL [746, 747] or, more generally, a graph neural network architecture combined with physical models for long-range interactions, such as GEMS [748] or SO3LR [749]. Another package that has been developed linking FHI-aims and MLIPs is the GKX package [750] and the FHI-vibes framework (see contribution 6.1). By using GKX, MLIPs trained on high-fidelity data generated with FHI-aims can be used to perform GPU-accelerated MD simulations for systems with thousands of atoms over timescales of nanoseconds. Such simulations can be used, for instance, to obtain converged thermal transport coefficients [541].

Despite the growing number of applications of MLIPs, concerns about their reliability arise when they are utilized to predict properties associated with configurations or chemical species that are significantly different from those in the training set [751]. Kang *et al.* developed ALMOMD (Active-Learning Machine-Operated Molecular Dynamics [536, 752]), a Python workflow package interfacing FHI-aims with the MLIP codes NequIP [753] and so3krates [754]. ALMOMD is designed to effectively train MLIP through an AL scheme with an automated framework that samples unfamiliar data, e.g., rare events, based on uncertainty estimates of MLIP predictions (Figure 8.1).

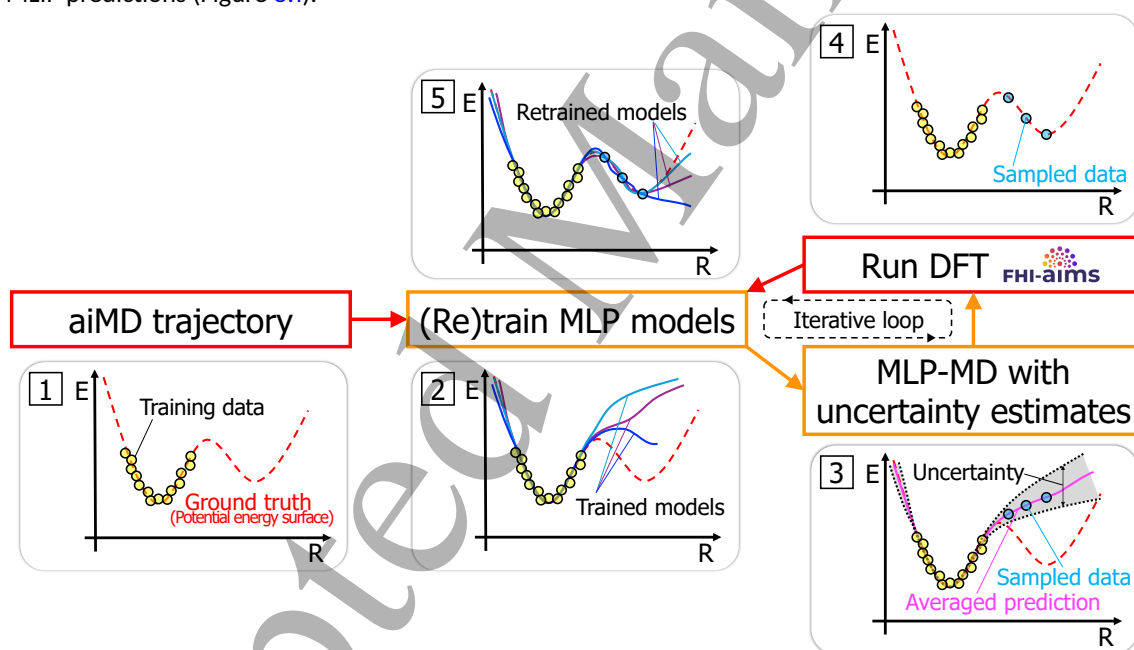


Figure 8.1: The overall iterative workflow of the ALMOMD. White boxes display indexed sequential steps for exploring the configurational space using MLIP-MD and sampling training data via uncertainty estimates.

While MLIPs hold great promise, many material problems are high-dimensional in nature and involve costly evaluation of an objective function. This can benefit from AL workflows that dramatically reduce sampling, such as those involving Bayesian optimization. The Bayesian Optimization Structure Search (BOSS) algorithm [755, 756] builds probabilistic N -dimensional surrogate models for materials energy or property landscapes,

Roadmap on Advancements of the FHI-aims Software Package

8.2. WORKFLOWS FOR ARTIFICIAL INTELLIGENCE

then refines them with smart sampling. The strategic acquisition strategy of blending data exploitation with design space exploration ensures fast identification of optimal solutions. BOSS is a Python tool for materials optimization and interoperable with FHI-aims and ASE.

Finally, high-quality materials data are typically sparse, demanding data-efficient AI approaches. Moreover, interpretability, i.e., the ability to inspect the model's inference is critical, highlighting the importance of using descriptor-based AI methods [757]. Nair *et al.* developed an AL workflow integrating the sure-independence screening and specifying operator (SISSO) approach with FHI-aims [758]. SISSO is a symbolic-regression method that utilizes compressed sensing to identify analytical expressions correlated with materials' properties or functions [759, 760]. It is a data-efficient method and offers better interpretability compared to widely used ML approaches in materials science. SISSO identifies analytical expressions that contain key physicochemical parameters, from many offered ones. The developed workflow utilizes an interface of FHI-aims with the high-throughput utility Taskblaster [715] for executing multiple tasks, such as geometry optimization, band-structure calculation, etc., for a large number of materials. Such workflows achieve efficient data acquisition and mitigate the issue of redundant data [761], making them suitable for generating information-rich materials databases (e.g., by including materials with high prediction uncertainty) which could be further used to develop AI models with enhanced generalizability.

Usability and Tutorials

This section illustrates the applications of the workflows with tutorials, that researchers can adapt to their specific projects. The ALMOMD framework is demonstrated in the context of atomistic simulations of strongly anharmonic materials. Incomplete MLIP training often happens due to the absence or insufficiency of data within training regions. For example, the MLIPs may be unable to predict rare dynamical events, like defect creations, that are not included in training data or regularized away in the learning as they are rare. Consequently, it leads to critical deviations in predictions for transport properties. The ALMOMD framework can actively learn these unfamiliar data missed during MLIP training and correct the potential erroneous predictions during molecular dynamics (MD) simulations. It also corrects for fake rare events that an MLIP may produce.

ALMOMD consists of two important steps: exploration and data-sampling. The efficient exploration of configurational space is achieved by explorative MD employing MLIPs (MLIP-MD). Uncertainty estimates serve as a warning indicating when MLIP-MD goes beyond its trained area, and thus, it can identify unfamiliar data that need to be sampled and retrained for MLIPs in subsequent steps. ALMOMD provides the user with an automated workflow environment and on-line tutorials [762].

BOSS was applied to study molecular conformers [763] and surface adsorbates [764, 765], thin film growth [766], solid-solid interfaces [767] and it can also combine multiple fidelity simulations. The computation workflow illustrated in Figure 8.2 relies on uncertainty-aware and interpretable surrogate model landscapes to extract optimal solution basins, from which structural optimization leads to final structures and associated functional properties. The BOSS website facilitates adaptations of this workflow to different use cases [756], with the code, manual and extensive tutorials available to the research community

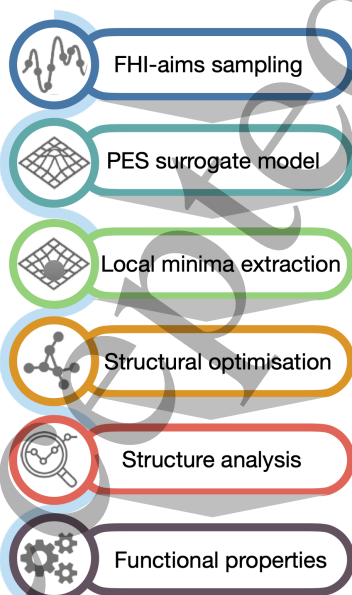


Figure 8.2: BOSS AI workflow: from surrogate models to optimal solutions.

8.2. WORKFLOWS FOR ARTIFICIAL INTELLIGENCE

[768, 769].

The SISSO-based AL workflow [758] is applied to the efficient discovery of acid-stable oxides for water splitting reaction from a large space of candidate materials, i.e., with a reduced number of calculations compared to high-throughput screening (Figure 8.3). Ensembles of SISSO models are used to obtain mean predictions as well as estimates of the prediction uncertainties. This opened the opportunity to use SISSO as a surrogate model in the aforementioned Bayesian optimization approach. DFT calculations were carried out for these materials by leveraging an efficient implementation of hybrid functionals [6]. A tutorial demonstrating the workflow can be found at ref [770]. The workflow has also been transformed into a user-friendly web app [771], featuring a step-by-step interactive demonstrator that guides users with limited AI experience in utilizing it effectively. Such a workflow reduces the risk of overlooking potentially interesting portions of the materials space that were disregarded in the initial training data and hence enables efficient materials discovery.

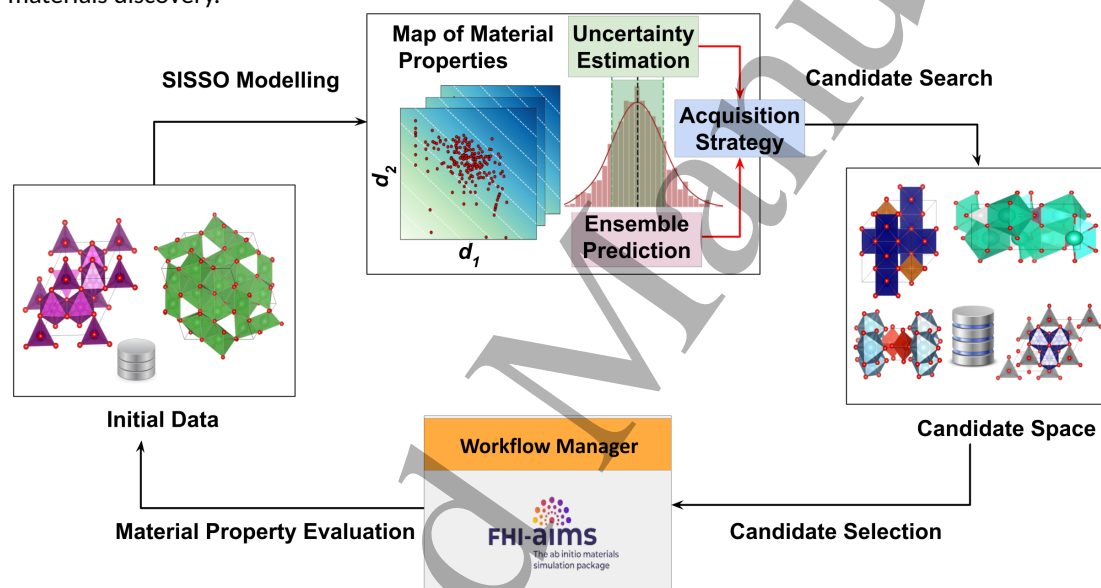


Figure 8.3: Schematic representation of the workflow integrating SISSO and FHI-aims. d_1 and d_2 represent the descriptors from the SISSO model constituting a materials map where the initially available data is labeled with red circles.

Future Plans and Challenges

Currently, the majority of MLIP-based methods which have been integrated into workflows to guide FHI-aims simulations, function as standalone packages. The next step is to enable direct execution of MLIPs within the FHI-aims environment, creating a more streamlined process that integrates MLIPs into the simulation workflow without the need for separate executions. For AL workflows, obtaining accurate and reliable uncertainty estimates is challenging, as overconfidence could lead to inefficient sampling of the materials or configuration spaces [772]. Additionally, for systems like strongly correlated materials, workflows need to be adapted to integrate advancements in beyond-DFT methods, such as GW or RPA.

*Roadmap on Advancements of the FHI-aims Software Package***8.2. WORKFLOWS FOR ARTIFICIAL INTELLIGENCE**

Apart from the use as a calculation engine for data acquisition, AI models could also be used to accelerate FHI-aims calculations or improve their accuracy. For example, promising research directions include AI-based initial guesses for wavefunctions based on learning Kohn-Sham matrices [773, 774], the prediction of electron density in 3D space [775] or the development of novel density functionals [776, 777]. The training data for such approaches could itself be generated by FHI-aims, allowing to improve AI models in a feedback loop. Such methods and their potential applications are discussed in greater detail in the contribution (8.3).

Acknowledgements

MS acknowledges support by his TEC1p Advanced Grant (the European Research Council (ERC) Horizon 2020 research and innovation programme, grant agreement No. 740233).

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

8.3 Machine Learning the Electronic Structure in a Real-Space and Atom-Centered Framework

Joseph W. Abbott¹, Michele Ceriotti², Andrea Grisafi², Wei Bin How¹, Alan Lewis³, Andrew J. Logsdail⁴, Zekun Lou⁵, Reinhard J. Maurer^{6,7,8}, *Mariana Rossi⁵, Pavel V. Stishenko⁴, Zechen Tang⁹, and Yong Xu^{9,10}

¹Institute of Materials, École Polytechnique Fédérale de Lausanne, Rte. Cantonale, Lausanne 1015, Switzerland

²Physicochimie des Électrolytes et Nanosystèmes Interfaciaux, Sorbonne Université, CNRS, F-75005 Paris, France

³Department of Chemistry, University of York, Heslington, York, YO10 5DD, UK

⁴Cardiff Catalysis Institute, School of Chemistry, Cardiff University, Park Place, Cardiff CF10 3AT, United Kingdom

⁵Max Planck Institute for the Structure and Dynamics of Matter, 22761 Hamburg, Germany

⁶Department of Chemistry, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

⁷Department of Physics, University of Warwick, Gibbet Hill Road, CV4 7AL Coventry, United Kingdom

⁸Faculty of Physics, University of Vienna, Vienna A-1090, Austria

⁹Department of Physics, Tsinghua University, Beijing 100084, P.R.China

¹⁰RIKEN Center for Emergent Matter Science (CEMS), Wako, Saitama 351-0198, Japan

*Coordinator of this contribution.

Summary

Machine-learning (ML) methods are driving a revolution in electronic structure simulations. Many approaches target the learning and prediction of energies, forces and stresses that are produced by a density functional theory (DFT) calculation, with the goal of training high-accuracy force-fields that can be used to investigate atomic structural and dynamical properties of materials (see 8.2). However, one drawback of such models is that they do not give access to electronic-structure information, contrary to explicit quantum calculations that, in addition to energies and forces, also provide a wealth of derived electronic structure properties.

A route to access these observables is to train machine learning models to learn a specific property, such as the electronic band gap, dipoles, surface work functions, and many others. However, a more transferable solution exists in methods that directly target the electronic-density or the Kohn-Sham Hamiltonian (in the framework of DFT). When successful, these approaches are advantageous because, with a single ML model, one can recover most electronic structure properties and, in principle, also calculate accurate energies and forces.

The field of ML for electronic structure is admittedly still in its infancy, despite a very fast pace of progress in recent years. Many different approaches have been reported in the literature, based on a diverse set of underlying ML models and basis representations. In this contribution, we focus on methods that are directly interfaced to FHI-aims, namely SALTED [83], rholearn [778], doslearn [778, 779], ACEhamiltonians.jl [780, 781] and DeepH [782-784].

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

FHI-aims offers unique benefits for ML surrogate models of electronic structure. The numeric atomic-orbital (NAO) basis offers a compact, accurate, and local representation of the electronic structure and yields compact Hamiltonian, overlap, and density matrices. The seamless treatment of periodic and non-periodic systems means that molecules and solids can be treated with the same infrastructure, making the training of models easier and more versatile. Furthermore, the density fitting (resolution-of-identity) infrastructure that has been developed for hybrid functionals within the NAO basis [62] provides the ingredients to represent various real-space quantities through a set of basis coefficients with known rotational symmetry properties. Finally, the Atomic Simulation Interface (ASI) [785], developed in FHI-aims, enables plain C and Python application programming interfaces, which are beneficial for integration with software packages and libraries as they allow in-memory data transfer with minimal latency and storage overheads.

Current Status of the Implementation

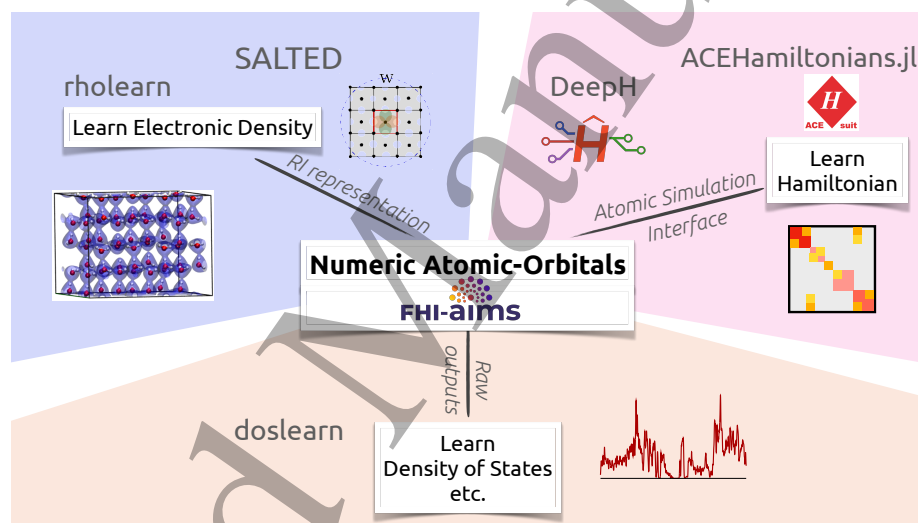


Figure 8.4: A sketch of the ecosystem of machine-learning (ML) methods for the electronic structure that are interfaced to FHI-aims. The raw outputs of the code, the resolution of the identity (RI) representation of the electronic density, and the atomic simulation interface (ASI) enable diverse types of ML algorithms to be used.

Machine-learning methods that interface with FHI-aims and target the electronic structure are, so far, all based on the real-space representation of the Kohn-Sham problem or the density from a quantum mechanical calculation. The challenge of learning these quantities reduces to representing atom-centered local density contributions or intra- and inter-atomic matrix blocks. Because of the common representation of the electronic density $n(\mathbf{r})$ in isolated and periodic systems, FHI-aims offers a seamless route to learn $n(\mathbf{r})$ for mixed datasets. For the Hamiltonian of periodic systems, once the real-space Hamiltonian in the extended crystal volume is known, the Hamiltonian at arbitrary k -points can be evaluated in a straightforward manner, similar to tight-binding techniques.

We start by summarizing methods that target $n(\mathbf{r})$. SALTED is a machine learning method based on a symmetry-adapted Gaussian process regression.[83] Using a resolution-of-the-identity (RI) expansion, $n(\mathbf{r})$ (from LDA, GGA, or hybrid-functionals) is first expressed as a linear combination of atom-centred basis func-

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

tions $\phi_{i\sigma}$:

$$n(\mathbf{r}) \approx \tilde{n}(\mathbf{r}) = \sum_{i,\sigma,\mathbf{U}} c_{i\sigma} \phi_{i\sigma}(\mathbf{r} - \mathbf{R}_i - \mathbf{T}(\mathbf{U})) \quad (8.1)$$

where i labels the atom on which the basis function is centered, \mathbf{T} is a translation vector, and $\sigma = (n\lambda\mu)$ is a compound label describing the radial R_n and spherical $Y_{\lambda\mu}$ part of the basis function. In FHI-aims, these basis functions can be chosen from the auxiliary basis introduced to accelerate the calculation of hybrid functionals.[62] The RI expansion coefficients $c_{i\sigma} \equiv c_{\sigma}(A_i)$ are then themselves approximated as a linear combination of a similarity measure $k_{\mu\mu'}^{\lambda}$ between the atomic environment A_i associated with the target coefficient and each reference atomic environment M_j , and their associated regression weights $b_{\mu j}^{\lambda}$ to be determined using SALTED:

$$c_{\sigma}(A_i) = \sum_j \sum_{\mu' \leq \lambda} b_{\mu j}^{\lambda} k_{\mu\mu'}^{\lambda}(A_i, M_j). \quad (8.2)$$

These similarity measures are symmetry adapted, being tensors of dimension $2\lambda + 1$ that transform as the λ -th irrep of the $O(3)$ group, since the basis functions and their associated coefficients transform in this way. In SALTED, atom-density based descriptors are computed with the featomic [786] library. These can be smooth-overlap of atomic position (SOAP) descriptors, or descriptors that contain information about long-range electrostatics (e.g., LODE descriptors).

For a full description of the calculation of the regression weight-vector \mathbf{b} , which lies at the core of the symmetry-adapted Gaussian process regression of the SALTED method, the reader is referred to Refs. [83, 787]. Here we highlight that, in order to calculate \mathbf{b} , two quantities are required for each structure in the training set: \mathbf{S} , the overlap matrix of the basis functions ϕ_i ; and the optimal expansion coefficients $\mathbf{c}^{\text{RI}} = \mathbf{S}^{-1}\mathbf{w}$, where \mathbf{w} is a vector containing the projection of the self-consistent electron density onto each basis function $\phi_{i\sigma}$.^a

The calculation of \mathbf{S} , \mathbf{w} and \mathbf{c}^{RI} has been implemented in FHI-aims, with the option to write these quantities to files for later use with SALTED to train a ML model. Additionally, SALTED has a number of FHI-aims specific “helper” functions to support integration between the two codes. These include functions to calculate the error introduced by the RI approximation, to help users find an accurate set of basis functions $\phi_{i\sigma}$, to convert the data output from FHI-aims into a suitable format to be read by SALTED, and to convert coefficients predicted using SALTED into a format that can be read by FHI-aims. In the latter case, FHI-aims has the capability to read in a set of expansion coefficients $c_{i\sigma}$ during the SCF initialisation, and construct from them $n(\mathbf{r})$ on the real-space integration grid used in the code. This approach replaces the default choice of the initial electron density (i.e., superposition of atomic densities). In preliminary tests with a SALTED model trained on a subset of around 9000 structures of the QM9 dataset [788] (see Ref. [787]) we have observed, on average, that this approach leads to a 35% reduction in the number of SCF steps needed for convergence.

The infrastructure built for SALTED in FHI-aims, including the RI representation for $n(\mathbf{r})$, can be re-used for ML schemes that use a neural network (NN) architecture rather than kernel regression. One such method is rholearn [778]. While the inputs (nuclear coordinates) and outputs (RI coefficients) of the prediction pipeline are the same, there are a few key differences compared to SALTED. The featomic [786] library is used to generate equivariant descriptors of arbitrary body order through Clebsch-Gordan products. These descriptor vectors, as opposed to the kernels used in SALTED, form the part of the typically (but not nec-

^aThe weights are $w_{i\sigma} = \sum_{\mathbf{U}}^{\text{cut}} \langle \phi_{i\sigma}(\mathbf{U}) | n \rangle_{\text{u.c.}}$. The subscript “u.c.” indicates that in FHI-aims the integral is performed only over the “central” unit cell in the case of periodic systems, with the sum over \mathbf{U} being truncated to only include unit cells that contain basis functions with some support in this central unit cell. The overlap matrix \mathbf{S} is evaluated in a similar way.

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

essarily) fixed transformations of the nuclear coordinates into an intermediate representation. Then, the training workflow uses the metatensor-torch library as its backbone. Native torch modules for tensor operations and learning utilities are wrapped to be compatible with TensorMap, the sparse data storage format of the metatensor [786] library. This allows arbitrarily complex equivariance-preserving NNs to be defined, and applied to the equivariant descriptors. Finally, the use of descriptor vectors, the ability to train by mini-batch gradient descent, and the use of sparse operations (most notably when evaluating the loss) reduce the memory overhead of rholearn.

Models that target learning the (Kohn-Sham) Hamiltonian matrix profit from a more tightly integrated interface with FHI-aims for AI/ML workflows. This interface is realised via ASI [785], which is a plain C interface that provides direct in-memory access to electronic structure quantities within FHI-aims, such as the Hamiltonian, and overlap matrices. It also allows to inject AI/ML predicted matrices into FHI-aims processes at runtime, providing new opportunities in modular software integration [781]. The integration of ACEhamiltonians.jl and DeepH to FHI-aims use this interface.

ACEhamiltonians.jl [780] uses atom- and bond-centred descriptors based on the Atomic Cluster Expansion (ACE) [789] to represent non-orthogonal Hamiltonian matrix blocks that transform equivariantly with respect to the full rotation group. ACEhamiltonians models are analytical and linear, and can produce accurate models with little data. The model can directly be trained on FHI-aims data that is outputted either via the keyword `output_rs_matrices` in human readable or HDF5 format, or through the ASI interface. Furthermore, the model is also integrated with FHI-aims through ASI to directly feed predicted quantities such as the Hamiltonian or the density matrix back into FHI-aims, e.g., to provide an improved initial guess of the density matrix for the SCF algorithm. We have also shown that ASI can be used as a bridge to train ACEhamiltonians models on FHI-aims data and then inject them into a different electronic structure code, such as DFTB+, during runtime. [781]

The deep-learning density functional theory Hamiltonian (DeepH) method [782–784] is an equivariant graph neural network approach for learning and predicting the electronic Hamiltonians for given material structures. Based on the predicted Hamiltonian, various physical properties can also be derived, including band structures, density of states, and the electric susceptibility, among others. Benefiting from the principle of nearsightedness, the neural networks of DeepH can infer the properties of large-scale material structures by learning from small ones. In the original version of DeepH developed in 2021 [782], a local-coordinates-based scheme was proposed to handle the rotational covariance of the DFT Hamiltonian matrix. A subsequent implementation, DeepH-E3 [783], incorporated an E(3)-equivariant neural network architecture to address the equivariance with respect to the E(3) group, vastly improving the prediction accuracy. Recently, more advanced equivariant tensor product techniques together with transformer architectures have been integrated into the DeepH framework, further enhancing both accuracy and efficiency [790]. The method is versatile and has also been extended to various deep-learning electronic structure schemes, including magnetic materials [784], density-functional perturbation theory [791], hybrid density functionals [792], among others. The current implementation of the DeepH interface for FHI-aims is based on the ASI package [785]. Structural information, Hamiltonian matrices and overlap matrices are extracted from the ASI plug-ins and converted to DeepH formats. The current interface has been tested on both molecular and periodic systems, in which the open source DeepH-E3 method [783] reaches sub-meV mean absolute error in Hamiltonian matrix elements across the test systems.

Finally, there are ML methods interfaced with FHI-aims that parse raw outputs of electronic-structure quantities from FHI-aims. doslearn targets the electronic density of states (DOS), employing a locality ansatz in which the total DOS of a structure is built as a sum of atom-centered contributions [793]. The machine learn-

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

ing pipeline directly takes in FHI-aims outputs, extracts the Kohn-Sham eigenvalues for each structure, and uses `scipy` to generate cubic Hermite splines of the DOS to support an adaptive energy reference, necessary for bulk periodic calculations. Model training is performed using the PyTorch package, with the adaptive reference framework supported *via* a modification of the loss function to concurrently optimize the energy reference during model training. Although the current `doslearn` implementation employs a simple feed forward network using invariant SOAP Power Spectrum descriptors generated from the `featome` [786] library as inputs, `doslearn` primarily focuses on the optimization procedure and is model agnostic, allowing it to be easily modified to deploy on any arbitrary model architecture of the user's choice.

Usability and Tutorials

The usability of the electronic-structure learning techniques described above together with FHI-aims is ensured by specific keywords, with examples and tutorials maintained either natively in the code or in dedicated repositories.

We begin by briefly describing the usability and tutorials of the electronic-density learning approaches. The SALTED code is written in Python and FORTRAN and it is available through a dedicated GitHub repository^b. `rholearn` is Python-based and is also available through a dedicated GitHub repository^c. Their installation and dependencies are described in their respective documentation.

Briefly, the FHI-aims keywords for `control.in` that trigger the outputs needed for SALTED and `rholearn` are:

- `ri_density_restart write` will create a file in the working directory containing the coefficients c^{RI} .
- `ri_density_restart read` will read the expansion coefficients contained in the file previously created (in the same directory) and replace the default initial density with the density obtained from these coefficients.
- `ri_full_output`, when used in conjunction with `ri_density_restart write`, will additionally output a file containing the overlap matrix S , and the file `ri_projections.out` containing the projections vector w .
- `ri_skip_scf` is used in conjunction with `ri_density_restart write` and allows RI decomposition of a previously-converged density as part of a two-step SCF + RI process. For instance, `elsi_restart write` can be used to save a converged solution to the SCF procedure, then `elsi_restart read` and `ri_skip_scf` can be used to fit directly to the corresponding converged density.

The keywords `prodbas_acc`, `max_l_prodbas`, and `wave_threshold` are not SALTED or `rholearn`-specific, but control the construction of the auxiliary basis that is used to expand the density. Specifying these keywords ensures a consistent basis is used, which is especially important if training on mixed periodic and non-periodic datasets.

The SALTED integration with FHI-aims is thoroughly described in an online tutorial^d. The tutorial guides the user on training a SALTED model for the electronic density of a water monomer, and predicting the density of

^b<https://github.com/andreagrisafi/SALTED>

^c<https://github.com/lab-cosmo/rholearn>

^d<https://fhi-aims-club.gitlab.io/tutorials/fhi-aims-with-salted>

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

water dimers. The rholearn integration with FHI-aims shares many of the features of the SALTED integration and is also described in [an online tutorial](#)^e. Data for a subset of the QM7 dataset [794, 795] is generated with the `aims-interface` in rholearn, and a descriptor-based equivariant neural network trained by minibatch gradient descent using a PyTorch- and metatensor [786]-based workflow.

The interfaces for Hamiltonian learning are based on ASI, as mentioned above. The ASI documentation is available via the library website [796]. The Python wrapper for ASI, `asi4py`, can be installed from the Python Package Index (PyPI) repository. Tutorials on how to train ACEhamiltonians models with example FHI-aims data for bulk Aluminium are provided on GitHub [797]. The tutorial covers how to parse and process FHI-aims Hamiltonian data generated with the `output_rs_matrices` keyword into the expected HDF5 database structure. It also covers essential keywords and hyperparameters for model construction, model fitting, and model prediction. ACEhamiltonians directly predicts the real-space representation of the Hamiltonian in the extended crystal volume. Matrices (and therefore eigenvalues and eigenvectors) during prediction can be constructed at arbitrary k-points.

Documentation on how to use ASI to provide an improved initialisation of the density matrix in FHI-aims during runtime is available in the `dm` submodule of `asi4py`^f. This submodule provides a prediction of the density matrix that can reduce the number of required self-consistent-field iterations, when compared to the standard initialization. The `ASI_register_dm_init_callback` function of the ASI API is used to register a user-provided callback function that returns an initial guess of the density matrix at the beginning of the SCF loop. The `asi4py.dm` module provides a few classes implementing density matrix prediction via ASI; these classes can be used directly, subclassed, or used as inspiration for custom density matrix predictors.

A tutorial for DeepH is available in Zenodo [798], which provides the source code for the interface with FHI-aims, along with demonstrations for generating datasets and two example datasets, each accompanied by configuration files for DeepH-E3 training. The source code for the interface is also distributed alongside FHI-aims. To utilize the interface, FHI-aims should be compiled as a shared library through ASI and the `asi4py` package should also be installed. The interface can be employed to extract all required data for DeepH training, utilizing the ASI API to access Hamiltonian and overlap matrices. The DeepH-E3 model [783] has been evaluated on water and graphene datasets provided in the Zenodo repository, achieving mean absolute errors of 0.31 and 0.28 meV, respectively, for the Hamiltonian matrix elements in the test data sets. The recently developed version DeepH-2 [790], which will be soon released, demonstrates significantly improved accuracy, with mean absolute errors of 0.08 and 0.04 meV for the water and graphene datasets mentioned above, respectively.

Finally, the doslearn integration with FHI-aims is available in the [the same repository](#)^g as rholearn, with an [an online tutorial](#) that covers the data processing pipeline and model training of a fully connected network using batch gradient descent. The keyword used to generate the energy eigenvalues necessary for doslearn is: `output postscf_eigenvalues`.

Future Plans and Challenges

The area of ML applied directly to the electronic-structure is growing at a fast pace. Many such methods, including the ones interfaced with FHI-aims discussed in this contribution, show high-accuracy in predicting the electronic-density and Hamiltonian matrix elements. The methods allow the prediction of electronic

^e<https://github.com/lab-cosmo/rholearn/tree/main/example/rholearn-aims-tutorial>

^fhttps://pvst.gitlab.io/asi/ml_dm.html

^g<https://github.com/lab-cosmo/rholearn>

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

structure properties of systems containing up to $\sim 10^5$ atoms when trained on much smaller structures. Efficiency in terms of speed and memory usage can still be bottlenecks, and these are currently being addressed by the community. It is exciting to see that such methods are already able to predict not only the ground-state electronic density and Hamiltonian, but also density linear-responses [469, 791, 799], for example. Remaining challenges in this area regard the proper and efficient handling of equivariant architectures for vector-field prediction.

It is further interesting to see the extension of methods such as DeepH to unsupervised learning [800], force fields for magnetic materials [801], and to train high-accuracy universal models covering over 20 elements in the periodic table and over 10,000 material structures [802]. For DeepH, despite its $O(N)$ scaling that enables highly efficient computation of large systems, there remain a few post-processing workflows for calculating physical properties that could be computationally expensive for studying large-size material systems. Future advances in efficient algorithms for property calculations, such as sparse-matrix techniques, Green's function methods, and Wannierization, will significantly enhance the practical applications of deep-learning electronic structure methods, allowing for the computation of a broader range of physical properties in large-scale material simulations.

Finally, to realise the full potential of ML for electronic structure, new infrastructure is necessary to integrate existing software with AI-enabling workflows. ML models and workflows to generate and manage data are increasingly available in FHI-aims. Data can be communicated inward and outward in raw or fitted forms, with the latter beneficial for reducing data quantities; Data can also be communicated to and from FHI-aims *via* file input/output (with minimal overheads), sockets-type interfaces, or through direct interfacing. Nevertheless, there is an ever growing need for flexible and robust interfaces that allow more user interaction with these calculations, and data extraction. This need motivates further efforts to support workflow connectivity and development of standardised data communication paradigms. For the direct access to in-memory data objects, development plans for ASI include more flexible and extendable interface that supports a wider range of data objects and diverse sparse matrix formats. Long-term aims include applying ASI to develop an infrastructure for integration with a broad range of ML models, which can then accelerate or refine electronic structure calculations.

Acknowledgements

We acknowledge fruitful discussions with Volker Blum, Ben Hourahine, Thomas Keal, Scott Woodley. We thank Adam McSloy and Connor L. Box for their contributions to Hamiltonian output functionality in FHI-aims.

AJL and PVS acknowledge funding by the UKRI Future Leaders Fellowship program (MR/To18372/1, MR/Yo34279/1). AJL, PVS, RJM, and MR acknowledge funding from the ARCHER2 eCSE Programme (eCSE03-10). RJM acknowledges support through the UKRI Future Leaders Fellowship programme (MR/So16023/1, MR/Xo23109/1), and a UKRI frontier research grant (EP/Xo14088/1). YX and ZT acknowledge funding from the Basic Science Center Project of NSFC (52388201), the Ministry of Science and Technology of China (2023YFA1406400), the National Natural Science Foundation of China (12334003, 12421004, and 12361141826), and the National Science Fund for Distinguished Young Scholars (12025405).

8.3. MACHINE LEARNING THE ELECTRONIC STRUCTURE IN A REAL-SPACE AND ATOM-CENTERED FRAMEWORK

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39
40
41
42
43
44
45
46
47
48
49
50
51
52
53
54
55
56
57
58
59
60

Accepted Manuscript

9 Appendix

9.1 Summary of the Delta-DFT test for the FHI-aims species defaults

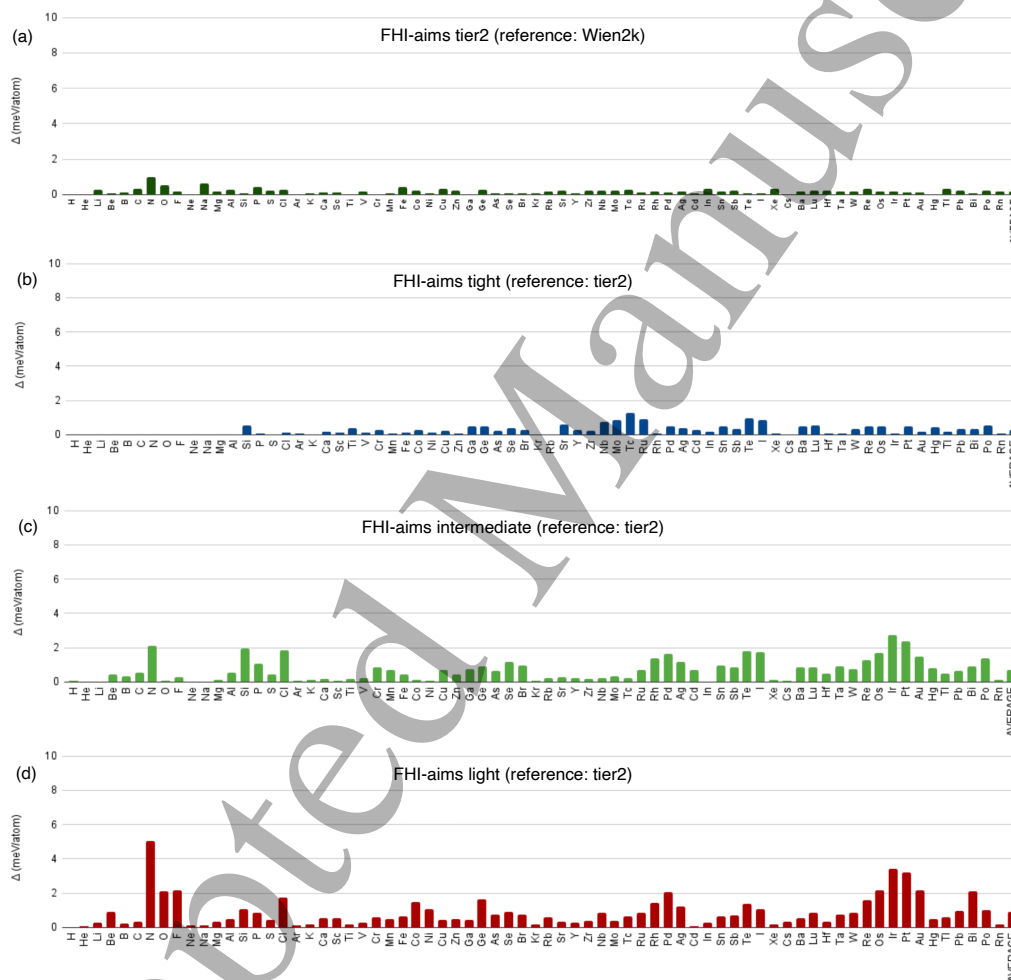


Figure 9.1: The Δ gauge [44] between (a) WIEN2k and FHI-aims with really tight settings and tier2 basis functions (tier2), (b) FHI-aims tight and tier2 (c) FHI-aims intermediate and tier2, and (d) FHI-aims light and tier2. The raw output data from the FHI-aims calculations is available at Ref.[47].

*Roadmap on Advancements of the FHI-aims Software Package**9.1. SUMMARY OF THE DELTA-DFT TEST FOR THE FHI-AIMS SPECIES DEFAULTS*

Accepted Manuscript

Bibliography

- [1] Volker Blum, Ryoji Asahi, Jochen Autschbach, Christoph Bannwarth, Gustav Bihlmayer, Stefan Blügel, Lori A Burns, T Daniel Crawford, William Dawson, and Wibe Albert et al. de Jong. Roadmap on methods and software for electronic structure based simulations in chemistry and materials. *Electronic Structure*, 6(4):042501, 2024.
- [2] Stefan Bauer, Peter Benner, Tristan Berau, Volker Blum, Mario Boley, Christian Carbogno, Richard Catlow, Gerhard Dehm, Sebastian Eibl, Ralph Ernstorfer, et al. Roadmap on data-centric materials science. *Modelling and Simulation in Materials Science and Engineering*, 2024.
- [3] Vikram Gavini, Stefano Baroni, Volker Blum, David R Bowler, Alexander Buccheri, James R Chelikowsky, Sambit Das, William Dawson, Pietro Delugas, Mehmet Dogan, et al. Roadmap on electronic structure codes in the exascale era. *Modelling and Simulation in Materials Science and Engineering*, 31(6):063301, 2023.
- [4] Andrew M Teale, Trygve Helgaker, Andreas Savin, Carlo Adamo, Bálint Aradi, Alexei V Arbuznikov, Paul W Ayers, Evert Jan Baerends, Vincenzo Barone, Patrizia Calaminici, et al. DFT exchange: sharing perspectives on the workhorse of quantum chemistry and materials science. *Physical chemistry chemical physics*, 24(47):28700–28781, 2022.
- [5] A Marek, V Blum, R Johanni, V Havu, B Lang, T Auckenthaler, A Heinecke, H-J Bungartz, and H Lederer. The elpa library: scalable parallel eigenvalue solutions for electronic structure theory and computational science. *Journal of Physics: Condensed Matter*, 26(21):213201, may 2014.
- [6] Sebastian Kokott, Florian Merz, Yi Yao, Christian Carbogno, Mariana Rossi, Ville Havu, Markus Rampp, Matthias Scheffler, and Volker Blum. Efficient all-electron hybrid density functionals for atomistic simulations beyond 10 000 atoms. *The Journal of Chemical Physics*, 161(2):024112, 07 2024.
- [7] Victor Wen zhe Yu, Carmen Campos, William Dawson, Alberto García, Ville Havu, Ben Hourahine, William P. Huhn, Mathias Jacquelin, Weile Jia, Murat Keçeli, Raul Laasner, Yingzhou Li, Lin Lin, Jianfeng Lu, Jonathan Moussa, Jose E. Roman, Álvaro Vázquez-Mayagoitia, Chao Yang, and Volker Blum. Elsi — an open infrastructure for electronic structure solvers. *Comput. Phys. Commun.*, 256:107459, 2020.
- [8] Volker Blum, Ralf Gehrke, Felix Hanke, Paula Havu, Ville Havu, Xinguo Ren, Karsten Reuter, and Matthias Scheffler. Ab initio molecular simulations with numeric atom-centered orbitals. *Computer Physics Communications*, 180(11):2175–2196, 2009.
- [9] F. W. Averill and D. E. Ellis. An efficient numerical multicenter basis set for molecular orbital calculations: Application to fecl4. *The Journal of Chemical Physics*, 59(12):6412–6418, 12 1973.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [10] Alex Zunger and A. J. Freeman. Self-consistent numerical-basis-set linear-combination-of-atomic-orbitals model for the study of solids in the local density formalism. *Phys. Rev. B*, 15:4716–4737, May 1977.
- [11] B. Delley and D. E. Ellis. Efficient and accurate expansion methods for molecules in local density models. *The Journal of Chemical Physics*, 76(4):1949–1960, 02 1982.
- [12] Bernard Delley. An all-electron numerical method for solving the local density functional for poly-atomic molecules. *The Journal of chemical physics*, 92(1):508–517, 1990.
- [13] G. te Velde and E. J. Baerends. Precise density-functional method for periodic structures. *Phys. Rev. B*, 44:7888–7903, Oct 1991.
- [14] Klaus Koepernik and Helmut Eschrig. Full-potential nonorthogonal local-orbital minimum-basis band-structure scheme. *Phys. Rev. B*, 59:1743–1757, Jan 1999.
- [15] Martin Fuchs Fuchs and Matthias Scheffler. Ab initio pseudopotentials for electronic structure calculations of poly-atomic systems using density-functional theory. *Computer Physics Communications*, 119(1):67–98, 1999.
- [16] Stefan Goedecker, 2016. private communication.
- [17] Ondřej Čertík, John E. Pask, and Jiří Vackář. dftatom: A robust and general schrödinger and dirac solver for atomic structure calculations. *Computer Physics Communications*, 184(7):1777–1791, 2013.
- [18] Javier Junquera, Óscar Paz, Daniel Sánchez-Portal, and Emilio Artacho. Numerical atomic orbitals for linear-scaling calculations. *Phys. Rev. B*, 64:235111, Nov 2001.
- [19] V. Havu, V. Blum, P. Havu, and M. Scheffler. Efficient integration for all-electron electronic structure calculation using numeric basis functions. *Journal of Computational Physics*, 228(22):8367–8379, December 2009.
- [20] Igor Ying Zhang, Xinguo Ren, Patrick Rinke, Volker Blum, and Matthias Scheffler. Numeric atom-centered-orbital basis sets with valence-correlation consistency from h to ar. *New J. Phys.*, 15(12):123033, dec 2013.
- [21] Dirk Andrae. Finite nuclear charge density distributions in electronic structure calculations for atoms and molecules. *Physics Reports*, 336(6):413–525, 2000.
- [22] C.W. De Jager, H. De Vries, and C. De Vries. Nuclear charge- and magnetization-density-distribution parameters from elastic electron scattering. *Atomic Data and Nuclear Data Tables*, 14(5):479–508, 1974. Nuclear Charge and Moment Distributions.
- [23] H. De Vries, C.W. De Jager, and C. De Vries. Nuclear charge-density-distribution parameters from elastic electron scattering. *Atomic Data and Nuclear Data Tables*, 36(3):495–536, 1987.
- [24] G. Fricke, C. Bernhardt, K. Heilig, L.A. Schaller, L. Schellenberg, E.B. Shera, and C.W. Dejager. Nuclear ground state charge radii from electromagnetic interactions. *Atomic Data and Nuclear Data Tables*, 60(2):177–285, 1995.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [25] Donal Day, Alan Brody, Bryan Lewis, and Stephen Washington. Nuclei charge density archive, 2014. <https://discovery.phys.virginia.edu/research/groups/ncd/download.html>. Last accessed on November 17, 2024.
- [26] Rundong Zhao, Victor Wen-zhe Yu, Kimberly Zhang, Yunlong Xiao, Yong Zhang, and Volker Blum. Quasi-four-component method with numeric atom-centered orbitals for relativistic density functional simulations of molecules and solids. *Phys. Rev. B*, 103:245144, Jun 2021.
- [27] John P Perdew, Kieron Burke, and Matthias Ernzerhof. Generalized gradient approximation made simple. *Phys. Rev. Lett.*, 77(18):3865, 1996.
- [28] Stig Rune Jensen, Santanu Saha, José A. Flores-Livas, William Huhn, Volker Blum, Stefan Goedecker, and Luca Frediani. The elephant in the room of density functional theory calculations. *The Journal of Physical Chemistry Letters*, 8(7):1449–1457, 2017. PMID: 28291362.
- [29] Multiresolution chemistry (mrchem) program package. <https://mrchem.readthedocs.io/en/latest/>.
- [30] P. Blaha, H. Hofstätter, O. Koch, R. Laskowski, and K. Schwarz. Iterative diagonalization in augmented plane wave based methods in electronic structure calculations. *Journal of Computational Physics*, 229(2):453–460, 2010.
- [31] R. Laasner, I. Mandzheeva, W.P. Huhn, J. Collet, V. Yu, W.S. Warren, T. Theis, and V. Blum. Molecular nmr shieldings, j-couplings, and magnetizabilities from numeric atom-centered orbital based density-functional calculations. *Electron. Struct.*, 2024.
- [32] T. H. Dunning. Gaussian basis sets for use in correlated molecular calculations. I. The atoms boron through neon and hydrogen. *J. Chem. Phys.*, 90(2):1007–1023, 1989.
- [33] Chi Liu, Jan Kloppenburg, Yi Yao, Xinguo Ren, Heiko Appel, Yosuke Kanai, and Volker Blum. All-electron ab initio bethe-salpeter equation approach to neutral excitations in molecules with numeric atom-centered orbitals. *The Journal of Chemical Physics*, 152(4), 2020.
- [34] Rick A. Kendall, Thom H. Dunning, and Robert J. Harrison. Electron affinities of the first-row atoms revisited. systematic basis sets and wave functions. *J. Chem. Phys.*, 96(9):6796–6806, 1992.
- [35] Yi Yao, Dorothea Golze, Patrick Rinke, Volker Blum, and Yosuke Kanai. All-Electron BSE@GW Method for K-Edge Core Electron Excitation Energies. *J. Chem. Theory Comput.*, 18(3):1569–1583, 2022.
- [36] X. Ren, F. Merz, H. Jiang, Y. Yao, M. Rapp, H. Lederer, V. Blum, and M. Scheffler. All-electron periodic g_0w_0 implementation with numerical atomic orbital basis functions: Algorithm and benchmarks. *Phys. Rev. Mater.*, 5:013807, 2021.
- [37] Ruiyi Zhou, Yi Yao, Volker Blum, Xinguo Ren, and Yosuke Kanai. All-electron bse@ gw method with numeric atom-centered orbitals for extended periodic systems. *Journal of Chemical Theory and Computation*, 2025.
- [38] Frank Jensen. The basis set convergence of spin-spin coupling constants calculated by density functional methods. *Journal of Chemical Theory and Computation*, 2(5):1360–1369, 2006. PMID: 26626843.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [39] F. Jensen. The optimum contraction of basis sets for calculating spin-spin coupling constants. *Theor. Chem. Acc.*, 126:371–382, 2010.
- [40] Benjamin P. Pritchard, Doaa Altarawy, Brett Didier, Tara D. Gibson, and Theresa L. Windus. New basis set exchange: An open, up-to-date resource for the molecular sciences community. *Journal of Chemical Information and Modeling*, 59(11):4814–4820, 2019. PMID: 31600445.
- [41] Michiel J. van Setten, Fabio Caruso, Sahar Sharifzadeh, Xinguo Ren, Matthias Scheffler, Fang Liu, Johannes Lischner, Lin Lin, Jack R. Deslippe, Steven G. Louie, Chao Yang, Florian Weigend, Jeffrey B. Neaton, Ferdinand Evers, and Patrick Rinke. GW100: Benchmarking GoWo for Molecular Systems. *J. Chem. Theory Comput.*, 11:5665–5687, 2015.
- [42] Sebastian Kokott, Iker Hurtado, Christian Vorwerk, Claudia Draxl, Volker Blum, and Matthias Scheffler. GIMS: graphical interface for materials simulations. *Journal of Open Source Software*, 6(57):2767, 2021.
- [43] Ask Hjorth Larsen, Jens Jørgen Mortensen, Jakob Blomqvist, Ivano E Castelli, Rune Christensen, Marcin Du lak, Jesper Friis, Michael N Groves, Bjørk Hammer, Cory Hargus, et al. The atomic simulation environment—a python library for working with atoms. *Journal of Physics: Condensed Matter*, 29(27):273002, 2017.
- [44] Kurt Lejaeghere, Gustav Bihlmayer, and Torbjörn Björkman *et al.* Reproducibility in density functional theory calculations of solids. *Science*, 351(6280):aad3000, 2016.
- [45] Peter Blaha, Karlheinz Schwarz, Fabien Tran, Robert Laskowski, Georg K. H. Madsen, and Laurence D. Marks. Wien2k: An apw+lo program for calculating the properties of solids. *The Journal of Chemical Physics*, 152(7):074101, 2020.
- [46] Andris Gulans, Stefan Kontur, Christian Meisenbichler, Dmitrii Nabok, Pasquale Pavone, Santiago Rigmonti, Stephan Sagmeister, Ute Werner, and Claudia Draxl. Exciting: a full-potential all-electron package implementing density-functional theory and many-body perturbation theory. *Journal of Physics: Condensed Matter*, 26(36):363202, 2014.
- [47] Fhi-aims delta test results, recomputed in 2020. <https://doi.org/10.17172/nomad.1yfb-ttj3>.
- [48] William P Huhn and Volker Blum. One-hundred-three compound band-structure benchmark of post-self-consistent spin-orbit coupling treatments in density functional theory. *Physical Review Materials*, 1(3):033803, 2017.
- [49] Lars Goerigk, Andreas Hansen, Christoph Bauer, Stephan Ehrlich, Asim Najibi, and Stefan Grimme. A look at the density functional theory zoo with the advanced GMTKN55 database for general main group thermochemistry, kinetics and noncovalent interactions. *Phys. Chem. Chem. Phys.*, 19:32184–32215, 2017.
- [50] E. R. Johnson. GMTKN55 inputs in FHI-aims format, 2024. <https://github.com/erin-r-johnson/gmtn55-fhiaims>.
- [51] Elisabeth Keller, Jack Morgenstein, Karsten Reuter, and Johannes T. Margraf. Small basis set density functional theory method for cost-efficient, large-scale condensed matter simulations. *The Journal of Chemical Physics*, 161(7):074104, 08 2024.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [52] Alberto García, Nick Papior, Arsalan Akhtar, Emilio Artacho, Volker Blum, Emanuele Bosoni, Pedro Brandimarte, Mads Brandbyge, J. I. Cerdá, Fabiano Corsetti, Ramón Cuadrado, Vladimir Dikan, Jaime Ferrer, Julian Gale, Pablo García-Fernández, V. M. García-Suárez, Sandra García, Georg Huhs, Sergio Illera, Richard Korytár, Peter Koval, Irina Lebedeva, Lin Lin, Pablo López-Tarifa, Sara G. Mayo, Stephan Mohr, Pablo Ordejón, Andrei Postnikov, Yann Pouillon, Miguel Pruneda, Roberto Robles, Daniel Sánchez-Portal, Jose M. Soler, Rafi Ullah, Victor Wen-zhe Yu, and Javier Junquera. Siesta: Recent developments and applications. *The Journal of Chemical Physics*, 152(20):204108, 05 2020.
- [53] Axel D Becke. A multicenter numerical integration scheme for polyatomic molecules. *The Journal of chemical physics*, 88(4):2547–2553, 1988.
- [54] Franz Knuth, Christian Carbogno, Viktor Atalla, Volker Blum, and Matthias Scheffler. All-electron formalism for total energy strain derivatives and stress tensor components for numeric atom-centered orbitals. *Computer Physics Communications*, 190:33–50, 2015.
- [55] Vyacheslav Ivanovich Lebedev. Values of the nodes and weights of quadrature formulas of gauss-markov type for a sphere from the ninth to seventeenth order of accuracy that are invariant with respect to an octahedron group with inversion. *Zhurnal Vychislitel'noi Matematiki i Matematicheskoi Fiziki*, 15(1):48–54, 1975.
- [56] Vyacheslav Ivanovich Lebedev. Quadratures on a sphere. *USSR Computational Mathematics and Mathematical Physics*, 16(2):10–24, 1976.
- [57] Vyacheslav Ivanovich Lebedev and Dimitri N Laikov. A quadrature formula for the sphere of the 131st algebraic order of accuracy. In *Doklady Mathematics*, volume 59, pages 477–481. Pleiades Publishing, Ltd., 1999.
- [58] Bernard Delley. High order integration schemes on the unit sphere. *Journal of computational chemistry*, 17(9):1152–1155, 1996.
- [59] Robert S Womersley. Efficient spherical designs with good geometric properties. In: *Dick J., Kuo F., Wozniakowski H. (eds) Contemporary Computational Mathematics - A Celebration of the 80th Birthday of Ian Sloan.*, pages 1243–1285, 2018.
- [60] R Eric Stratmann, Gustavo E Scuseria, and Michael J Frisch. Achieving linear scaling in exchange-correlation density functional quadratures. *Chemical physics letters*, 257(3-4):213–223, 1996.
- [61] Christoph Freysoldt, Jörg Neugebauer, and Chris G Van de Walle. Fully ab initio finite-size corrections for charged-defect supercell calculations. *Physical review letters*, 102(1):016402, 2009.
- [62] Xinguo Ren, Patrick Rinke, Volker Blum, Jürgen Wieferink, Alexandre Tkatchenko, Andrea Sanfilippo, Karsten Reuter, and Matthias Scheffler. Resolution-of-identity approach to hartree-fock, hybrid density functionals, rpa, mp2 and gw with numeric atom-centered orbital basis functions. *New Journal of Physics*, 14(5):053020, 2012.
- [63] Arvid Conrad Ihrig, Jürgen Wieferink, Igor Ying Zhang, Matti Ropo, Xinguo Ren, Patrick Rinke, Matthias Scheffler, and Volker Blum. Accurate localized resolution of identity approach for linear-scaling hybrid density functionals and for many-body perturbation theory. *New Journal of Physics*, 17(9):093020, 2015.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [64] Sergey V Levchenko, Xinguo Ren, Jürgen Wieferink, Rainer Johanni, Patrick Rinke, Volker Blum, and Matthias Scheffler. Hybrid functionals for large periodic systems in an all-electron, numeric atom-centered basis framework. *Computer Physics Communications*, 192:60–69, 2015.
- [65] Wiley S Morgan, John E Christensen, Parker K Hamilton, Jeremy J Jorgensen, Branton J Campbell, Gus LW Hart, and Rodney W Forcade. Generalized regular k-point grid generation on the fly. *Computational Materials Science*, 173:109340, 2020.
- [66] Atsushi Togo, Kohei Shinohara, and Isao Tanaka. Spglib: a software library for crystal symmetry search. *Science and Technology of Advanced Materials: Methods*, August 2024.
- [67] William P. Huhn, Björn Lange, Victor W. z. Yu, Mina Yoon, and Volker Blum. Gpu acceleration of all-electron electronic structure theory using localized numeric atom-centered basis functions. *Computer Physics Communications*, 254:107314, 2020.
- [68] M. N. Tahir, T. Zhu, H. Shang, J. Li, V. Blum, and X. Ren. Localized resolution of identity approach to the analytical gradients of random-phase approximation ground-state energy: Algorithm and benchmarks. *J. Chem. Theory Comput.*, 18:5297, 2022.
- [69] Jorge Nocedal and Stephen J. Wright. *Numerical Optimization*. Springer, New York, 1999.
- [70] Maja-Olivia Lenz, Thomas AR Purcell, David Hicks, Stefano Curtarolo, Matthias Scheffler, and Christian Carbogno. Parametrically constrained geometry relaxations for high-throughput materials science. *npj Computational Materials*, 5(1):123, 2019.
- [71] C. G. BROYDEN. The Convergence of a Class of Double-rank Minimization Algorithms. *IMA Journal of Applied Mathematics*, 6(3):222–231, 1970.
- [72] R. Fletcher. A new approach to variable metric algorithms. *The Computer Journal*, 13(3):317–322, 1970.
- [73] Donald Goldfarb. A family of variable-metric methods derived by variational means. *Mathematics of Computation*, 24(109):23–26, 1970.
- [74] D. F. Shanno. Conditioning of quasi-Newton methods for function minimization. *Mathematics of Computation*, 24(111):647–656, 1970.
- [75] Roland Lindh, Anders Bernhardsson, Gunnar Karlström, and Per-Åke Malmqvist. On the use of a Hessian model function in molecular geometry optimizations. *Chemical Physics Letters*, 241(4):423–428, 1995.
- [76] Bernd G. Pfrommer, Michel Côté, Steven G. Louie, and Marvin L. Cohen. Relaxation of Crystals with the Quasi-Newton Method. *Journal of Computational Physics*, 131(1):233–240, 1997.
- [77] E. B. Tadmor, G. S. Smith, N. Bernstein, and E. Kaxiras. Mixed finite element and atomistic formulation for complex crystals. *Physical Review B*, 59(1):235–245, 1999.
- [78] Florian Knoop. *Heat transport in strongly anharmonic solids from first principles*. PhD thesis, Humboldt-Universität zu Berlin, 2022. <https://doi.org/10.18452/24244>.
- [79] HANNES JÓNSSON, GREG MILLS, and KARSTEN W. JACOBSEN. *Nudged elastic band method for finding minimum energy paths of transitions*, pages 385–404. World Scientific, 1998.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [80] Graeme Henkelman and Hannes Jónsson. Improved tangent estimate in the nudged elastic band method for finding minimum energy paths and saddle points. *The Journal of Chemical Physics*, 113(22):9978–9985, 12 2000.
- [81] Weinan E, Weiqing Ren, and Eric Vanden-Eijnden. Simplified and improved string method for computing the minimum energy paths in barrier-crossing events. *The Journal of Chemical Physics*, 126(16):164103, 04 2007.
- [82] Baron Peters, Andreas Heyden, Alexis T. Bell, and Arup Chakraborty. A growing string method for determining transition states: Comparison to the nudged elastic band and string methods. *The Journal of Chemical Physics*, 120(17):7877–7886, 05 2004.
- [83] Alan M. Lewis, Andrea Grisafi, Michele Ceriotti, and Mariana Rossi. Learning electron densities in the condensed phase. *Journal of Chemical Theory and Computation*, 17(11):7203–7214, 2021.
- [84] Evgeny Posenitskiy, Vijay Gopal Chilkuri, Abdallah Ammar, Micha l Hapka, Katarzyna Pernal, Ravindra Shinde, Edgar Josué Landinez Borda, Claudia Filippi, Kosuke Nakano, Otto Kohulák, Sandro Sorella, Pablo de Oliveira Castro, William Jalby, Pablo López Riós, Ali Alavi, and Anthony Scemama. TREXIO: A File Format and Library for Quantum Chemistry. *The Journal of Chemical Physics*, 158(17):174801, 2023.
- [85] Yann Garniron, Thomas Applencourt, Kevin Gasperich, Anouar Benali, Anthony Ferté, Julien Paquier, Barthélémy Pradines, Roland Assaraf, Peter Reinhardt, Julien Toulouse, Pierrette Barbaresco, Nicolas Renon, Grégoire David, Jean-Paul Malrieu, Mickaël Véril, Michel Caffarel, Pierre-François Loos, Emmanuel Giner, and Anthony Scemama. Quantum Package 2.0: An Open-Source Determinant-Driven Suite of Programs. *J. Chem. Theory Comput.*, 15(6):3591–3609, 2019.
- [86] Kai Guther, Robert J. Anderson, Nick S. Blunt, Nikolay A. Bogdanov, Deidre Cleland, Nike Dattani, Werner Dobrautz, Khaldoon Ghanem, Peter Jeszenszki, Niklas Liebermann, Giovanni Li Manni, Alexander Y. Lozovoi, Hongjun Luo, Dongxia Ma, Florian Merz, Catherine Overy, Markus Rampf, Pradipta Kumar Samanta, Lauretta R. Schwarz, James J. Shepherd, Simon D. Smart, Eugenio Vitale, Oskar Weser, George H. Booth, and Ali Alavi. NECI: N -Electron Configuration Interaction with an emphasis on state-of-the-art stochastic methods. *The Journal of Chemical Physics*, 153(3):034107, 2020.
- [87] Anthony Scemama, Michel Caffarel, Emmanuel Oseret, and William Jalby. Quantum Monte Carlo for large chemical systems: Implementing efficient strategies for petascale platforms and beyond. *J Comput Chem*, 34(11):938–951, 2013.
- [88] Anthony Scemama, Thomas Applencourt, Emmanuel Giner, and Michel Caffarel. Quantum Monte Carlo with very large multideterminant wavefunctions. *J. Comput. Chem.*, 37(20):1866–1875, 2016.
- [89] Peter J. Knowles and Nicholas C. Handy. A determinant based full configuration interaction program. *Computer Physics Communications*, 54(1):75–83, 1989.
- [90] Levi Keller, Volker Blum, Patrick Rinke, and Dorothea Golze. Relativistic correction scheme for core-level binding energies from GW . *J. Chem. Phys.*, 153(11):114110, 2020.
- [91] Ian P. Grant. *Relativistic Quantum Theory of Atoms and Molecules*. Springer New York, NY, April 2007.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [92] D.A. Rehn, J.M. Wills, T.E. Battelle, and A.E. Mattsson. Dirac's equation and its implications for density functional theory based calculations of materials containing heavy elements. *Phys. Rev. B*, 101:085114, 2020.
- [93] E. van Lenthe, E. Baerends, and J. Snijders. Relativistic regular two-component hamiltonians. *J. Chem. Phys.*, 99(6):4597–4610, 1993.
- [94] E. van Lenthe, E. Baerends, and J. Snijders. Relativistic total energy using regular approximations. *J. Chem. Phys.*, 101(11):9783–9792, 1994.
- [95] D. Peng, W. Liu, Y. Xiao, and L. Cheng. Making four- and two-component relativistic density functional methods fully equivalent based on the idea of "from atoms to molecule". *J. Chem. Phys.*, 127(10):104106, 2007.
- [96] Werner Kutzelnigg and Wenjian Liu. Quasirelativistic theory equivalent to fully relativistic theory. *J. Chem. Phys.*, 123(24):241102, 2005.
- [97] Ondřej Čertík, John E. Pask, and Jiří Vackář. Dftatom: A robust and general schrödinger and dirac solver for atomic structure calculations. *Comput. Phys. Commun.*, 184(7):1777, 2013.
- [98] Manoj K Jana, Ruyi Song, Haoliang Liu, Dipak Raj Khanal, Svenja M Janke, Rundong Zhao, Chi Liu, Z Valy Vardeny, Volker Blum, and David B Mitzi. Organic-to-inorganic structural chirality transfer in a 2d hybrid perovskite and impact on rashba-dresselhaus spin-orbit coupling. *Nature communications*, 11(1):4699, 2020.
- [99] S. Kim, M. Lee, C. Hong, Y. Yoon, H. An, D. Lee, W. Jeong, D. Yoo, Y. Kang, Y. Youn, and S. Hang. A band-gap database for semiconducting inorganic materials calculated with hybrid functional. *Sci. Data*, 7:387, 2020.
- [100] Walter Kohn and Lu Jeu Sham. Self-consistent equations including exchange and correlation effects. *Physical review*, 140(4A):A1133, 1965.
- [101] A. Seidl, A. Görling, P. Vogl, J. A. Majewski, and M. Levy. Generalized kohn-sham schemes and the band-gap problem. *Phys. Rev. B*, 53:3764–3774, Feb 1996.
- [102] <https://wordpress.elsi-interchange.org/>.
- [103] Victor Wen zhe Yu, Fabiano Corsetti, Alberto García, William P. Huhn, Mathias Jacquelin, Weile Jia, Björn Lange, Lin Lin, Jianfeng Lu, Wenhui Mi, Ali Seifitokaldani, Álvaro Vázquez-Mayagoitia, Chao Yang, Haizhao Yang, and Volker Blum. Elsi: A unified software interface for kohn–sham electronic structure solvers. *Comput. Phys. Commun.*, 222:267, 2018.
- [104] B. Hourahine, B. Aradi, V. Blum, F. Bonafé, A. Buccheri, C. Camacho, C. Cevallos, M. Y. Deshayé, T. Dumitrică, and A. et al. Dominguez. Dftb+, a software package for efficient approximate density functional theory based atomistic simulations. *The Journal of Chemical Physics*, 152(12):124101, 03 2020.
- [105] Nwchemex community. <https://github.com/NWChemEx>.
- [106] <https://www.netlib.org/lapack/>.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [107] E. Anderson, Z. Bai, C. Bischof, L. S. Blackford, J. Demmel, J. Dongarra, J. Du Croz, A. Greenbaum, S. Hammarling, A. McKenney, and D. Sorensen. *LAPACK Users' Guide*. Society for Industrial and Applied Mathematics, third edition, 1999.
- [108] <https://elpa.mpcdf.mpg.de/>.
- [109] A. Marek, P. Karpov, and T. Melson. Elpa manual: User's guide and best practices. https://elpa.mpcdf.mpg.de/elpa_userguide.pdf, 2024.
- [110] Lin Lin, Jianfeng Lu, Lexing Ying, Roberto Car, and Weinan E. Fast algorithm for extracting the diagonal of the inverse matrix with application to the electronic structure analysis of metallic systems. *Communications in Mathematical Sciences*, 7(3):755-777, 2009.
- [111] Lin Lin, Mohan Chen, Chao Yang, and Lixin He. Accelerating atomic orbital-based electronic structure calculation via pole expansion and selected inversion. *Journal of Physics: Condensed Matter*, 25(29):295501, June 2013.
- [112] William Dawson and Takahito Nakajima. Massively parallel sparse matrix function calculations with ntpoly. *Computer Physics Communications*, 225:154-165, 2018.
- [113] Toshiyuki Imamura, Susumu YAMADA, and Masahiko Machida. Development of a high-performance eigensolver on a peta-scale next-generation supercomputer system. *Progress in Nuclear Science and Technology*, 2:643-650, 10 2011.
- [114] Jack Dongarra, Mark Gates, Azzam Haidar, Jakub Kurzak, Piotr Luszczek, Stanimire Tomov, and Ichitaro Yamazaki. *Accelerating Numerical Dense Linear Algebra Calculations with GPUs*, pages 3-28. 06 2014.
- [115] Jan Winkelmann, Paul Springer, and Edoardo Di Napoli. Chase: Chebyshev accelerated subspace iteration eigensolver for sequences of hermitian eigenvalue problems. *ACM Trans. Math. Softw.*, 45(2), April 2019.
- [116] Xinzhe Wu, Davor Davidović, Sebastian Achilles, and Edoardo Di Napoli. Chase: a distributed hybrid cpu-gpu eigensolver for large-scale hermitian eigenvalue problems. In *Proceedings of the Platform for Advanced Scientific Computing Conference, PASC '22*, New York, NY, USA, 2022. Association for Computing Machinery.
- [117] Murat Keçeli, Fabiano Corsetti, Carmen Campos, Jose E. Roman, Hong Zhang, Álvaro Vázquez-Mayagoitia, Peter Zapol, and Albert F. Wagner. Siesta-sips: Massively parallel spectrum-slicing eigensolver for an ab initio molecular dynamics package. *Journal of Computational Chemistry*, 39(22):1806-1814, 2018.
- [118] Fabiano Corsetti. The orbital minimization method for electronic structure calculations with finite-range atomic basis sets. *Computer Physics Communications*, 185(3):873-883, 2014.
- [119] Meiyue Shao, Felipe H. da Jornada, Chao Yang, Jack Deslippe, and Steven G. Louie. Structure preserving parallel algorithms for solving the bethe-salpeter eigenvalue problem. *Linear Algebra and its Applications*, 488:148-167, 2016.
- [120] Distributed linear algebra from the future. <https://github.com/eth-cscs/DLA-Future>.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [121] Raffaele Solcà, Mikael Simberg, Rocco Meli, Alberto Invernizzi, Auriane Reverdell, and John Biddiscombe. Dla-future: A task-based linear algebra library which provides a gpu-enabled distributed eigensolver. In Patrick Diehl, Joseph Schuchart, Pedro Valero-Lara, and George Bosilca, editors, *Asynchronous Many-Task Systems and Applications*, pages 135–141, Cham, 2024. Springer Nature Switzerland.
- [122] X. Gonze, B. Amadon, G. Antonius, F. Arnardi, L. Baguet, J.-M. Beuken, J. Bieder, F. Bottin, J. Bouchet, E. Bousquet, et al. The abinitproject: Impact, environment and recent developments. *Computer Physics Communications*, 248:107042, 03 2020.
- [123] J. Deslippe, G. Samsonidze, D. A. Strubbe, M. Jain, M. L. Cohen, and S. G. Louie. Berkeleygw: A massively parallel computer package for the calculation of the quasiparticle and optical properties of materials and nanostructures. *Comput. Phys. Commun.*, 183:1269, 2012.
- [124] Thomas D. Kühne, Marcella Iannuzzi, Mauro Del Ben, Vladimir V. Rybkin, Patrick Seewald, Frederick Stein, Teodoro Laino, Rustam Z. Khaliullin, Ole Schütt, Florian Schiffmann, Dorothea Golze, Jan Wilhelm, Sergey Chulkov, Mohammad Hossein Bani-Hashemian, Valéry Weber, Urban Borštnik, Mathieu Taillefumier, Alice Shoshana Jakobovits, Alfio Lazzaro, Hans Pabst, Tiziano Müller, Robert Schade, Manuel Guidon, Samuel Andermatt, Nico Holmberg, Gregory K. Schenter, Anna Hehn, Augustin Bussy, Fabian Belleflamme, Gloria Tabacchi, Andreas Glöß, Michael Lass, Iain Bethune, Christopher J. Mundy, Christian Plessl, Matt Watkins, Joost VandeVondele, Matthias Krack, and Jürg Hutter. Cp2k: An electronic structure and molecular dynamics software package - quickstep: Efficient and accurate electronic structure calculations. *The Journal of Chemical Physics*, 152(19), May 2020.
- [125] Tobias Klöffel, Gerald Mathias, and Bernd Meyer. Integrating state of the art compute, communication, and autotuning strategies to multiply the performance of ab initio molecular dynamics on massively parallel multi-core supercomputers. *Computer Physics Communications*, 260:107745, 2021.
- [126] J Enkovaara, C Rostgaard, JJ Mortensen, J Chen, M Du lak, L Ferrighi, J Gavnholt, C Glinsvad, V Haikola, H A Hansen, et al. Electronic structure calculations with gpaw: a real-space implementation of the projector augmented-wave method. *Journal of Physics: Condensed Matter*, 22(25):253202, jun 2010.
- [127] E. Aprà, E. J. Bylaska, W. A. de Jong, N. Govind, K. Kowalski, T. P. Straatsma, M. Valiev, H. J. J. van Dam, Y. Alexeev, J. Anchell, et al. Nwchem: Past, present, and future. *The Journal of Chemical Physics*, 152(18):184102, 05 2020.
- [128] Nicolas Tancogne-Dejean, Micael J. T. Oliveira, Xavier Andrade, Heiko Appel, Carlos H. Borca, Guillaume Le Breton, Florian Buchholz, Alberto Castro, Stefano Corni, Correa, et al. Octopus, a computational framework for exploring light-driven phenomena and quantum dynamics in extended and finite systems. *The Journal of Chemical Physics*, 152(12):124119, 03 2020.
- [129] Søren Smidstrup, Troels Markussen, Pieter Vancaerfeld, Jess Wellendorff, Julian Schneider, Tue Gunst, Brecht Verstichel, Daniele Stradi, Petr A Khomyakov, Ulrik G Vej-Hansen, et al. Quantumatk: an integrated platform of electronic and atomic-scale modelling tools. *Journal of Physics: Condensed Matter*, 32(1):015901, oct 2019.
- [130] Paolo Giannozzi, Stefano Baroni, Nicola Bonini, Matteo Calandra, Roberto Car, Carlo Cavazzoni, Davide Ceresoli, Guido L Chiarotti, Matteo Cococcioni, Ismaila Dabo, et al. Quantum espresso: a modular and open-source software project for quantum simulations of materials. *Journal of Physics: Condensed Matter*, 21(39):395502, sep 2009.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [131] P Giannozzi, O Andreussi, T Brumme, O Bunau, M Buongiorno Nardelli, M Calandra, R Car, C Cavazzoni, D Ceresoli, M Cococcioni, et al. Advanced capabilities for materials modelling with quantum espresso. *Journal of Physics: Condensed Matter*, 29(46):465901, oct 2017.
- [132] G. Kresse and J. Furthmüller. Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set. *Phys. Rev. B*, 54:11169–11186, Oct 1996.
- [133] Pavel Kůs, Hermann Lederer, and Andreas Marek. Gpu optimization of large-scale eigenvalue solver. In Florin Adrian Radu, Kundan Kumar, Inga Berre, Jan Martin Nordbotten, and Iuliu Sorin Pop, editors, *Numerical Mathematics and Advanced Applications ENUMATH 2017*, pages 123–131, Cham, 2019. Springer International Publishing.
- [134] Victor Wen zhe Yu, Jonathan Moussa, Pavel Kůs, Andreas Marek, Peter Messmer, Mina Yoon, Hermann Lederer, and Volker Blum. Gpu-acceleration of the elpa2 distributed eigensolver for dense symmetric and hermitian eigenproblems. *Computer Physics Communications*, 262:107808, 2021.
- [135] Gabriel Wlaz lowski, Michael McNeil Forbes, Saptarshi Rajan Sarkar, Andreas Marek, and Maciej Szpindler. Fermionic quantum turbulence: Pushing the limits of high-performance computing. *PNAS Nexus*, 3(5):pgae160, 04 2024.
- [136] Scalapack. <https://www.netlib.org/scalapack/>.
- [137] L. S. Blackford, J. Choi, A. Cleary, E. D’Azevedo, J. Demmel, I. Dhillon, J. Dongarra, S. Hammarling, G. Henry, A. Petitet, K. Stanley, D. Walker, and R. C. Whaley. *ScaLAPACK Users’ Guide*. Society for Industrial and Applied Mathematics, 1997.
- [138] Nccl. <https://developer.nvidia.com/nccl>.
- [139] Rccl. <https://rocm.docs.amd.com/projects/rccl>.
- [140] Sycl. <https://www.intel.com/content/www/us/en/developer/tools/oneapi/data-parallel-c-plus-plus.html>.
- [141] onemkl. <https://www.intel.com/content/www/us/en/developer/tools/oneapi/onemkl.html>.
- [142] Bruno Lang. A parallel algorithm for reducing symmetric banded matrices to tridiagonal form. *SIAM Journal on Scientific Computing*, 14(6):1320–1338, 1993.
- [143] C. Bischof, Xiaobai Sun, and B. Lang. Parallel tridiagonalization through two-step band reduction. In *Proceedings of IEEE Scalable High Performance Computing Conference*, pages 23–27, 1994.
- [144] T. Auckenthaler, V. Blum, H.-J. Bungartz, T. Huckle, R. Johanni, L. Krämer, B. Lang, H. Lederer, and P.R. Willems. Parallel solution of partial symmetric eigenvalue problems from electronic structure calculations. *Parallel Computing*, 37(12):783–794, 2011. 6th International Workshop on Parallel Matrix Algorithms and Applications (PMAA’10).
- [145] P. Kůs, A. Marek, S.S. Köcher, H.-H. Kowalski, C. Carbogno, Ch. Scheurer, K. Reuter, M. Scheffler, and H. Lederer. Optimizations of the eigensolvers in the elpa library. *Parallel Computing*, 85:167–177, 2019.
- [146] R. A. Van De Geijn and J. Watts. Summa: scalable universal matrix multiplication algorithm. *Concurrency: Practice and Experience*, 9(4):255–274, 1997.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [147] Almadena Chtchelkanova, John Gunnels, Greg Morrow, James Overfelt, and Robert A. van de Geijn. Parallel implementation of blas: general techniques for level 3 blas. *Concurrency: Practice and Experience*, 9(9):837–857, 1997.
- [148] Lynn Elliot Cannon. *A cellular computer to implement the kalman filter algorithm*. PhD thesis, Carnegie-Mellon University, USA, 1969.
- [149] Nvidia mps. <https://docs.nvidia.com/deploy/mps>.
- [150] Raven. <https://docs.mpcdf.mpg.de/doc/computing/raven-user-guide.html>.
- [151] P. Karpov. ScaLAPACK + ELPA tutorial. <https://github.com/karpov-peter/elpa-tutorial>, 2023. GitHub repository.
- [152] Apu. <https://www.amd.com/de/products/accelerators/instinct/mi300/mi300a.html>.
- [153] Vpu. <https://github.com/riscvarchive/riscv-v-spec>.
- [154] R Grima Torres, P Vizcaíno, F Mantovani, and JJ Gutiérrez Moreno. Co-designing ab initio electronic structure methods on a risc-v vector architecture. *Open Research Europe*, 4:165, 10 2024.
- [155] Ryan Pederson, John Kozłowski, Ruyi Song, Jackson Beall, Martin Ganahl, Markus Hauru, Adam G. M. Lewis, Yi Yao, Shrestha Basu Mallick, Volker Blum, and Guifre Vidal. Large scale quantum chemistry with tensor processing units. *Journal of Chemical Theory and Computation*, 19:25–32, 2023.
- [156] Victor Wen-zhe Yu, Jonathan Moussa, and Volker Blum. Accurate frozen core approximation for all-electron density-functional theory. *The Journal of Chemical Physics*, 154(22):224107, 2021.
- [157] Timothy Hughbanks and Roald Hoffmann. Chains of trans-edge-sharing molybdenum octahedra: metal-metal bonding in extended systems. *Journal of the American Chemical Society*, 105(11):3528–3537, 1983.
- [158] Izumi Takahara, Kiyuu Shibata, and Teruyasu Mizoguchi. Crystal orbital overlap population based on all-electron ab initio simulation with numeric atom-centered orbitals and its application to chemical-bonding analysis in li-intercalated layered materials. *Modelling and Simulation in Materials Science and Engineering*, 32(5):055028, 2024.
- [159] Yoyo Hinuma, Giovanni Pizzi, Yu Kumagai, Fumiyasu Oba, and Isao Tanaka. Band structure diagram paths based on crystallography. *Computational Materials Science*, 128:140–184, 2017.
- [160] Robert S Mulliken. Electronic population analysis on lcao–mo molecular wave functions. i. *The Journal of chemical physics*, 23(10):1833–1840, 1955.
- [161] R Mulliken. Overlap populations, bond orders and covalent bond energies. *J. chem. Phys*, 23:1841–1846, 1955.
- [162] Fred L Hirshfeld. Bonded-atom fragments for describing molecular charge densities. *Theoretica chimica acta*, 44:129–138, 1977.
- [163] Patrick Bultinck, Christian Van Alsenoy, Paul W Ayers, and Ramon Carbó-Dorca. Critical analysis and extension of the hirshfeld atoms in molecules. *The Journal of chemical physics*, 126(14), 2007.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [164] RFW Bader. Atoms in molecules: A quantum theory, 438 p, 1990.
- [165] Miroslav Kohout. Electron localizability indicator and bonding analysis with dgrid. *Chapt*, 4:75–112, 2021.
- [166] Raffaele Resta. Macroscopic polarization in crystalline dielectrics: the geometric phase approach. *Reviews of modern physics*, 66(3):899, 1994.
- [167] Christian Carbogno, Nikita Rybin, Sara Panahian Jand, Alaa Akkoush, Carlos Mera Acosta, Zhenkun Yuan, and Mariana Rossi. Polarisation, born effective charges, and topological invariants via a berry-phase approach, 2025.
- [168] Axel D Becke and Kenneth E Edgecombe. A simple measure of electron localization in atomic and molecular systems. *The Journal of chemical physics*, 92(9):5397–5403, 1990.
- [169] A Savin, B Silvi, and F Colonna. Topological analysis of the electron localization function applied to delocalized bonds. *Canadian journal of chemistry*, 74(6):1088–1096, 1996.
- [170] M Kohout and A Savin. Atomic shell structure and electron numbers. *International journal of quantum chemistry*, 60(4):875–882, 1996.
- [171] Definition of the CUBE file format. <https://h5cube-spec.readthedocs.io/en/latest/cubeformat.html>.
- [172] Jerry Tersoff and Donald R Hamann. Theory of the scanning tunneling microscope. *Physical Review B*, 31(2):805, 1985.
- [173] AH MacDonald, SH Vosko, and PT Coleridge. Extensions of the tetrahedron method for evaluating spectral properties of solids. *Journal of Physics C: Solid State Physics*, 12(15):2991, 1979.
- [174] Dimitris Zaharioudakis. Tetrahedron methods for brillouin zone integration. *Computer physics communications*, 157(1):17–31, 2004.
- [175] Ruyi Song and Rundong Zhao. Density functional theory study of two-dimensional hybrid organic-inorganic perovskites: frontier level alignment and chirality-induced spin splitting. *AAPPS Bulletin*, 34(1):1–15, 2024.
- [176] Carlos Campaña, Bastien Mussard, and Tom K Woo. Electrostatic potential derived atomic charges for periodic systems using a modified error functional. *Journal of Chemical Theory and Computation*, 5(10):2866–2878, 2009.
- [177] De-Li Chen, Abraham C Stern, Brian Space, and J Karl Johnson. Atomic charges derived from electrostatic potentials for molecular and periodic systems. *The Journal of Physical Chemistry A*, 114(37):10225–10233, 2010.
- [178] Susi Lehtola and Hannes Jónsson. Unitary optimization of localized molecular orbitals. *Journal of chemical theory and computation*, 9(12):5365–5372, 2013.
- [179] János Pipek and Paul G Mezey. A fast intrinsic localization procedure applicable for ab initio and semiempirical linear combination of atomic orbital wave functions. *The Journal of Chemical Physics*, 90(9):4916–4926, 1989.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [180] Yang Guo, Wei Li, and Shuhua Li. An efficient linear scaling procedure for constructing localized orbitals of large molecules based on the one-particle density matrix. *The Journal of chemical physics*, 135(13), 2011.
- [181] Peter E Blöchl, Ove Jepsen, and Ole Krogh Andersen. Improved tetrahedron method for brillouin-zone integrations. *Physical Review B*, 49(23):16223, 1994.
- [182] Arieh Warshel and Michael Levitt. Theoretical studies of enzymic reactions: dielectric, electrostatic and steric stabilization of the carbonium ion in the reaction of lysozyme. *Journal of Molecular Biology*, 103(2):227–249, 1976.
- [183] Hans Martin Senn and Walter Thiel. Qm/mm methods for biomolecular systems. *Angewandte Chemie International Edition*, 48(7):1198–1229, 2009.
- [184] Stefan Ringe, Nicolas G. Hörmann, Harald Oberhofer, and Karsten Reuter. Implicit solvation methods for catalysis at electrified interfaces. *Chemical Reviews*, 122(12):10777–10820, 2022.
- [185] Katja-Sophia Csizi and Markus Reiher. Universal qm/mm approaches for general nanoscale applications. *Wiley Interdisciplinary Reviews: Computational Molecular Science*, 13(4):e1656, 2023.
- [186] Gabriel Adrian Bramley, Owain Tomos Beynon, Pavel Viktorovich Stishenko, and Andrew James Logsdail. The application of qm/mm simulations in heterogeneous catalysis. *Physical Chemistry Chemical Physics*, 25(9):6562–6585, 2023.
- [187] Daniel Berger, Andrew J Logsdail, Harald Oberhofer, Matthew R Farrow, C Richard A Catlow, Paul Sherwood, Alexey A Sokol, Volker Blum, and Karsten Reuter. Embedded-cluster calculations in a numeric atomic orbital density-functional theory framework. *The Journal of Chemical Physics*, 141(2):024105, 2014.
- [188] You Lu, Matthew R Farrow, Pierre Fayon, Andrew J Logsdail, Alexey A Sokol, C Richard A Catlow, Paul Sherwood, and Thomas W Keal. Open-source, python-based redevelopment of the chemshell multiscale qm/mm environment. *Journal of Chemical Theory and Computation*, 15(2):1317–1328, 2018.
- [189] Norina A Richter, Sabrina Siculo, Sergey V Levchenko, Joachim Sauer, and Matthias Scheffler. Concentration of vacancies at metal-oxide surfaces: Case study of mgo (100). *Physical Review Letters*, 111(4):045502, 2013.
- [190] Thomas Stecher, Karsten Reuter, and Harald Oberhofer. First-principles free-energy barriers for photoelectrochemical surface reactions: Proton abstraction at tio 2 (110). *Physical Review Letters*, 117(27):276001, 2016.
- [191] Matthias Kick, Karsten Reuter, and Harald Oberhofer. Intricacies of DFT+U, Not Only in a Numeric Atom Centered Orbital Framework. *J. Chem. Theory Comput.*, 15(3):1705–1718, mar 2019.
- [192] You Lu, Kakali Sen, Chin Yong, David SD Gunn, John A Purton, Jingcheng Guan, Alec Desmoutier, Jamal Abdul Nasir, Xingfan Zhang, Lei Zhu, et al. Multiscale qm/mm modelling of catalytic systems with chemshell. *Physical Chemistry Chemical Physics*, 25(33):21816–21835, 2023.
- [193] Oliviero Andreussi, Ismaila Dabo, and Nicola Marzari. Revised self-consistent continuum solvation in electronic-structure calculations. *The Journal of Chemical Physics*, 136(6):064102, February 2012.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [194] Jakob Filser, Edan Bainglass, Karsten Reuter, and Oliviero Andreussi. Coupling all-electron full-potential density functional theory with grid-based continuum embeddings. *The Journal of Chemical Physics*, 163(16):164103, 10 2025.
- [195] Jakob Filser, Karsten Reuter, and Harald Oberhofer. Piecewise multipole-expansion implicit solvation for arbitrarily shaped molecular solutes. *Journal of Chemical Theory and Computation*, 18(1):461–478, 2021.
- [196] A. Klamt and G. Schüürmann. Cosmo: a new approach to dielectric screening in solvents with explicit expressions for the screening energy and its gradient. *J. Chem. Soc., Perkin Trans. 2*, (5):799–805, 1993.
- [197] Darrin M York and Martin Karplus. A smooth solvation potential based on the conductor-like screening model. *The Journal of Physical Chemistry A*, 103(50):11060–11079, 1999.
- [198] Opium - pseudopotential generation project. <https://opium.sourceforge.net/>. Accessed: 2024-10-01.
- [199] Using fhi-aims for embedded cluster simulations with chemshell. <https://fhi-aims-club.gitlab.io/tutorials/using-fhi-aims-for-embedded-cluster-simulations-in-chemshell/>. Accessed: 2024-10-01.
- [200] Pavel V Stishenko, Jakob Filser, Gabriel Bramley, Reinhard Maurer, Volker Blum, Harald Oberhofer, and Andrew Logsdail. Improvements in periodic representation of solvated systems with fhi-aims (1.0). 2024.
- [201] Stefan Ringe. *First-Principles Modeling in Poisson-Boltzmann Electrolytes*. PhD thesis, Technische Universität München, 2017. <https://mediatum.ub.tum.de/doc/1353874/document.pdf>.
- [202] C Dupont, O Andreussi, and N Marzari. Self-consistent continuum solvation (SCCS): the case of charged systems. *The Journal of Chemical Physics*, 139(21):214110, December 2013.
- [203] Christoph Hille, Stefan Ringe, Martin Deimel, Christian Kunkel, William E Acree, Karsten Reuter, and Harald Oberhofer. Generalized molecular solvation in non-aqueous solutions by a single parameter implicit solvation scheme. *The Journal of Chemical Physics*, 150(4):041710, January 2019.
- [204] Stefan Ringe, Harald Oberhofer, Christoph Hille, Sebastian Matera, and Karsten Reuter. Function-space-based solution scheme for the size-modified Poisson–Boltzmann equation in full-potential DFT. *Journal of Chemical Theory and Computation*, 12(8):4052–4066, 2016.
- [205] Stefan Ringe, Harald Oberhofer, and Karsten Reuter. Transferable ionic parameters for first-principles poisson-boltzmann solvation calculations: Neutral solutes in aqueous monovalent salt solutions. *The Journal of Chemical Physics*, 146(13):134103, 2017.
- [206] Chen Huang and Emily A Carter. Potential-functional embedding theory for molecules and materials. *The Journal of chemical physics*, 135(19), 2011.
- [207] Sebastian JR Lee, Matthew Welborn, Frederick R Manby, and Thomas F Miller III. Projection-based wavefunction-in-DFT embedding. *Accounts of chemical research*, 52(5):1359–1368, 2019.
- [208] Pierre Hohenberg and Walter Kohn. Inhomogeneous electron gas. *Physical review*, 136(3B):B864, 1964.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [209] John P. Perdew. Jacob's ladder of density functional approximations for the exchange-correlation energy. In *AIP Conference Proceedings*, volume 577, page 1–20. AIP, 2001.
- [210] Michael Seidl, Sara Giarrusso, Stefan Vuckovic, Eduardo Fabiano, and Paola Gori-Giorgi. Communication: Strong-interaction limit of an adiabatic connection in hartree-fock theory. *J. Chem. Phys.*, 149(24):241101, 2018.
- [211] Timothy J Daas, Eduardo Fabiano, Fabio Della Sala, Paola Gori-Giorgi, and Stefan Vuckovic. Noncovalent interactions from models for the møller-plesset adiabatic connection. *J. Phys. Chem. Lett.*, 12(20):4867–4875, 2021.
- [212] Lucian A. Constantin, Eduardo Fabiano, and Fabio Della Sala. Nonempirical adiabatic connection correlation functional from hartree-fock orbitals. *The Journal of Physical Chemistry Letters*, 16(13):3378–3388, 2025.
- [213] Timothy J. Daas, Derk P. Kooi, Arthur J. A. F. Grooteman, Michael Seidl, and Paola Gori-Giorgi. Gradient expansions for the large-coupling strength limit of the møller-plesset adiabatic connection. *J. Chem. Theory Comput.*, 18(3):1584–1594, 2022.
- [214] Susi Lehtola, Conrad Steigemann, Micael JT Oliveira, and Miguel AL Marques. Recent developments in libxc—a comprehensive library of functionals for density functional theory. *SoftwareX*, 7:1–5, 2018.
- [215] Wolfram Koch and Max C Holthausen. *A chemist's guide to density functional theory*. John Wiley & Sons, 2015.
- [216] David S Sholl and Janice A Steckel. *Density functional theory: a practical introduction*. John Wiley & Sons, 2022.
- [217] Luis A Agapito, Stefano Curtarolo, and Marco Buongiorno Nardelli. Reformulation of DFT+ U as a Pseudohybrid Hubbard Density Functional for Accelerated Materials Discovery. *Phys. Rev. X*, 5(1):11006, jan 2015.
- [218] V I Anisimov, J Zaanen, and O K Andersen. Band theory and Mott insulators: Hubbard U instead of Stoner I . *Phys. Rev. B*, 44(3):943–954, jul 1991.
- [219] S L Dudarev, G A Botton, S Y Savrasov, C J Humphreys, and A P Sutton. Electron-energy-loss spectra and the structural stability of nickel oxide: An LSDA+ U study. *Phys. Rev. B*, 57(3):1505–1509, jan 1998.
- [220] Luca M. Ghiringhelli, Carsten Baldauf, Tristan Berau, Sandor Brockhauser, Christian Carbogno, Javad Chamanara, Stefano Cozzini, Stefano Curtarolo, Claudia Draxl, Shyam Dwaraknath, Ádám Fekete, James Kermode, Christoph T. Koch, Markus Kühbach, Alvin Noe Ladines, Patrick Lambrix, Maja-Olivia Himmer, Sergey V. Levchenko, Micael Oliveira, Adam Michalchuk, Ronald E. Miller, Berk Onat, Pasquale Pavone, Giovanni Pizzi, Benjamin Regler, Gian-Marco Rignanese, Jörg Schaarschmidt, Markus Scheidgen, Astrid Schneidewind, Tatyana Sheveleva, Chuanxun Su, Denis Usyat, Omar Valsion, Christof Wöll, and Matthias Scheffler. Shared metadata for data-centric materials science. *Sci. Data*, 10:626, 2023.
- [221] Anders Brakestad, Peter Wind, Stig Rune Jensen, Luca Frediani, and Kathrin Helen Hopmann. Multiwavelets applied to metal-ligand interactions: Energies free from basis set errors. *J. Chem. Phys.*, 154:214302, 2021.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [222] Susi Lehtola and Miguel A. L. Marques. Reproducibility of density functional approximations: How new functionals should be reported. *J. Chem. Phys.*, 159:114116, 2023.
- [223] Robin Strange, FR Manby, and PJ Knowles. Automatic code generation in density functional theory. *Computer Physics Communications*, 136(3):310–318, 2001.
- [224] Max Dion, Henrik Rydberg, Elsebeth Schröder, David C Langreth, and Bengt I Lundqvist. Van der Waals density functional for general geometries. *Physical review letters*, 92(24):246401, 2004.
- [225] Andris Gulans, Martti J Puska, and Risto M Nieminen. Linear-scaling self-consistent implementation of the van der Waals density functional. *Physical Review B—Condensed Matter and Materials Physics*, 79(20):201105, 2009.
- [226] John A Pople, Peter M W Gill, and Benny G Johnson. Kohn-Sham density-functional theory within a finite basis set. *Chem. Phys. Lett.*, 199(6):557–560, 1992.
- [227] RT Sharp and GK Horton. A variational approach to the unipotential many-electron problem. *Physical Review*, 90(2):317, 1953.
- [228] JB Krieger, Yan Li, and GJ Iafrate. Derivation and application of an accurate kohn-sham potential with integer discontinuity. *Physics Letters A*, 146(5):256–260, 1990.
- [229] Alexei V Arbuznikov and Martin Kaupp. The self-consistent implementation of exchange-correlation functionals depending on the local kinetic energy density. *Chemical physics letters*, 381(3-4):495–504, 2003.
- [230] Ralf Neumann, Ross H Nobes, and Nicholas C Handy. Exchange functionals and potentials. *Molecular Physics*, 87(1):1–36, 1996.
- [231] Jianwei Sun, Martijn Marsman, Gábor I Csonka, Adrienn Ruzsinszky, Pan Hao, Yoon-Suk Kim, Georg Kresse, and John P Perdew. Self-consistent meta-generalized gradient approximation within the projector-augmented-wave method. *Physical Review B—Condensed Matter and Materials Physics*, 84(3):035117, 2011.
- [232] Hans-Joachim Bungartz, Christian Carbogno, Martin Galgon, Thomas Huckle, Simone Köcher, Hagen-Henrik Kowalski, Pavel Kus, Bruno Lang, Hermann Lederer, Valeriy Manin, et al. Elpa: A parallel solver for the generalized eigenvalue problem 1. In *Parallel Computing: Technology Trends*, pages 647–668. IOS Press, 2020.
- [233] Fritz Haber Institute *ab initio* molecular simulations: FHI-aims, 2025. [Online; accessed 11-Dec-2025].
- [234] Burak Himmetoglu, Andrea Floris, Stefano De Gironcoli, and Matteo Cococcioni. Hubbard-corrected DFT energy functionals: The LDA+U description of correlated systems. *International Journal of Quantum Chemistry*, 114(1):14–49, 2014.
- [235] Libxc - a library of exchange-correlation functionals for density-functional theory. <https://libxc.gitlab.io/functionals/>. Accessed: 2024-10-01.
- [236] Suhwan Song, Stefan Vuckovic, Eunji Sim, and Kieron Burke. Density-corrected DFT explained: Questions and answers. *Journal of chemical theory and computation*, 18(2):817–827, 2022.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [237] Neil Qiang Su and Xin Xu. The xyg3 type of doubly hybrid density functionals. *Wiley Interdisciplinary Reviews: Computational Molecular Science*, 6(6):721–747, 2016.
- [238] Yizhen Wang, Yajing Li, Jun Chen, Igor Ying Zhang, and Xin Xu. Doubly hybrid functionals close to chemical accuracy for both finite and extended systems: implementation and test of xyg3 and xygios. *JACS Au*, 1(5):543–549, 2021.
- [239] Steven E Wheeler and KN Houk. Integration grid errors for meta-gga-predicted reaction energies: Origin of grid errors for the mo6 suite of functionals. *Journal of Chemical Theory and Computation*, 6(2):395–404, 2010.
- [240] Yi Yao and Yosuke Kanai. Plane-wave pseudopotential implementation and performance of scan meta-gga exchange-correlation functional for extended systems. *The Journal of chemical physics*, 146(22):224105, 2017.
- [241] Albert P Bartók and Jonathan R Yates. Regularized scan functional. *The Journal of chemical physics*, 150(16), 2019.
- [242] Susi Lehtola and Miguel A. L. Marques. Many recent density functionals are numerically ill-behaved. *J. Chem. Phys.*, 157:174114, 2022.
- [243] Jeremy P Allen and Graeme W Watson. Occupation matrix control of d- and f-electron localisations using DFT+U. *Phys. Chem. Chem. Phys.*, 16(39):21016–21031, 2014.
- [244] Boris Dorado, Bernard Amadon, Michel Freyss, and Marjorie Bertolus. DFT + U calculations of the ground state and metastable states of uranium dioxide. *Phys. Rev. B*, 79:235125, Jun 2009.
- [245] Matthias Kick, Cristina Grosu, Markus Schuderer, Christoph Scheurer, and Harald Oberhofer. Mobile small polarons qualitatively explain conductivity in lithium titanium oxide battery electrodes. *The Journal of Physical Chemistry Letters*, 11(7):2535–2540, 2020.
- [246] Matthias Kick, Christoph Scheurer, and Harald Oberhofer. Formation and stability of small polarons at the lithium-terminated li4ti5o12 (lto) (111) surface. *The Journal of Chemical Physics*, 153(14):144701, 10 2020.
- [247] Matthias Kick, Christoph Scheurer, and Harald Oberhofer. Polaron-assisted charge transport in li-ion battery anode materials. *ACS Applied Energy Materials*, 4(8):8583–8591, 2021.
- [248] Daniel Mejía-Rodríguez and SB Trickey. Deorbitalized meta-gga exchange-correlation functionals in solids. *Physical Review B*, 98(11):115161, 2018.
- [249] Tomasz A Wesolowski, Sapana Shedje, and Xiuwen Zhou. Frozen-density embedding strategy for multilevel simulations of electronic structure. *Chemical reviews*, 115(12):5891–5928, 2015.
- [250] Matteo Cococcioni and Stefano De Gironcoli. Linear response approach to the calculation of the effective interaction parameters in the LDA+U method. *Phys. Rev. B*, 71(3):35105, 2005.
- [251] Vivaldo Leiria Campo and Matteo Cococcioni. Extended DFT+U+V method with on-site and inter-site electronic interactions. *J. Phys. Cond. Matter*, 22(5):55602, 2010.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [252] Haipeng Lu, Gabrielle Koknat, Yi Yao, Ji Hao, Xixi Qin, Chuanxiao Xiao, Ruyi Song, Florian Merz, Markus Rampp, Sebastian Kokott, Christian Carbogno, Tianyang Li, Glenn Teeter, Matthias Scheffler, Joseph J. Berry, David B. Mitzi, Jeffrey L. Blackburn, Volker Blum, and Matthew C. Beard. Electronic impurity doping of a 2d hybrid lead iodide perovskite by Bi and Sn. *PRX Energy*, 2:023010, Jun 2023.
- [253] Juana Moreno and José M Soler. Optimal meshes for integrals in real-and reciprocal-space unit cells. *Physical Review B*, 45(24):13891, 1992.
- [254] Matthias Ernzerhof and Gustavo E Scuseria. Assessment of the perdew–burke–ernzerhof exchange–correlation functional. *The Journal of chemical physics*, 110(11):5029–5036, 1999.
- [255] Carlo Adamo and Vincenzo Barone. Toward reliable density functional methods without adjustable parameters: The pbeo model. *The Journal of chemical physics*, 110(13):6158–6170, 1999.
- [256] Jorge M del Campo, José L Gázquez, SB Trickey, and Alberto Vela. Non-empirical improvement of pbe and its hybrid pbeo for general description of molecular properties. *The Journal of chemical physics*, 136(10), 2012.
- [257] Philip J Stephens, Frank J Devlin, Cary F Chabalowski, and Michael J Frisch. Ab initio calculation of vibrational absorption and circular dichroism spectra using density functional force fields. *The Journal of physical chemistry*, 98(45):11623–11627, 1994.
- [258] Jochen Heyd, Gustavo E Scuseria, and Matthias Ernzerhof. Hybrid functionals based on a screened coulomb potential. *The Journal of chemical physics*, 118(18):8207–8215, 2003.
- [259] Jochen Heyd, Gustavo E. Scuseria, and Matthias Ernzerhof. Erratum: “Hybrid functionals based on a screened Coulomb potential” [J. Chem. Phys. 118, 8207 (2003)]. *The Journal of Chemical Physics*, 124(21), 06 2006. 219906.
- [260] Oleg A. Vydrov and Gustavo E. Scuseria. Assessment of a long-range corrected hybrid functional. *The Journal of Chemical Physics*, 125(23):234109, 12 2006.
- [261] Yan Zhao and Donald G Truhlar. The mo6 suite of density functionals for main group thermochemistry, thermochemical kinetics, noncovalent interactions, excited states, and transition elements: two new functionals and systematic testing of four mo6-class functionals and 12 other functionals. *Theoretical chemistry accounts*, 120:215–241, 2008.
- [262] Yan Zhao and Donald G Truhlar. Exploring the limit of accuracy of the global hybrid meta density functional for main-group thermochemistry, kinetics, and noncovalent interactions. *Journal of Chemical Theory and Computation*, 4(11):1849–1868, 2008.
- [263] Roberto Peverati and Donald G Truhlar. Improving the accuracy of hybrid meta-gga density functionals by range separation. *The Journal of Physical Chemistry Letters*, 2(21):2810–2817, 2011.
- [264] Michael Lorke, Peter Deák, and Thomas Frauenheim. Koopmans-compliant screened exchange potential with correct asymptotic behavior for semiconductors. *Physical Review B*, 102(23):235168, 2020.
- [265] Danish Khan, Alastair JA Price, Bing Huang, Maximilian L Ach, and O Anatole von Lilienfeld. Adapting hybrid density functionals with machine learning. *Science Advances*, 11(5):eadt7769, 2025.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [266] Adaptive hybrid density functionals tutorial. <https://adaptive-hybrid-density-functionals-6101eb.gitlab.io/>, 2025. Accessed: 2025-12-19.
- [267] Sebastian Kokott, Volker Blum, and Matthias Scheffler. Efficient computation of the long-range exact exchange using an extended screening function. *The Journal of Chemical Physics*, 162(22), 2025.
- [268] libxc-X. libxc-X gitlab repository. <https://gitlab.com/FHI-aims-club/libxc-x>, 2024. [Online; accessed 26-Sep-2024].
- [269] A. Tkatchenko and M. Scheffler. Accurate molecular van der Waals interactions from ground-state electron density and free-atom reference data. *Phys. Rev. Lett.*, 102:073005, 2009.
- [270] A. Tkatchenko, R. A. DiStasio Jr., R. Car, and M. Scheffler. Accurate and efficient method for many-body van der Waals interactions. *Phys. Rev. Lett.*, 108:236402, 2012.
- [271] Alberto Ambrosetti, Anthony M Reilly, Robert A DiStasio Jr., and Alexandre Tkatchenko. Long-range correlation energy calculated from coupled atomic response functions. *The Journal of chemical physics*, 140(18), 2014.
- [272] J. Hermann and A. Tkatchenko. Density functional model for van der Waals interactions: Unifying many-body atomic approaches with nonlocal functionals. *Phys. Rev. Lett.*, 124:146401, 2020.
- [273] E. R. Johnson and A. D. Becke. A post-hartree-fock model of intermolecular interactions: Inclusion of higher-order corrections. *J. Chem. Phys.*, 124:174104, 2006.
- [274] E. R. Johnson. The exchange-hole dipole moment dispersion model. In G. A. DiLabio and A. Otero de la Roza, editors, *Non-covalent interactions in quantum chemistry and physics*, volume 5, pages 169–194. Elsevier, 2017.
- [275] A. J. A. Price, A. Otero de la Roza, and E. R. Johnson. XDM-corrected hybrid DFT with numerical atomic orbitals predicts molecular crystal lattice energies with unprecedented accuracy. *Chem. Sci.*, 14:1252–1262, 2023.
- [276] J. Hermann, M. Stöhr, S. Göger, S. Chaudhuri, B. Aradi, R. J. Maurer, and A. Tkatchenko. libMBD: A general-purpose package for scalable quantum many-body dispersion calculations. *J. Chem. Phys.*, 159:174802, 2023.
- [277] N. Ferri, R. A. DiStasio Jr., A. Ambrosetti, R. Car, and A. Tkatchenko. Electronic properties of molecules and surfaces with a self-consistent interatomic van der Waals density functional. *Phys. Rev. Lett.*, 114:176802, 2015.
- [278] N. Ferri, A. Ambrosetti, and A. Tkatchenko. Electronic charge rearrangement at metal/organic interfaces induced by weak van der Waals interactions. *Phys. Rev. Mat.*, 1:026003, 2017.
- [279] Stefan Grimme, Jens Antony, Stephan Ehrlich, and Helge Krieg. A consistent and accurate ab initio parametrization of density functional dispersion correction (dft-d) for the 94 elements h-pu. *The Journal of chemical physics*, 132(15), 2010.
- [280] Stefan Grimme, Stephan Ehrlich, and Lars Goerigk. Effect of the damping function in dispersion corrected density functional theory. *Journal of computational chemistry*, 32(7):1456–1465, 2011.
- [281] simple dftd3. <https://github.com/dftd3/simple-dftd3>.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [282] Sebastian Ehlert. Simple dft-d3: Library first implementation of the d3 dispersion correction. *Journal of Open Source Software*, 9:7169, 2024.
- [283] Kyle Bryenton and Erin Johnson. Dispersion-Corrected DFT with XDM. <https://fhi-aims-club.gitlab.io/tutorials/xdm-tutorial/>, 2024. [Online; accessed 28-Oct-2024].
- [284] T. Björkman, A. Gulans, A. V. Krasheninnikov, and R. M. Nieminen. Are we van der Waals ready? *J. Phys.: Condens. Matter*, 24:424218, 2012.
- [285] A. J. A. Price, K. R. Bryenton, and E. R. Johnson. Requirements for an accurate dispersion-corrected density functional. *J. Chem. Phys.*, 154:230902, 2021.
- [286] J. F. Dobson and A. Ambrosetti. MBD+C: how to incorporate metallic character into atom-based dispersion energy schemes. *J. Chem. Theory Comput.*, 19:6434–6451, 2023.
- [287] A. Otero de la Roza, L. M. LeBlanc, and E. R. Johnson. What is “many-body” dispersion and should I worry about it? *Phys. Chem. Chem. Phys.*, 22:8266–8276, 2020.
- [288] K. R. Bryenton and E. R. Johnson. Many-body dispersion in model systems and the sensitivity of self-consistent screening. *J. Chem. Phys.*, 158:204110, 2023.
- [289] T. Bučko, S. Lebègue, J. G. Ángyán, and J. Hafner. Extending the applicability of the Tkatchenko-Scheffler dispersion correction via iterative Hirshfeld partitioning. *J. Chem. Phys.*, 141:034114, 2014.
- [290] T. Gould, S. Lebègue, J. G. Ángyán, and T. Bučko. A fractionally ionic approach to polarizability and van der Waals many-body dispersion calculations. *J. Chem. Theory Comput.*, 12:5920–5930, 2016.
- [291] Axel D. Becke. A remarkably simple dispersion damping scheme and the dh24 double hybrid density functional. *The Journal of Chemical Physics*, 160(20):204118, 05 2024.
- [292] S. Góger, A. Khabibrakhmanov, O Vaccarelli, D. V. Fedorov, and A. Tkatchenko. Optimized quantum drude oscillators for atomic and molecular response properties. *J. Phys. Chem. Lett.*, 14:6217–6223, 2023.
- [293] D. Massa, A. Ambrosetti, and P. L. Silvestrelli. Beyond-dipole van der Waals contributions within the many-body dispersion framework. *Elec. Struct.*, 3:044002, 2021.
- [294] X. Ren, P. Rinke, C. Joas, and M. Scheffler. Random-phase approximation and its applications in computational chemistry and materials science. *J. Mater. Sci.*, 47:7447, 2012.
- [295] H. Eshuis, J. E. Bates, and F. Furche. Electron correlation methods based on the random phase approximation. *Theor. Chem. Acc.*, 131:1084, 2012.
- [296] X. Ren, A. Tkatchenko, P. Rinke, and M. Scheffler. Beyond the random-phase approximation for the electron correlation energy: The importance of single excitations. *Phys. Rev. Lett.*, 106:153003, 2011.
- [297] A. Grüneis, M. Marsman, J. Harl, L. Schimka, and G. Kresse. Making the random phase approximation to electronic correlation accurate. *J. Chem. Phys.*, 131:154115, 2009.
- [298] X. Ren, P. Rinke, G. Scuseria, and M. Scheffler. Renormalized second-order perturbation theory for the electron correlation energy: Concept, implementation, and benchmarks. *Phys. Rev. B*, 88:035120, 2013.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [299] X. Ren, P. Rinke, and M. Scheffler. Exploring the random phase approximation: Application to co-adsorbed on cu(111). *Phys. Rev. B*, 80:045402, 2009.
- [300] Muhammad N Tahir, Honghui Shang, Jia Li, and Xinguo Ren. Efficient structural relaxation based on the random phase approximation: Applications to water clusters. *The Journal of Physical Chemistry A*, 2024.
- [301] R. Shi, M.-Y. Zhang, P. Lin, L. He, and X. Ren. Subquadratic-scaling real-space random phase approximation correlation energy calculations for periodic systems with numerical atomic orbitals. *Phys. Rev. B*, 109:035103, 2024.
- [302] Igor Ying Zhang, Rong Shi, and Xinguo Ren. Introduction to the fifth-rung density functional approximations: concept, formulation, and applications. *Electronic Structure*, 7(4):043002, 2025.
- [303] H. Shang, C. Carbogno, P. Rinke, and M. Scheffler. Lattice dynamics calculations based on density-functional perturbation theory in real space. *Comput. Phys. Commun.*, 215:26, 2017.
- [304] Sixian Yang and Xinguo Ren. Phase stability of the argon crystal: first-principles study based on random phase approximation plus renormalized single excitation corrections. *New Journal of Physics*, 24(3):033049, 2022.
- [305] M. Azizi, J. Wilhelm, D. Golze, M. Giantomassi, R. L. Panadés-Barrueta, F. A. Delesma, A. Buccheri, A. Gulans, P. Rinke, C. Draxl, and X. Gonze. Time-frequency component of the greenx library: minimax grids for efficient rpa and gw calculations. *J. Open Source Softw.*, 8:5570, 2023.
- [306] M. Azizi, J. Wilhelm, D. Golze, F. A. Delesma, R. L. Panadés-Barrueta, P. Rinke, M. Giantomassi, and X. Gonze. Validation of the greenx library time-frequency component for efficient gw and rpa calculations. *Phys. Rev. B*, 109:245101, 2024.
- [307] Rong Shi, Min-Ye Zhang, Peize Lin, Lixin He, and Xinguo Ren. Librpa: A software package for low-scaling first-principles calculations of random phase approximation electron correlation energy based on numerical atomic orbitals. *Computer Physics Communications*, 309:109496, 2025.
- [308] Muhammad N. Tahir, Honghui Shang, and Xinguo Ren. Analytical gradients of random-phase approximation plus corrections from renormalized single excitations. *Journal of Chemical Theory and Computation*, 21(21):10822–10841, 2025.
- [309] Fhi-aims tutorial: Rpa and gw for molecules and solids. <https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids/Part-2>.
- [310] Sixian Yang, Igor Ying Zhang, and Xinguo Ren. Developing correlation-consistent numeric atom-centered orbital basis sets for krypton: Applications in RPA-based correlated calculations. *J. Chem. Phys.*, 160(2):024112, 01 2024.
- [311] Igor Ying Zhang and Andreas Grüneis. Coupled cluster theory in materials science. *Front. Mater.*, 6:123, 2019.
- [312] Evgeny Moerman, Felix Hummel, Andreas Grüneis, Andreas Irmeler, and Matthias Scheffler. Interface to high-performance periodic coupled-cluster theory calculations with atom-centered, localized basis functions. *J. Open Source Softw.*, 7:4040, 2022.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [313] Felix Hummel, Theodoros Tsatsoulis, and Andreas Grüneis. Low rank factorization of the coulomb integrals for periodic coupled cluster theory. *J. Chem. Phys.*, 146:124105, 2017.
- [314] Evgeny Moerman. Fhi-aims coupled cluster tutorial. <https://periodic-cc-methods-in-fhi-aims-fhi-aims-club-tu-9d435dcb75ed8d.gitlab.io/>, 2024. [Online; accessed 09-Oct-2024].
- [315] Evgeny Moerman, Alejandro Gallo, Andreas Irmeler, Tobias Schäfer, Felix Hummel, Andreas Grüneis, and Matthias Scheffler. Finite-size effects in periodic eom-ccsd for ionization energies and electron affinities: Convergence rate and extrapolation to the thermodynamic limit. *Journal of Chemical Theory and Computation*, 21(4):1865–1878, 2025.
- [316] Divya Tripathi and Achintya Kumar Dutta. The performance of approximate eom-ccsd for ionization potential and electron affinity of genetic material subunits: A benchmark investigation. *Int. J. Quantum Chem.*, 122:e26918, 2022.
- [317] Tonghao Shen, Zhenyu Zhu, Igor Ying Zhang, and Matthias Scheffler. Massive-parallel implementation of the resolution-of-identity coupled-cluster approaches in the numeric atom-centered orbital framework for molecular systems. *J. Chem. Theory Comput.*, 15:4721–4734, 2019.
- [318] R. J. Hunt, B. Monserrat, V. Zólyomi, and N. D. Drummond. Diffusion quantum monte carlo and gw study of the electronic properties of monolayer and bulk hexagonal boron nitride. *Phys. Rev. B*, 101:205115, 2020.
- [319] Krzysztof Rościszewski, Beate Paulus, Peter Fulde, and Hermann Stoll. Ab initio calculation of ground-state properties of rare-gas crystals. *Phys. Rev. B*, 60:7905, 1999.
- [320] Yasmine S. Al-Hamdani, Péter R. Nagy, Andrea Zen, Dennis Barton, Mihály Kállay, Jan Gerit Brandenburg, and Alexandre Tkatchenko. Interactions between large molecules pose a puzzle for reference quantum mechanical methods. *Nat. Commun.*, 12:3927, 2021.
- [321] Evgeny Moerman, Henrique Miranda, Alejandro Gallo, Andreas Irmeler, Tobias Schäfer, Felix Hummel, Manuel Engel, Georg Kresse, Matthias Scheffler, and Andreas Grüneis. Exploring the accuracy of the equation-of-motion coupled-cluster band gap of solids. *Phys. Rev. B*, 111:L121202, Mar 2025.
- [322] Per-Olov Löwdin. Quantum theory of many-particle systems. i. physical interpretations by means of density matrices, natural spin-orbitals, and convergence problems in the method of configurational interaction. *Phys. Rev.*, 97:1474, 1955.
- [323] Andreas Grüneis, George H. Booth, Martijn Marsman, James Spencer, Ali Alavi, and Georg Kresse. Natural orbitals for wave function based correlated calculations using a plane wave basis set. *J. Chem. Theory Comput.*, 7:2780–2785, 2011.
- [324] Florian Libisch, Chen Huang, and Emily A Carter. Embedded correlated wavefunction schemes: Theory and applications. *Accounts of chemical research*, 47(9):2768–2775, 2014.
- [325] Benjamin X Shi, Andrew S Rosen, Tobias Schäfer, Andreas Grüneis, Venkat Kapil, Andrea Zen, and Angelos Michaelides. An accurate and efficient framework for modelling the surface chemistry of ionic materials. *Nature Chemistry*, pages 1–8, 2025.
- [326] Tobias Schäfer, Alejandro Gallo, Andreas Irmeler, Felix Hummel, and Andreas Grüneis. Surface science using coupled cluster theory via local wannier functions and in-rpa-embedding: The case of water on graphitic carbon nitride. *The Journal of Chemical Physics*, 155(24), 2021.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [327] Péter R Nagy, Gyula Samu, and Mihály Kállay. Optimization of the linear-scaling local natural orbital ccSD (t) method: Improved algorithm and benchmark applications. *Journal of Chemical Theory and Computation*, 14(8):4193–4215, 2018.
- [328] Dimitrios G Liakos, Manuel Sparta, Manoj K Kesharwani, Jan ML Martin, and Frank Neese. Exploring the accuracy limits of local pair natural orbital coupled-cluster theory. *Journal of chemical theory and computation*, 11(4):1525–1539, 2015.
- [329] Hong-Zhou Ye and Timothy C Berkelbach. Periodic local coupled-cluster theory for insulators and metals. *Journal of Chemical Theory and Computation*, 20(20):8948–8959, 2024.
- [330] Péter R Nagy and Mihály Kállay. Approaching the basis set limit of ccSD (t) energies for large molecules with local natural orbital coupled-cluster methods. *Journal of Chemical Theory and Computation*, 15(10):5275–5298, 2019.
- [331] Isolde Sandler, Junbo Chen, Mackenzie Taylor, Shaleen Sharma, and Junming Ho. Accuracy of dlpno-ccSD (t): Effect of basis set and system size. *The Journal of Physical Chemistry A*, 125(7):1553–1563, 2021.
- [332] P Bernát Szabó, József Csóka, Mihály Kállay, and Péter R Nagy. Linear-scaling local natural orbital ccSD (t) approach for open-shell systems: Algorithms, benchmarks, and large-scale applications. *Journal of Chemical Theory and Computation*, 19(22):8166–8188, 2023.
- [333] Lars Hedin. New method for calculating the one-particle green's function with application to the electron-gas problem. *Phys. Rev.*, 139(3A):A796, 1965.
- [334] Dorothea Golze, Marc Dvorak, and Patrick Rinke. The GW Compendium: A Practical Guide to Theoretical Photoemission Spectroscopy. *Front. Chem.*, 7:377, 2019.
- [335] Dorothea Golze, Jan Wilhelm, Michiel J. van Setten, and Patrick Rinke. Core-Level Binding Energies from GW: An Efficient Full-Frequency Approach within a Localized Basis. *J. Chem. Theory Comput.*, 14:4856–4869, 2018.
- [336] Dorothea Golze, Levi Keller, and Patrick Rinke. Accurate Absolute and Relative Core-Level Binding Energies from GW. *J. Phys. Chem. Lett.*, 11(5):1840–1847, 2020.
- [337] Jiachen Li, Ye Jin, Patrick Rinke, Weitao Yang, and Dorothea Golze. Benchmark of gw methods for core-level binding energies. *J. Chem. Theory Comput.*, 18(12):7570–7585, 2022.
- [338] F. Aryasetiawan and S. Biermann. Generalized hedin's equations for quantum many-body systems with spin-dependent interactions. *Phys. Rev. Lett.*, 100:116402, Mar 2008.
- [339] Qinglong Liu and Dorothea Golze. All-electron two-component *GW* implementation with numeric atomic centered orbitals. *to be published*, 2024.
- [340] F. Caruso, P. Rinke, X. Ren, M. Scheffler, and A. Rubio. Unified description of ground and excited states of finite systems: The self-consistent *gw* approach. *Phys. Rev. B*, 86:081102, Aug 2012.
- [341] Fabio Caruso, Patrick Rinke, Xinguo Ren, Angel Rubio, and Matthias Scheffler. Self-consistent *gw*: All-electron implementation with localized basis functions. *Phys. Rev. B*, 88:075105, Aug 2013.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [342] Ramón L. Panadés-Barrueta and Dorothea Golze. Accelerating Core-Level GW Calculations by Combining the Contour Deformation Approach with the Analytic Continuation of W. *J. Chem. Theory Comput.*, 19(16):5450–5464, 2023.
- [343] Jannis Kockläuner, Majid Shaker, Maximilian Muth, Simon Steinbach, Christoph Oleszak, Ole Lytken, Hans-Peter Steinrück, and Dorothea Golze. Decoding shake-up satellites in xps through large-scale ab initio simulations: Spectral signatures of ring fusion in porphyrins. *arXiv:2509.26057*, 2025.
- [344] O. Vahtras, J. Almlöf, and M.W. Feyereisen. Integral approximations for LCAO-SCF calculations. *Chem. Phys. Lett.*, 213:514–518, 1993.
- [345] Xinguo Ren, Noa Marom, Fabio Caruso, Matthias Scheffler, and Patrick Rinke. Beyond the *gw* approximation: A second-order screened exchange correction. *Phys. Rev. B*, 92:081104, Aug 2015.
- [346] Yanyong Wang, Patrick Rinke, and Xinguo Ren. Assessing the $g_0w_0\gamma_0^{(1)}$ approach: Beyond *gowo* with hedin’s full second-order self-energy contribution. *J. Chem. Theory Comput.*, 17(8):5140–5154, 2021. PMID: 34319724.
- [347] Jannis Kockläuner and Dorothea Golze. Gw plus cumulant approach for predicting core-level shakeup satellites in large molecules. *J. Chem. Theory Comput.*, 21(6):3101–3119, 2025.
- [348] Fabien Bruneval and Arno Förster. Fully dynamic *g3w2* self-energy for finite systems: Formulas and benchmark. *J. Chem. Theory Comput.*, 20(8):3218–3230, 2024. PMID: 38603811.
- [349] Ferdi Aryasetiawan, Lars Hedin, and Krister Karlsson. Multiple plasmon satellites in na and al spectral functions from ab initio cumulant expansion. *Phys. Rev. Lett.*, 77(11):2268, 1996.
- [350] Matteo Guzzo, Giovanna Lani, Francesco Sottile, Pina Romaniello, Matteo Gatti, Joshua J Kas, John J Rehr, Mathieu G Silly, Fausto Sirotti, and Lucia Reining. Valence electron photoemission spectrum of semiconductors: Ab initio description of multiple satellites. *Phys. Rev. Lett.*, 107(16):166401, 2011.
- [351] Matteo Gatti and Matteo Guzzo. Dynamical screening in correlated metals: Spectral properties of srvo_3 in the *gw* approximation and beyond. *Phys. Rev. B*, 87:155147, Apr 2013.
- [352] Fabio Caruso and Feliciano Giustino. Spectral fingerprints of electron-plasmon coupling. *Phys. Rev. B*, 92:045123, Jul 2015.
- [353] Jianqiang Sky Zhou, J. J. Kas, Lorenzo Sponza, Igor Reshetnyak, Matteo Guzzo, Christine Giorgetti, Matteo Gatti, Francesco Sottile, J. J. Rehr, and Lucia Reining. Dynamical effects in electron spectroscopy. *J. Chem. Phys.*, 143(18):184109, 2015.
- [354] Jianqiang Sky Zhou, Matteo Gatti, JJ Kas, JJ Rehr, and Lucia Reining. Cumulant green’s function calculations of plasmon satellites in bulk sodium: Influence of screening and the crystal environment. *Phys. Rev. B*, 97(3):035137, 2018.
- [355] Gw eigenvalues and theoretical spectroscopy. <https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids/Part-1/GWEigenvaluesAndPhotoemission/>, 2024. Accessed: 2024-09-18.
- [356] Core levels with gw. <https://fhi-aims-club.gitlab.io/tutorials/rpa-and-gw-for-molecules-and-solids/Part-1/CoreLevels/>, 2024. Accessed: 2024-09-18.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [357] Annika Stuke, Christian Kunkel, Dorothea Golze, Milica Todorović, Johannes T Margraf, Karsten Reuter, Patrick Rinke, and Harald Oberhofer. Atomic structures and orbital energies of 61,489 crystal-forming organic molecules. *Sci. Data*, 7(1):1–11, 2020.
- [358] Dorothea Golze, Markus Hirvensalo, Patricia Hernández-León, Anja Aarva, Jarkko Etula, Toma Susi, Patrick Rinke, Tomi Laurila, and Miguel A. Caro. Accurate Computational Prediction of Core-Electron Binding Energies in Carbon-Based Materials: A Machine-Learning Model Combining Density-Functional Theory and GW. *Chem. Mater.*, 34(14):6240–6254, 2022.
- [359] N. Marom, F. Caruso, X. Ren, O. T. Hofmann, T. Körzdörfer, J. R. Chelikowsky, A. Rubio, M. Scheffler, and P. Rinke. Benchmark of *GW* methods for azabenzenes. *Phys. Rev. B*, 86:245127, Dec 2012.
- [360] Fabio Caruso, Matthias Dauth, Michiel J. van Setten, and Patrick Rinke. Benchmark of GW Approaches for the GW100 Test Set. *J. Chem. Theory Comput.*, 12(10):5076–5087, September 2016.
- [361] Mia Schambeck, Dorothea Golze, and Jan Wilhelm. Solving multipole challenges in the *gw*100 benchmark enables precise low-scaling *gw* calculations. *Phys. Rev. B*, 110:125146, Sep 2024.
- [362] Jan Wilhelm, Patrick Seewald, and Dorothea Golze. Low-Scaling GW with Benchmark Accuracy and Application to Phosphorene Nano-sheets. *J. Chem. Theory Comput.*, 17(3):1662–1677, 2021.
- [363] A. K. Wilson, T. van Mourik, and T. H. Dunning. Gaussian basis sets for use in correlated molecular calculations. VI. Sextuple zeta correlation consistent basis sets for boron through neon. *J. Mol. Struct. (Theochem)*, 388:339–349, 1996.
- [364] Florian Weigend and Reinhart Ahlrichs. Balanced basis sets of split valence, triple zeta valence and quadruple zeta valence quality for h to rn: Design and assessment of accuracy. *Phys. Chem. Chem. Phys.*, 7:3297–3305, 2005.
- [365] Maximilien A Ambroise, Andreas Dreuw, and Frank Jensen. Probing basis set requirements for calculating core ionization and core excitation spectra using correlated wave function methods. *J. Chem. Theory Comput.*, 17(5):2832–2842, 2021.
- [366] Daniel Mejia-Rodriguez, Alexander Kunitsa, Edoardo Aprà, and Niranjana Govind. Basis set selection for molecular core-level gw calculations. *J. Chem. Theory Comput.*, 18(8):4919–4926, 2022.
- [367] Hao Peng, Sixian Yang, Hong Jiang, Hongming Weng, and Xinguo Ren. Basis-set-error-free random-phase approximation correlation energies for atoms based on the sternheimer equation. *J. Chem. Theory Comput.*, 19(20):7199–7214, 2023. PMID: 37811855.
- [368] Francisco A. Delesma, Moritz Leucke, Dorothea Golze, and Patrick Rinke. Benchmarking the accuracy of the separable resolution of the identity approach for correlated methods in the numeric atom-centered orbitals framework. *J. Chem. Phys.*, 160(2):024118, 01 2024.
- [369] M. S. Hybertsen and S. G. Louie. Electron correlation in semiconductors and insulators: Band gaps and quasiparticle energies. *Phys. Rev. B*, 34:5390, 1986.
- [370] Lucia Reining. The GW approximation: Content, successes and limitations. *WIREs Comput Mol Sci*, 8(3):e1344, 2018.
- [371] H. N. Rojas, R. W. Godby, and R. J. Needs. Space-time method for ab initio calculations of self-energies and dielectric response functions of solids. *Phys. Rev. Lett.*, 74:1827, 1995.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [372] M. Shishkin and G. Kresse. Implementation and performance of the frequency-dependent gw method within the paw framework. *Phys. Rev. B*, 74:035101, 2006.
- [373] C. Friedrich, A. Schindlmayr, and S. Blügel. Efficient implementation of the gw approximation within the all-electron flapw method. *Phys. Rev. B*, 81:125102, 2010.
- [374] H. Jiang, R. I. Gómez-Abal, X.-Z. Li, C. Meisenbichler, C. Ambrosch-Draxl, and M. Scheffler. Fhi-gap: A gw code based on the all-electron augmented plane wave method. *Comp. Phys. Commun.*, 184:348, 2013.
- [375] T. Zhu and G. K.-L. Chan. All-electron gaussian-based gowo for valence and core excitation energies of periodic systems. *J. Chem. Theory Comput.*, 17:727, 2021.
- [376] H. Jiang and P. Blaha. Gw with linearized augmented plane waves extended by high-energy local orbitals. *Phys. Rev. B*, 93:115203, 2016.
- [377] Maryam Azizi, Francisco A. Delesma, Matteo Giantomassi, Davis Zavickis, Mikael Kuisma, Kristian Thyghesen, Dorothea Golze, Alexander Buccheri, Min-Ye Zhang, Patrick Rinke, Claudia Draxl, Andris Gulans, and Xavier Gonze. Precision benchmarks for solids: GoWo calculations with different basis sets. *Comput. Mater. Sci.*, 250:113655, 2025.
- [378] F. Giustino. Electron-phonon interactions from first principles. *Rev. Mol. Phys.*, 89:015003, 2017.
- [379] C. Pellegrini and A. Sanna. Ab initio methods for superconductivity. *Nat. Rev. Phys.*, 6:509, 2024.
- [380] Ruiyi Zhou, Yi Yao, Volker Blum, Xinguo Ren, and Yosuke Kanai. All-electron *bse@gw* method with numeric atom-centered orbitals for extended systems. *J. Chem. Theory Comput.*, 21:291–306, 2025.
- [381] Michael Rohlfing and Steven G Louie. Electron-hole excitations and optical spectra from first principles. *Physical Review B*, 62(8):4927, 2000.
- [382] G Strinati. Effects of dynamical screening on resonances at inner-shell thresholds in semiconductors. *Physical Review B*, 29(10):5718, 1984.
- [383] Marc Dvorak and Patrick Rinke. Dynamical configuration interaction: Quantum embedding that combines wave functions and green's functions. *Phys. Rev. B*, 99:115134, Mar 2019.
- [384] Marc Dvorak, Dorothea Golze, and Patrick Rinke. Quantum embedding theory in the screened coulomb interaction: Combining configuration interaction with *gw*/BSE. *Phys. Rev. Mater.*, 3:070801, Jul 2019.
- [385] Mark E. Casida. *Time-Dependent Density Functional Response Theory for Molecules*, page 155–192. WORLD SCIENTIFIC, November 1995.
- [386] Giovanni Onida, Lucia Reining, and Angel Rubio. Electronic excitations: density-functional versus many-body green's-function approaches. *Reviews of modern physics*, 74(2):601, 2002.
- [387] Xavier Blase, Ivan Duchemin, and Denis Jacquemin. The bethe–salpeter equation in chemistry: relations with td-dft, applications and challenges. *Chem. Soc. Rev.*, 47(3):1022–1043, 2018.
- [388] David E. Woon and Thom H. Dunning. Gaussian basis sets for use in correlated molecular calculations. iv. calculation of static electrical response properties. *J. Chem. Phys.*, 100(4):2975–2988, 1994.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [389] Marc Dvorak and Patrick Rinke. A balanced treatment of static and dynamic correlation in free- and mg-porphyrin, 2020.
- [390] Sampreeti Bhattacharya, Jiachen Li, Weitao Yang, and Yosuke Kanai. Bse@gw prediction of charge transfer exciton in molecular complexes: Assessment of self-energy and exchange-correlation dependence. *The Journal of Physical Chemistry A*, 128(29):6072–6083, 2024. PMID: 39011742.
- [391] David Kammerlander, Silvana Botti, Miguel A. L Marques, Andrea Marini, and Claudio Attaccalite. Speeding up the solution of the bethe-salpeter equation by a double-grid method and wannier interpolation. *Phys. Rev. B*, 86:125203, Sep 2012.
- [392] Tobias Sander, Emanuele Maggio, and Georg Kresse. Beyond the tamm-dancoff approximation for extended systems using exact diagonalization. *Physical Review B*, 92(4):045209, 2015.
- [393] Jack Deslippe, Georgy Samsonidze, David A Strubbe, Manish Jain, Marvin L Cohen, and Steven G Louie. Berkeleygw: A massively parallel computer package for the calculation of the quasiparticle and optical properties of materials and nanostructures. *Computer Physics Communications*, 183(6):1269–1289, 2012.
- [394] Ph. Ghosez, X. Gonze, and R. W. Godby. Long-wavelength behavior of the exchange-correlation kernel in the kohn-sham theory of periodic systems. *Phys. Rev. B*, 56:12811–12817, Nov 1997.
- [395] Young-Moo Byun, Jiuyu Sun, and Carsten A Ullrich. Time-dependent density-functional theory for periodic solids: assessment of excitonic exchange-correlation kernels. *Electronic Structure*, 2(2):023002, apr 2020.
- [396] Zeng-hui Yang, Francesco Sottile, and Carsten A. Ullrich. Simple screened exact-exchange approach for excitonic properties in solids. *Phys. Rev. B*, 92:035202, Jul 2015.
- [397] Dahvyd Wing, Jonah B. Haber, Roy Noff, Bradford Barker, David A. Egger, Ashwin Ramasubramaniam, Steven G. Louie, Jeffrey B. Neaton, and Leeor Kronik. Comparing time-dependent density functional theory with many-body perturbation theory for semiconductors: Screened range-separated hybrids and the *gw* plus bethe-salpeter approach. *Phys. Rev. Mater.*, 3:064603, Jun 2019.
- [398] Lucia Reining, Valerio Olevano, Angel Rubio, and Giovanni Onida. Excitonic effects in solids described by time-dependent density-functional theory. *Phys. Rev. Lett.*, 88:066404, Jan 2002.
- [399] Xiaosong Li, Niranjana Govind, Christine Isborn, A. Eugene DePrince, and Kenneth Lopata. Real-time time-dependent electronic structure theory. *Chemical Reviews*, 120(18):9951–9993, 2020.
- [400] Erich Runge and E. K. U. Gross. Density-functional theory for time-dependent systems. *Phys. Rev. Lett.*, 52:997–1000, 1984.
- [401] Jianhang Xu, Thomas E. Carney, Ruiyi Zhou, Christopher Shepard, and Yosuke Kanai. Real-time time-dependent density functional theory for simulating nonequilibrium electron dynamics. *Journal of the American Chemical Society*, 146(8):5011–5029, 2024. PMID: 38362887.
- [402] Christopher Shepard, Ruiyi Zhou, Dillon C. Yost, Yi Yao, and Yosuke Kanai. Simulating electronic excitation and dynamics with real-time propagation approach to TDDFT within plane-wave pseudopotential formulation. *The Journal of Chemical Physics*, 155(10):100901, 2021.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [403] Alina Kononov, Cheng-Wei Lee, Tatiane Pereira dos Santos, Brian Robinson, Yifan Yao, Yi Yao, Xavier Andrade, Andrew David Baczewski, Emil Constantinescu, Alfredo A. Correa, Yosuke Kanai, Normand Modine, and André Schleife. Electron dynamics in extended systems within real-time time-dependent density-functional theory. *MRS Communications*, 12(6), 2022.
- [404] Takehiro Yonehara, Kota Hanasaki, and Kazuo Takatsuka. Fundamental approaches to nonadiabaticity: Toward a chemical theory beyond the born-oppenheimer paradigm. *Chemical Reviews*, 112(1):499–542, 2012. PMID: 22077497.
- [405] John Crank and Phyllis Nicolson. A practical method for numerical evaluation of solutions of partial differential equations of the heat-conduction type. In *Mathematical Proceedings of the Cambridge Philosophical Society*, volume 43, pages 50–67. Cambridge University Press, 1947.
- [406] Ying Zhu and John M. Herbert. Self-consistent predictor/corrector algorithms for stable and efficient integration of the time-dependent kohn-sham equation. *The Journal of Chemical Physics*, 148(4):044117, 2018.
- [407] Sharon Hammes-Schiffer. Nuclear-electronic orbital methods: Foundations and prospects. *J. Chem. Phys.*, 155(3):030901, Jul 2021.
- [408] Jianhang Xu, Ruiyi Zhou, Volker Blum, Tao E Li, Sharon Hammes-Schiffer, and Yosuke Kanai. First-principles approach for coupled quantum dynamics of electrons and protons in heterogeneous systems. *Phys. Rev. Lett.*, 131(23):238002, 2023.
- [409] Jianhang Xu, Ruiyi Zhou, Zhen Tao, Christopher Malbon, Volker Blum, Sharon Hammes-Schiffer, and Yosuke Kanai. Nuclear-electronic orbital approach to quantization of protons in periodic electronic structure calculations. *J. Chem. Phys.*, 156(22):224111, 2022.
- [410] André Schleife, Erik W. Draeger, Yosuke Kanai, and Alfredo A. Correa. Plane-wave pseudopotential implementation of explicit integrators for time-dependent kohn-sham equations in large-scale simulations. *The Journal of Chemical Physics*, 137(22):22A546, 2012.
- [411] André Schleife, Erik W. Draeger, Victor M. Anisimov, Alfredo A. Correa, and Yosuke Kanai. Quantum dynamics simulation of electrons in materials on high-performance computers. *Computing in Science & Engineering*, 16(5):54–60, 2014.
- [412] Adam Bruner, Daniel LaMaster, and Kenneth Lopata. Accelerated broadband spectra using transition dipole decomposition and padé approximants. *J. Chem. Theo. Comput.*, 12(8):3741, 2016.
- [413] Matthias Kick, Ezra Alexander, Anton Beiersdorfer, and Troy Van Voorhis. Super-resolution techniques to simulate electronic spectra of large molecular systems. *Nature Communications*, 15(1):8001, sep 2024.
- [414] Joscha Hekele. *Implementation and Application of Density Functional Theory Methods for Electron and Nuclear Dynamics*. PhD thesis, University of Duisburg-Essen, 2021. <https://doi.org/10.17185/duetpublico/74983>.
- [415] Kushal Ramakrishna, Mani Lokamani, and Attila Cangi. Electrical conductivity of warm dense hydrogen from ohm's law and time-dependent density functional theory. *Electronic Structure*, 6(4):045008, nov 2024.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [416] Joscha Hekele, Yi Yao, Yosuke Kanai, Volker Blum, and Peter Kratzer. All-electron real-time and imaginary-time time-dependent density functional theory within a numeric atom-centered basis function framework. *J. Chem. Phys.*, 155:154801, 2021.
- [417] P. S. Bagus. Self-consistent-field wave functions for hole states of some ne-like and ar-like ions. *Phys. Rev.*, 139(A):A619, 1965.
- [418] Nicholas A. Besley, Andrew T. B. Gilbert, and Peter M. W. Gill. Self-consistent-field calculations of core excited states. *J. Chem. Phys.*, 130:124308, 2009.
- [419] Samuel J. Hall, Benedikt P. Klein, and Reinhard J. Maurer. Characterizing molecule-metal surface chemistry with ab initio simulation of x-ray absorption and photoemission spectra. *J. Phys. Chem. C*, 127:1870, 2023.
- [420] Delano P. Chong. Density-functional calculation of core-electron binding energies of c, n, o, and f. *J. Chem. Phys.*, 103:1842, 1995.
- [421] Marcus Annegarn, Juhan Matthias Kahk, and Johannes Lischner. Combining time-dependent density functional theory and the δ SCF approach for accurate core-electron spectra. *J. Chem. Theory Comput.*, 18:7620, 2022.
- [422] Felix Uhl and Volker Staemmler. An ab initio study of the o1s and mg1s, mg2s, mg2p core electron binding energies in bulk mgo. *J. Electron Spectros. Relat. Phenomena*, 233:90, 2019.
- [423] Nicholas A. Besley. Modeling of the spectroscopy of core electrons with density functional theory. *WIREs Comput. Mol. Sci.*, 11:e1527, 2021.
- [424] J. Matthias Kahk and Johannes Lischner. Predicting core electron binding energies in elements of the first transition series using the δ -self-consistent-field method. *Faraday Discuss.*, 236:373, 2022.
- [425] Francisco A. Delesma, Maxime Van den Bossche, Henrik Grönbeck, Patrizia Calaminici, Andreas M. Köster, and Lars G. M. Pettersson. A chemical view on x-ray photoelectron spectroscopy: the esca molecule and surface-to-bulk xps shifts. *ChemPhysChem*, 19:169, 2018.
- [426] Paul S. Bagus, Eugene Ilton, and Connie J. Nelin. Extracting chemical information from xps spectra: A perspective. *Catal. Lett.*, 148:1785, 2018.
- [427] Misa Nishino, Kouji Inagaki, Yoshitada Morikawa, Kazuya Yamamura, and Yuji Ohkubo. Identification of chemical species on plasma-treated polytetrafluoroethylene surface by ab-initio calculations of core-energy-level shift in x-ray photoelectron spectra. *Appl. Surf. Sci.*, 655:159369, 2024.
- [428] Weitao Yang and Paul W. Ayers. Foundation for the δ SCF approach in density functional theory, 2024.
- [429] Tom Ziegler, Michael Seth, Mykhaylo Krykunov, Jochen Autschbach, and Fan Wang. On the relation between time-dependent and variational density functional theory approaches for the determination of excitation energies and transition moments. *J. Chem. Phys.*, 130:154102, 2009.
- [430] John Cullen, Mykhaylo Krykunov, and Tom Ziegler. The formulation of a self-consistent constricted variational density functional theory for the description of excited states. *Chem. Phys.*, 391:11, 2011.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [431] Tom Ziegler, Mykhaylo Krykunov, and John Cullen. The application of constricted variational density functional theory to excitations involving electron transitions from occupied lone-pair orbitals to virtual π^* orbitals. *J. Chem. Theory Comput.*, 7:2485, 2011.
- [432] Tom Ziegler, Mykhaylo Krykunov, and John Cullen. The implementation of a self-consistent constricted variational density functional theory for the description of excited states. *J. Chem. Phys.*, 136:124107, 2012.
- [433] Andreas Görling. Density-functional theory beyond the hohenberg-kohn theorem. *Phys. Rev. A*, 59:3359, 1999.
- [434] Paul W. Ayers and Mel Levy. Time-independent (static) density-functional theories for pure excited states: Extensions and unification. *Phys. Rev. A*, 80:012508, 2009.
- [435] Benedikt P Klein, Samuel J Hall, and Reinhard J Maurer. The nuts and bolts of core-hole constrained ab initio simulation for k-shell x-ray photoemission and absorption spectra. *J. Phys.: Condens. Matter*, 33:154005, 2021.
- [436] Shayantan Chaudhuri, Samuel J. Hall, Benedikt P. Klein, Marc Walker, Andrew J. Logsdail, Julie V. Macpherson, and Reinhard J. Maurer. Coexistence of carbonyl and ether groups on oxygen-terminated (110)-oriented diamond surfaces. *Commun. Mater.*, 3(6), 2022.
- [437] J. Matthias Kahk and Johannes Lischner. Accurate absolute core-electron binding energies of molecules, solids, and surfaces from first-principles calculations. *Phys. Rev. Materials*, 3:100801, 2019.
- [438] J. Matthias Kahk, Georg S. Michelitsch, Reinhard J. Maurer, Karsten Reuter, and Johannes Lischner. Core electron binding energies in solids from periodic all-electron δ -self-consistent-field calculations. *J. Phys. Chem. Lett.*, 12:9353, 2021.
- [439] J. Matthias Kahk and Johannes Lischner. Combining the δ -self-consistent-field and gw methods for predicting core electron binding energies in periodic solids. *J. Chem. Theory Comput.*, 19:3276, 2023.
- [440] Thomas C. Taucher, Oliver T. Hofmann, and Egbert Zojer. Final-state simulations of core-level binding energies at metal-organic hybrid interfaces: Artifacts caused by spurious collective electrostatic effects. *ACS Omega*, 5:25868, 2020.
- [441] Reinhard J. Maurer and Karsten Reuter. Assessing computationally efficient isomerization dynamics: δ scf density-functional theory study of azobenzene molecular switching. *J. Chem. Phys.*, 135:224303, 2011.
- [442] Helene Lefebvre-Brion. Cecam-orsay documentation for molecular calculations - 1970, 2021.
- [443] Andrew T. B. Gilbert, Nicholas A. Besley, and Peter M. W. Gill. Self-consistent field calculations of excited states using the maximum overlap method (mom). *J. Phys. Chem. A*, 112:13164, 2008.
- [444] Samuel J. Hall, Dylan B. Morgan, and Reinhard J. Maurer. Archer2-ecse04-3: Relativistic all-electron orbital-constrained density functional theory to simulate x-ray photoemission and absorption spectroscopy. *Zenodo*, 2023.
- [445] Georg S. Michelitsch and Karsten Reuter. Efficient simulation of near-edge x-ray absorption fine structure (nexafs) in density-functional theory: Comparison of core-level constraining approaches. *J. Chem. Phys.*, 150:074104, 2019.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [446] Georg Siegmund Michelitsch. *Ab Initio simulation of core-level spectroscopies and correlated materials*. PhD thesis, Technische Universität München, 2019. <https://mediatum.ub.tum.de/doc/1472571/document.pdf>.
- [447] J. M. Foster and S. F. Boys. Canonical configurational interaction procedure. *Rev. Mod. Phys.*, 32:300, 1960.
- [448] Mariana Rossi, Michele Ceriotti, and David E. Manolopoulos. Nuclear quantum effects in h^+ and oh^- diffusion along confined water wires. *J. Phys. Chem. Lett.*, 7:3001, 2016.
- [449] G. Makov and M. C. Payne. Periodic boundary conditions in ab initio calculations. *Phys. Rev. B*, 51:4014, 1995.
- [450] S. Baroni, S. de Gironcoli, A. D. Corso, and P. Giannozzi. Phonons and related crystal properties from density-functional perturbation theory. *Rev. Mod. Phys.*, 73:515–562, 2001.
- [451] P. Giannozzi, S. de Gironcoli, P. Pavone, and S. Baroni. Ab initio calculation of phonon dispersions in semiconductors. *Phys. Rev. B*, 43:7231–7242, 1991.
- [452] G. D. Mahan. Modified sternheimer equation for polarizability. *Phys. Rev. A*, 22:1780–1785, 1980.
- [453] X. Gonze and C. Lee. Dynamical matrices, born effective charges, dielectric permittivity tensors, and interatomic force constants from density-functional perturbation theory. *Phys Rev B*, 55:10355–10368, 1997.
- [454] N. Raimbault, V. Athavale, and M. Rossi. Anharmonic effects in the low-frequency vibrational modes of aspirin and paracetamol crystals. *Phys. Rev. Mater.*, 3:053605, 2019.
- [455] A. Putrino, D. Sebastiani, and M. Parrinello. Generalized variational density functional perturbation theory. *J. Chem. Phys.*, 113:7102–7109, 2000.
- [456] F. Giustino, M. L. Cohen, and S. G. Louie. Electron-phonon interaction using wannier functions. *Phys. Rev. B*, 76:165108, 2007.
- [457] C. L. Box, W. G. Stark, and R. J. Maurer. Ab initio calculation of electron-phonon linewidths and molecular dynamics with electronic friction at metal surfaces with numeric atom-centred orbitals. *Electron. Struct.*, 5:035005, 2023.
- [458] H. Shang, C. Carbogno, P. Rinke, and M. Scheffler. Lattice dynamics calculations based on density-functional perturbation theory in real space. *Comput. Phys. Commun.*, 215:26–46, 2017.
- [459] H. Shang, N. Raimbault, P. Rinke, M. Scheffler, M. Rossi, and C. Carbogno. All-electron, real-space perturbation theory for homogeneous electric fields: theory, implementation, and application within DFT. *New J. Phys.*, 20:073040, 2018.
- [460] Connor L. Box, Andrew J. Logsdail, Volker Blum, Mariana Rossi, Christian Carbogno, and Reinhard J. Maurer. Archer2-ecse01-16 report achieving the sustainability and scalability of numeric-atomic-orbital-based linear response and electron-phonon functionality in fhi-aims, 2021.
- [461] Florian Knoop, Thomas A R Purcell, Matthias Scheffler, and Christian Carbogno. FHI-vibes: Ab initio vibrational simulations. *Journal of Open Source Software*, 5(56):2671, December 2020.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [462] R. M. Sternheimer. Electronic polarizabilities of ions from the hartree-fock wave functions. *Phys Rev*, 96:951–968, 1954.
- [463] M.A. Marques, M.J. Oliveira, and T. Burnus. Libxc: A library of exchange and correlation functionals for density functional theory. *Comput. Phys. Commun.*, 183:2272–2281, 2012.
- [464] H. Shang et al. Extreme-scale ab initio quantum raman spectra simulations on the leadership hpc system in china. In *Proceedings of the International Conference for High Performance Computing, Networking, Storage and Analysis*, 2021.
- [465] Honghui Shang, WanZhen Liang, Yunquan Zhang, and Jinlong Yang. Efficient parallel linear scaling method to get the response density matrix in all-electron real-space density-functional perturbation theory. *Computer Physics Communications*, 258:107613, 2021.
- [466] N. Raimbault, A. Grisafi, M. Ceriotti, and M. Rossi. Using gaussian process regression to simulate the vibrational raman spectra of molecular crystals. *New J. Phys.*, 21:105001, 2019.
- [467] Y. Zhang, R. J. Maurer, and B. Jiang. Symmetry-adapted high dimensional neural network representation of electronic friction tensor of adsorbates on metals. *J. Phys. Chem. C*, 124:186–195, 2020.
- [468] H. Shang and H. Wang. Anharmonic raman spectra simulation of crystals from deep neural networks. *AIP Adv.*, 11:035105, 2021.
- [469] A. M. Lewis, P. Lazzaroni, and M. Rossi. Predicting the electronic density response of condensed-phase systems to electric field perturbations. *J. Chem. Phys.*, 159:014103, 2023.
- [470] Paul Hodgkinson and Josep Sauri. *Nuclear Magnetic Resonance: Volume 50*. Royal Society of Chemistry, 11 2024.
- [471] A.G. Biternas, G.T.P. Charnock, and Ilya Kuprov. A standard format and a graphical user interface for spin system specification. *J. Magn. Reson.*, 240:124–131, 2014.
- [472] H.J. Hogben, M. Krzystyniak, G.T.P. Charnock, P.J. Hore, and Ilya Kuprov. Spinach - a software library for simulation of spin dynamics in large spin systems. *J. Magn. Reson.*, 208(2):179–194, 2011.
- [473] M. Bühl, M. Kaupp, O.L. Malkina, and V.G. Malkin. The dft route to nmr chemical shifts. *J. Comput. Chem.*, 20(1):91–105, 1999.
- [474] R. Ditchfield. Molecular orbital theory of magnetic shielding and magnetic susceptibility. *The Journal of Chemical Physics*, 56(11):5688–5691, 06 1972.
- [475] Caspar Jonas Schattenberg and Martin Kaupp. Extended benchmark set of main-group nuclear shielding constants and nmr chemical shifts and its use to evaluate modern dft methods. *Journal of Chemical Theory and Computation*, 17(12):7602–7621, 2021. PMID: 34797677.
- [476] Susi Lehtola, Maria Dimitrova, Heike Fliegl, and Dage Sundholm. Benchmarking magnetizabilities with recent density functionals. *Journal of Chemical Theory and Computation*, 17(3):1457–1468, 2021. PMID: 33599491.
- [477] J. San Fabián, J. García de la Vega, R. Suardiáz, M. Fernández-Oliva, C. Pérez, R. Crespo-Otero, and R. Contreras. Computational nmr coupling constants: shifting and scaling factors for evaluating $^1j_c/h$. *Magnetic Resonance in Chemistry*, 51(12):775–787.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [478] Frank Jensen. Basis set convergence of nuclear magnetic shielding constants calculated by density functional methods. *Journal of Chemical Theory and Computation*, 4(5):719–727, 2008. PMID: 26621087.
- [479] <https://fhi-aims-club.gitlab.io/tutorials/tutorials-overview/>.
- [480] Chemical Book. https://www.chemicalbook.com/SpectrumEN_64-17-5_1HNMR.htm. [Online reference data (subscription required); last accessed: 2025/01/19].
- [481] Giovanni Bussi, Davide Donadio, and Michele Parrinello. Canonical sampling through velocity rescaling. *The Journal of Chemical Physics*, 126(1):014101, 2007.
- [482] Iuliia Mandzhieva, Franziska Theiss, Xingtao He, Adam Ortmeier, Anuja Koirala, Stephen J. McBride, Stephen J. DeVience, Matthew S. Rosen, Volker Blum, and Thomas Theis. Zero-field nmr and millitesla-slic spectra for > 200 molecules from density functional theory and spin dynamics. Submitted for publication (2025).
- [483] Francesco Mauri and Steven G. Louie. Magnetic susceptibility of insulators from first principles. *Phys. Rev. Lett.*, 76:4246–4249, May 1996.
- [484] Francesco Mauri, Bernd G. Pfrommer, and Steven G. Louie. Ab initio theory of nmr chemical shifts in solids and liquids. *Phys. Rev. Lett.*, 77:5300–5303, Dec 1996.
- [485] Jochen Autschbach. Analyzing molecular properties calculated with two-component relativistic methods using spin-free natural bond orbitals: Nmr spin-spin coupling constants. *The Journal of Chemical Physics*, 127(12):124106, 09 2007.
- [486] Jochen Autschbach. Analyzing nmr shielding tensors calculated with two-component relativistic methods using spin-free localized molecular orbitals. *The Journal of Chemical Physics*, 128(16):164112, 04 2008.
- [487] Jochen Autschbach. Magnitude of finite-nucleus-size effects in relativistic density functional computations of indirect nmr nuclear spin-spin coupling constants. *ChemPhysChem*, 10(13):2274–2283, 2009.
- [488] A. Bansil, Hsin Lin, and Tanmoy Das. Colloquium: Topological band theory. *Reviews of Modern Physics*, 88(2), June 2016.
- [489] Liang Fu and C. L. Kane. Time reversal polarization and a z_2 adiabatic spin pump. *Physical Review B*, 74(19), November 2006.
- [490] R. Resta. Theory of the electric polarization in crystals. *Ferroelectrics*, 136(1):51–55, November 1992.
- [491] R. D. King-Smith and David Vanderbilt. Theory of polarization of crystalline solids. *Physical Review B*, 47(3):1651–1654, January 1993.
- [492] D. J. Thouless. Quantization of particle transport. *Physical Review B*, 27(10):6083–6087, May 1983.
- [493] E.I. Blount. *Formalisms of Band Theory*, page 305–373. Elsevier, 1962.
- [494] Nicola Marzari and David Vanderbilt. Maximally localized generalized wannier functions for composite energy bands. *Physical review B*, 56(20):12847, 1997.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [495] Nicola Marzari, Arash A. Mostofi, Jonathan R. Yates, Ivo Souza, and David Vanderbilt. Maximally localized wannier functions: Theory and applications. *Reviews of Modern Physics*, 84(4):1419–1475, October 2012.
- [496] David Vanderbilt. *Berry Phases in Electronic Structure Theory: Electric Polarization, Orbital Magnetization and Topological Insulators*. Cambridge University Press, October 2018.
- [497] Nicola A. Spaldin. A beginner's guide to the modern theory of polarization. *Journal of Solid State Chemistry*, 195:2–10, November 2012.
- [498] Alexey A. Soluyanov and David Vanderbilt. Computing topological invariants without inversion symmetry. *Physical Review B*, 83(23):235401, June 2011.
- [499] Rui Yu, Xiao Liang Qi, Andrei Bernevig, Zhong Fang, and Xi Dai. Equivalent expression of \mathbb{Z}_2 topological invariant for band insulators using the non-abelian berry connection. *Physical Review B*, 84(7):075119, August 2011.
- [500] Chen Si, Junwei Liu, Yong Xu, Jian Wu, Bing-Lin Gu, and Wenhui Duan. Functionalized germanene as a prototype of large-gap two-dimensional topological insulators. *Physical Review B*, 89(11):115429, 2014.
- [501] L. Matthes, S. Küfner, J. Furthmüller, and F. Bechstedt. Intrinsic spin hall conductivity in one-, two-, and three-dimensional trivial and topological systems. *Physical Review B*, 94(8):085410, August 2016.
- [502] Liang Fu, C. L. Kane, and E. J. Mele. Topological insulators in three dimensions. *Physical Review Letters*, 98(10):106803, 2007.
- [503] Guohua Cao, Runhai Ouyang, Luca M. Ghiringhelli, Matthias Scheffler, Huijun Liu, Christian Carbogno, and Zhenyu Zhang. Artificial intelligence for high-throughput discovery of topological insulators: The example of alloyed tetradymites. *Physical Review Materials*, 4(3):034204, March 2020.
- [504] Yair Litman, Venkat Kapil, Yotam M. Y. Feldman, Davide Tisi, Tomislav Begušić, Karen Fidanyan, Guillaume Fraux, Jacob Higer, Matthias Kellner, Tao E. Li, Eszter S. Pócs, Elia Stocco, George Trenins, Barak Hirshberg, Mariana Rossi, and Michele Ceriotti. i-pi 3.0: A flexible and efficient framework for advanced atomistic simulations. *The Journal of Chemical Physics*, 161(6):062504, Aug 2024.
- [505] Elia Stocco, Christian Carbogno, and Mariana Rossi. Electric-Field Driven Nuclear Dynamics of Liquids and Solids from a Multi-Valued Machine-Learned Dipolar Model. *arXiv*, 2502.02413, 2025.
- [506] Linfeng Zhang, Mohan Chen, Xifan Wu, Han Wang, Weinan E, and Roberto Car. Deep neural network for the dielectric response of insulators. *Physical Review B*, 102(4):041121, 2020.
- [507] Koji Shimizu, Ryuji Otsuka, Masahiro Hara, Emi Minamitani, and Satoshi Watanabe. Prediction of Born effective charges using neural network to study ion migration under electric fields: applications to crystalline and amorphous Li_3PO_4 . *Science and Technology of Advanced Materials: Methods*, 3(1):2253135, 2023.
- [508] Nicolas Bergmann, Nicéphore Bonnet, Nicola Marzari, Karsten Reuter, and Nicolas G. Hörmann. Machine Learning the Energetics of Electrified Solid-Liquid Interfaces. *Physical Review Letters*, 135(14):146201, 2025.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [509] Giovanni Pizzi, Valerio Vitale, Ryotaro Arita, Stefan Blügel, Frank Freimuth, Guillaume Granton, Marco Gibertini, Dominik Gresch, Charles Johnson, Takashi Koretsune, Julen Ibaez-Azpiroz, Hyungjun Lee, Jae-Mo Lihm, Daniel Marchand, Antimo Marrazzo, Yuriy Mokrousov, Jamal I. Mustafa, Yoshiro Nohara, Yusuke Nomura, Lorenzo Paulatto, Samuel Ponc, Thomas Ponweiser, Junfeng Qiao, Florian Thäle, Stepan S. Tsirkin, Ma Igorzata Wierzbowska, Nicola Marzari, David Vanderbilt, Ivo Souza, Arash A. Mostofi, and Jonathan R. Yates. Wannier90 as a community code: new features and applications. *Journal of Physics: Condensed Matter*, 32(16):165902, 2020.
- [510] S. Ponce, E. R. Margine, C. Verdi, and F. Giustino. Epw: Electron-phonon coupling, transport and superconducting properties using maximally localized wannier functions. *Computer Physics Communications*, 209:116–133, December 2016.
- [511] Jin-Jian Zhou, Jinsoo Park, I-Te Lu, Ivan Maliyov, Xiao Tong, and Marco Bernardi. Perturbo: A software package for ab initio electron-phonon interactions, charge transport and ultrafast dynamics. *Computer Physics Communications*, 264:107970, 2021.
- [512] M. Born and K. Huang. *Dynamical Theory of Crystal Lattices*. Clarendon Press, Oxford, 1954.
- [513] Günther Leibfried and Wolfgang Ludwig. Theory of anharmonic effects in crystals. In *Solid state physics*, volume 12, pages 275–444. Elsevier, 1961.
- [514] Paul G Klemens. Thermal conductivity and lattice vibrational modes. In *Solid state physics*, volume 7, pages 1–98. Elsevier, 1958.
- [515] M. Dove. *Introduction to Lattice Dynamics*. Cambridge University Press, 1993.
- [516] B. Fultz. Progress in materials science. *Prog. Mater. Sci.*, 55:247, 2010.
- [517] Lorenzo Monacelli, Raffaello Bianco, Marco Cherubini, Matteo Calandra, Ion Errea, and Francesco Mauri. The stochastic self-consistent harmonic approximation: calculating vibrational properties of materials with full quantum and anharmonic effects. *Journal of Physics: Condensed Matter*, 33(36):363001, 2021.
- [518] Fredrik Eriksson, Erik Fransson, and Paul Erhart. The hiphive package for the extraction of high-order force constants by machine learning. *Advanced Theory and Simulations*, 2(5):1800184, May 2019.
- [519] Olle Hellman, Igor A Abrikosov, and Sergei I Simak. Lattice dynamics of anharmonic solids from first principles. *Physical Review B—Condensed Matter and Materials Physics*, 84(18):180301, 2011.
- [520] Ambroise van Roekeghem, Jesús Carrete, and Natalio Mingo. Quantum self-consistent ab-initio lattice dynamics. *Computer Physics Communications*, 263:107945, 2021.
- [521] T Tadano, Y Gohda, and S Tsuneyuki. Anharmonic force constants extracted from first-principles molecular dynamics: applications to heat transfer simulations. *Journal of Physics: Condensed Matter*, 26(22):225402, 2014.
- [522] Atsushi Togo, Laurent Chaput, Terumasa Tadano, and Isao Tanaka. Implementation strategies in phonopy and phono3py. *Journal of Physics: Condensed Matter*, 35(35):353001, 2023.
- [523] Atsushi Togo, Laurent Chaput, and Isao Tanaka. Distributions of phonon lifetimes in brillouin zones. *Physical Review-Section B-Condensed Matter*, 91(9):094306, March 2015.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [524] Florian Knoop, Nina Shulumba, Alois Castellano, J P Alvarinhas Batista, Roberta Farris, Matthieu J Verstraete, Matthew Heine, David Broido, Dennis S Kim, Johan Klarbring, Igor A Abrikosov, Sergei I Simak, and Olle Hellman. Tdep: Temperature dependent effective potentials. *Journal of Open Source Software*, 9(94):6150, 2024.
- [525] Florian Knoop, Thomas A R Purcell, Matthias Scheffler, and Christian Carbogno. Anharmonicity measure for materials. *Phys. Rev. Mater.*, 4(8):083809, August 2020.
- [526] Anubhav Jain, Shyue Ping Ong, Wei Chen, Bharat Medasani, Xiaohui Qu, Michael Kocher, Miriam Brafman, Guido Petretto, Gian-Marco Rignanese, Geoffroy Hautier, Daniel Gunter, and Kristin A. Persson. Fireworks: a dynamic workflow system designed for high-throughput applications. *Concurrency and Computation: Practice and Experience*, 27(17):5037–5059, 2015. CPE-14-0307.R2.
- [527] K Esfarjani and H T Stokes. Method to extract anharmonic force constants from first principles calculations. *Physical Review B*, 77:144112, 2008.
- [528] K Parlinski, ZQ Li, and Y Kawazoe. First-principles determination of the soft mode in cubic ZrO_2 . *Physical Review Letters*, 78(21):4063, 1997.
- [529] Slawomir Biernacki and Matthias Scheffler. Negative thermal expansion of diamond and zinc-blende semiconductors. *Physical review letters*, 63(3):290, 1989.
- [530] S. Rasti and J. Meyer. Thermodynamic properties of complex molecular systems. *J. Chem. Phys.*, 150:234504, 2019.
- [531] S. Rasti, E. O. Jonsson, H. Jonsson, and J. Meyer. Kinetic mechanisms in molecular systems: An advanced perspective. *J. Phys. Chem. Lett.*, 13:11831, 2022.
- [532] D West and SK Estreicher. First-Principles Calculations of Vibrational Lifetimes and Decay Channels: Hydrogen-Related Modes in Si. *Physical review letters*, 96(11):115504, 2006.
- [533] Florian Knoop, Thomas A. R. Purcell, Matthias Scheffler, and Christian Carbogno. Anharmonicity in thermal insulators: An analysis from first principles. *Physical Review Letters*, 130(23):236301, 2023.
- [534] Thomas AR Purcell, Matthias Scheffler, Luca M Ghiringhelli, and Christian Carbogno. Accelerating materials-space exploration for thermal insulators by mapping materials properties via artificial intelligence. *npj computational materials*, 9(1):112, 2023.
- [535] Jingkai Quan, Christian Carbogno, and Matthias Scheffler. Carrier mobility of strongly anharmonic materials from first principles. *Physical Review B*, 110(23):235202, 2024.
- [536] Kisung Kang, Thomas AR Purcell, Christian Carbogno, and Matthias Scheffler. Accelerating the training and improving the reliability of machine-learned interatomic potentials for strongly anharmonic materials through active learning. *Physical Review Materials*, 9(6):063801, 2025.
- [537] Zezhu Zeng, Xingchen Shen, Ruihuan Cheng, Olivier Perez, Niuchang Ouyang, Zheyong Fan, Pierrick Lemoine, Bernard Raveau, Emmanuel Guilmeau, and Yue Chen. Pushing thermal conductivity to its lower limit in crystals with simple structures. *Nature Communications*, 15(1):3007, 2024.
- [538] Yixin Xu and Yanguang Zhou. Metric for quantifying elastic and inelastic thermal transport at interfaces. *Physical Review B*, 110(11):115305, 2024.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [539] Owain T Beynon, Alun Owens, Christian Carbogno, and Andrew J Logsdail. Evaluating the role of anharmonic vibrations in zeolite β materials. *The Journal of Physical Chemistry C*, 127(32):16030–16040, 2023.
- [540] Xavier Gonze and Changyol Lee. Dynamical matrices, born effective charges, dielectric permittivity tensors, and interatomic force constants from density-functional perturbation theory. *Physical Review B*, 55(16):10355, 1997.
- [541] Marcel F. Langer, Florian Knoop, Christian Carbogno, Matthias Scheffler, and Matthias Rupp. Heat flux for semilocal machine-learning potentials. *Physical Review B*, 108(10):L100302, 2023.
- [542] Roberto Car and M Parrinello. Unified approach for molecular dynamics and density-functional theory. *Physical Review Letters*, 55(22):2471 – 2474, 1985.
- [543] Mariana Rossi, Volker Blum, Peter Kupser, Gert von Helden, Frauke Bierau, Kevin Pagel, Gerard Meijer, and Matthias Scheffler. Secondary structure of ac-ala-lysh+ polyalanine peptides ($n = 5, 10, 15$) in vacuo: Helical or not? *The Journal of Physical Chemistry Letters*, 1(24):3465–3470, 2010.
- [544] Elizabeth C. Beret, Luca M. Ghiringhelli, and Matthias Scheffler. Free gold clusters: beyond the static, monostructure description. *Faraday Discuss.*, 152:153–167, 2011.
- [545] Yuanyuan Zhou, Matthias Scheffler, and Luca M Ghiringhelli. Determining surface phase diagrams including anharmonic effects. *Physical Review B*, 100(17):174106, 2019.
- [546] M Born and R Oppenheimer. On the quantum theory of molecules. *Annalen der Physik*, 84:457 – 484, 1927.
- [547] L Verlet. Computer "Experiments" on classical fluids. *Physical Review*, 159(1):98, 1967.
- [548] Daan Frenkel and Berend Smit. Understanding molecular simulation: from algorithms to applications. page 638, 2002.
- [549] H C Andersen. Molecular dynamics simulations at constant pressure and/or temperature. *The Journal of Chemical Physics*, 72:2384, 1980.
- [550] W G Hoover. Canonical dynamics: Equilibrium phase-space distributions. *Physical Review A*, 31(3):1695 – 1697, 03 1985.
- [551] Michele Ceriotti, Giovanni Bussi, and Michele Parrinello. Langevin Equation with Colored Noise for Constant-Temperature Molecular Dynamics Simulations. *Physical Review Letters*, 102(2):020601, 2009.
- [552] Karsten Reuter and Matthias Scheffler. Composition and structure of the $\text{RuO}_2(110)$ surface in an O_2 and CO environment: Implications for the catalytic formation of CO_2 . *Phys. Rev. B*, 68:045407, Jul 2003.
- [553] Qiliang Yan and Juan J de Pablo. Hyper-parallel tempering monte carlo: Application to the lennard-jones fluid and the restricted primitive model. *The Journal of chemical physics*, 111(21):9509–9516, 1999.
- [554] Roland Faller, Qiliang Yan, and Juan J de Pablo. Multicanonical parallel tempering. *The Journal of chemical physics*, 116(13):5419–5423, 2002.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [555] Enzo Marinari and Giorgio Parisi. Simulated tempering: a new monte carlo scheme. *Europhysics letters*, 19(6):451, 1992.
- [556] Yuji Sugita and Yuko Okamoto. Replica-exchange molecular dynamics method for protein folding. *Chemical physics letters*, 314(1-2):141–151, 1999.
- [557] Yuanyuan Zhou, Chunye Zhu, Matthias Scheffler, and Luca M Ghiringhelli. Ab initio approach for thermodynamic surface phases with full consideration of anharmonic effects: The example of hydrogen at si (100). *Physical Review Letters*, 128(24):246101, 2022.
- [558] Michael R Shirts and John D Chodera. Statistically optimal analysis of samples from multiple equilibrium states. *The Journal of chemical physics*, 129(12), 2008.
- [559] Che Ting Chan, Klaus Peter Bohnen, and KM Ho. Accelerating the convergence of force calculations in electronic-structure computations. *Physical Review B*, 47(8):4771, 1993.
- [560] M. Parrinello and A. Rahman. Study of an F center in molten KCl. *The Journal of Chemical Physics*, 80(2):860–867, Jan 1984.
- [561] Venkat Kapil, Mariana Rossi, Ondrej Marsalek, Riccardo Petraglia, Yair Litman, Thomas Spura, Bingqing Cheng, Alice Cuzzocrea, Robert H Meißner, David M Wilkins, et al. i-PI 2.0: A universal force engine for advanced molecular simulations. *Computer Physics Communications*, 236:214–223, 2019.
- [562] Simon P. Webb, Tzvetelin Iordanov, and Sharon Hammes-Schiffer. Multiconfigurational nuclear-electronic orbital approach: Incorporation of nuclear quantum effects in electronic structure calculations. *J. Chem. Phys.*, 117(9):4106–4118, 2002.
- [563] Fabijan Pavošević, Tanner Culpitt, and Sharon Hammes-Schiffer. Multicomponent quantum chemistry: Integrating electronic and nuclear quantum effects via the nuclear-electronic orbital method. *Chem. Rev.*, 120(9):4222–4253, May 2020.
- [564] Xi Xu and Yang Yang. Full-quantum descriptions of molecular systems from constrained nuclear-electronic orbital density functional theory. *J. Chem. Phys.*, 153(7):074106, Aug 2020.
- [565] Michael V. Pak, Arindam Chakraborty, and Sharon Hammes-Schiffer. Density functional theory treatment of electron correlation in the nuclear-electronic orbital approach. *J. Phys. Chem. A*, 111(20):4522–4526, May 2007.
- [566] Arindam Chakraborty, Michael V. Pak, and Sharon Hammes-Schiffer. Development of electron-proton density functionals for multicomponent density functional theory. *Phys. Rev. Lett.*, 101(15):153001, Oct 2008.
- [567] T. Kreibich and E. K. U. Gross. Multicomponent Density-Functional Theory for Electrons and Nuclei. *Phys. Rev. Lett.*, 86(14):2984–2987, April 2001.
- [568] Ondrej Marsalek and Thomas E. Markland. Ab initio molecular dynamics with nuclear quantum effects at classical cost: Ring polymer contraction for density functional theory. *The Journal of Chemical Physics*, 144(5):054112, Feb 2016.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [569] V. Kapil, J. VandeVondele, and M. Ceriotti. Accurate molecular dynamics and nuclear quantum effects at low cost by multiple steps in real and imaginary time: Using density functional theory to accelerate wavefunction methods. *The Journal of Chemical Physics*, 144(5):054111, Feb 2016.
- [570] Yair Litman, Davide Donadio, Michele Ceriotti, and Mariana Rossi. Decisive role of nuclear quantum effects on surface mediated water dissociation at finite temperature. *The Journal of Chemical Physics*, 148:102320, 2018.
- [571] Igor Poltavsky, Robert A. DiStasio Jr., and Alexandre Tkatchenko. Perturbed path integrals in imaginary time: Efficiently modeling nuclear quantum effects in molecules and materials. *The Journal of Chemical Physics*, 148(10):102325, Dec 2017.
- [572] Stuart C. Althorpe. Path integral simulations of condensed-phase vibrational spectroscopy. *Annual Review of Physical Chemistry*, 75:397–420, 2024.
- [573] Benjamin J. Sutherland, William H. D. Moore, and David E. Manolopoulos. Nuclear quantum effects in thermal conductivity from centroid molecular dynamics. *The Journal of Chemical Physics*, 154(17):174104, May 2021.
- [574] William H. Miller. Semiclassical limit of quantum mechanical transition state theory for nonseparable systems. *The Journal of Chemical Physics*, 62(5):1899–1906, Mar 1975.
- [575] Jeremy O. Richardson and Stuart C. Althorpe. Ring-polymer molecular dynamics rate-theory in the deep-tunneling regime: Connection with semiclassical instanton theory. *The Journal of Chemical Physics*, 131(21):214106, Dec 2009.
- [576] Jeremy O. Richardson. Ring-polymer instanton theory. *International Reviews in Physical Chemistry*, 37(2):171–216, 2018.
- [577] Jianhang Xu, Ruiyi Zhou, and Zhenhong Liu. Nuclear quantum effects in molecular dynamics simulations. *Nature Reviews Chemistry*, 6(4):164–182, 2022.
- [578] Yang Yang, Kurt R. Brorsen, Tanner Culpitt, Michael V. Pak, and Sharon Hammes-Schiffer. Development of a practical multicomponent density functional for electron-proton correlation to produce accurate proton densities. *J. Chem. Phys.*, 147(11):114113, Sep 2017.
- [579] Kurt R. Brorsen, Yang Yang, and Sharon Hammes-Schiffer. Multicomponent density functional theory: Impact of nuclear quantum effects on proton affinities and geometries. *J. Phys. Chem. Lett.*, 8(15):3488–3493, Aug 2017.
- [580] Qi Yu, Fabijan Pavošević, and Sharon Hammes-Schiffer. Development of nuclear basis sets for multicomponent quantum chemistry methods. *J. Chem. Phys.*, 152(24):244123, Jun 2020.
- [581] Jianhang Xu, Ruiyi Zhou, Tao E. Li, Sharon Hammes-Schiffer, and Yosuke Kanai. Lagrangian formulation of nuclear-electronic orbital ehrenfest dynamics with real-time TDDFT for extended periodic systems, 2024.
- [582] Xi Xu and Yang Yang. Constrained nuclear-electronic orbital density functional theory: Energy surfaces with nuclear quantum effects. *The Journal of Chemical Physics*, 152(8), 2020.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [583] Songrui Liu, Jianhang Xu, and Yosuke Kanai. Constrained nuclear–electronic orbital method for periodic density functional theory: Application to h₂ chemisorption on si (001) surfaces. *The Journal of Chemical Physics*, 163(8), 2025.
- [584] Xi Xu, Zehua Chen, and Yang Yang. Molecular dynamics with constrained nuclear electronic orbital density functional theory: Accurate vibrational spectra from efficient incorporation of nuclear quantum effects. *J. Am. Chem. Soc.*, 144(9):4039–4046, 2022.
- [585] Rafael Ramírez and Telesforo López-Ciudad. Phase-space formulation of thermodynamic and dynamical properties of quantum particles. *Phys. Rev. Lett.*, 83:4456–4459, Nov 1999.
- [586] Karen Fidanyan, Ikutaro Hamada, and Mariana Rossi. Quantum nuclei at weakly bonded interfaces: The case of cyclohexane on Rh(111). *Advanced Theory and Simulations*, 4(4):2000241, 2021.
- [587] Yair Litman and Mariana Rossi. Multidimensional hydrogen tunneling in supported molecular switches: The role of surface interactions. *Physical Review Letters*, 125:216001, Nov 2020.
- [588] Karen Fidanyan, Guoyuan Liu, and Mariana Rossi. Ab initio study of water dissociation on a charged Pd(111) surface. *The Journal of Chemical Physics*, 158(9):094707, 03 2023.
- [589] Mariana Rossi, Piero Gasparotto, and Michele Ceriotti. Anharmonic and quantum fluctuations in molecular crystals: A first-principles study of the stability of paracetamol. *Physical Review Letters*, 117:115702, Sep 2016.
- [590] Venkat Kapil and Edgar A. Engel. A complete description of thermodynamic stabilities of molecular crystals. *Proceedings of the National Academy of Sciences*, 119(6):e2111769119, 2022.
- [591] David P. Jelenfi, Anita Schneiker, Attila Tajti, Gabor Magyarfalvi, and Gyorgy Tarczay. Polyaromatic hydrocarbons with an imperfect aromatic system as catalysts of interstellar h₂ formation. *Molecular Physics*, 121(11-12):e2142168, 2023.
- [592] Yair Litman, Jeremy O. Richardson, Takashi Kumagai, and Mariana Rossi. Elucidating the nuclear quantum dynamics of intramolecular double hydrogen transfer in porphycene. *Journal of the American Chemical Society*, 141(6):2526–2534, 2019.
- [593] Yair Litman, Jörg Behler, and Mariana Rossi. Temperature dependence of the vibrational spectrum of porphycene: A qualitative failure of classical-nuclei molecular dynamics. *Faraday Discussions*, 221:526–546, 2020.
- [594] Y. Litman, E. S. Pócs, C. L. Box, R. Martinazzo, R. J. Maurer, and M. Rossi. Dissipative tunneling rates through the incorporation of first-principles electronic friction in instanton rate theory. ii. benchmarks and applications. *The Journal of Chemical Physics*, 156(19):194107, 05 2022.
- [595] Oliver Bridge, Paolo Lazzaroni, Rocco Martinazzo, Mariana Rossi, Stuart C. Althorpe, and Yair Litman. Quantum rates in dissipative systems with spatially varying friction. *The Journal of Chemical Physics*, 161(2):024110, 07 2024.
- [596] Mariana Rossi, Wei Fang, and Angelos Michaelides. Stability of complex biomolecular structures: van der Waals, hydrogen bond cooperativity, and nuclear quantum effects. *The Journal of Physical Chemistry Letters*, 6(21):4233–4238, 2015.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [597] Gregory Fonseca, Igor Poltavsky, Valentin Vassilev-Galindo, and Alexandre Tkatchenko. Improving molecular force fields across configurational space by combining supervised and unsupervised machine learning. *The Journal of Chemical Physics*, 154(12):124102, 03 2021.
- [598] Matheus Jacobs, Karen Fidanyan, Mariana Rossi, and Caterina Cocchi. Impact of nuclear effects on the ultrafast dynamics of an organic/inorganic mixed-dimensional interface. *Electronic Structure*, 6(2):025006, May 2024.
- [599] Ruji Zhao, Peiwei You, and Sheng Meng. Ring polymer molecular dynamics with electronic transitions. *Physical Review Letters*, 130:166401, Apr 2023.
- [600] O. Bünermann, H. Jiang, Y. Dorenkamp, A. Kandratsenka, S. M. Janke, D. J. Auerbach, and A. M. Wodtke. Electron-hole pair excitation determines the mechanism of hydrogen atom adsorption. *Science*, 350:1346–1349, 2015.
- [601] M. Head-Gordon and J. C. Tully. Molecular dynamics with electronic frictions. *Journal of Chemical Physics*, 103:10137–10145, 1995.
- [602] R. J. Maurer, M. Askerka, V. S. Batista, and J. C. Tully. Ab initio tensorial electronic friction for molecules on metal surfaces: Nonadiabatic vibrational relaxation. *Physical Review B*, 94:115432, 2016.
- [603] R. J. Maurer, B. Jiang, H. Guo, and J. C. Tully. Mode specific electronic friction in dissociative chemisorption on metal surfaces: H₂ on ag (111). *Physical Review Letters*, 118:256001, 2017.
- [604] C. L. Box, Y. Zhang, R. Yin, B. Jiang, and R. J. Maurer. Determining the effect of hot electron dissipation on molecular scattering experiments at metal surfaces. *JACS Au*, 1:164–173, 2020.
- [605] M. Sachs, W. G. Stark, R. J. Maurer, and C. Ortner. Equivariant representation of configuration-dependent friction tensors in langevin heatbaths. *arXiv preprint arXiv:2407.13935*, 2024.
- [606] M. Head-Gordon and J. C. Tully. Molecular-orbital calculations of the lifetimes of the vibrational modes of co on cu (100). *Physical Review B*, 46:1853, 1992.
- [607] Y. Litman, E. S. Pósz, C. L. Box, R. Martinazzo, R. J. Maurer, and M. Rossi. Dissipative tunneling rates through the incorporation of first-principles electronic friction in instanton rate theory. i. theory. *Journal of Chemical Physics*, 156, 2022.
- [608] Y. Litman, V. Kapil, Y. M. Y. Feldman, D. Tisi, T. Begušić, K. Fidanyan, G. Fraux, J. Higer, M. Kellner, T. E. Li, E. S. Pósz, E. Stocco, G. Trenins, B. Hirshberg, M. Rossi, and M. Ceriotti. i-pi 3.0: a flexible, efficient framework for advanced atomistic simulations. *arXiv preprint arXiv:2405.15224*, 2024.
- [609] Feliciano Giustino, Steven G. Louie, and Marvin L. Cohen. Electron-phonon renormalization of the direct band gap of diamond. *Physical Review Letters*, 105(26):265501, 2010.
- [610] E. Cannuccia and A. Marini. Zero point motion effect on the electronic properties of diamond, trans-polyacetylene and polyethylene. *The European Physical Journal B*, 85(9):320, 2012.
- [611] S Ponce, G Antonius, P Boulanger, E Cannuccia, A Marini, M Cote, and X Gonze. Verification of first-principles codes: Comparison of total energies, phonon frequencies, electron-phonon coupling and zero-point motion correction to the gap between abinit and qe/yambo. *Computational Materials Science*, 83:341–348, 2014.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [612] Marios Zacharias and Feliciano Giustino. Theory of the special displacement method for electronic structure calculations at finite temperature. *Physical Review Research*, 2(1):013357, 2020.
- [613] Jesse Noffsinger, Emmanouil Kioupakis, Chris G. Van de Walle, Steven G. Louie, and Marvin L. Cohen. Phonon-assisted optical absorption in silicon from first principles. *Physical Review Letters*, 108(16):167402, 2012.
- [614] Marios Zacharias, Christopher E. Patrick, and Feliciano Giustino. Stochastic approach to phonon-assisted optical absorption. *Physical Review Letters*, 115(17):177401, 2015.
- [615] Jamal I. Mustafa, Marco Bernardi, Jeffrey B. Neaton, and Steven G. Louie. Ab initio electronic relaxation times and transport in noble metals. *Physical Review B*, 94(15):155105, 2016.
- [616] Samuel Ponce, Wenbin Li, Sven Reichardt, and Feliciano Giustino. First-principles calculations of charge carrier mobility and conductivity in bulk semiconductors and two-dimensional materials. *Reports on Progress in Physics*, 83(3):036501, 2020.
- [617] Marios Zacharias, Matthias Scheffler, and Christian Carbogno. Fully anharmonic nonperturbative theory of vibronically renormalized electronic band structures. *Physical Review B*, 102(4):045126, July 2020.
- [618] Jean Paul Nery and Francesco Mauri. Nonperturbative green's function method to determine the electronic spectral function due to electron-phonon interactions: Application to a graphene model from weak to strong coupling. *Physical Review B*, 105(24):245120, 2022.
- [619] Nikita Rybin. *Band structure renormalization at finite temperatures from first principles*. PhD thesis, Humboldt-Universität zu Berlin, 2023. <https://doi.org/10.18452/27063>.
- [620] Jingkai Quan, Nikita Rybin, Matthias Scheffler, and Christian Carbogno. Efficient band structure unfolding with atom-centered orbitals: General theory and application. *Phys. Rev. B*, 113:085112, Feb 2026.
- [621] Timothy B Boykin and Gerhard Klimeck. Practical application of zone-folding concepts in tight-binding calculations. *Physical Review B*, 71(11):115215, March 2005.
- [622] Wei Ku, Tom Berlijn, and Chi-Cheng Lee. Unfolding first-principles band structures. *Physical Review Letters*, 104(21):216401, May 2010.
- [623] Voicu Popescu and Alex Zunger. Extracting e versus k-effective band structure from supercell calculations on alloys and impurities. *Physical Review B*, 85(8):085201, 2012.
- [624] P B Allen, T Berlijn, D A Casavant, and J M Soler. Recovering hidden bloch character: Unfolding electrons, phonons, and slabs. *Physical Review B*, 87(8):085322, February 2013.
- [625] Xie Zhang, Jun Kang, and Su-Huai Wei. Profiling electronic and phononic band structures of semiconductors at finite temperatures: Methods and applications. *Chinese Physics Letters*, 41(2):026301, 2024.
- [626] G. Antonius, S. Poncé, P. Boulanger, M. Côté, and X. Gonze. Many-body effects on the zero-point renormalization of the band structure. *Physical Review Letters*, 112(21):215501, May 2014.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [627] Ferenc Karsai, Manuel Engel, Espen Flage-Larsen, and Georg Kresse. Electron-phonon coupling in semiconductors within the gw approximation. *New Journal of Physics*, 20(12):123008, December 2018.
- [628] Jörg Behler and Gábor Csányi. Machine learning potentials for extended systems: a perspective. *The European Physical Journal B*, 94(7):142, 2021.
- [629] Liwei Zhang, Berk Onat, Geneviève Dusson, Adam McSloy, G. Anand, Reinhard J. Maurer, Christoph Ortner, and James R. Kermode. Equivariant analytical mapping of first principles hamiltonians to accurate and transferable materials models. *npj Computational Materials*, 8(1):158, 2022.
- [630] Michele Lazzeri and Francesco Mauri. Nonadiabatic kohn anomaly in a doped graphene monolayer. *Physical Review Letters*, 97(26):266407, 2006.
- [631] Anna Miglio, Véronique Brousseau-Couture, Emile Godbout, Gabriel Antonius, Yang-Hao Chan, Steven G. Louie, Michel Côté, Matteo Giantomassi, and Xavier Gonze. Predominance of non-adiabatic effects in zero-point renormalization of the electronic band gap. *npj Computational Materials*, 6(1):167, 2020.
- [632] N W Ashcroft and N D Mermin. *Solid State Physics*. Saunders College Publishing, 1976.
- [633] R Peierls. Zur kinetischen theorie der wärmeleitung in kristallen. *Annalen der Physik*, 395(8):1055-1101, 1929.
- [634] Florian Knoop, Matthias Scheffler, and Christian Carbogno. Ab initio green-kubo simulations of heat transport in solids: Method and implementation. *Physical Review B*, 107(22):224304, 2023.
- [635] A. F. Ioffe and A. R. Regel. Non-crystalline, amorphous and liquid electronic semiconductors. *Prog. Semicond.*, 4(89):237-291, 1960.
- [636] Michele Simoncelli, Nicola Marzari, and Francesco Mauri. Unified theory of thermal transport in crystals and glasses. *Nature Physics*, 15(8):809-813, 2019.
- [637] D. A. Broido, M. Malorny, G. Birner, Natalio Mingo, and D. A. Stewart. Intrinsic lattice thermal conductivity of semiconductors from first principles. *Applied Physics Letters*, 91(23):231922, 2007.
- [638] Tianli Feng and Xiulin Ruan. Quantum mechanical prediction of four-phonon scattering rates and reduced thermal conductivity of solids. *Physical Review B*, 93(4):045202, 2016.
- [639] Navaneetha K Ravichandran and David Broido. Unified first-principles theory of thermal properties of insulators. *Physical Review-Section B-Condensed Matter*, 98(8):085205, August 2018.
- [640] Jivtesh Garg, Nicola Bonini, Boris Kozinsky, and Nicola Marzari. Role of disorder and anharmonicity in the thermal conductivity of silicon-germanium alloys: A first-principles study. *Physical Review Letters*, 106(4):045901, 2011.
- [641] Olle Hellman and I A Abrikosov. Temperature-dependent effective third-order interatomic force constants from first principles. *Physical Review-Section B-Condensed Matter*, 88(14):144301, October 2013.
- [642] S Stackhouse, L Stixrude, and BB Karki. Thermal conductivity of periclase (mgo) from first principles. *Physical Review Letters*, 104(20):208501, 2010.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [643] Florian Müller-Plathe. A simple nonequilibrium molecular dynamics method for calculating the thermal conductivity. *The Journal of Chemical Physics*, 106(14):6082–6085, 1997.
- [644] T. M. Gibbons and S. K. Estreicher. Erratum: Impact of impurities on the thermal conductivity of semiconductor nanostructures: First-principles theory [phys. rev. lett. 102, 255502 (2009)]. *Physical Review Letters*, 103(9):099904, August 2009.
- [645] Marcello Puligheddu, François Gygi, and Giulia Galli. First-principles simulations of heat transport. *Physical Review Materials*, 1(6):060802, November 2017.
- [646] Bingqing Cheng and Daan Frenkel. Computing the heat conductivity of fluids from density fluctuations. *Physical Review Letters*, 125(13):130602, September 2020.
- [647] Enrico Drigo, Maria Grazia Izzo, and Stefano Baroni. Heat conductivity from energy-density fluctuations. *The Journal of Chemical Physics*, 159(18):184107, 2023.
- [648] Ryogo Kubo. Statistical-mechanical theory of irreversible processes. i. general theory and simple applications to magnetic and conduction problems. *Journal of the Physical Society of Japan*, 12(6):570–586, 1957.
- [649] Olle Hellman and David A. Broido. Phonon thermal transport in Bi_2Te_3 from first principles. *Physical Review B*, 90(13):134309, 2014.
- [650] Christian Carbogno, Rampi Ramprasad, and Matthias Scheffler. Ab initio green-kubo approach for the thermal conductivity of solids. *Phys. Rev. Lett.*, 118(17):175901, April 2017.
- [651] P C Howell. Comparison of molecular dynamics methods and interatomic potentials for calculating the thermal conductivity of silicon. *The Journal of Chemical Physics*, 137(22):224111, 2012.
- [652] Loris Ercole, Aris Marcolongo, Paolo Umari, and Stefano Baroni. Gauge invariance of thermal transport coefficients. *Journal of Low Temperature Physics*, 185(1-2):79–86, April 2016.
- [653] Marcel F. Langer, J. Thorben Frank, and Florian Knopp. Stress and heat flux via automatic differentiation. *The Journal of Chemical Physics*, 159(17):174105, 2023.
- [654] Jie Chen, Xiangfan Xu, Jun Zhou, and Baowen Li. Interfacial thermal resistance: Past, present, and future. *Reviews of Modern Physics*, 94(2):025002, 2022.
- [655] T J Scheidemantel, C Ambrosch-Draxl, T Thonhauser, J V Badding, and J O Sofo. Transport coefficients from first-principles calculations. *Physical Review-Section B-Condensed Matter*, 68(12):125210, September 2003.
- [656] Samuel Poncé, Elena R Margine, and Feliciano Giustino. Towards predictive many-body calculations of phonon-limited carrier mobilities in semiconductors. *Physical Review-Section B-Condensed Matter*, 97(12):121201, March 2018.
- [657] Jin-Jian Zhou, Olle Hellman, and Marco Bernardi. Electron-phonon scattering in the presence of soft modes and electron mobility in SrTiO_3 perovskite from first principles. *Physical Review Letters*, 121(22):226603, June 2018.
- [658] Andrea Cepellotti and Boris Kozinsky. Interband tunneling effects on materials transport properties using the first principles wigner distribution. *Materials Today Physics*, 19:100412, 2021.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [659] D A Greenwood. The boltzmann equation in the theory of electrical conduction in metals. *Proceedings of the Physical Society*, 71(4):585, 1958.
- [660] Bastian Holst, Martin French, and Ronald Redmer. Electronic transport coefficients from ab initio simulations and application to dense liquid hydrogen. *Phys. Rev. B*, 83(23):235120, June 2011.
- [661] Florian Fiebig. Assessing electronic transport in solid materials via the fluctuation-dissipation theorem. Master thesis. <https://doi.org/10.14279/depositonce-21491>, 2023.
- [662] Zhenkun Yuan. *Electrical conductivity from first principles*. PhD thesis, Humboldt-Universität zu Berlin, 2022. <https://doi.org/10.18452/24219>.
- [663] Chenmu Zhang, Ruoyu Wang, Himani Mishra, and Yuanyue Liu. Two-dimensional semiconductors with high intrinsic carrier mobility at room temperature. *Physical Review Letters*, 130(8):087001, 2023.
- [664] Georg K.H. Madsen, Jesús Carrete, and Matthieu J. Verstraete. BoltzTraP2, a program for interpolating band structures and calculating semi-classical transport coefficients. *Computer Physics Communications*, 231:140–145, 2018.
- [665] Tao Fan and Artem R Oganov. Aicon2: A program for calculating transport properties quickly and accurately. *Computer Physics Communications*, 266:108027, 2021.
- [666] G Jeffrey Snyder and Eric S Toberer. Complex thermoelectric materials. *Nat. Mater.*, 7(2):105–114, 2008.
- [667] Matteo Rinaldi, Matthias Kick, Karsten Reuter, and Christian Carbogno. Advancing our Understanding of Optoionic Effects for the Design of Solar Batteries: A Theoretical Perspective. *arXiv*, 2503.07460, 2025.
- [668] Rotraut Merkle, Maximilian F. Hoedl, Giulia Raimondi, Reihaneh Zohourian, and Joachim Maier. Oxides with mixed protonic and electronic conductivity. *Annual Review of Materials Research*, 51(1):461–493, 2021.
- [669] T. A. Su, M. Neupane, M. L. Steigerwald, L. Venkataraman, and C. Nuckolls. Chemical principles of single-molecule electronics. *Nature Reviews Materials*, 1:16002, 2016.
- [670] M. Thoss and F. Evers. Perspective: Theory of quantum transport in molecular junctions. *Journal of Chemical Physics*, 148:030901, 2018.
- [671] F. Evers, R. Korytár, S. Tewari, and J. M. van Ruitenbeek. Advances and challenges in single-molecule electron transport. *Reviews of Modern Physics*, 92:035001, 2020.
- [672] J. C. Cuevas and E. Scheer. *Molecular Electronics*. World Scientific, 2nd edition, 2010.
- [673] M. Camarasa-Gómez, D. Hernangómez-Pérez, M. S. Inkpen, G. Lovat, E.-D. Fung, X. Roy, L. Venkataraman, and F. Evers. Mechanically tunable quantum interference in ferrocene-based single-molecule junctions. *Nano Letters*, 20:6381, 2020.
- [674] V. Meded, A. Bagrets, K. Fink, R. Chandrasekar, M. Ruben, F. Evers, A. Bernard-Mantel, J. S. Seldenthuis, A. Beukman, and H. S. J. van der Zant. Electrical control over the fe(ii) spin crossover in a single molecule: Theory and experiment. *Physical Review B*, 83:245415, 2011.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [675] M. Elbing, R. Ochs, M. Koentopp, M. Fischer, C. von Hänisch, F. Weigend, F. Evers, H. B. Weber, and M. Mayor. A single-molecule diode. *Proceedings of the National Academy of Sciences*, 102:8815, 2005.
- [676] F. Evers, A. Aharony, N. Bar-Gill, O. Entin-Wohlman, P. Hedegård, O. Hod, P. Jelinek, G. Kamieniarz, M. Lemeshko, K. Michaeli, V. Mujica, R. Naaman, Y. Paltiel, S. Refaely-Abramson, O. Tal, J. Thijssen, M. Thoss, J. M. van Ruitenbeek, L. Venkataraman, D. H. Waldeck, B. Yan, and L. Kronik. Theory of chirality induced spin selectivity: Progress and challenges. *Advanced Materials*, 34:2106629, 2022.
- [677] P. Havu, M. J. Hashemi, M. Kaukonen, E. T. Seppälä, and R. M. Nieminen. Effect of gating and pressure on the electronic transport properties of crossed nanotube junctions: Formation of a schottky barrier. *Journal of Physics: Condensed Matter*, 23:112203, 2011.
- [678] P. Havu, A. Sillanpää, N. Runeberg, J. Tarus, E. T. Seppälä, and R. M. Nieminen. Effects of chemical functionalization on electronic transport in carbon nanobuds. *Physical Review B*, 85:115446, 2012.
- [679] T. Ketolainen, V. Havu, and M. J. Puska. Conductivity of aucl_4 -functionalized carbon nanotube networks. *Journal of Physical Chemistry C*, 121:4627, 2017.
- [680] A. Arnold, F. Weigend, and F. Evers. Quantum chemistry calculations for molecules coupled to reservoirs: Formalism, implementation, and application to benzenedithiol. *Journal of Chemical Physics*, 126:174101, 2007.
- [681] M. Camarasa-Gómez, D. Hernangómez-Pérez, and F. Evers. Spin-orbit torque in single-molecule junctions from ab initio. *Journal of Physical Chemistry Letters*, 15:5747, 2024.
- [682] A. Bagrets. Spin-polarized electron transport across metal-organic molecules: A density functional theory approach. *Journal of Chemical Theory and Computation*, 9:2801, 2013.
- [683] J. Wilhelm, M. Walz, M. Stendel, A. Bagrets, and F. Evers. Ab initio simulations of scanning-tunneling-microscope images with embedding techniques and application to c_{58} -dimers on $\text{au}(111)$. *Physical Chemistry Chemical Physics*, 15:6684, 2013.
- [684] H. J. W. Haug and A.-P. Jauho. *Quantum Kinetics in Transport and Optics of Semiconductors*. Springer-Verlag, 2nd edition, 2008.
- [685] M. Di Ventra. *Electrical Transport in Nanoscale Systems*. Cambridge University Press, 1st edition, 2008.
- [686] M. Camarasa-Gómez. *Ab Initio Electronic Transport in Single-Molecule Junctions: Quantum Interference Effects and Spin-Orbit Torque*. PhD thesis, Universität Regensburg, 2021. <https://epub.uni-regensburg.de/46138/>.
- [687] Y. Meir and N. S. Wingreen. Landauer formula for the current through an interacting electron region. *Physical Review Letters*, 68:2512, 1992.
- [688] F. Evers and A. Arnold. Molecular conductance from ab initio calculations: Self energies and absorbing boundary conditions. In C. Röthig, G. Schön, and M. Vojta, editors, *CFN Lectures on Functional Nanostructures Vol. 2*, volume 820 of *Lecture Notes in Physics*, pages 27–53. Springer, 2011.
- [689] P.-O. Löwdin. On the non-orthogonality problem connected with the use of atomic wave functions in the theory of molecules and crystals. *Journal of Chemical Physics*, 18:365, 1950.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [690] M. A. García-Blázquez, W. Dednam, and J. J. Palacios. Nonequilibrium magneto-conductance as a manifestation of spin filtering in chiral nanojunctions. *Journal of Physical Chemistry Letters*, 14:7931, 2023.
- [691] S. Naskar, V. Mujica, and C. Herrmann. Chiral-induced spin selectivity and non-equilibrium spin accumulation in molecules and interfaces: A first-principles study. *Journal of Physical Chemistry Letters*, 14:694, 2023.
- [692] M. Walz, J. Wilhelm, and F. Evers. Current patterns and orbital magnetism in mesoscopic dc transport. *Physical Review Letters*, 113:136602, 2014.
- [693] M. Walz, A. Bagrets, and F. Evers. Local current density calculations for molecular films from ab initio. *Journal of Chemical Theory and Computation*, 11:5161, 2015.
- [694] J. Wilhelm, M. Walz, and F. Evers. Ab initio spin-flip conductance of hydrogenated graphene nanoribbons: Spin-orbit interaction and scattering with local impurity spins. *Physical Review B*, 92:014405, 2015.
- [695] Harald Oberhofer, Karsten Reuter, and Jochen Blumberger. Charge transport in molecular materials: An assessment of computational methods. *Chemical Reviews*, 117:10319–10357, 2017.
- [696] R. Hoffmann. A chemical and theoretical way to look at bonding on surfaces. *Reviews of Modern Physics*, 60:601–628, 1988.
- [697] B. Hammer and J. K. Nørskov. Electronic factors determining the reactivity of metal surfaces. *Surface Science*, 343:211–220, 1995.
- [698] D. News. Self-consistent model of hydrogen chemisorption. *Physical Review*, 178:1123–1135, 1969.
- [699] J. Tersoff and D. Hamann. Theory and application for the scanning tunneling microscope. *Physical Review Letters*, 50:1998–2001, 1983.
- [700] E. Gull, A. Millis, A. Lichtenstein, A. Rubtsov, M. Troyer, and P. Werner. Continuous-time monte carlo methods for quantum impurity models. *Reviews of Modern Physics*, 83:349–404, 2011.
- [701] Alexander Neef, Sebastian Hammer, Yuxuan Yao, Shubham Sharma, Samuel Beaulieu, Shuo Dong, Tommaso Pincelli, Maximilian Frank, Martin Wolf, Mariana Rossi, Harald Oberhofer, Laurenz Rettig, Jens Pflaum, and Ralph Ernstorfer. Frontier orbitals control dynamical disorder in molecular semiconductors, 2024.
- [702] T. Van Voorhis, T. Kowalczyk, B. Kaduk, L.-P. Wang, C.-L. Cheng, and Q. Wu. The diabatic picture of electron transfer, reaction barriers, and molecular dynamics. *Annual Review of Physical Chemistry*, 61:149–170, 2010.
- [703] Vitaly A. Rassolov, Mark A. Ratner, John A. Pople, Paul C. Redfern, and Larry A. Curtiss. 6-31g* basis set for third-row atoms. *Journal of Computational Chemistry*, 22(9):976–984, 2001.
- [704] Christoph Schober, Karsten Reuter, and Harald Oberhofer. Virtual screening for high carrier mobility in organic semiconductors. *The journal of physical chemistry letters*, 7(19):3973–3977, 2016.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [705] Anthony M Reilly, Richard I Cooper, Claire S Adjiman, Saswata Bhattacharya, A Daniel Boese, Jan Gerit Brandenburg, Peter J Bygrave, Rita Bylsma, Josh E Campbell, Roberto Car, et al. Report on the sixth blind test of organic crystal structure prediction methods. *Acta Crystallographica Section B: Structural Science, Crystal Engineering and Materials*, 72(4):439–459, 2016.
- [706] Christoph Schober, Karsten Reuter, and Harald Oberhofer. Critical analysis of fragment-orbital DFT schemes for the calculation of electronic coupling values. *The Journal of chemical physics*, 144(5), 2016.
- [707] K Senthilkumar, FC Grozema, FM Bickelhaupt, and LDA Siebbeles. Charge transport in columnar stacked triphenylenes: Effects of conformational fluctuations on charge transfer integrals and site energies. *The Journal of chemical physics*, 119(18):9809–9817, 2003.
- [708] Harald Oberhofer and Jochen Blumberger. Revisiting electronic couplings and incoherent hopping models for electron transport in crystalline c 60 at ambient temperatures. *Physical Chemistry Chemical Physics*, 14(40):13846–13852, 2012.
- [709] I. Kondov, M. Čížek, C. Benesch, H. Wang, and M. Thoss. Quantum dynamics of photoinduced electron-transfer reactions in dye-semiconductor systems: First-principles description and application to coumarin 343-tio₂. *Journal of Physical Chemistry C*, 111:11970–11981, 2007.
- [710] Zdenek Futera and Jochen Blumberger. Electronic couplings for charge transfer across molecule/metal and molecule/semiconductor interfaces: Performance of the projector operator-based diabatization approach. *Journal of Physical Chemistry C*, 121:19677–19689, 2017.
- [711] S. Ghan, C. Kunkel, K. Reuter, and H. Oberhofer. Improved projection-operator diabatization schemes for the calculation of electronic coupling values. *Journal of Chemical Theory and Computation*, 12:7431–7443, 2020.
- [712] S. Ghan, E. Diesen, C. Kunkel, K. Reuter, and H. Oberhofer. Interpreting ultrafast electron transfer on surfaces with a converged first-principles newns-anderson chemisorption function. *Journal of Chemical Physics*, 158:234103, 2023.
- [713] Shyue Ping Ong, William Davidson Richards, Anubhav Jain, Geoffroy Hautier, Michael Kocher, Shreyas Cholia, Dan Gunter, Vincent L Chevrier, Kristin A Persson, and Gerbrand Ceder. Python materials genomics (pymatgen): A robust, open-source python library for materials analysis. *Computational Materials Science*, 68:314–319, 2013.
- [714] Alex Ganose, Hrushikesh Sahasrabudde, Mark Asta, Kevin Beck, Tathagata Biswas, Alexander Bonkowski, Joana Bustamante, Xin Chen, Yuan Chiang, and Daryl et al. Chrzan. Atomate2: Modular workflows for materials science. *Digital Discovery*, 4:1944–1973, 2025.
- [715] Ask Hjorth Larsen, Mikael J Kuisma, Tara M Boland, Fredrik A Nilsson, and Kristian S Thygesen. Taskblaster: A generic framework for automated computational workflows. *Digital Discovery*, 4(9):2512–2520, 2025.
- [716] Sebastiaan P Huber, Spyros Zoupanos, Martin Uhrin, Leopold Talirz, Leonid Kahle, Rico Häuselmann, Dominik Gresch, Tiziano Müller, Aliaksandr V Yakutovich, Casper W Andersen, et al. Aiida 1.0, a scalable computational infrastructure for automated reproducible workflows and data provenance. *Scientific data*, 7(1):300, 2020.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [717] Koichi Momma and Fujio Izumi. Vesta: A three-dimensional visualization system for electronic and structural analysis. *Journal of Applied Crystallography*, 41:653–658, 06 2008.
- [718] Anton Kokalj. Xcrysden — a new program for displaying crystalline structures and electron densities. *Journal of Molecular Graphics and Modelling*, 17(3):176–179, 1999.
- [719] William Humphrey, Andrew Dalke, and Klaus Schulten. VMD – Visual Molecular Dynamics. *Journal of Molecular Graphics*, 14:33–38, 1996.
- [720] Jmol: an open-source Java viewer for chemical structures in 3D.
- [721] Sebastian Kokott, Iker Hurtado, Christian Vorwerk, Claudia Draxl, Volker Blum, and Matthias Scheffler. GIMS: graphical interface for materials simulations. *Journal of Open Source Software*, 6(57):2767, 2021.
- [722] Andrew S. Rosen, Max Gallant, Janine George, Janosh Riebesell, Hrushikesh Sahasrabudde, Jimmy-Xuan Shen, Mingjian Wen, Matthew L. Evans, Guido Petretto, David Waroquiers, Gian-Marco Rignanesi, Kristin A. Persson, Anubhav Jain, and Alex M. Ganose. Jobflow: Computational workflows made simple. *Journal of Open Source Software*, 9(93):5995, 2024.
- [723] Thomas Purcell and Andrei Sobolev. FHI-aims with ASE. <https://fhi-aims-club.gitlab.io/tutorials/fhi-aims-with-ase/>, 2024. [Online; accessed 26-Sep-2024].
- [724] Andrei Sobolev and Uthpala Herath. High-throughput workflows with FHI-aims and atomate2. <https://workflows-with-atomate2-fhi-aims-club-tutorials-a0acc2efc2fba83.gitlab.io/>, 2024. [Online; accessed 26-Sep-2024].
- [725] Akhil Nair. Working in FHI-aims with TaskBlaster: Tutorial. <https://gitlab.com/FHI-aims-club/tutorials/fhi-aims-with-taskblaster>, 2024. [Online; accessed 26-Sep-2024].
- [726] Andrei Sobolev. Working with FHI-aims and AiiDA. <https://fhi-aims-club.gitlab.io/tutorials/fhi-aims-with-aiida/>, 2024. [Online; accessed 26-Sep-2024].
- [727] Alexander Knoll. Managing Research Data with AiiDA and FHI-aims. <https://fhi-aims-club.gitlab.io/tutorials/fhi-aims-and-aiida/>, 2021. [Online; accessed 26-Sep-2024].
- [728] GIMS stable release version. Available at: <https://gims.ms1p.org>.
- [729] GIMS development version. Available at: <https://gims-dev.ms1p.org>.
- [730] Hagen-Henrik Kowalski, Sebastian Kokott, and Andrei Sobolev. GIMS Manual. <https://gims-developers.gitlab.io/gims/>, 2024. [Online; accessed 26-Sep-2024].
- [731] Ask Hjorth Larsen, Jens Jørgen Mortensen, Jakob Blomqvist, Ivano E Castelli, Rune Christensen, Marcin Dułak, Jesper Friis, Michael N Groves, Bjørk Hammer, Cory Hargus, Eric D Hermes, Paul C Jennings, Peter Bjerre Jensen, James Kermode, John R Kitchin, Esben Leonhard Kolsbjerg, Joseph Kubal, Kristen Kaasbjerg, Steen Lysgaard, Jón Bergmann Maronsson, Tristan Maxson, Thomas Olsen, Lars Pastewka, Andrew Peterson, Carsten Rostgaard, Jakob Schiøtz, Ole Schütt, Mikkel Strange, Kristian S Thygesen, Tejs Vegge, Lasse Vilhelmsen, Michael Walter, Zhenhua Zeng, and Karsten W Jacobsen. The atomic simulation environment—a python library for working with atoms. *Journal of Physics: Condensed Matter*, 29(27):273002, 2017.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [732] Stefan Bauer, Peter Benner, Tristan Bereau, Volker Blum, Mario Boley, Christian Carbogno, C Richard A Catlow, Gerhard Dehm, Sebastian Eibl, Ralph Ernstorfer, et al. Roadmap on data-centric materials science. *Modelling and Simulation in Materials Science and Engineering*, 32(6):063301, 2024.
- [733] Jörg Behler and Michele Parrinello. Generalized neural-network representation of high-dimensional potential-energy surfaces. *Physical Review Letters*, 98:146401, 2007.
- [734] Volker L Deringer, Albert P Bartók, Noam Bernstein, David M Wilkins, Michele Ceriotti, and Gábor Csányi. Gaussian process regression for materials and molecules. *Chemical Reviews*, 121(16):10073–10141, 2021.
- [735] Ilyes Batatia, David P Kovacs, Gregor Simm, Christoph Ortner, and Gábor Csányi. Mace: Higher order equivariant message passing neural networks for fast and accurate force fields. *Advances in Neural Information Processing Systems*, 35:11423–11436, 2022.
- [736] Simon Wengert, Gábor Csányi, Karsten Reuter, and Johannes T. Margraf. Data-efficient machine learning for molecular crystal structure prediction. *Chem. Sci.*, 12:4536–4546, 2021.
- [737] Carsten G. Staacke, Hendrik H. Heenen, Christoph Scheurer, Gábor Csányi, Karsten Reuter, and Johannes T. Margraf. On the role of long-range electrostatics in machine-learned interatomic potentials for complex battery materials. *ACS Applied Energy Materials*, 4(11):12562–12569, Nov 2021.
- [738] Sina Stocker, Hyunwook Jung, Gábor Csányi, C. Franklin Goldsmith, Karsten Reuter, and Johannes T. Margraf. Estimating free energy barriers for heterogeneous catalytic reactions with machine learning potentials and umbrella integration. *Journal of Chemical Theory and Computation*, 19(19):6796–6804, Oct 2023.
- [739] Hyunwook Jung, Lena Sauerland, Sina Stocker, Karsten Reuter, and Johannes T. Margraf. Machine-learning driven global optimization of surface adsorbate geometries. *npj Computational Materials*, 9(1):114, Jun 2023.
- [740] Elena Gelžinytė, Simon Wengert, Tamás K. Stenczel, Hendrik H. Heenen, Karsten Reuter, Gábor Csányi, and Noam Bernstein. wfl Python toolkit for creating machine learning interatomic potentials and related atomistic simulation workflows. *The Journal of Chemical Physics*, 159(12):124801, 09 2023.
- [741] Libatoms. <https://libatoms.github.io/workflow/index.html>.
- [742] Jiequn Han, Linfeng Zhang, Roberto Car, et al. Deep potential: A general representation of a many-body potential energy surface. *arXiv preprint arXiv:1707.01478*, 2017.
- [743] Tongqi Wen, Linfeng Zhang, Han Wang, David J Srolovitz, et al. Deep potentials for materials science. *Materials Futures*, 1(2):022601, 2022.
- [744] Roman Kempf, Agnieszka Kuc, Thomas Brumme, and Thomas Heine. Edge conductivity in ptse2 nanostructures. *Small Structures*, 5(2):2300222, 2024.
- [745] Tsz Wai Ko, Jonas A. Finkler, Stefan Goedecker, and Jörg Behler. General-purpose machine learning potentials capturing nonlocal charge transfer. *Acc. Chem. Res.*, 54:808–817, 2021.
- [746] Stefan Chmiela, Valentin Vassilev-Galindo, Oliver T Unke, Adil Kabylda, Huziel E Sauceda, Alexandre Tkatchenko, and Klaus-Robert Müller. Accurate global machine learning force fields for molecules with hundreds of atoms. *Science Advances*, 9(2):eadf0873, 2023.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [747] Huziel E Saucedo, Luis E Gálvez-González, Stefan Chmiela, Lauro Oliver Paz-Borbón, Klaus-Robert Müller, and Alexandre Tkatchenko. Bigdml—towards accurate quantum machine learning force fields for materials. *Nature communications*, 13(1):3733, 2022.
- [748] Oliver T Unke, Martin Stöhr, Stefan Ganscha, Thomas Unterthiner, Hartmut Maennel, Sergii Kashubín, Daniel Ahlin, Michael Gastegger, Leonardo Medrano Sandonas, Joshua T Berryman, et al. Biomolecular dynamics with machine-learned quantum-mechanical force fields trained on diverse chemical fragments. *Science Advances*, 10(14):eadn4397, 2024.
- [749] J Thorben Frank, Oliver T Unke, Klaus-Robert Müller, and Stefan Chmiela. A euclidean transformer for fast and stable machine learned force fields. *Nature Communications*, 15(1):6539, 2024.
- [750] Gkx. <https://github.com/sirmarcel/gkx>.
- [751] Leonid Kahle and Federico Zipoli. Quality of uncertainty estimates from neural network potential ensembles. *Physical Review E*, 105(1):015311, 2022.
- [752] Available at:.
- [753] Simon Batzner, Albert Musaelian, Lixin Sun, Mario Geiger, Jonathan P Mailoa, Mordechai Kornbluth, Nicola Molinari, Tess E Smidt, and Boris Kozinsky. E(3)-equivariant graph neural networks for data-efficient and accurate interatomic potentials. *Nature Communications*, 13(1):2453, 2022.
- [754] Thorben Frank, Oliver Unke, and Klaus-Robert Müller. So3krates: Equivariant attention for interactions on arbitrary length-scales in molecular systems. In S. Koyejo, S. Mohamed, A. Agarwal, D. Belgrave, K. Cho, and A. Oh, editors, *Advances in Neural Information Processing Systems*, volume 35, pages 29400–29413. Curran Associates, Inc., 2022.
- [755] Milica Todorović, Michael U. Gutmann, Jukka Corander, and Patrick Rinke. Bayesian inference of atomistic structure in functional materials. *npj Computational Materials*, 5(1), 2019.
- [756] Boss public website. <https://sites.utu.fi/boss>.
- [757] Felipe Oviedo, Juan Lavista Ferres, Tonio Buonassisi, and Keith T Butler. Interpretable and explainable machine learning for materials science and chemistry. *Accounts of Materials Research*, 3(6):597–607, 2022.
- [758] Akhil S Nair, Lucas Foppa, and Matthias Scheffler. Materials-discovery workflow guided by symbolic regression for identifying acid-stable oxides for electrocatalysis. *npj Computational Materials*, 11(1):150, 2025.
- [759] Runhai Ouyang, Stefano Curtarolo, Emre Ahmetcik, Matthias Scheffler, and Luca M Ghiringhelli. Sisso: A compressed-sensing method for identifying the best low-dimensional descriptor in an immensity of offered candidates. *Physical Review Materials*, 2(8):083802, 2018.
- [760] Thomas A. R. Purcell, Matthias Scheffler, and Luca M. Ghiringhelli. Recent advances in the SISO method and their implementation in the SISO++ code. *The Journal of Chemical Physics*, 159(11):114110, 09 2023.
- [761] Kangming Li, Daniel Persaud, Kamal Choudhary, Brian DeCost, Michael Greenwood, and Jason Hattrick-Simpers. Exploiting redundancy in large materials datasets for efficient machine learning with less data. *Nature Communications*, 14(1):7283, 2023.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [762] Kisung Kang. ALMOMD. <https://gitlab.com/FHI-aims-club/ALmoMD>, 2024. [Online; accessed 26-Sep-2024].
- [763] Lincan Fang, Esko Makkonen, Milica Todorović, Patrick Rinke, and Xi Chen. Efficient Amino Acid Conformer Search with Bayesian Optimization. *Journal of Chemical Theory and Computation*, 17(3):1955–1966, 2021.
- [764] Jari Järvi, Patrick Rinke, and Milica Todorović. Detecting stable adsorbates of (1S)-camphor on Cu(111) with Bayesian optimization. *Beilstein Journal of Nanotechnology*, 11(1):1577–1589, 2020.
- [765] Jari Järvi, Benjamin Alldritt, Ondřej Krejčí, Milica Todorović, Peter Liljeroth, and Patrick Rinke. Integrating Bayesian Inference with Scanning Probe Experiments for Robust Identification of Surface Adsorbate Configurations. *Advanced Functional Materials*, 31(32), 2021.
- [766] Alexander T. Egger, Lukas Hörmann, Andreas Jeindl, Michael Scherbela, Veronika Obersteiner, Milica Todorović, Patrick Rinke, and Oliver T. Hofmann. Charge Transfer into Organic Thin Films: A Deeper Insight through Machine-Learning-Assisted Structure Search. *Advanced Science*, 7(15), 2020.
- [767] Azimatu Fangnon, Marc Dvorak, Ville Havu, Milica Todorović, Jingrui Li, and Patrick Rinke. Protective Coating Interfaces for Perovskite Solar Cell Materials: A First-Principles Study. *ACS Applied Materials & Interfaces*, 14(10):12758–12765, 2022.
- [768] Boss code. <https://gitlab.com/cest-group/boss>.
- [769] Boss documentation. <https://cest-group.gitlab.io/boss>.
- [770] Sequential learning with sisso. <https://gitlab.com/FHI-aims-club/tutorials/tutorial-sl-sisso>.
- [771] Available at: <https://gims.ms1p.org/static/index.html>.
- [772] Yujin Hwang, Won Jo, Juyoung Hong, and Yukyung Choi. Overcoming overconfidence for active learning. *IEEE Access*, 2024.
- [773] Kristof T Schütt, Michael Gastegger, Alexandre Tkatchenko, K-R Müller, and Reinhard J Maurer. Unifying machine learning and quantum chemistry with a deep neural network for molecular wavefunctions. *Nature communications*, 10(1):5024, 2019.
- [774] Oliver Unke, Mihail Bogojeski, Michael Gastegger, Mario Geiger, Tess Smidt, and Klaus-Robert Müller. Se (3)-equivariant prediction of molecular wavefunctions and electronic densities. *Advances in Neural Information Processing Systems*, 34:14434–14447, 2021.
- [775] Xiang Fu, Andrew Rosen, Kyle Bystrom, Rui Wang, Albert Musaelian, Boris Kozinsky, Tess Smidt, and Tommi Jaakkola. A recipe for charge density prediction. *arXiv preprint arXiv:2405.19276*, 2024.
- [776] John C Snyder, Matthias Rupp, Katja Hansen, Klaus-Robert Müller, and Kieron Burke. Finding density functionals with machine learning. *Physical review letters*, 108(25):253002, 2012.
- [777] James Kirkpatrick, Brendan McMorrow, David HP Turban, Alexander L Gaunt, James S Spencer, Alexander GDG Matthews, Annette Obika, Louis Thiry, Meire Fortunato, David Pfau, et al. Pushing the frontiers of density functionals by solving the fractional electron problem. *Science*, 374(6573):1385–1389, 2021.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [778] Joseph W. Abbott, Wei Bin How, Guillaume Fraux, and Michele Ceriotti. lab-cosmo/rholearn: rholearn vo.1.0. Zenodo, October 2024.
- [779] Wei Bin How, Sanggyu Chong, Federico Grasselli, Kevin K. Huguenin-Dumittan, and Michele Ceriotti. Adaptive energy reference for machine-learning models of the electronic density of states, July 2024.
- [780] Liwei Zhang, Berk Onat, Geneviève Dusson, Adam McSloy, G. Anand, Reinhard J. Maurer, Christoph Ortner, and James R. Kermode. Equivariant analytical mapping of first principles Hamiltonians to accurate and transferable materials models. *npj Computational Materials*, 8(1):1–14, July 2022.
- [781] Pavel Stishenko, Adam McSloy, Berk Onat, Ben Hourahine, Reinhard J. Maurer, James R. Kermode, and Andrew Logsdail. Integrated workflows and interfaces for data-driven semi-empirical electronic structure calculations. *The Journal of Chemical Physics*, 161(1):012502, July 2024.
- [782] He Li, Zun Wang, Nianlong Zou, Meng Ye, Runzhang Xu, Xiaoxun Gong, Wenhui Duan, and Yong Xu. Deep-learning density functional theory Hamiltonian for efficient ab initio electronic-structure calculation. *Nature Computational Science*, 2(6):367–377, June 2022.
- [783] Xiaoxun Gong, He Li, Nianlong Zou, Runzhang Xu, Wenhui Duan, and Yong Xu. General framework for E(3)-equivariant neural network representation of density functional theory Hamiltonian. *Nature Communications*, 14(1):2848, May 2023.
- [784] He Li, Zechen Tang, Xiaoxun Gong, Nianlong Zou, Wenhui Duan, and Yong Xu. Deep-learning electronic-structure calculation of magnetic superstructures. *Nature Computational Science*, 3(4):321–327, April 2023.
- [785] Pavel V. Stishenko, Thomas W. Keal, Scott M. Woodley, Volker Blum, Benjamin Hourahine, Reinhard J. Maurer, and Andrew J. Logsdail. Atomic Simulation Interface (ASI): Application programming interface for electronic structure codes. *Journal of Open Source Software*, 8(85):5186, May 2023.
- [786] Filippo Bigi, Joseph W. Abbott, Philip Loche, Arslan Mazitov, Davide Tisi, Marcel F. Langer, Alexander Goscinski, Paolo Pegolo, Sanggyu Chong, Rohit Goswami, Sofiiia Chorna, Matthias Kellner, Michele Ceriotti, and Guillaume Fraux. Metatensor and metatomic: foundational libraries for interoperable atomistic machine learning, 2025.
- [787] Andrea Grisafi, Alan M. Lewis, Mariana Rossi, and Michele Ceriotti. Electronic-Structure Properties from Atom-Centered Predictions of the Electron Density. *Journal of Chemical Theory and Computation*, 19(14):4451–4460, July 2023.
- [788] Raghunathan Ramakrishnan, Pavlo O. Dral, Matthias Rupp, and O. Anatole von Lilienfeld. Quantum chemistry structures and properties of 134 kilo molecules. *Scientific Data*, 1(1):140022, August 2014.
- [789] Ralf Drautz. Atomic cluster expansion for accurate and transferable interatomic potentials. *Phys. Rev. B*, 99:014104, January 2019.
- [790] Yuxiang Wang, He Li, Zechen Tang, Honggeng Tao, Yanzhen Wang, Zilong Yuan, Zezhou Chen, Wenhui Duan, and Yong Xu. DeepH-2: Enhancing deep-learning electronic structure via an equivariant local-coordinate transformer, January 2024.
- [791] He Li, Zechen Tang, Jingheng Fu, Wen-Han Dong, Nianlong Zou, Xiaoxun Gong, Wenhui Duan, and Yong Xu. Deep-Learning Density Functional Perturbation Theory. *Physical Review Letters*, 132(9):096401, February 2024.

Roadmap on Advancements of the FHI-aims Software Package

BIBLIOGRAPHY

- [792] Zechen Tang, He Li, Peize Lin, Xiaoxun Gong, Gan Jin, Lixin He, Hong Jiang, Xinguo Ren, Wenhui Duan, and Yong Xu. A deep equivariant neural network approach for efficient hybrid density functional calculations. *Nature Communications*, 15(1):8815, October 2024.
- [793] Chiheb Ben Mahmoud, Andrea Anelli, Gábor Csányi, and Michele Ceriotti. Learning the electronic density of states in condensed matter. *Physical Review B*, 102(23):235130, December 2020.
- [794] Lorenz C. Blum and Jean-Louis Reymond. 970 Million Druglike Small Molecules for Virtual Screening in the Chemical Universe Database GDB-13. *Journal of the American Chemical Society*, 131(25):8732–8733, July 2009.
- [795] Matthias Rupp, Alexandre Tkatchenko, Klaus-Robert Müller, and O. Anatole von Lilienfeld. Fast and Accurate Modeling of Molecular Atomization Energies with Machine Learning. *Physical Review Letters*, 108(5):058301, January 2012.
- [796] Atomic Simulation Interface (ASI) API. <https://pvst.gitlab.io/asi>.
- [797] ACEhamiltonians.jl. <https://acesuit.github.io/ACEhamiltonians.jl/dev>.
- [798] Zechen Tang. Code and tutorial for DeepH@FHI-aims interface, October 2024. <https://zenodo.org/records/13943187>.
- [799] Andrea Grisafi, Augustin Bussy, Mathieu Salanne, and Rodolphe Vuilleumier. Predicting the charge density response in metal electrodes. *Physical Review Materials*, 7(12):125403, December 2023.
- [800] Yang Li, Zechen Tang, Zezhou Chen, Minghui Sun, Boheng Zhao, He Li, Honggeng Tao, Zilong Yuan, Wenhui Duan, and Yong Xu. Neural-network Density Functional Theory Based on Variational Energy Minimization. *Physical Review Letters*, 133(7):076401, August 2024.
- [801] Zilong Yuan, Zhiming Xu, He Li, Xinle Cheng, Honggeng Tao, Zechen Tang, Zhiyuan Zhou, Wenhui Duan, and Yong Xu. Equivariant neural network force fields for magnetic materials. *Quantum Frontiers*, 3(1):8, April 2024.
- [802] Yuxiang Wang, Yang Li, Zechen Tang, He Li, Zilong Yuan, Honggeng Tao, Nianlong Zou, Ting Bao, Xinghao Liang, Zezhou Chen, Shanghua Xu, Ce Bian, Zhiming Xu, Chong Wang, Chen Si, Wenhui Duan, and Yong Xu. Universal materials model of deep-learning density functional theory Hamiltonian. *Science Bulletin*, 69(16):2514–2521, August 2024.