Lawrence Berkeley National Laboratory
Recent Work

Title
MOLGW 1: Many-body perturbation theory software for atoms, molecules, and clusters

Permalink
https://escholarship.org/uc/item/6280604q

Authors
Bruneval, F
Rangel, T
Hamed, SM
et al.

Publication Date
2016-11-01

DOI
10.1016/j.cpc.2016.06.019

Peer reviewed
MOLGW 1: Many-body perturbation theory software for atoms, molecules, and clusters

Article in Computer Physics Communications · July 2016
DOI: 10.1016/j.cpc.2016.06.019

CITATIONS
4

READS
62

6 authors, including:

Tonatiuh Rangel
Lawrence Berkeley National Laboratory
30 PUBLICATIONS 1,109 CITATIONS
SEE PROFILE

Meiyue Shao
Lawrence Berkeley National Laboratory
22 PUBLICATIONS 78 CITATIONS
SEE PROFILE

Chao Yang
Shanghai Institutes for Biological Sciences
130 PUBLICATIONS 1,657 CITATIONS
SEE PROFILE

Some of the authors of this publication are also working on these related projects:

NUCLEI View project

Excited state properties of photovoltaic materials from many-body perturbation theory View project

All content following this page was uploaded by Tonatiuh Rangel on 31 October 2016.
The user has requested enhancement of the downloaded file. All in-text references underlined in blue are added to the original document and are linked to publications on ResearchGate, letting you access and read them immediately.
**molgw 1: many-body perturbation theory software for atoms, molecules, and clusters**

Fabien Bruneval\(^a\)^b,c,*, Tonatiuh Rangel\(^b\)^c, Samia M. Hamed\(^b\)^c,d,e, Meiyue Shao\(^f\), Chao Yang\(^f\), Jeffrey B. Neaton\(^b\)^c,e

\(^a\)CEA, DEN, Service de Recherches de Métallurgie Physique, F-91191 Gif-sur-Yvette, France
\(^b\)Molecular Foundry, Lawrence Berkeley National Laboratory, Berkeley, California 94720, United States
\(^c\)Department of Physics, University of California, Berkeley, California 94720, United States
\(^d\)Department of Chemistry, University of California, Berkeley, California 94720, United States
\(^e\)Kavli Energy Nanosciences Institute at Berkeley, Berkeley, California 94720, United States
\(^f\)Computational Research Division, Lawrence Berkeley National Laboratory, Berkeley, CA 94720, United States

**Abstract**

We summarize the molgw code that implements density-functional theory and many-body perturbation theory in a Gaussian basis set. The code is dedicated to the calculation of the many-body self-energy within the GW approximation and the solution of the Bethe-Salpeter equation. These two types of calculations allow the user to evaluate physical quantities that can be compared to spectroscopic experiments. Quasiparticle energies, obtained through the calculation of the GW self-energy, can be compared to photoemission or transport experiments, and neutral excitation energies and oscillator strengths, obtained via solution of the Bethe-Salpeter equation, are measurable by optical absorption. The implementation choices outlined here have aimed at the accuracy and robustness of calculated quantities with respect to measurements. Furthermore, the algorithms implemented in molgw allow users to consider molecules or clusters containing up to 100 atoms with rather accurate basis sets, and to choose whether or not to apply the resolution-of-the-identity approximation. Finally, we demonstrate the parallelization efficacy of the molgw code over several hundreds of processors.

**Keywords:** Electronic structure of molecules; Many-body perturbation theory; GW approximation; Bethe-Salpeter equation

**PROGRAM SUMMARY**

Program Title: molgw

Journal Reference:  
Catalogue identifier:  
Licensing provisions: GPL v3.0 [1]

Programming language: Fortran 2003 with a few C subroutines, Python scripts

Number of processors used: 1-1024

Keywords: Electronic structure of molecules; Many-body perturbation theory; GW approximation; Bethe-Salpeter equation

Classification:

External routines/libraries: libint [2], libxc [3]

Nature of problem:  
Prediction of the electronic structure of atoms, molecules, clusters with a particular interest in their spectroscopic features, such as quasiparticle energies and optical spectra.

Solution method:

Using the GW approximation to many-body perturbation theory, molgw calculates total-energies, quasiparticle energies, and optical excitations.

Running time:  
From 30 seconds to a few hours


1. General presentation

Over the years, many-body perturbation theory (MBPT) [1, 2, 3] has been demonstrated to lead to meaningful approximations for a variety of interesting electronic properties, but especially for those that involve excitations. Excited electronic states arise naturally when one connects calculations to, for example,
experimental measurements of photoemission, inverse-photoemission, charge transport, or optical absorption.

In the field of theoretical chemistry, the celebrated Möller-Plesset perturbation theory [4] is most commonly derived in increasing orders with respect to the bare Coulomb interaction \( v \). However, for bulk metallic systems, the expansion with respect to \( v \) gives rise to infinite terms, and is therefore inadequate. The use of the screened Coulomb interaction \( W \), instead of the bare Coulomb interaction \( v \), remedies this issue [2] and consequently, a comprehensive perturbative approach based on \( W \) can be derived [5]. Its first-order approximation, namely the \( GW \) approximation, has been used with great success for the last 30 years in extended systems to calculate the band gaps of solids [6, 7, 8, 9]. In the past, the application of the \( GW \) approximation to finite systems was rather infrequent [10, 11, 12, 13]. But there has been much recent increase in interest [14, 15, 16, 17, 18, 19, 20, 21, 22, 23] in the application of the \( GW \) approximation to problems involving atoms, molecules, and clusters, in part driven by the quest to develop efficient techniques to address mixed systems such as molecular junctions [24, 25, 26, 27, 28].

Several codes that implement MBPT in its \( GW \) approximation for periodic systems are well-established and regularly utilized. To name a few of them, Abinit [29], Yambo [30], sax [31], vasp [32], and BerkeleyGW [33] produce \( GW \) quasiparticle energies and optical spectra from the Bethe-Salpeter equation (BSE). A few newer implementations have the advantage of a specific representation of \( W \) to target large but still periodic systems [34, 35].

Implementations of the \( GW \) approximation of MBPT for finite systems are much rarer, but their numbers have increased during the last few years and include kg-wbs [13], fhi-aims [36], Fiesta [15], Turbomol [37], and molgw. While molgw has been employed in several previous studies [18, 23, 38, 39, 40], it has never been fully described in a single text.

In this article, we present version 1 of the molgw code. This software is particularly dedicated to the study of excited state properties of finite systems with MBPT using a Gaussian basis. molgw is a self-contained code that is able to perform both the starting self-consistent field calculation in the generalized Kohn-Sham (gKS) framework and the MBPT post-treatment. The code can produce \( GW \) quasiparticle energies and optical spectra through solution of the BSE.

The article is organized as follows: in Section 2, we quickly review the theoretical foundation of the \( GW \) approximation and of the BSE; in Section 3, we detail the technical implementation; in Section 4, we describe the code organization; in Section 5, we present a few validation studies; and in Section 6, we exhibit the parallelization capabilities of molgw.

In this article, Hartree atomic units will be used in all equations; wavefunctions will be considered real and consequently all conjugations will be dropped; state indexes \( m \) and \( n \) run over all states, empty and occupied; indexes \( i \) and \( j \) are limited to occupied states; and indexes \( a \) and \( b \) are limited to empty states.

2. Theoretical background

In most cases, an MBPT calculation is performed as a post-processing step based on a previous self-consistent gKS run, as shown in Fig. 1. The optimal choice of gKS starting point, whether Hartree-Fock, a semi-local approximation, or a hybrid functional within density-functional theory (DFT), is not trivial for a given system and is currently an active area of research [41, 39, 42, 38, 43, 44]. In any case, the wavefunctions and eigenvalues that result from the gKS calculation are used to produce the screened Coulomb interaction \( W \) which is an ingredient in both the \( GW \) approximation to the self-energy and in the BSE. The \( GW \) self-energy is meant to produce accurate quasiparticle energies \( \epsilon_{q,i}^{W,\sigma} \) that can be compared to photoemission and inverse-photoemission spectroscopy, and these quasiparticle energies are in turn required input for the BSE solution. Finally, the solution to the BSE yields neutral excitation energies \( \Omega \), and oscillator strengths \( f \), that can be compared to optical absorption experiments.

Bearing in mind the different stages of the MBPT calculations, we are now prepared to introduce a few useful physical quantities. We do not intend to review all of

![Figure 1: Typical workflow of perturbative GW and BSE calculations.](image-url)
the developments necessary for a complete derivation of MBPT. Rather, we sketch a rapid overview of the minimal knowledge required for the implementation of the GW approximation to the self-energy and for the BSE solution. For a more complete understanding, we refer the reader to the review articles of Strinati [45], Aryasetiawan and Gunnarsson [8], and Onida et al. [46].

2.1. GW self-energy

The time-ordered Green’s function is the central object in MBPT. It is a compact representation of additional electrons (for positive times) or missing electrons (for negative times) in the system.

In the specific case of electrons in a mean-field potential, which can be described with one-electron wavefunctions $\varphi_{\sigma}^{m}(\mathbf{r})$ and one-electron energies $\epsilon_{\sigma}^{m}$, the one-particle Green’s function reduces to

$$G_{\sigma}(\mathbf{r}, \mathbf{r}', \omega) = \sum_{i} \frac{\varphi_{\sigma}^{m}(\mathbf{r}) \varphi_{\sigma}^{m}(\mathbf{r}')}{\omega - \epsilon_{\sigma}^{m} + i\eta},$$

where the index $i$ runs over the occupied one-electron states, the index $a$ runs over the empty one-electron states, $\sigma$ is the spin channel, and $\eta$ is a small positive real number.

With this Green’s function, one can build the non-interacting polarization $\chi_{0}$,

$$\chi_{0}(\mathbf{r}, \mathbf{r}', \omega) = -\frac{i}{2\pi} \sum_{\sigma} \int d\omega_{1} G_{\sigma}(\mathbf{r}, \mathbf{r}', \omega+\omega_{1}) G_{\sigma}(\mathbf{r}', \mathbf{r}, \omega_{1}),$$

and the dielectric function $\varepsilon$ within the random-phase approximation (RPA)

$$\varepsilon(\mathbf{r}, \mathbf{r}', \omega) = \delta(\mathbf{r} - \mathbf{r}') - \int d\mathbf{r} \nu(\mathbf{r} - \mathbf{r}_{1}) \chi_{0}(\mathbf{r}_{1}, \mathbf{r}, \omega).$$

The screened Coulomb interaction $W$ is then obtained as

$$W(\mathbf{r}, \mathbf{r}', \omega) = \int d\mathbf{r} \nu^{-1}(\mathbf{r}, \mathbf{r}_{1}, \omega) \nu(\mathbf{r}_{1} - \mathbf{r}).$$

The $GW$ approximation is the first-order expansion of the self-energy in $W$. It reads

$$\Sigma_{\sigma}(\mathbf{r}, \mathbf{r}', \omega) = \frac{i}{2\pi} \int d\omega_{1} e^{i\omega_{1}a} G_{\sigma}(\mathbf{r}, \mathbf{r}', \omega+\omega_{1}) W(\mathbf{r}', \mathbf{r}, \omega_{1}),$$

where the factor $e^{i\omega_{1}a}$ enforces the correct time-ordered form of the self-energy.

The $GW$ self-energy is most commonly evaluated within a perturbative approach, called $G_{0}W_{0}$ or alternatively “one-shot $GW$”. In such a case, the $GW$ quasiparticle energies $\epsilon_{n}^{GW\sigma}$ are obtained through

$$\epsilon_{n}^{GW\sigma} = \epsilon_{n}^{\sigma} + \langle \varphi_{n}^{\sigma} | \Sigma_{\sigma}^{GW} | \varphi_{n}^{\sigma} \rangle,$$

Note that the self-energy is evaluated at the quasiparticle energy $\epsilon_{n}^{GW\sigma}$, therefore Eq. (6) must be solved numerically.

Some authors [42, 43, 47] advocate for iterating further the $GW$ equations by updating the eigenvalues in Eq. (1) to those from Eq. (6), while keeping the wavefunction fixed to the mean-field result. This procedure, usually called “ev$GW$”, appears as simple and robust.

Alternatively, Faleev et al. [48, 49, 50] proposed that the Hermitian part of a static approximation to the $GW$ self-energy be retained to produce new orthogonal wavefunctions that could be iterated as well. This approach is referred to as “QSGW” for quasiparticle self-consistent $GW$.

Nowadays, some groups [51, 14, 21] are exploring the possibility of fully self-consistent $GW$ calculations. As $mol$ does not have this capability, we will not mention this any further.

2.2. Bethe-Salpeter equation

The BSE is an equation for a contraction of the two-particle Green’s function $L$ that permits the calculation of polarizabilities that readily include the electron-hole interaction.

The BSE is usually assuming a static approximation of the $GW$ self-energy, which simplifies the equation into

$$L_{0}^{\sigma\sigma'}(\mathbf{r}_{1}, \mathbf{r}_{2}; \mathbf{r}_{1}', \mathbf{r}_{2}'; \omega) = L_{0}^{\sigma\sigma'}(\mathbf{r}_{1}, \mathbf{r}_{2}; \mathbf{r}_{1}', \mathbf{r}_{2}'; \omega)$$

$$+ \int d\mathbf{r}_{3} d\mathbf{r}_{4} d\mathbf{r}_{5} d\mathbf{r}_{6} L_{0}^{\sigma\sigma'}(\mathbf{r}_{1}, \mathbf{r}_{4}; \mathbf{r}_{1}', \mathbf{r}_{5})$$

$$\times (\nu(\mathbf{r}_{3} - \mathbf{r}_{4}) \nu(\mathbf{r}_{5} - \mathbf{r}_{6})$$

$$- iW(\mathbf{r}_{1}, \mathbf{r}_{4}, \omega = 0) \delta(\mathbf{r}_{5} - \mathbf{r}_{3}) \delta(\mathbf{r}_{4} - \mathbf{r}_{6}) \delta_{\sigma\sigma'})$$

$$\times L_{0}^{\sigma\sigma'}(\mathbf{r}_{6}, \mathbf{r}_{2}; \mathbf{r}_{5}, \mathbf{r}_{2}'; \omega),$$

where $L_{0}$ stands for the non-interacting counterpart of $L$.

$$L_{0}^{\sigma\sigma'}(\mathbf{r}_{1}, \mathbf{r}_{2}; \mathbf{r}_{1}', \mathbf{r}_{2}'; \omega) = \frac{1}{2\pi} \int d\omega_{1} G_{\sigma}(\mathbf{r}_{1}, \mathbf{r}_{2}; \omega + \omega_{1})$$

$$\times G_{\sigma}(\mathbf{r}_{1}', \mathbf{r}_{2}'; \omega).$$

The electron-hole interaction arises from the statically screened Coulomb interaction $W(\mathbf{r}_{3}, \mathbf{r}_{4}, \omega = 0)$, which is often named the BSE kernel $\Xi$.

A contraction of the indexes of $L_{0}$ gives back the usual non-interacting polarizability introduced in Eq. (2):

$$\chi_{0}(\mathbf{r}_{1}, \mathbf{r}_{2}, \omega) = -\sum_{\sigma} L_{0}^{\sigma\sigma}(\mathbf{r}_{1}, \mathbf{r}_{2}; \mathbf{r}_{1}, \mathbf{r}_{2}; \omega).$$
The fully interacting polarizability is obtained with the same index contraction:
\[ \chi(r_1, r_2, \omega) = -i \sum_{\sigma} L^{\sigma e}(r_1, r_2; r_1, r_2; \omega). \] 

(10)

3. Technical implementation

In this section, we will describe how the general equations shown in the previous section translate when a finite basis set is introduced. The equations derived here closely match the notation and the forms used in the practical implementation of molgw and its source code.

3.1. Gaussian basis sets

In molgw, one-electron wavefunctions are expanded as linear combinations of atom-centered orbitals \( \phi_\mu \):
\[ \psi_{\sigma}(r) = \sum_\mu C_{\sigma \mu} \phi_\mu(r), \] 

(11)

For numerical ease, the radial components of the atom-centered orbitals are chosen to be linear combinations of Gaussian orbitals or “contracted Gaussians” [3]. The angular component can be either “Cartesian”
\[ \phi_\mu(r) = x^\mu_1 y^\mu_2 z^\mu_3 \sum_b C_{\mu b} e^{-\alpha_b r^2}, \] 

(12)
or “Pure”
\[ \phi_\mu(r) = Y_{\mu \lambda}(\hat{r}) r^\lambda \sum_b C_{\mu b} e^{-\alpha_b r^2}, \] 

(13)

where \( Y_{\mu \lambda}(\hat{r}) \) are the real spherical harmonics. The choice of the angular part type or of the coefficients is dictated by the basis set selection. The transformations from Cartesian to pure are tabulated, for instance, in Ref. [52].

3.2. Self-consistent field

In practice, the GW self-energy and the BSE solution require the knowledge of a mean-field starting point. With a gKS approach to obtain one-electron wavefunctions and energies, molgw solves the Roothan-Hall equation for spin restricted calculations and the Pople-Nesbet equations for those that are spin unrestricted [3]. This framework encompasses Hartree-Fock and semi-local approximations to DFT, as well as the hybrid functionals and range-separated hybrids.

The Roothan-Hall and Pople-Nesbet equations require a self-consistent solution to calculate the wavefunction coefficients \( C_{\sigma} \) introduced in Eq. (11) and the one-electron energies \( e_{\sigma} \) (which are subsequently stored in a diagonal matrix). These equations comprise the generalized diagonalization problem
\[ H_{\sigma} C_{\sigma} = S C_{\sigma} e_{\sigma}, \] 

(14)

where \( S \) is the basis function overlap matrix,
\[ S_{\mu \nu} = \int d\mathbf{r} \phi_\mu(\mathbf{r}) \phi_\nu(\mathbf{r}), \] 

(15)

and \( H_{\sigma} \) is the spin-dependent hamiltonian,
\[ H_{\mu \nu}^{\sigma} = T_{\mu \nu} + V_{\text{ext} \mu \nu} + J_{\mu \nu} - \alpha K_{\mu \nu}^{\sigma} - \beta K_{\mu \nu}^{\sigma} + (V_{\text{xc} \sigma})_{\mu \nu}. \] 

(16)

We show in Fig. 2 how the different components of the exchange are taken into account within the range-separated hybrid family of functionals, either calculated with the Fock exchange or with a DFT semi-local functional. The \( \gamma \) parameter (an inverse length) governs the crossover between short and long range. The value of the \( \alpha \) parameter is a measure of the amount of Fock exact-exchange. In Fig. 2, this fraction of the exchange is present at all ranges and it is graphically represented by the lower gray area. The \( \beta \) parameter dictates the proportion of long-range Fock exchange. In Fig. 2, because of positive \( \beta \), the amount of exact-exchange increases with interelectron distance. It is graphically represented by the central blue area. The DFT semi-local exchange contribution complements the Fock exchange.
so that a full account of the exchange is recovered at any interelectron separation. Therefore, the amount of DFT semi-local exchange decreases with interelectron distance for positive \( \beta \). The DFT exchange contribution consists of two parts: a standard exchange part, which is constant for any distance and which is represented by the upper red area in Fig. 2; and a short-range part, which precisely complements the long-range exact-exchange Fock contribution. This latter contribution is proportional to \( \beta \) and is represented by the central green area in Fig. 2. Note that the standard DFT exchange-correlation potential \( V_{\text{xc}} \) is a function of \( \alpha \), \( \beta \), and \( \gamma \), even though it is not explicitly noted.

Functional with no range separation, such as B3LYP [53] or PBE0 [54], can be obtained by setting \( \beta \) or \( \gamma \) to 0. The functionals with the correct asymptotics [55] enforce \( \alpha + \beta = 1 \). The short-range only functionals, such as HSE06 [56, 57], use \( \beta = -\alpha \). With this framework, we have then the full flexibility to account for most current hybrid functionals on the market.

Let us detail now the terms in the Hamiltonian in Eq. (16). It consists of a kinetic energy term

\[
T_{\mu\nu} = -\frac{1}{2} \int dr \lim_{r \to 0} \phi_{\mu}(r) \nabla_{\mu}^{2} \phi_{\nu}(r),
\]

an external potential energy term

\[
V_{\text{ext}} = -\sum_{\alpha} \int dr \phi_{\alpha}(r) \frac{Z_{\alpha}}{|r - R_{\alpha}|} \phi_{\alpha}(r),
\]

where \( Z_{\alpha} \) and \( R_{\alpha} \) are the charge and the position of the nuclei, and electron-electron interaction terms.

First, the Hartree term accounts for the classical electron-electron repulsion. It reads

\[
J_{\mu\nu} = \sum_{\lambda \tau} \langle \mu \nu | \lambda \tau \rangle \sum_{\sigma} P_{\lambda\sigma}^{T},
\]

where the density matrix \( P \) is introduced as

\[
P_{\mu\nu} = \sum_{n} f_{n}^{\sigma} C_{\mu n}^{\sigma} C_{\nu n}^{\sigma},
\]

with \( f_{n}^{\sigma} \) the occupation number, as well as the electron repulsion integrals in Mulliken notation:

\[
\langle \mu \nu | \lambda \tau \rangle = \int dr dr' \phi_{\mu}(r) \phi_{\nu}(r) \frac{1}{|r - r'|} \phi_{\lambda}(r') \phi_{\tau}(r').
\]

In order to account for hybrid functionals (and range-separated hybrids), we have explicitly written the all-range exchange matrix, rescaled with a factor \( \alpha \) in Eq. (16),

\[
K_{\mu\nu}^{x} = \sum_{\lambda \tau} P_{\lambda\tau}^{x}(\mu \lambda | r v) \frac{\epsilon_{\tau}(r)}{1 + \gamma_{\tau\sigma} r^{-\gamma}}
\]

and the long-range only exchange matrix, rescaled with a factor \( \beta \) in Eq. (16),

\[
K_{\mu\nu}^{y} = \sum_{\lambda \tau} P_{\lambda\tau}^{y}(\mu \lambda | r v) \frac{\epsilon_{\tau}(r)}{1 + \gamma_{\tau\sigma} r^{-\gamma}}
\]

where the long-range electron repulsion integrals are defined as

\[
(\mu \nu | \lambda \tau) = \int dr dr' \phi_{\mu}(r) \phi_{\nu}(r) \frac{\epsilon_{\tau}(r)}{1 + \gamma_{\tau\sigma} r^{-\gamma}} \phi_{\lambda}(r') \phi_{\tau}(r').
\]

In practice, the electron repulsion integrals are evaluated using recursion formulas [52] as implemented in the \textsc{libxc} library [53]. The range-separated integrals are obtained simply by changing the initial term of the recursion [56].

A priori integral screening with the Schwarz inequality

\[
(\mu \nu | \lambda \tau) \leq \sqrt{\mu \nu} \sqrt{\lambda \tau} \sqrt{\lambda \tau}
\]

can drastically reduce the number of 4-center integrals to be calculated [60], and we make use of this here.

The density-functional exchange-correlation potential \( V_{\text{xc}} \) is then evaluated with the expression [61]

\[
(V_{\text{xc}}^{\mu\nu})_{\mu\nu} = \langle \mu | V_{\text{xc}}(\rho^{\mu\nu}(r), \nabla_{\tau} \rho^{\tau\gamma}(r)) | \nu \rangle
\]

\[
= \int dr \left( \frac{\partial \epsilon_{\text{xc}}}{\partial \rho^{\mu\tau}(r)} \phi_{\mu}(r) \phi_{\nu}(r) + \frac{2}{\partial \gamma_{\tau\gamma}} \frac{\partial \epsilon_{\text{xc}}}{\partial \gamma_{\tau\gamma}} \nabla_{\tau} \rho^{\tau\gamma}(r) \right)
\]

\[
\times \nabla_{\tau} (\phi_{\mu}(r) \phi_{\nu}(r)),
\]

where \( \epsilon_{\text{xc}} \) is the exchange-correlation energy density (per unit volume), and where

\[
\gamma_{\tau\sigma} = \nabla_{\tau} \rho^{\tau\sigma}(r) \cdot \nabla_{\tau} \rho^{\tau\sigma}(r).
\]

The corresponding expression for \( V_{\text{xc}}^{\mu\nu} \) is analogous. In \textsc{molgw}, the values of \( \epsilon_{\text{xc}} \) and its derivatives are obtained from the external library \textsc{libxc} [62].

The density and the density gradient are obtained from the density matrix as

\[
\rho^{\mu\tau}(r) = \sum_{\nu} P_{\nu\tau}^{\mu}(\mu \nu | r \nu) \phi_{\nu}(r)
\]

\[
\nabla \rho^{\mu\tau}(r) = \sum_{\nu} P_{\nu\tau}^{\mu} \nabla_{\nu} (\phi_{\nu}(r) \phi_{\nu}(r)).
\]
As is standard in quantum chemistry, the integration in Eq. (27) for the evaluation of the matrix elements \( (V^\alpha_{\sigma})_{ij} \) is performed by numerical quadrature. First, the space is partitioned in smooth regions around each atom using Becke [63] or SSF schemes [64]. Then, each atom carries its own quadrature points in spherical coordinates combining an angular Lebedev-Laikov mesh [65] and a radial mesh [66].

Finally, the Roothan-Hall or Pople-Nesbet equations should be solved self-consistently because of the dependence of the terms \( J, K, K', V_{xc} \) on the density matrix. The method due to Pulay, extremely efficient [67, 68] and rather robust, is used here.

### 3.3. Auxiliary basis set

In order to reduce the computational and the memory costs associated with the 4-center integrals \( \langle \mu \nu | I \tau \rangle \), the introduction of an auxiliary basis set to represent the orbital products \( \phi_i(r) \phi_j(r) \) has been proposed [69, 70, 71]. This method, known as the resolution-of-the-identity method, allows one to approximate the 4-center integrals in terms of 2- and 3-center integrals, as

\[
(\mu \nu | I \tau) \approx \sum Q (\mu \nu | P) (P | Q)_{-1} (Q | I \tau)
\]  
(31)

where the 3-center integrals read

\[
(\mu \nu | P) = \int d \mathbf{r} d \mathbf{r}' \phi_i(r) \phi_j(r') \frac{1}{|r - r'|} \phi_k(r')
\]  
(32)

and the 2-center integrals are

\[
(P | Q) = \int d \mathbf{r} d \mathbf{r}' \phi_i(r) \frac{1}{|r - r'|} \phi_j(r').
\]  
(33)

The index running over the auxiliary basis are labeled with capital letters \( P, Q, \) etc.

The quality of the approximation in Eq. (31) depends on the quality of the auxiliary basis set. The basis sets proposed by Weigend [71] permits an accurate expansion of the orbital products that are used in both the Hartree term \( J \) and in the Fock exchange term \( K \). This method is called “RI-JK”.

In practice, the 2-center integral matrix \( (P | Q) \) is positive definite and its square-root can be calculated so as to transform Eq. (31) into

\[
(\mu \nu | I \tau) \approx \sum P M_{\mu \nu}^P M_{I \tau}^P
\]  
(34)

where

\[
M_{\mu \nu}^P = \sum_Q (\mu \nu | Q) (Q | P)_{-1/2}.
\]  
(35)

This is computationally simpler to evaluate \( (N \) operations instead of \( N^2 \)).

In \textsc{molgw}, one can choose whether or not to evaluate the Hartree term \( J \) and the Fock term \( K \) with the resolution-of-the-identity (RI) approximation.

### 3.4. Polarizability in the state product basis

Both the \( GW \) calculations and the BSE require the calculation of the polarizability. In standard \( GW \) calculations, it is obtained within the random phase approximation (RPA), whereas in the BSE, one needs to introduce the additional kernel \( \Xi \) described in Section 2.2. In \textsc{molgw}, whatever the approximation, the calculation of the polarizability follows the same procedure. The advantage of this is that the spectral decomposition of the polarizability may be obtained so that its evaluation at any frequency is exact and, as a consequence, the \( GW \) self-energy convolution can be calculated analytically.

The BSE equation in Eq. (7) can be written in terms of a product basis of occupied to unoccupied states (and the converse). This transformation is precisely the same one frequently used to represent time-dependent Hartree-Fock (TD-HF) and time-dependent density-functional theory (TD-DFT) within linear response [72, 73, 74]. Due to the symmetry between resonant (occupied to empty) and anti-resonant (empty to occupied) transitions, these equations have a very specific block form:

\[
\begin{pmatrix}
A & B \\
-\overline{B} & -\overline{A}
\end{pmatrix}
\begin{pmatrix}
X_i \\
Y_i
\end{pmatrix} = \Omega_i
\]  
(36)

where \( \Omega_i \) are the neutral excitation energies and \( \{X_i, Y_i\} \) are the corresponding eigenvectors.

The only difference between RPA, TD-HF, TD-DFT and BSE lies in the specific expression of the matrix elements in \( A \) and \( B \). Let \( i \) and \( j \) indexes stand for occupied states and \( a \) and \( b \) for unoccupied states. The expressions for these elements read

\[
A_{\text{RPA}}^{\text{bor}} = (\epsilon_i - \epsilon_r) \delta_{ij} \delta_{ab} \delta_{\sigma' \sigma''}
\]
\[
-(i \sigma r | j \sigma' r') + \Xi_{\text{bor}}^{\text{bor}}
\]  
(37a)

\[
B_{\text{RPA}}^{\text{bor}} = -(i \sigma r | j \sigma' r') + \Xi_{\text{bor}}^{\text{bor}}
\]  
(37b)

where the operator \( \Xi \) stands for the kernel of the selected approximation. In the specific case of RPA, \( \Xi = 0 \). The expression of \( \Xi \) in BSE will be addressed further below.

The molecular orbital electron repulsion integrals are defined as

\[
(i \sigma r | j \sigma' r') = \int d \mathbf{r} d \mathbf{r}' \phi_i^\sigma(r) \phi_j^{\sigma'}(r') \frac{1}{|r - r'|} \phi_j^{\sigma'}(r) \phi_i^\sigma(r')
\]  
(38)
and are obtained by the canonical transform

$$(ia\sigma|j\sigma r') = \sum_{\mu\nu} C_{\mu i}^{*} C_{\nu a}^{*} C_{\nu b}^{*} C_{\mu j}(\nu|\mu r).$$

(39)

When the resolution-of-the-identity approximation is used, the scaling of the transform above can be improved:

$$(ia\sigma|j\sigma r') \approx \sum_{\mu} \left( \sum_{\nu} C_{\mu i}^{*} C_{\nu a}^{*} M_{\mu}^{p} \right) \times \left( \sum_{\nu} C_{\nu b}^{*} C_{\nu b}^{*} M_{\nu}^{p} \right).$$

(40)

Due to the peculiar block structure of Eq. (36), which induces many symmetries, it can be recast into a smaller matrix diagonalization, after some algebra [73]:

$$CZ' = Z'\Omega^{2},$$

(41)

where $C = (A - B)^{1/2}(A + B)(A - B)^{-1/2}$ is a symmetric matrix that is half the size of the initial problem in Eq. (36). The above expression assumes matrix $(A - B)$ to be positive definite. From the knowledge of an eigenvector $Z'$, one can recover both $X'$ and $Y'$ as

$$X' = \frac{1}{2} \left[ \Omega_{s}^{-1/2}(A - B)^{1/2} + \Omega_{s}^{1/2}(A - B)^{-1/2} \right] Z',$$

(42a)

$$Y' = \frac{1}{2} \left[ \Omega_{s}^{-1/2}(A - B)^{1/2} - \Omega_{s}^{1/2}(A - B)^{-1/2} \right] Z'.

(42b)

The equations for $X'$ and $Y'$ imply normalization conditions that can be recast into a matrix form:

$$\begin{pmatrix} X & -Y' \\ -Y & X \end{pmatrix} \begin{pmatrix} X & Y \\ Y & X \end{pmatrix} = I,$$

(43)

where the left-hand matrix contains the left eigenvectors and the right-hand matrix contains the right eigenvectors of Eq. (36). A more efficient way of solving Eq. (36) that avoids the computation of the matrix square-root will be discussed in Section 6.2.

Once the eigenvalues $\Omega_{s}$ and the eigenvectors $(X', Y')$ have been obtained through the diagonalization of Eq. (41) and the matrix multiplications in Eqs. (42a-42b), the spectral representation of the polarizability follows thanks to Eq. (10). Then, using the definition of the screened Coulomb interaction in Eq. (4), $W$ reads

$$W^{s}_{mnra}(\omega) = (mn\sigma|\sigma r') + \sum_{s} w^{s}_{mnr} w^{\sigma r'} + \sum_{s} w^{s}_{mnra} \left( \frac{1}{\omega - \Omega_{s} - i\eta} - \frac{1}{\omega + \Omega_{s} + i\eta} \right),$$

(44)

where the residues $w^{s}_{mnr}$ are defined as

$$w^{s}_{mnr} = \sum_{s} (mn\sigma|\sigma r')(X'_{mnr} + Y'_{mnr}).$$

(45)

Note that once again the resonant/antiresonant symmetry has been used in Eq. (44) to produce a more compact expression.

3.5. Many-body total energies

The GW approximation has been mentioned so far with the aim of obtaining self- and quasiparticle energies, but it can also be used to calculate total energies [5, 75, 12, 21].

There are several expressions for the GW total energy that all match for a self-consistent solution of the Green’s function $G$. However, they differ when evaluated with a perturbative $G$ obtained from a gKS starting point [12, 76]. One of these expressions, the Klein functional [77], is particularly interesting, since it yields a stationary expression for the total energy and it matches the RPA total energy obtained in the DFT framework [78, 79, 80].

In practice, when calculating the Klein functional from a gKS potential, Furch [81] proposed a very compact formula that avoids the numerical integration over the coupling constant and over frequencies. The expression only involves the results already calculated for the polarizability anyway:

$$E_{c}^{RPA} = -\frac{1}{2} \sum_{s} \Omega_{s} + \frac{1}{2} \sum_{s} \epsilon_{s} - \epsilon_{i} - (ia\sigma|j\sigma r).$$

(46)

This last equation is nothing more than the difference between the eigenvalues and the diagonal terms of Eq. (36).

3.6. GW self-energy

With the spectral decomposition of $W$ at hand, and using the residue theorem, it is possible to give an analytical expression for the GW self-energy convolution in Eq. (5).

The first term on the right-hand side of Eq. (44) gives the usual Fock exchange term. The rest yields the correlation part of the self-energy $\Sigma$, whose diagonal matrix elements read

$$\langle \phi_{i}^{\sigma}|\Sigma_{i}^{\sigma}(\omega)|\phi_{j}^{\sigma}\rangle = \sum_{s} \frac{w^{s}_{mnr} w^{s}_{mnr} \left( \omega - \epsilon_{s} + \Omega_{s} - i\eta \right)}{\omega - \epsilon_{s} - \Omega_{s} + i\eta} + \sum_{as} \frac{w^{s}_{mnr} w^{s}_{mnr} \left( \omega - \epsilon_{s} + \Omega_{s} + i\eta \right)}{\omega - \epsilon_{s} + \Omega_{s} + i\eta}.$$
and the graphical solution to the quasiparticle equation.

When the poles of the self-energy are distant from the
\( \epsilon \) and hence, the \( GW \) reduces to
\[ \Sigma_{\omega} = \Sigma_{\omega}^{\text{linear}} \]

The first term in Eq. (48) is the pure screened part of
\[ \langle \phi_i^\sigma | \Sigma^{\sigma} | \phi_a^\alpha \rangle = 2 \sum_{ij} \frac{w_{ia}}{\Omega_s} \frac{w_{aj}}{\Omega_s} \sum_{nm} \frac{w_{nm}^{\sigma} w_{mn}^{\alpha}}{\Omega_{ij}}. \]

The symbol \( \tilde{x} \) is the position operator along the \( x \) direction in space.

The oscillator strengths should add up to the number of electrons in the system \( N_e \) according to the Thomas-
Reiche-Kuhn sum rule [73]
\[ \sum_{ia} (f^\prime)^2 \Omega_s = N_e. \]

However this relation is fulfilled only in the limit of a complete basis set.

Finally, we would like to mention that the so-called Tamam-Danoff approximation can be used within BSE. This approximation consists in setting the \( B \) block to zero in Eq. (36). The BSE for a spin-triplet final state can be evaluated as shown, for instance in Ref. [38].
4. Code description

molgw is being developed mainly in the Fortran 2003 language. In general, molgw prioritizes simplicity of implementation to facilitate new developments. As a consequence, symmetry operations are not used. All the Coulomb integrals are calculated once and then stored in memory (a so-called in-core algorithm). molgw’s reliance on several external libraries eases the code’s maintenance. 2-, 3-, and 4-center Coulomb integrals are all obtained from the library libint [59], while the exchange-correlation energy density, potential, and kernel are calculated with the library libxc [62].

molgw is parallelized using Message Passing Interface (MPI) for the communications and ScaLAPACK for the linear algebra.

The accuracy of molgw for the self-consistent field has been tested against existing quantum chemistry codes with total energies and one-electron energies that agree within 10^{-6} Ha. Recently, we have also tested the accuracy against the fiesta code for the GW quasiparticle energies [87]. The agreement is better than 1 meV.

Regression tests are present that certify about 150 critical check points using 30 different input files.

Many basis sets are available with molgw. However, new ones can be obtained, for instance from the EMSL basis set exchange website [88, 89, 90]. A script is available in molgw that transforms the basis sets in nwchem [91] format into the internal format of molgw.

5. Accuracy tests

5.1. Auxiliary basis set convergence

Whereas the RI approximation has been assessed for MP2 calculations [93], its quality for GW calculations is still rather unclear. While the RI for GW has been used by several groups [94, 15, 41], molgw offers a unique opportunity to test this approximation precisely, since it is capable of turning it on or off.

As the GW calculations without RI quickly become unfeasible, we have tested the approximation on a small molecule, namely silane SiH₄. Silane is a relevant test example since the natures of the HOMO and LUMO differ significantly: the LUMO is unbound. In Table 1, we provide the results for the Dunning family of basis sets with and without diffuse functions [95, 96] in conjunction with the auxiliary basis set developed by Weigend and coworkers [92]. The error induced by the RI approximation appears as insensitive to the state nature (bound or unbound) as to the presence of diffuse function (augmented or not). The error is at most 11 meV, which is excellent compared to the other sources of error, such as the wavefunction basis set. We note that the RI error seems to decrease when going to more complete basis sets.

In our experience, the accuracy of the RI approximation shown here is typical of other small molecules. We conclude that the RI approximation can be safely used for production. It is used in all further shown calculations.

5.2. Basis set convergence

In several previous works [18, 39], we have already illustrated slow convergence of the GW quasiparticle energies as a function of the basis set. We have shown that the slow convergence is not unexpected for a correlated method and the GW convergence rate is not very different between MP2 and the coupled-cluster method CCSD(T).

Figure 4 shows a typical example of convergence for the famous-as-an-electron-acceptor tetracyanoethylene molecule (TCNE). We use the Dunning basis set families (cc-pVnZ and aug-cc-pVnZ) [95, 96] that are precisely designed to smoothly reach the complete basis set. The convergence is indeed smooth, but very slow. We propose an extrapolating function with 2 parameters \( a \) and \( \epsilon_m^\infty \) for the quasiparticle energy as a function of the basis set,

\[
\epsilon_m(l_{\text{max}}) = \epsilon_m^\infty + \frac{a}{l_{\text{max}}^2},
\]

Figure 4: Basis set convergence of the LUMO energy of tetracyanoethylene (TCNE) with the Dunning basis sets without diffuse functions (blue squares) or with diffuse functions (red circles). The continuous curves are a fit to the calculated data with the function given in Eq. (54).
Table 1: HOMO and LUMO quasiparticle energy within GW@PBE0 for silane SiH₄ for different Dunning basis sets using or not the RI approximation in eV. The difference is given in meV. We use here the auxiliary basis sets developed Weigend [92].

<table>
<thead>
<tr>
<th></th>
<th>cc-pVDZ</th>
<th>cc-pVTZ</th>
<th>cc-pVQZ</th>
<th>aug-cc-pVDZ</th>
<th>aug-cc-pVTZ</th>
<th>aug-cc-pVQZ</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOMO no RI (eV)</td>
<td>-12.3566</td>
<td>-12.6027</td>
<td>-12.6879</td>
<td>-12.3426</td>
<td>-12.6018</td>
<td>-12.6958</td>
</tr>
<tr>
<td>RI (eV)</td>
<td>-12.3553</td>
<td>-12.6012</td>
<td>-12.6871</td>
<td>-12.3407</td>
<td>-12.6002</td>
<td>-12.6952</td>
</tr>
<tr>
<td>Diff. (meV)</td>
<td>1.3</td>
<td>1.5</td>
<td>0.8</td>
<td>1.9</td>
<td>1.6</td>
<td>0.6</td>
</tr>
<tr>
<td>LUMO no RI (eV)</td>
<td>3.9687</td>
<td>3.2614</td>
<td>2.6863</td>
<td>0.9958</td>
<td>0.8431</td>
<td>0.7516</td>
</tr>
<tr>
<td>RI (eV)</td>
<td>3.9800</td>
<td>3.2578</td>
<td>2.6864</td>
<td>0.9999</td>
<td>0.8420</td>
<td>0.7516</td>
</tr>
<tr>
<td>Diff. (meV)</td>
<td>11.3</td>
<td>-3.6</td>
<td>0.0</td>
<td>4.1</td>
<td>-1.1</td>
<td>-0.1</td>
</tr>
</tbody>
</table>

where \( l_{\text{max}} \) is the highest angular momentum in the basis set (cc-pVDZ has \( l_{\text{max}} = 2 \), cc-pVTZ has \( l_{\text{max}} = 3 \), etc.).

The form of the extrapolating function has been empirically selected. The squared exponent in the denominator yields the best match with the explicitly-calculated points. Note that Truhlar [97] obtained a rather close exponent of 2.2 for the MP2 correlation energies. With this function, we have obtained the complete basis set value and hence we can define the basis set error. The error is as large as 1.1 eV for cc-pVDZ and even the most complete basis set available, i.e. aug-cc-pV5Z, still experiences a 0.15 eV deviation.

From all the examples we have calculated so far, the convergence of the quasiparticle energies as a function of the basis set size is always from above. As the convergence is indeed very slow, we advocate for the systematic use of an extrapolation technique when comparison to experiment is made.

We also provide the example of the convergence of an optical spectrum in Fig. 5. It shows the photoabsorption cross section of pyrene, \( \text{C}_{16}\text{H}_{10} \), for different basis sets with and without diffuse functions. The whole calculation workflow (PBE0, then GW, then BSE) uses the same basis set. In contrast with the previous statement for \( \text{GW} \), here the optical excitations converge smoothly with the basis set, especially for augmented basis sets. This demonstrates that the slow convergence of the quasiparticle energies observed above compensates, so that their differences converge more quickly. For instance, the first bright excitation at 3.42 eV is evaluated with an accuracy of 50 meV already with the aug-cc-pDZ basis. Just the higher energy lineshape around 7-9 eV seems to be more sensitive to the basis set.

In conclusion, the convergence of the optical spectrum can in cases be achieved with unconverged absolute quasiparticle energies.

![Figure 5: Basis set convergence of the photoabsorption cross section \( \sigma \) of pyrene, \( \text{C}_{16}\text{H}_{10} \), within BSE with with the Dunning basis sets without diffuse functions (upper panel) or with diffuse functions (lower panel). This calculation is based on PBE0.](image-url)
Figure 6: Convergence of the GW@PBE0 HOMO and LUMO quasiparticle energies as a function of the number of states explicitly used in the calculation of the RPA polarizability.

5.3. Single Pole Approximation

The strategy retained to evaluate the polarizability in molgw possesses the unique feature of being exact in the chosen basis set. Its drawback is that it scales poorly with system size. Indeed, the transition space is spanned by the product of orbitals $|ia\sigma\rangle$, where $i$ is occupied and $a$ is empty. This space scales as $N^2$, if $N$ is a measure of the system size (for instance the number of electrons). Then the matrix $C$ to be diagonalized in Eq. (41) occupies memory as $N^4$ and the diagonalization time itself scales as $N^6$. For large systems this can indeed be problematic. Consider buckminster fullerene C60 in the cc-pVTZ basis set. There are 1800 basis functions and, among them, 120 constitute the occupied states (considering that the C 1s are frozen). The $C$ matrix is then a 194400x194400 real symmetric matrix that requires 281 Tb of storage. It is then necessary to have a procedure to reduce the size of the matrix while keeping reasonable accuracy.

A simple truncation of the number of empty states in the state product basis $|ia\sigma\rangle$ is not much help. We exemplify this statement with the convergence behavior of the HOMO and LUMO GW@PBE0 quasiparticle energies as shown in Fig. 6. When the product basis is limited to the empty states below a given state index $N_{\text{states}}$, both the HOMO and LUMO converge very slowly to the value when all the empty states are included ($N_{\text{states}} = 1800$). $N_{\text{states}} = 300$ yields quasiparticle energies that deviate by 0.7 eV from the converged values within this basis set.

In the recent past, several techniques have been designed to account for the states that are neglected in the context of plane-wave approaches [13, 98, 99]. We propose to introduce the single pole approximation (SPA) [73] for the high energy states to reduce the $C$ matrix dimension. In this approximation, the RPA matrix elements when $a > N_{\text{states}}$ are considered diagonal:

$$A_{ib\sigma'}^{a\sigma} \approx \delta_{ij} \delta_{ab} \delta_{\sigma\sigma'} \left[ \epsilon_{ib} - \epsilon_{ia} - \langle i\sigma|a\sigma'\rangle \alpha_{\text{SPA}} \right]$$

$$B_{ib\sigma'}^{a\sigma} \approx -\delta_{ij} \delta_{ab} \delta_{\sigma\sigma'} \langle i\sigma|a\sigma'\rangle \alpha_{\text{SPA}}.$$  

(55a) (55b)

Thus, the $C$ matrix has a dense square block up to $a = N_{\text{states}}$ and is diagonal above. Then, the diagonalization needs only to be done for the dense block.

The performance of this procedure is shown with the green line in Fig. 6. The approximation clearly overshoots the targeted value for both the HOMO and the LUMO. As a consequence, we propose that the diagonal terms above $N_{\text{states}}$ be multiplied by an empirical parameter $\alpha_{\text{SPA}} = 1.5$. This empirical factor does not contain much physical meaning and it is just meant to reduce the overshooting of the SPA. However, it yields a substantial accuracy gain in the case of the C60 molecule, as can be observed from the red line in Fig. 6. Now an accuracy of 0.05 eV can be reached for both the HOMO and the LUMO with $N_{\text{states}} = 600$, therefore reducing the matrix dimension by a factor of 4 and hence dividing the computational time by 64! This parameter $\alpha_{\text{SPA}}$ is meant to be tuned, however the values ranging from 1.0 to 1.5 are quite reasonable across all the molecules we have calculated so far.

The use of the SPA allows one to significantly reduce computational time and memory consumption by changing the scaling prefactor, however keeping an overall $N^6$ scaling.

6. Performance and parallelization

To access the GW quasiparticle energies and the BSE optical spectra of larger molecules, the parallelization of the code becomes unavoidable. Since the medium to large molecules can be calculated within a reasonable
time only when the RI approximation is used, the parallelization effort has been focusing on the RI calculations in molgw. Two types of operations benefit from the parallelization: the construction of the operators, such as the Hamiltonian in Eq. (16) or the polarizability matrix in Eqs. (37a), (37b), and the large eigenvalue problem described in Eq. (36).

### 6.1. Parallelization for the construction of the operators

As the parallelization is only meant to be used in conjunction with the RI approximation, the parallelization of the data and of the computation tasks most often relies on the distribution over the auxiliary basis functions $P$. This choice is validated by the observation that the number of basis functions $P$ in the auxiliary basis is in general much larger than the number of functions in the wave function basis $\mu$. Furthermore, distributing over the auxiliary basis functions allows us to calculate many operators with only a few changes from the serial version of the code.

Let us exemplify the parallelization with the calculation of the exchange operator $K$. The auxiliary basis functions $P$ are divided among the processors. Except for a subset of $P$, each processor owns in memory the 3-center Coulomb integral $M^P_{\mu\nu}$ for all $\mu$ and $\nu$. Upon introduction of the RI approximation [See Eq. (31)], and the square root of the density matrix $P^{1/2}$, Eq. (22) can be cast into

$$K^{\sigma\tau}_{\mu\nu} = \sum_P \sum_{\kappa} \left( \sum_A M^P_{\mu\lambda} P^{1/2}_{\lambda\kappa} \right) \left( \sum_{\tau} M^P_{\tau\kappa} P^{1/2}_{\kappa\nu} \right).$$  \hspace{1cm} (56)

where $\kappa$ only runs over the non-vanishing eigenvalues of $P$, i.e. the occupied states. As can be observed from Eq. (56), the sum over $P$ is only performed at the end of the calculation of $K^{\sigma\tau}_{\mu\nu}$. Therefore, each processor can calculate its workload independently and only at the end, an overall reduction is carried out.

Another example is that of the well-known atomic orbital to molecular orbital transforms that are necessary to evaluate the integrals ($ia\sigma$/$jbc\tau$) in Eqs. (37a), (37b). In this case, the parallelization is completely independent. Each processor performs the transforms

$$M^P_{i\alpha\sigma} = \sum_{\mu\nu} C^\sigma_{\mu\nu} C^{ia}_{\nu\alpha} M^P_{\mu\nu}$$ \hspace{1cm} (57)

for its subset of auxiliary basis functions $P$. Then the integrals ($ia\sigma$/$jbc\tau$) are calculated following the same spirit as in Eq. (56).

In Fig. 7, we show the performance of the parallelization of molgw for the gKS part for a medium molecule with almost 100 atoms, namely $\beta$-Carotene $\beta$-Carotene using the cc-pVTZ basis set has 1984 basis functions. The corresponding auxiliary basis has 4920 functions. The speed-up has been calculated with respect to the smallest core number that fits in memory, i.e. 16 cores. The scaling of the computational time is rather good as long as the number of auxiliary basis functions per core is not too low, as demonstrated in the upper panel of Fig. 7. A lower limit of 30-50 auxiliary basis functions per core is reasonable. It should be stressed that anyway, the computational time is not large: even with just 16 cores, the calculation lasts only roughly 3 hours.

Concerning the memory consumption shown in the lower panel of Fig. 7, the parallelization of the data is efficient. This is not surprising since, for the in-core algorithm used in molgw, most of the memory is used to store the 3-center Coulomb integrals $M^P_{\mu\nu}$ that are perfectly distributed among the cores.

![Figure 7: Parallelization performance of the gKS part of molgw for the $\beta$-Caroten molecule C30H56 within the rather complete basis set cc-pVTZ. The upper panel represents the computational time speed-up as a function of the number of cores. The reference for speed-up is taken for 16 cores. The lower panel shows the memory consumption per core. The actual results (blue circles) are compared to the ideal parallelization (black line).](image)
In general, the bottleneck of the calculation is not the gKS step. Rather, it is the calculation of the polarizability, either to prepare the GW self-energy evaluation or to evaluate the optical spectrum within BSE.

6.2. Diagonalizing the BSE Hamiltonian in parallel

Within the semi-local approximations to TD-DFT and within RPA, the matrix $(A - B)$ entering the expression of $C$ in Eq. (41) is purely diagonal and therefore its square-root can straightforwardly be calculated.

On the contrary, for hybrid functionals or for BSE, the matrix $(A - B)$ is dense and the direct application of Eq. (41) would require two diagonalizations. To avoid this, we use the Cholesky decomposition to replace the matrix square root. Our approach is summarized as follows [100]:

1. Compute the Cholesky factorization $A + B = LL^T$.
2. Form $W = L^T(A - B)L$.
3. Diagonalize $W = ZΩZ^T$.
4. Set $X = (L^T ZΩ^{1/2} + LZΩ^{1/2})/2$.
5. Set $Y = (L^T ZΩ^{1/2} - LZΩ^{1/2})/2$.

All of these steps make use of existing functionalities from the ScaLAPACK software package [101]. The eigenvectors are normalized as $X^TX - Y^TY = I$ so that both right and left eigenvectors can be readily used without further scaling.

Figure 8 exemplifies the performance of the BSE solver in molgw with an oligomer, namely polyacetylene $C_{2n}H_{2n+2}$. The scaling with system size is made apparent. The number of transitions from occupied to empty states scales as $n^2$. The BSE matrix is $n^2 \times n^2$. The time necessary to construct it (upper panel) and the memory necessary to store it (lower panel) indeed scale as $n^4$. Then the structure preserving algorithm presented above is dominated by the diagonalization in its step 3. This operation scales as the cube of the matrix dimension, which yields an overall scaling of $n^6$. The measured slopes in Fig. 8 match rather well with the scalings we have just anticipated.

As the calculation complexity increases quickly with the size of the problem, parallelization of the code is highly necessary. molgw uses a ScaLAPACK distribution of the BSE matrix to distribute the memory and workload for the diagonalization. It can be observed from the lower panel of Fig. 8 that by increasing the number of cores the memory consumption can be kept below 2 Gb per core, which is a reasonable target for modern computers. The middle panel shows how well the structure preserving algorithm behaves as a function of the number of cores. The construction of the BSE

Figure 8: Numerical performance of the BSE solution in molgw for polyacetylene oligomer $C_{2n}H_{2n+2}$ within the aug-cc-pVDZ basis with increasing length $n$. The upper panel shows the wall clock time necessary to build the BSE. The middle panel shows the wall clock time used in the structure preserving solution of the BSE. The lower panel shows the peak memory used in a BSE run. The thin dotted lines highlight the anticipated scaling of the computational load. The calculations have been performed with a number of cores ranging from 1 to 144.
matrix in the upper panel is more problematic for the parallelization. Indeed, as in the rest of the code, the calculation of the BSE matrix elements relies on the distribution of the auxiliary basis. Then this distribution of the data must be transposed into the ScaLAPACK distribution, which induces some intensive communications.

The performance as a function of the number of cores is perhaps better appreciated from Fig. 9. For this small example, we have managed to multiply the number of cores up to the parallelization breakdown. The BSE matrix building experiences an efficiency collapse beyond 64 cores, whereas the structure preserving diagonalization based on ScaLAPACK allows one to reach 144 cores without breakdown. It is remarkable that the speed-up is rather close to the ideal one over two orders of magnitude of core counts.

Of course, molgw offers the possibility to limit the BSE matrix size by removing the highest empty states from the transition basis. This approximation should be tested as an extra convergence parameter.

7. Summary

In this paper, we have presented version 1 of the molgw code. This code is dedicated to MBPT calculations in the GW approximation for atoms, molecules, and clusters. The focus is to calculate i) the GW self-energy to obtain accurate quasiparticle energies that may be compared to photoemission spectroscopy and ii) the solutions to the BSE, to evaluate precise optical spectra that include the electron-hole interaction. molgw relies on Gaussian type orbitals and therefore make use of all of the past developments in quantum chemistry.

We have shown the most central equations used in practice in molgw, starting from the self-consistent field within gKS, to the GW self-energy and BSE solution. molgw is a unique code, since it implements the equations with or without the RI approximation. It also has the advantage of calculating directly the spectral decomposition of the screened Coulomb interaction \( W \) and hence is able to calculate the GW self-energy at any frequency \( \omega \) with no further technical approximation. Incidentally, the RPA total energy is obtained for free. molgw employs a recently developed structure preserving parallel algorithm to perform the diagonalization of the BSE Hamiltonian.

This paper was also an opportunity to demonstrate the validity of the RI by directly comparing calculations with and without this approximation. Furthermore, we introduced a technique, based on the single pole approximation, to speed-up the GW calculations and save memory.

In conclusion, we stress that molgw version 1 is enclosed with this paper. However the forthcoming updates will be directly and freely obtained on a versioning server hosted on the web [102].

Acknowledgement

F.B. acknowledges the Enhanced Eurotalent program and the France Berkeley Fund for supporting his sabbatical leave in UC Berkeley. This work is supported by the U.S. Department of Energy, Office of Basic Energy Sciences and of Advanced Scientific Computing Research through the SciDAC Program on Excited State Phenomena. S.M.H. is supported by the Chemical Sciences, Geosciences, and Biosciences Division in Office of Basic Energy Sciences of the U.S. Department of Energy. Portions of this work took place at the Molecular Foundry, supported by the U.S. Department of Energy, Office of Basic Energy Sciences. This work was performed using HPC resources from GENCI-CCRT-TGCC (Grants No. 2015-096018).
