CONVERGENCE ANALYSIS OF DIRECT MINIMIZATION AND SELF-CONSISTENT ITERATIONS*

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Abstract. This article is concerned with the numerical solution of subspace optimization problems, consisting of minimizing a smooth functional over the set of orthogonal projectors of fixed rank. Such problems are encountered in particular in electronic structure calculation (Hartree–Fock and Kohn–Sham density functional theory models). We compare from a numerical analysis perspective two simple representatives, the damped self-consistent field (SCF) iterations and the gradient descent algorithm, of the two classes of methods competing in the field: SCF and direct minimization methods. We derive asymptotic rates of convergence for these algorithms and analyze their dependence on the spectral gap and other properties of the problem. Our theoretical results are complemented by numerical simulations on a variety of examples, from toy models with tunable parameters to realistic Kohn–Sham computations. We also provide an example of chaotic behavior of the simple SCF iterations for a nonquadratic functional.

Key words. mathematical quantum chemistry, electronic structure, density matrices, SCF, direct minimization

AMS subject classifications. 65H17, 65K10

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1. Introduction. This paper is concerned with the convergence behavior of algorithms to solve the *subspace optimization problem*

(1.1)
$$\min \left\{ E(P) \mid P \in \mathbb{R}^{N_b \times N_b}, \ P^2 = P = P^*, \ \text{Tr}(P) = N \right\}$$

consisting of optimizing a C^2 function $E: \mathbb{R}^{N_b \times N_b} \to \mathbb{R}$ over the set of rank-N orthogonal projectors P. Here P^* denotes the adjoint (transpose) of P. This problem can also be reformulated as

(1.2)
$$\min \left\{ E\left(\sum_{i=1}^{N} \phi_i \phi_i^*\right) \middle| \phi_i \in \mathbb{R}^{N_b}, \ \phi_i^* \phi_j = \delta_{ij} \quad \forall \ i, j \in \{1, \dots, N\} \right\},$$

using an orthonormal basis $(\phi_i)_{i=1,\dots,N}$ for the subspace $\operatorname{Ran}(P)$. This problem is of interest in a number of contexts, such as matrix approximation, computer vision [1], and electronic structure theory [12, 26, 37, 38, 43, 53], the latter of which is the main motivation for this work.

Let $H(P) = \nabla E(P)$. The first-order conditions for problem (1.1) are

$$PH(P)(1-P) = (1-P)H(P)P = 0.$$

Up to an appropriate choice for the orthonormal basis $(\phi_i)_{i=1,...,N}$ of Ran(P), this yields

$$(1.3) H(P)\phi_i = \varepsilon_i \phi_i,$$

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which reveals an alternative interpretation of this problem as a nonlinear eigenvector problem (to be distinguished from nonlinear eigenvalue problems of the form $A(\varepsilon)\phi = 0$, where $A : \mathbb{R} \to \mathbb{R}^{N_b \times N_b}$). In the case when $E(P) = \text{Tr}(H_0P)$ for a fixed symmetric matrix H_0 , one recovers the classical eigenvalue problem $H_0\phi_i = \varepsilon_i\phi_i$. At a minimizer of (1.1), the $(\varepsilon_i)_{i=1,...,N}$ are the lowest eigenvalues of H_0 , counting multiplicities.

Problems of the form (1.1) are found in the Hartree–Fock and Kohn–Sham theories of electronic structure [26, 43], both approximations of the many-body Schrödinger equation. In this context, the ϕ_i are (discretized) orbitals, the projector P is the density matrix, and the energy E(P) includes linear contributions from the kinetic and external potential energy of the electrons, as well as nonlinear terms arising from electron-electron interaction. Another notable problem of this form is the nonlinear Schrödinger or Gross–Pitaevskii equation for Bose–Einstein condensates [6], where N=1. In all these cases, the first-order condition (1.3) is interpreted as a self-consistent or mean-field equation: the particles behave as independent particles in an effective Hamiltonian H(P) (also known as the Fock matrix) involving the mean-field they create. In the rest of this paper, we will work on the formulation (1.1) without specifying E for generality.

The minimization problem (1.1) is compact but nonconvex: there exists at least one minimizer, but the minimizer might not be unique, and local minima might not be global ones. Solving this optimization problem is of considerable practical interest, and algorithms for doing so date back to the early days of quantum mechanics [24]. The first introduced and still most popular approach is the *self-consistent field* (SCF) method, which, in its original version [45, 51], works as follows: if P^k is the current iterate of the algorithm, P^{k+1} is found by solving (1.3) for the fixed matrix $H(P^k)$:

$$H(P^k)\phi_i^k = \varepsilon_i^k \phi_i^k, \quad (\phi_i^k)^* \phi_i^k = \delta_{ij}$$

with the ε_i^k sorted in nondecreasing order, and building P^{k+1} as

$$P^{k+1} = \sum_{i=1}^{N} \phi_i^k (\phi_i^k)^*.$$

This algorithm assumes the Aufbau property, which is that at a minimum P_* we have $P_* = \sum_{i=1}^N \phi_i \phi_i^*$ with ϕ_i a system of orthogonal eigenvectors associated with the lowest N eigenvalues of $H(P_*)$. This property holds for the (spin-unconstrained) Hartree–Fock model [4], and the Gross–Pitaevskii model without magnetic field [10]. It usually holds for molecular systems in the Kohn–Sham model but does not hold in general for the Gross–Pitaevskii model with strong magnetic fields.

This basic procedure converges for systems where the nonlinearity is weak but fails to converge otherwise (see [14] for a comprehensive mathematical analysis of this behavior when the functional E is a sum of a linear and a quadratic term in P, which is the case for the Hartree–Fock model). A solution is to damp this procedure, and mix the iterates to accelerate convergence. This gives rise to a variety of SCF algorithms, among which are Broyden-like and Anderson-like mixing algorithms [31, 42, 49, 54], the direct inversion in the iterative space (DIIS) algorithm [34, 47, 48], the optimal damping algorithm [13] (ODA), and the energy-DIIS algorithm combining the latter two approaches [35].

A second class of algorithms solves the minimization problem (1.1) directly. The minimization set $\{P \in \mathbb{R}^{N_b \times N_b}, P^2 = P^* = P, \text{Tr } P = N\}$ is diffeomorphic to the Grassmann manifold of the N-dimensional vector subspaces of \mathbb{R}^{N_b} . This set

is naturally equipped with the structure of a Riemannian manifold, and this allows the use of Riemann optimization algorithms [1, 20]. Direct minimization algorithms are preferred for the Gross-Pitaevskii model with magnetic fields [3, 18, 25, 27], for which the Aufbau principle is not satisfied in general. Gradient-type [2, 17, 46, 57, 60], Newton-type [5, 15, 62], and trust-region methods have also been designed to solve (1.1) for larger values of N. At the time of writing, direct minimization algorithms are less popular than SCF algorithms in electronic structure calculation, where N can be very large, but it is not clear whether this is for sound scientific reasons or because SCF algorithms have been implemented and optimized for decades in the main production codes, which has not been the case for direct minimization algorithms.

While the convergence of several SCF and direct minimization algorithms has been analyzed from a mathematical point of view (see, e.g., [16, 36, 40, 50, 56, 59] and references therein), the two approaches have not been compared in a systematic way to our knowledge. The purpose of this paper is to contribute to filling this gap, by focusing on very simple representatives of each class, namely, the damped SCF iteration and the gradient descent. We emphasize that neither of these two algorithms is a practical choice as is. The SCF iteration should be accelerated (for instance, using the Anderson acceleration technique), and the gradient information in direct minimization methods should rather be used as part of a quasi-Newton method (such as the limited-memory BFGS algorithm [1]). Depending on the exact problem at hand, all these methods should be preconditioned to avoid issues related to small mesh sizes (which leads to a divergence of the kinetic energy term) and/or large computational domains (which can lead to a divergence of the Coulomb energy, or the confining potential). We refer to [58] for a recent review in the context of the Kohn-Sham equations for solids. Rather, in this paper, we aim to focus on the very simplest representative of each general strategy (SCF and direct minimization). The investigation of these two basic algorithms is informative on the strengths and weaknesses of the two classes and is a first step in the analysis of more complex methods.

The paper is organized as follows. In section 2, we recall some results about optimization on Grassmann manifolds, in particular the first- and second-order optimality conditions, and prove preparatory lemmas. In section 3, we present the two algorithms that are in the scope of this paper: a fixed-step gradient descent and a damped SCF algorithm. We prove their local convergence as long as the step is small enough and we derive convergence rates. We find that the convergence rates depend on the spectral radius of operators (acting on $\mathbb{R}^{N_b \times N_b}$) of the form $1 - \beta J$ with β the fixed step and $J = \Omega_* + K_*$ for the gradient descent, $J = 1 + \Omega_*^{-1} K_*$ for the SCF algorithm, where the operators Ω_* and K_* are specified in the next section. Let us just mention at this stage that the lowest eigenvalue of Ω_* is equal to the spectral gap between the Nth and (N+1)st eigenvalues of $H(P_*)$, allowing us to analyze the convergence rates of the algorithms in terms of natural quantities of the problem. This also shows that the damped SCF algorithm can be seen as a matrix splitting of the fixed-step gradient descent algorithm.

In section 4, we compare the two algorithms on several test problems. First, we focus on a toy model for which we can easily tune the gap and observe some fundamental differences between SCF and direct minimization algorithms, in agreement with the mathematical results established in section 3. We also provide an example of chaos in SCF iterations, complementing the results of [14, 36] in the case of a nonquadratic objective functional E. Then, we analyze a one-dimensional (1D) Gross-Pitaevskii

model (N=1) and its fermionic counterpart for N=2, for which we investigate the behavior of the algorithms when the gap closes. We conclude with an example from electronic structure calculation: a silicon crystal, in the framework of Kohn–Sham density functional theory (DFT), where we show in particular that accelerated SCF algorithms are less sensitive to small gaps than the simple damped SCF. Finally, in section 5 we draw conclusions and outline perspectives for future work.

2. Optimization on Grassmann manifolds. We focus in this paper on the case of real symmetric matrices, but the study can be easily extended to complex Hermitian matrices. Let $\mathcal{H} := \mathbb{R}_{\text{sym}}^{N_b \times N_b}$ be the vector space of $N_b \times N_b$ real symmetric matrices endowed with the Frobenius inner product $\langle A, B \rangle_{\text{F}} := \text{Tr}(AB)$. Let

$$\mathcal{M} := \{ P \in \mathcal{H} \mid P^2 = P \} \text{ and } \mathcal{M}_N := \{ P \in \mathcal{H} \mid P^2 = P, \operatorname{Tr}(P) = N \}.$$

From a geometrical point of view, \mathcal{M} is a compact subset of \mathcal{H} with N_b+1 connected components \mathcal{M}_N , $N=0,\ldots,N_b$, each of them being characterized by the value of Tr(P), namely, the rank of the orthogonal projector P, and being diffeomorphic to the Grassmann manifold $\text{Grass}(N,N_b)$ [1]. From now on, we fix the number of electrons N and we seek the local minimizers of the problem

(2.1)
$$\min_{P \in \mathcal{M}_N} E(P),$$

where $E:\mathcal{H}\to\mathbb{R}$ is a discretized energy functional, for which some examples are given below.

Example 2.1. As an example, we study a discrete Gross-Pitaevskii model in subsection 4.4. Other models from electronic structure can be considered, such as the discretized Hartree-Fock or Kohn-Sham models, where the energy is of the form

$$E(P) := \operatorname{Tr}(H_0 P) + E_{\operatorname{nl}}(P)$$

with H_0 being the core Hamiltonian (representing the kinetic energy and the external potential) and $E_{\rm nl}$ a nonlinear energy functional depending on the model (representing the interaction between electrons). For instance, for the Hartree–Fock model,

$$E_{\mathrm{nl}}(P) := \frac{1}{2} \operatorname{Tr}(G(P)P), \quad \text{where} \quad (G(P))_{ij} := \sum_{k,l=1}^{N_b} A_{ijkl} P_{kl} \quad \forall \ i,j=1,\ldots,N_b,$$

with A a symmetric tensor of order 4. For more details on these models or electronic structure in general, we refer to [12, 38, 53].

In plane-wave, finite differences, finite elements, or wavelets electronic structure calculation codes, the size N_b of the discretized space is in practice much larger than the number N of electrons. Therefore, it is not practical to store and manipulate the (dense) matrix P. Instead, algorithms work on the orbitals $(\phi_i)_{i=1,...,N}$ introduced in (1.3). The density matrix P is then recovered as

$$P = \sum_{i=1}^{N} \phi_i \phi_i^*.$$

All the results in this article are presented in the density matrix framework. However, the algorithms we study can be expressed in a way that avoids ever forming the density matrix. We refer to [58] for details.

We will need two assumptions for our results.

Assumption 2.2. The energy functional $E: \mathcal{H} \to \mathbb{R}$ is of class C^2 (twice continuously differentiable).

Assumption 2.2 is true for Hartree–Fock models. For Kohn–Sham models, it is true when the density $\rho = \sum_{i=1}^N |\phi_i|^2$ is uniformly bounded away from zero, which is the case, for instance, in condensed phase systems. Most of the results presented in this article are local in nature, and therefore this assumption can be relaxed to local regularity.

Assumption 2.3. $P_* \in \mathcal{M}_N$ is a nondegenerate local minimizer of (2.1) in the sense that there exists some $\eta > 0$ such that, for $P \in \mathcal{M}_N$ in a neighborhood of P_* , we have

$$E(P) \geqslant E(P_*) + \eta \|P - P_*\|_{F}^{2}$$
.

It is very hard in most practical situations to check this assumption, but it seems to be verified in practice. Notable exceptions are systems invariant with respect to continuous symmetry groups, in which case $E(P) = E(P_*)$ for all P in the orbit of P_* along the symmetry group. In this case, the assumption cannot be true, and $||P - P_*||_F$ must be replaced by the distance from P to the orbit of P_* . Our results can be extended to this case up to quotienting \mathcal{H} by the symmetry group.

Throughout the paper, we will use the following notation:

- $H(P) := \nabla E(P)$ is the gradient, and $H_* := H(P_*)$;
- $K(P) := \Pi_P \nabla^2 E(P) \Pi_P$ is the Hessian projected onto the tangent space at P, and $K_* := K(P_*)$ (the projection Π_P is defined below in Proposition 2.4).
- **2.1. First-order condition.** To study the first-order optimality conditions, we start by recalling some classical results about the geometry of the manifold \mathcal{M}_N .

PROPOSITION 2.4. \mathcal{M}_N is a smooth real manifold and its tangent space $T_P \mathcal{M}_N$ at $P \in \mathcal{M}_N$ is given by

$$T_P \mathcal{M}_N = \{ X \in \mathcal{H} \mid PX + XP = X, \text{ Tr}(X) = 0 \}$$

= $\{ X \in \mathcal{H} \mid PXP = (1 - P)X(1 - P) = 0 \}$.

The orthogonal projection Π_P on $T_P\mathcal{M}_N$ for the Frobenius inner product is

(2.2)
$$\forall X \in \mathcal{H}, \quad \Pi_P(X) = PX(1-P) + (1-P)XP = [P, [P, X]],$$

where [A, B] := AB - BA.

This classical result is proved in, e.g., [1, section 3.4]. Using the fact that $\mathcal{H} = \operatorname{Ran}(P) \oplus \operatorname{Ran}(1-P)$ and the induced decomposition of $P \in \mathcal{M}_N$ and $X \in \mathcal{H}$ as

(2.3)
$$P = \begin{bmatrix} I_N & 0 \\ 0 & 0 \end{bmatrix}, \quad X = \begin{bmatrix} (X)_{\text{oo}} & (X)_{\text{ov}} \\ (X)_{\text{vo}} & (X)_{\text{vv}} \end{bmatrix},$$

the projection Π_P is given by

$$\Pi_P(X) = \begin{bmatrix} 0 & (X)_{\text{ov}} \\ (X)_{\text{vo}} & 0 \end{bmatrix}.$$

Here the subscript "o" (resp., "v") stands for *occupied* (resp., *virtual*). The first-order optimality condition at P_* is $\Pi_{P_*}(H_*) = 0$, which can be formulated as follows:

(2.4) First-order optimality condition:
$$P_*H_*(1-P_*)=(1-P_*)H_*P_*=0$$
.

Note that this condition can be rewritten as $[H_*, P_*] = 0$, showing that H_* and P_* can be codiagonalized. Let $(\phi_k)_{1 \leqslant k \leqslant N_b}$ be an orthonormal basis of eigenvectors of H_* associated with the eigenvalues $(\varepsilon_k)_{1 \leqslant k \leqslant N_b}$ sorted in ascending order. Then $P = \sum_{i \in \mathcal{I}} \phi_i \phi_i^*$, where $\mathcal{I} \subset \{1, \ldots, N_b\}$, $|\mathcal{I}| = N$ is the set of occupied orbitals. The minimizer P_* is said to satisfy

- the Aufbau principle if $\mathcal{I} = \{1, \dots, N\}$;
- the strong Aufbau principle if $\mathcal{I} = \{1, ..., N\}$ and if in addition $\varepsilon_N < \varepsilon_{N+1}$, in which case $P_* = \sum_{i=1}^N \phi_i \phi_i^*$.
- **2.2. Second-order condition.** We derive here the second-order optimality condition from the nondegeneracy of the minimum (Assumption 2.3).

Let $X \in T_{P_*}\mathcal{M}_N$, I be a real interval containing 0 and $\gamma: I \to \mathcal{M}_N$ be a smooth path such that $\gamma(0) = P_*$ and $\dot{\gamma}(0) = X$. An example of a possible γ is given in subsection 4.1. We expand

$$E\left(\gamma(t)\right) = E(P_*) + t \left\langle H_*, X \right\rangle_{\mathcal{F}} + \frac{t^2}{2} \left(\left\langle H_*, \ddot{\gamma}(0) \right\rangle_{\mathcal{F}} + \left\langle X, \nabla^2 E(P_*) X \right\rangle_{\mathcal{F}} \right) + o(t^2)$$

$$= E(P_*) + \frac{t^2}{2} \left(\left\langle H_*, \ddot{\gamma}(0) \right\rangle_{\mathcal{F}} + \left\langle X, K_* X \right\rangle_{\mathcal{F}} \right) + o(t^2)$$

as H_* is orthogonal to $T_{P_*}\mathcal{M}_N$ at the minimum. Differentiating the relation $\gamma(t)^2 = \gamma(t)$ at t = 0, we get

$$P_* \ddot{\gamma}(0) + \ddot{\gamma}(0)P_* + 2X^2 = \ddot{\gamma}(0),$$

from which we obtain the following two relations on the diagonal blocks of $\ddot{\gamma}(0)$ in the decomposition (2.3):

$$\frac{1}{2} \left(\ddot{\gamma}(0) \right)_{\mathrm{oo}} = - \left(X^2 \right)_{\mathrm{oo}} = - \left(X \right)_{\mathrm{ov}} \left(X \right)_{\mathrm{vo}}, \qquad \frac{1}{2} \left(\ddot{\gamma}(0) \right)_{\mathrm{vv}} = \left(X^2 \right)_{\mathrm{vv}} = \left(X \right)_{\mathrm{vo}} \left(X \right)_{\mathrm{ov}}.$$

Thus, since $(H_*)_{vo} = (H_*)_{ov}^* = 0$ at the minimum, we have

$$\begin{split} \langle H_*, \ddot{\gamma}(0) \rangle_{\mathrm{F}} &= \mathrm{Tr} \left(\begin{bmatrix} (H_*)_{\mathrm{oo}} & 0 \\ 0 & (H_*)_{\mathrm{vv}} \end{bmatrix} \begin{bmatrix} (\ddot{\gamma}(0))_{\mathrm{oo}} & (\ddot{\gamma}(0))_{\mathrm{ov}} \\ (\ddot{\gamma}(0))_{\mathrm{vo}} & (\ddot{\gamma}(0))_{\mathrm{vv}} \end{bmatrix} \right) \\ &= 2 \, \mathrm{Tr} \left(- \left(H_* \right)_{\mathrm{oo}} \left(X \right)_{\mathrm{ov}} \left(X \right)_{\mathrm{vo}} \right) + 2 \, \mathrm{Tr} \left(\left(H_* \right)_{\mathrm{vv}} \left(X \right)_{\mathrm{vo}} \left(X \right)_{\mathrm{ov}} \right) \\ &= 2 \, \mathrm{Tr} \left(- \left(X \right)_{\mathrm{vo}} \left(H_* \right)_{\mathrm{oo}} \left(X \right)_{\mathrm{ov}} \right) + 2 \, \mathrm{Tr} \left(\left(X \right)_{\mathrm{ov}} \left(H_* \right)_{\mathrm{vv}} \left(X \right)_{\mathrm{vo}} \right) \\ &= \mathrm{Tr} \left(X \left(\Omega_* X \right) \right), \end{split}$$

where the operator $\Omega_*: T_{P_*}\mathcal{M}_N \to T_{P_*}\mathcal{M}_N$ is defined as

$$\Omega_* X := P_* X (1 - P_*) H_* - H_* P_* X (1 - P_*) + \text{sym}$$

$$= \begin{bmatrix} 0 & (X)_{\text{ov}} (H_*)_{\text{vv}} - (H_*)_{\text{oo}} (X)_{\text{ov}} \\ (H_*)_{\text{vv}} (X)_{\text{vo}} - (X)_{\text{vo}} (H_*)_{\text{oo}} & 0 \end{bmatrix},$$

where "sym" stands for the transpose of the previous expression. Introducing the operator

$$(2.6) \Omega_* + K_* : T_{P_*} \mathcal{M}_N \to T_{P_*} \mathcal{M}_N,$$

one gets in the end

$$E\left(\gamma(t)\right) = E(P_*) + \frac{t^2}{2} \left\langle X, (\Omega_* + K_*)X \right\rangle_{\mathcal{F}} + o(t^2).$$

At the critical point P_* , the second-order expansion of $E(\gamma(t))$ only depends on $X = \dot{\gamma}(0)$, a common feature in constrained optimization. The operator $\Omega_* + K_*$ can be interpreted as the Hessian of the energy on the manifold, or alternatively as the partial Hessian of the Lagrangian on \mathcal{H} . The operator Ω_* represents the influence of the curvature of the manifold on the Hessian of E.

As P_* is a nondegenerate minimum in the sense of Assumption 2.3, we have the

(2.7) Second-order optimality condition:
$$\forall X \in T_{P_*} \mathcal{M}_N, \quad \langle X, (\Omega_* + K_*) X \rangle_F \geqslant \eta \|X\|_F^2.$$

Remark 2.5 (structure of Ω_* and link with the Aufbau principle). Let P_* be a nondegenerate minimizer of (2.1) in the sense of Assumption 2.3. Denoting by A_{kl} the component along $\phi_k \phi_l^*$ of the matrix $A \in \mathcal{H}$, the operator Ω_* defined in (2.5) can alternatively be defined by

$$\forall X \in T_{P_*} \mathcal{M}_N, \quad \begin{cases} (\Omega_* X)_{ia} = (\varepsilon_a - \varepsilon_i) X_{ia}, \\ (\Omega_* X)_{ai} = (\varepsilon_a - \varepsilon_i) X_{ai} \end{cases} \text{ for } i \in \mathcal{I}, \ a \notin \mathcal{I},$$

where we have used the standard notation in chemistry of using the subscripts i for occupied and a for virtual orbitals (\mathcal{I} is the set of occupied orbitals).

In the case when $E(D)=\operatorname{Tr}(HD)$ for some fixed symmetric matrix $H\in\mathcal{H}$ (linear eigenvalue problem), then $K_*=0$ and so (2.7) is equivalent to the Aufbau principle. This equivalence does not hold true in general for nonlinear models: (2.7) is independent of the Aufbau principle, and η is unrelated to the gap $\nu=\min_{a\notin\mathcal{I}}\varepsilon_a-\max_{i\in\mathcal{I}}\varepsilon_i$ (equal to the lowest eigenvalue of the operator Ω_*). However, in specific cases, such as the reduced Hartree–Fock or Gross–Pitaevskii model, where $K_*\geqslant 0$, we have $\eta\geqslant \nu$ and a positive gap is a sufficient (but not necessary) condition for optimality.

Remark 2.6 (link with the Liouvillian). Another way to understand Ω_* is to use the Liouvillian \mathcal{L}_{H_*} associated to H_* , which is defined by

$$\forall A \in \mathcal{H}, \quad \mathcal{L}_{H_*} A := [H_*, A].$$

The action of \mathcal{L}_{H_*} has a simple expression in the eigenvector decomposition $(\varepsilon_k, \phi_k)_{1 \leqslant k \leqslant N_b}$ of H_* :

$$(2.8) \forall 1 \leq k, l \leq N_b, \quad \mathcal{L}_{H_*}(\phi_k \phi_l^*) = (\varepsilon_k - \varepsilon_l) \phi_k \phi_l^*.$$

Thus, we have

$$\forall i \in \mathcal{I}, \ a \notin \mathcal{I}, \quad \Omega_*(\phi_i \phi_a^* + \phi_a \phi_i^*) = (\varepsilon_a - \varepsilon_i)(\phi_i \phi_a^* + \phi_a \phi_i^*).$$

Hence, one can easily check that, using again the decomposition (2.3), we have

$$(2.9) \forall X \in T_{P_*} \mathcal{M}_N, \quad \Omega_* X = -[P_*, \mathcal{L}_{H_*} X] = -[P_*, [H_*, X]].$$

This definition also provides a canonical way to extend the operator Ω_* , originally defined on $T_{P_*}\mathcal{M}_N$, to the whole space \mathcal{H} .

2.3. Fixed-point on a manifold. Finally, we state a general abstract result that we will use to study the convergence of optimization algorithms on manifolds.

LEMMA 2.7. Let \mathcal{M} be a smooth finite-dimensional Riemannian manifold. Let $P_* \in \mathcal{M}$ and $f: U \to \mathcal{M}$ be a continuously differentiable mapping on a neighborhood U of P_* such that $f(P_*) = P_*$. Let $\mathrm{d}f(P_*): T_{P_*}\mathcal{M} \to T_{P_*}\mathcal{M}$ be the derivative of f at P_* . If $\mathrm{d}f(P_*)$ verifies $r(\mathrm{d}f(P_*)) < 1$ where $r(\mathrm{d}f(P_*))$ is the spectral radius of $\mathrm{d}f(P_*)$, then, for P^0 close enough to P_* , the fixed point iteration $P^{k+1} = f(P^k)$ linearly converges to P_* with asymptotic rate $r(\mathrm{d}f(P_*))$, in the sense that for all $\theta > 0$ there exists $C_\theta > 0$ such that, for all P^0 close enough to P_* ,

$$||P^k - P_*|| \le C_\theta \left(r(\mathrm{d}f(P_*)) + \theta\right)^k ||P^0 - P_*||.$$

Proof. We use the notation presented in [8, Chapter 1]. Up to a restriction of U to a smaller neighborhood of P_* , there exists a neighborhood V of 0 in $T_{P_*}\mathcal{M}$ and $g:V\to U$ a diffeomorphic parametrization of the manifold such that $g(0)=P_*$ and $\mathrm{d}g(0)=\mathrm{Id}$ (take, for instance, the restriction to V of the exponential map). Therefore, as f is continuously differentiable, there exists a neighborhood $\widetilde{V}\subset V$ of 0 in $T_{P_*}\mathcal{M}$ such that $F:=g^{-1}\circ f\circ g:\widetilde{V}\to V$ is a continuously differentiable map with fixed-point 0 and $\mathrm{d}F(0)=\mathrm{d}f(P_*)$. As $r(\mathrm{d}F(0))=r(\mathrm{d}f(P_*))<1$, we can find a neighborhood $V'\subset V$ of 0 in $T_{P_*}\mathcal{M}$ such that F is a contraction in V' for some norm $\|\cdot\|_{\theta}$, with contraction factor $r(\mathrm{d}f(P_*))+\theta$, $\theta>0$ (see [30] for more details). Therefore, we can apply the Banach fixed-point theorem to F and we get that, for x^0 close enough to 0, $x^{k+1}=F(x^k)$ converges to 0. Finally, for $P^0=g(x^0)$, $P^{k+1}=g(x^{k+1})=g(F(x^k))=f(g(x^k))=f(P^k)$ converges to $P_*=g(0)$, with asymptotic rate $r(\mathrm{d}f(P_*))$.

3. Algorithms and analysis of convergence.

- **3.1. Direct minimization.** The gradient descent algorithm consists in following the steepest descent direction with a fixed step β at each iteration point. As the iterations are constrained to stay on the manifold, we have to
 - 1. project the gradient on the tangent space with Π_{P^k} to bring the steepest descent line $P^k \beta \Pi_{P^k}(\nabla E(P^k))$ back to the manifold at first order;
 - 2. retract the steepest descent line defined in the tangent space onto the manifold \mathcal{M}_N by a nonlinear retraction R mapping a neighborhood of \mathcal{M}_N in \mathcal{H} to \mathcal{M}_N .

An example of retraction is given in subsection 4.1 and we will assume that the retraction R satisfies the following.

Assumption 3.1. $R: \mathcal{H} \to \mathcal{H}$ is of class C^2 and for all $P \in \mathcal{M}_N$ and $X \in \mathcal{H}$ small enough,

$$R(P+X) \in \mathcal{M}_N \text{ and } R(P+X) = P + \Pi_P(X) + O(X^2).$$

These two successive operations are sketched in Figure 1 and the gradient descent algorithm is presented in Algorithm 3.1.

Algorithm 3.1 Gradient descent.

```
P^0 \in \mathcal{M}_N

while convergence not reached do

P^{k+1} := R\left(P^k - \beta \Pi_{P^k}\left(\nabla E(P^k)\right)\right)

end while
```

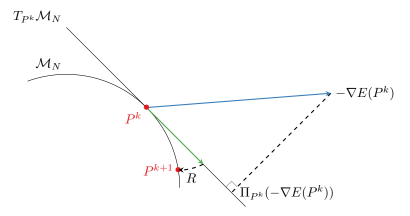


Fig. 1. Projection on the tangent space for the gradient descent, and retraction to the manifold.

At the continuous level, this algorithm can be seen as the discretization of the flow $\dot{P} = -\Pi_P \nabla E(P)$. Note that, by the use of the retraction R and Assumption 3.1, the projection step has no influence on the convergence of the algorithm for β small. Indeed, by Assumption 3.1,

$$\forall P \in \mathcal{M}_N, \quad R(P - \beta \Pi_P(\nabla E(P))) = P - \beta \Pi_P(\Pi_P(\nabla E(P))) + O(\beta^2)$$
$$= P - \beta \Pi_P(\nabla E(P)) + O(\beta^2)$$

and thus the first-order expansion is the same with or without the projection step. The reason we use this projection step is that it is convenient to interpret $\Pi_{P^k}\nabla E(P^k)$ as a residual.

The following theorems state that, for β small enough, Algorithm 3.1 globally converges in the sense that $\Pi_{P^k}\nabla E(P^k)\to 0$ and locally converges in the sense that $P^k\to P_*$ if P^0 is close enough to P_* .

THEOREM 3.2. Let $E: \mathcal{H} \to \mathbb{R}$ satisfy Assumption 2.2 and $R: \mathcal{H} \to \mathcal{H}$ satisfy Assumption 3.1. There exists $\beta_0 > 0$ such that for all $0 < \beta \leqslant \beta_0$ and all $P^0 \in \mathcal{M}_N$, the iterations

$$P^{k+1} := R \left(P^k - \beta \Pi_{P^k} \left(\nabla E(P^k) \right) \right)$$

satisfy the following properties:

- 1. $(E(P^k))_{k\in\mathbb{N}}$ is a nonincreasing sequence converging to some critical value E_c of E on \mathcal{M}_N ;
- 2. when k goes to infinity, $\Pi_{P^k}\nabla E(P^k) \to 0$, $\|P^{k+1}-P^k\|_F \to 0$, and $d(P^k, A_c) \to 0$, where A_c is one of the connected components of $C(E_c) := \{P \in \mathcal{M}_N \mid E(P) = E_c \text{ and } \Pi_P(\nabla E(P)) = 0\}$.

Proof. As $E: \mathcal{H} \to \mathbb{R}$ and $R: \mathcal{H} \to \mathcal{H}$ are C^2 , and \mathcal{M}_N is compact, we can use the expansion of Assumption 3.1 and obtain that there exists a constant $C \in \mathbb{R}_+$ such that for all $0 \le \beta \le 1$,

$$\forall \ k \in \mathbb{N}, \quad E(P^{k+1}) \leqslant E(P^k) - \beta \left\| \Pi_{P^k} \nabla E(P^k) \right\|_{\mathrm{F}}^2 + C\beta^2 \left\| \Pi_{P^k} \nabla E(P^k) \right\|_{\mathrm{F}}^2.$$

Therefore, we have for $\beta > 0$ small enough,

$$\forall \ k \in \mathbb{N}, \quad E(P^{k+1}) \leqslant E(P^k) - \frac{\beta}{2} \left\| \Pi_{P^k} \nabla E(P^k) \right\|_{\mathrm{F}}^2.$$

This shows that the sequence $(E(P^k))_{k\in\mathbb{N}}$ is nonincreasing. As E is continuous on the compact set \mathcal{M}_N , $(E(P^k))_{k\in\mathbb{N}}$ is bounded and hence converges to some $E_c \in \mathbb{R}$. Moreover,

$$\sum_{k \in \mathbb{N}} \left\| \Pi_{P^k} \nabla E(P^k) \right\|_{\mathcal{F}}^2 < \infty,$$

which implies that $\Pi_{P^k} \nabla E(P^k) \to 0$ when $k \to \infty$. It follows that $\|P^{k+1} - P^k\|_F \to 0$ when $k \to \infty$.

Let B be the nonempty compact set of accumulation points of $(P^k)_{k\in\mathbb{N}}$. By continuity of E and $P \mapsto \Pi_P \nabla E(P)$, it follows that $B \subset C(E_c)$. Assuming that $d(P^k, B)$ does not go to zero, we can extract a subsequence at finite distance of B which converges to a point in B, a contradiction. Assume that B is disconnected: it is then the union of two compact subsets B_1 and B_2 at positive distance from each other. Since $P^{k+1} - P^k \to 0$, there is an infinite number of points in $(P^k)_{k\in\mathbb{N}}$ at distance greater or equal to $\eta > 0$ from both B_1 and B_2 , from which we can extract a point in B, a contradiction. It follows that B is connected, hence the result.

This result implies in particular the convergence of the sequence $(P^k)_{k\in\mathbb{N}}$ in the generic case where critical points are isolated. If this is not the case but E and R are analytic, convergence can be shown following the approach in [36] based on Lojasiewicz inequality.

THEOREM 3.3. Let $E: \mathcal{H} \to \mathbb{R}$ satisfy Assumptions 2.2 and 2.3 with P_* a local minimizer of (2.1). Let $R: \mathcal{H} \to \mathcal{H}$ satisfy Assumption 3.1. Then, if $P^0 \in \mathcal{M}_N$ is close enough to P_* , the iterations

$$P^{k+1} := R \left(P^k - \beta \Pi_{P^k} \left(\nabla E(P^k) \right) \right)$$

linearly converge to P_* for $\beta > 0$ small enough, with asymptotic rate $r(1 - \beta J_{\rm grad})$ where $J_{\rm grad} \coloneqq \Omega_* + K_*$.

Proof. In order to prove convergence, one can apply Lemma 2.7 to the function $f: \mathcal{M}_N \to \mathcal{M}_N$ defined by

$$f(P) := R(P - \beta \Pi_P (\nabla E(P))),$$

for which we know by the first-order optimality condition that P_* is a fixed-point.

We compute explicitly $df(P_*)$ using the second-order optimality condition (2.7). To this end, take $X \in T_{P_*}\mathcal{M}_N$ and a smooth path $\gamma: I \to \mathcal{M}_N$ defined on a real interval I containing 0 such that $\gamma(0) = P_*$ and $\dot{\gamma}(0) = X$. We want to expand to the first order in t the following expression:

$$f(\gamma(t)) = R \left(\gamma(t) - \beta \Pi_{\gamma(t)} \left(\nabla E(\gamma(t)) \right) \right).$$

First, we focus on the projection of $H(\gamma(t))$ on $T_{\gamma(t)}\mathcal{M}_N$:

$$\begin{split} \Pi_{\gamma(t)}H(\gamma(t)) &= \gamma(t)H(\gamma(t))(1-\gamma(t)) + \text{sym} \\ &= (P_* + tX) \left(H_* + t \left(\nabla^2 E(P_*)X \right) \right) (1-P_* - tX) + \text{sym} + O(t^2) \\ &= t \left[P_* \left(\nabla^2 E(P_*)X \right) (1-P_*) + \text{sym} \right] \\ &+ t \left[X H_* (1-P_*) - P_* H_* X + \text{sym} \right] + O(t^2) \\ &= t (K_* + \Omega_*) X + O(t^2). \end{split}$$

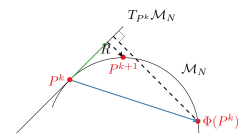


Fig. 2. Retraction for the damped SCF algorithm.

Inserting this into the expansion of $f(\gamma(t))$ and using Assumption 3.1 and the fact that $\Pi_{\gamma(t)}X = \Pi_{P_*}X + O(t^2)$ gives

$$f(\gamma(t)) = R(\gamma(t) - \beta t(\Omega_* + K_*)X + O(t^2)) = P_* + t(X - \beta(\Omega_* + K_*)X) + O(t^2).$$

Therefore.

$$df(P_*)X = (1 - \beta(\Omega_* + K_*))X.$$

As the second-order optimality condition (2.7) shows that $\Omega_* + K_*$ is positive definite on $T_{P_*}\mathcal{M}_N$, for β small enough, the spectral radius $r(\mathrm{d}f(P_*))$ of the derivative $\mathrm{d}f(P_*)$ is less than 1, which concludes the proof.

- **3.2.** Damped self-consistent field. The damped SCF algorithm is a damped version of the Roothaan algorithm [14, 36] and is presented in Algorithm 3.2, under the assumption that the strong Aufbau principle is satisfied, and represented in Figure 2. Note that it is well defined only if $\varepsilon_N^k < \varepsilon_{N+1}^k$ for all $k \in \mathbb{N}$. We introduce the nonlinear operators:
 - 1. $A(H) := \mathbf{1}_{(-\infty,\varepsilon_N(H)]}(H)$ with $\varepsilon_N(H)$ the lowest Nth eigenvalue of H and where we recall that $\mathbf{1}_{(-\infty,\mu]}(H) \coloneqq \sum_{\varepsilon_i \leq \mu} \phi_i \phi_i^*$, the ϕ_i 's being orthonormal eigenvectors of H associated to the eigenvalues ε_i ; 2. $\Phi(P) = A(H(P))$ or, equivalently, $\Phi(P) \coloneqq \sum_{i=1}^N \phi_i \phi_i^*$, where the ϕ_i 's are
 - orthonormal eigenvectors associated to the lowest N eigenvalues of H(P).

Algorithm 3.2 Damped SCF algorithm.

$$\begin{split} &P^0 \in \mathcal{M}_N \\ &\textbf{while} \text{ convergence not reached } \textbf{do} \\ &\text{solve} \begin{cases} H(P^k)\phi_i^k = \varepsilon_i^k\phi_i^k, \ \varepsilon_1^k \leqslant \cdots \leqslant \varepsilon_N^k < \varepsilon_{N+1}^k \leq \cdots \leq \varepsilon_{N_b}^k, \\ \left\langle \phi_i^k, \phi_j^k \right\rangle = \delta_{ij}, \end{cases} \\ &\Phi(P^k) \coloneqq \sum_{i=1}^N \phi_i^k \left(\phi_i^k \right)^* \\ &P^{k+1} \coloneqq R \left(P^k + \beta \Pi_{P^k} \left(\Phi(P^k) - P^k \right) \right) \\ &\textbf{end while} \end{split}$$

The following theorem states that, under the condition that there is a gap between the smallest Nth and (N+1)st eigenvalues of the Hamiltonian H_* , Algorithm 3.2 locally converges for β small enough.

THEOREM 3.4. Let $E: \mathcal{H} \to \mathbb{R}$ and $P_* \in \mathcal{M}_N$ satisfy Assumptions 2.2 and 2.3 and $R: \mathcal{H} \to \mathcal{H}$ satisfy Assumption 3.1. Assume that P_* satisfies the strong Aufbau

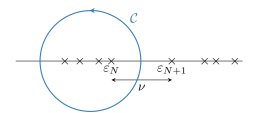


Fig. 3. Definition of A and graphical interpretation of the Aufbau principle and the existence of a gap.

principle

$$\Phi(P_*) = P_* \text{ and } \nu := \varepsilon_{N+1} - \varepsilon_N > 0,$$

where $(\varepsilon_i)_{1 \leq i \leq N_b}$ are the eigenvalues of H_* ranked in nondecreasing order. Then, for $\beta > 0$ small enough and $P^0 \in \mathcal{M}_N$ close enough to P_* , the iterations

$$P^{k+1} \coloneqq R \left(P^k + \beta \Pi_{P^k} \left(\Phi(P^k) - P^k \right) \right)$$

are well-defined and P^k linearly converges to P_* , with asymptotic rate $r(1 - \beta J_{SCF})$, where $J_{SCF} := 1 + \Omega_*^{-1} K_*$.

Proof. In order to prove convergence, we apply Lemma 2.7 to the function $f:\mathcal{M}_N\to\mathcal{M}_N$ defined by

$$f(P) := R(P + \beta \Pi_P (\Phi(P) - P)),$$

for which P_* is a fixed-point.

First, we compute the derivative of $\Phi = A \circ H$ at the minimizer P_* to get

$$d\Phi(P_*) = dA(H_*)\nabla^2 E(P_*).$$

Now, to compute $dA(H_*)$, note that, as there is a gap $\varepsilon_{N+1} > \varepsilon_N$ at the minimum, we can find a contour \mathcal{C} in the complex plane enclosing the lowest N eigenvalues of H_* (Figure 3) such that

(3.1)
$$A(H_*) = \frac{1}{2\pi i} \oint_{\mathcal{C}} \frac{1}{z - H_*} dz$$

(see [29, 32] for more details on spectral calculus, contour integrals, and perturbation theory for functions of matrices). By continuity, we also have

$$A(H) = \frac{1}{2\pi i} \oint_{\mathcal{C}} \frac{1}{z - H} \, \mathrm{d}z$$

for H in a neighborhood of H_* . Then, one can use the expression (3.1) of A and the expansion for H in a neighborhood of H_*

$$\forall z \in \mathcal{C}, \quad \frac{1}{z-H} = \frac{1}{z-H_*} (H - H^*) \frac{1}{z-H_*} + O(\|H - H_*\|_F^2)$$

to get

$$\forall h \in \mathcal{H}, \quad dA(H_*)h = \frac{1}{2\pi i} \oint_{\mathcal{C}} \frac{1}{z - H_*} h \frac{1}{z - H_*} dz$$
$$= \sum_{k=1}^{N_b} \sum_{l=1}^{N_b} \left(\frac{1}{2\pi i} \oint_{\mathcal{C}} \frac{1}{z - \varepsilon_k} h_{kl} \frac{1}{z - \varepsilon_l} dz \right) \phi_k \phi_l^*,$$

where $h_{kl} = \phi_k^* h \phi_l$. Now, let us denote by $1 \leq i \leq N$ the occupied orbitals (ε_i is inside \mathcal{C}) and by $N+1 \leq a \leq N_b$ the virtual ones (ε_a is outside \mathcal{C}). Then,

$$\oint_{\mathcal{C}} \frac{1}{z - \varepsilon_i} \frac{1}{z - \varepsilon_a} \, dz = \begin{cases} \frac{1}{\varepsilon_i - \varepsilon_a} & \text{if } 1 \leqslant i \leqslant N < a \leqslant N_b; \\ 0 & \text{otherwise.} \end{cases}$$

Thus, the sum becomes

$$dA(H_*)h = \sum_{i=1}^{N} \sum_{a=N+1}^{N_b} \frac{1}{\varepsilon_i - \varepsilon_a} \Big(h_{ia} \phi_i \phi_a^* + h_{ai} \phi_a \phi_i^* \Big) = -\Omega_*^{-1} \Pi_{P_*} h,$$

and we finally get

$$\forall X \in T_{P_*} \mathcal{M}_N, \quad d\Phi(P_*) X = -\Omega_*^{-1} K_* X.$$

Now, we compute the derivative of f at point P_* . Let $X \in T_{P_*}\mathcal{M}_N$ and $\gamma: I \to \mathcal{M}_N$ be a smooth path defined on a real interval I containing 0 such that $\gamma(0) = P_*$ and $\dot{\gamma}(0) = X$. First, we expand $\gamma(t)$ around 0 and Φ around $\gamma(0) = P_*$ to obtain

$$f(\gamma(t)) = P_* + t\Pi_{P_*} ((1-\beta) + \beta d\Phi(P_*)) X + O(t^2).$$

Thus, for $X \in T_{P_*}\mathcal{M}_N$,

$$df(P_*)X = ((1-\beta) - \beta\Omega_*^{-1}K_*)X$$

To conclude this proof, we compute the spectral radius of

$$df(P_*) = (1 - \beta) - \beta \Omega_*^{-1} K_* = 1 - \beta \left(1 + \Omega_*^{-1} K_*\right).$$

First, notice that

$$1 + {\Omega_*}^{-1} K_* = 1 + {\Omega_*}^{-1/2} \left({\Omega_*}^{-1/2} K_* {\Omega_*}^{-1/2} \right) {\Omega_*}^{1/2}$$

and thus, $1 + \Omega_*^{-1}K_*$ and the symmetric operator $1 + \Omega_*^{-1/2}K_*\Omega_*^{-1/2}$ have the same eigenvalues. Moreover, using the second-order optimality condition (2.7), with $X = \Omega_*^{-1/2}Y$, we get

$$(3.2) \qquad \forall Y \in T_{P_*} \mathcal{M}_N, \quad \left\langle Y, \left(1 + \Omega_*^{-1/2} K_* \Omega_*^{-1/2}\right) Y \right\rangle_{\mathcal{F}} \geqslant \eta \left\langle Y, \Omega_*^{-1} Y \right\rangle_{\mathcal{F}}$$

$$\geqslant \frac{\eta}{\|\Omega_*\|_{\text{op}}} \|Y\|_{\mathcal{F}}^2$$

with $\|\Omega_*\|_{\text{op}}$ the operator norm associated to $\|\cdot\|_{\text{F}}$. Thus, all the eigenvalues of $1 + \Omega_*^{-1/2}K_*\Omega_*^{-1/2}$, hence of $1 + \Omega_*^{-1}K$, are real and positive. Consequently, for β small enough, the spectral radius $r\left(\mathrm{d}f(P_*)\right)$ is less than 1 and we conclude by applying Lemma 2.7.

Remark 3.5 (case when the Aufbau principle is not satisfied). In the case when the minimizer P_* does not verify the Aufbau principle but does satisfy the condition that the eigenvalues of $1 + {\Omega_*}^{-1} K_*$ are positive (note that Ω_* is not positive when the Aufbau principle is not verified, but $1 + {\Omega_*}^{-1} K_*$ might still have only positive eigenvalues), the damped SCF still converges locally to P_* for $\beta > 0$ small enough if we change the way we select the occupied orbitals to build $\Phi(P)$ (in this case, we do not pick those associated to the smallest N eigenvalues of H(P), but those corresponding to the occupied orbitals of P_*).

We conclude this section by proving the local convergence of the nonretracted variant of Algorithm 3.2.

Algorithm 3.3 Nonretracted damped SCF algorithm.

$$\begin{split} P^0 &\in \mathcal{M}_N \\ \text{while convergence not reached do} \\ &\text{solve} \begin{cases} H(P^k) \phi_i^k = \varepsilon_i^k \phi_i^k, \ \varepsilon_1^k \leqslant \cdots \leqslant \varepsilon_N^k < \varepsilon_{N+1}^k \leqslant \cdots \leqslant \varepsilon_{N_b}^k, \\ \left\langle \phi_i^k, \phi_j^k \right\rangle = \delta_{ij}, \end{cases} \\ \Phi(P^k) &\coloneqq \sum_{i=1}^N \phi_i^k \left(\phi_i^k \right)^* \\ P^{k+1} &\coloneqq P^k + \beta \Pi_{P^k} \left(\Phi(P^k) - P^k \right) \\ \text{end while} \end{split}$$

THEOREM 3.6. Let $E: \mathcal{H} \to \mathbb{R}$ and P_* satisfy Assumptions 2.2 and 2.3. Moreover, assume that

$$\Phi(P_*) = P_*$$
 and $\nu := \varepsilon_{N+1} - \varepsilon_N > 0$ (strong Aufbau principle),

where $(\varepsilon_i)_{1 \leq i \leq N_b}$ are the eigenvalues of H_* ranked in nondecreasing order.

Then, for $\beta > 0$ small enough and $P^0 \in \mathcal{H}$ close enough to P_* and with trace N, the iterations

$$(3.3) P^{k+1} := P^k + \beta \left(\Phi(P^k) - P^k \right)$$

are well-defined and P^k linearly converges to $P_* \in \mathcal{M}_N$, with asymptotic rate $\max(r(1-\beta J_{SCF}), 1-\beta)$, where $J_{SCF} := 1 + \Omega_*^{-1} K_*$.

Note that the iterates P^k defined by (3.3) have trace N but do not lay on the manifold \mathcal{M}_N in general.

Proof. The proof follows that of Theorem 3.4. This time, we need to compute the Jacobian matrix of $f: \mathcal{H} \ni P \mapsto P + \beta (\Phi(P) - P) \in \mathcal{H}$ at the minimizer $P_* \in \mathcal{M}_N$. As we work in the whole space \mathcal{H} , the Jacobian matrix has the form, in the decomposition $\mathcal{H} = T_{P_*} \mathcal{M}_N \oplus (T_{P_*} \mathcal{M}_N)^{\perp}$,

$$df(P_*) = \begin{bmatrix} 1 - \beta J_{SCF} & \times \\ 0 & 1 - \beta \end{bmatrix},$$

where $J_{\text{SCF}} = 1 + \Omega_*^{-1} K_*$ has been computed in the proof of Theorem 3.4. Hence, this time the algorithm converges to $P_* \in \mathcal{M}_N$ as long as β is such that $\max(r(1 - \beta J_{\text{SCF}}), 1 - \beta) < 1$.

Within the local density approximation (LDA), or the generalized gradient approximation (GGA), the mean-field (Kohn–Sham) Hamiltonian H(P) is actually a function $\widetilde{H}(\rho_P)$ of the density ρ_P associated with the density matrix P. Since the map $P \mapsto \rho_P$ is linear, (3.3) can be rewritten as

$$\rho^{k+1} = (1 - \beta)\rho^k + \beta\Psi(\rho^k),$$

where $\Psi(\rho) = \rho_{A(\widetilde{H}(\rho))}$. We can therefore interpret (3.3) as the equivalent density matrix formulation of this density mixing algorithm.

- **3.3. Comparison.** In this section, we proved the local convergence of Algorithms 3.1 and 3.2. and we obtained asymptotic convergence rates. On the tangent space, both Jacobian matrices are of the form $1 \beta J$, where J has positive real spectrum and
 - for the gradient descent, $J_{\text{grad}} = K_* + \Omega_*$, which is self-adjoint for the Frobenius inner product;
 - for the damped SCF algorithm, if the strong Aufbau principle is satisfied at P_* , $J_{\text{SCF}} = 1 + \Omega_*^{-1} K_*$, which is self-adjoint for the inner product $\langle \cdot, \cdot \rangle_{\Omega_*} := \langle \Omega_* \cdot, \cdot \rangle_{\text{F}}$.

One can notice that, in the linear regime, the SCF iterations correspond to a matrix splitting of the gradient iterations. Whether this results in a faster method or not depends not only on the relative conditioning of the iteration matrices but also on the relative cost of each step.

To have the fastest convergence, we want the eigenvalues of $1 - \beta J$ to be as close to 0 as possible. If we denote by λ_1 (resp., λ_N) the smallest (resp., largest) eigenvalue of J, the optimal step β_* is the minimizer of $\min_{\beta} \max\{|1 - \beta \lambda_1|, |1 - \beta \lambda_N|\}$, which is given by

$$\beta_* = \frac{2}{\lambda_1 + \lambda_N}.$$

Then, the rate of convergence is, with $\kappa := \lambda_N/\lambda_1$ the spectral condition number of J.

$$r = \frac{\kappa - 1}{\kappa + 1}.$$

Now, we can evaluate the conditioning of J for the two algorithms:

• For the gradient descent, we have

(3.4)
$$\kappa(J_{\text{grad}}) \leqslant \frac{\|\Omega_*\|_{\text{op}} + \|K_*\|_{\text{op}}}{\eta},$$

where η is the coercivity constant in the nondegeneracy Assumption 2.3. First, the smaller η is, the more difficult the convergence. Note, however, that there is no relationship in general between η and the gap ν . Second, the bigger $\|\Omega_*\|_{\text{op}} = \varepsilon_{N_b} - \varepsilon_1$ is, the more difficult the convergence. In particular, for models arising from the discretization of partial differential equations, $\varepsilon_{N_b} - \varepsilon_1 \to \infty$ when the discretization is refined. In practice, this issue is solved by preconditioning (see Remark 3.7).

• For the damped SCF algorithm, a naive bound would be

$$\kappa(J_{SCF}) \leqslant \|\Omega_*\|_{op} \frac{1 + \nu^{-1} \|K_*\|_{op}}{\eta}.$$

In this bound the right-hand side diverges when $\|\Omega_*\|_{op} \to \infty$ as above, whereas the left-hand side may actually remain bounded. For instance, under the uniform coercivity assumption [11],

$$\forall X \in T_{P_*} \mathcal{M}_N, \quad \langle X, (\Omega_* + K_*) X \rangle_{\mathcal{F}} \geqslant \widetilde{\eta} \langle \Omega_* X, X \rangle_{\mathcal{F}},$$

with $\tilde{\eta}$ independent of N_b (which is often the case in practice), we have

$$\kappa(J_{\text{SCF}}) \leqslant \frac{1 + \nu^{-1} \|K_*\|_{\text{op}}}{\widetilde{\eta}}.$$

In contrast with the bound (3.4), we can see that the smaller the gap ν is, the slower the convergence.

As a special case, if we consider the case where the Hessian $\nabla^2 E \equiv 0$, i.e., a linear eigenvalue problem, then the SCF algorithm converges in one iteration, which is consistent with $J_{\text{SCF}} = 1$. The gradient descent with optimal step locally converges with asymptotic rate $r = \frac{\kappa - 1}{\kappa + 1}$, where $\kappa = \frac{\varepsilon_{N_b} - \varepsilon_1}{\varepsilon_{N_{+1}} - \varepsilon_N}$.

The convergence rates we derived in Theorems 3.3 and 3.4 are consistent with well-known convergence issues, for instance, the failure of the simple damped SCF algorithm to converge for systems with small gaps [52] (although subsection 4.5 shows this is not necessarily true for more sophisticated acceleration methods).

Remark 3.7 (preconditioning). We discuss here the extension of Theorem 3.3 to the preconditioned gradient descent:

$$P^{k+1} \coloneqq R \left(P^k - \beta \Pi_{P^k} B \Pi_{P^k} \left(\nabla E(P^k) \right) \right)$$

with $B: \mathcal{H} \to \mathcal{H}$ a symmetric positive definite preconditioner. If we denote by $\widetilde{B}_* := \Pi_{P_*}B\Pi_{P_*}$ its restriction to the tangent plane, the Jacobian matrix of the gradient becomes $1 - \beta \widetilde{B}_*(\Omega_* + K_*)$, where $(\Omega_* + K_*)$ is positive definite (under Assumption 2.3) and the proof of local convergence for β small enough follows exactly in the same way, using the positive definiteness of \widetilde{B}_* to show that $\widetilde{B}_*(\Omega_* + K_*)$ has real positive spectrum. The same analysis holds true for the preconditioned SCF algorithm. In practice, preconditioning is a crucial tool to accelerate iterations, in particular in order to achieve mesh- and domain-size independence of the number of iterations for discretized partial differential equations. However, we are interested here in the intrinsic aspects of each algorithm (direct minimization vs. SCF) and the influence of physical parameters (e.g., the gap ν), so that the study of preconditioned algorithms is not in the scope of this paper.

Remark 3.8 (dielectric operator). In the context of KS-DFT, the operator $(1 + K_*\Omega_*^{-1})^{-1}$, the transpose of the inverse of the Jacobian of the simple SCF mapping, is known as the dielectric operator: it represents the infinitesimal change in the self-consistent Hamiltonian $H(P_*)$ in response to a change in the energy functional. Our results show that this operator is well-defined and has real positive spectrum, with no assumption on the sign of Hartree-exchange-correlation kernel K_* , recovering in an algebraic framework the results of [19, 23] obtained using a different variational principle.

4. Numerical tests. We present here some numerical experiments to illustrate our theoretical results, explore their limits, and investigate the global behavior of the algorithms. First, we start by specifying the retraction R that we use in our

numerical tests. In subsection 4.2, we use a simple toy model for which we can control the gap and analytically compute the exact minimizer: this allows us to study the impact of the gap on the convergence of Algorithms 3.1 and 3.2. In subsection 4.3, we show that simple (nondamped) SCF iterations can exhibit chaotic behavior for some nonlinearities. Then, in subsection 4.4, we report numerical tests for a 1D Gross–Pitaevskii model (N=1) and its fermionic version for N=2. Finally, in subsection 4.5, we present results obtained with a more realistic case: silicon in the framework of the Kohn–Sham DFT.

4.1. The retraction. We choose the following algorithm: for a given symmetric matrix \widetilde{P} close to \mathcal{M}_N with eigendecomposition $\widetilde{P} = V\widetilde{D}V^*$ with \widetilde{D} diagonal and V orthogonal, we set the diagonal matrix D as

$$D_{ii} = \begin{cases} 1 & \text{if } \widetilde{D}_{ii} > 0.5, \\ 0 & \text{otherwise} \end{cases}$$

and $R(\widetilde{P}) = VDV^*$. When \widetilde{P} is close to \mathcal{M}_N , its eigenvalues are close to either 0 or 1. Given a contour \mathcal{C} enclosing only the eigenvalues close to 1, R has the following explicit expression:

(4.1)
$$R(P) = \frac{1}{2\pi i} \oint_{\mathcal{C}} \frac{1}{z - P} dz.$$

Therefore, it follows from arguments similar to those used in the proof of Theorem 3.4 that R is analytic and satisfies Assumption 3.1.

4.2. A toy model with tunable spectral gap. We work here in the very simple framework of real density matrices of order 2, i.e., the 2×2 real matrices P such that $P^* = P$, $P^2 = P$ and Tr(P) = 1. Then, we consider the following minimization problem, with a parameter $\varepsilon \ge 0$: energy functional

$$E_{\varepsilon}(P) := \operatorname{Tr}\left(\left(P - \begin{bmatrix} 1 & \varepsilon \\ \varepsilon & 0 \end{bmatrix}\right)^2\right)$$

for parameters $\varepsilon \geqslant 0$. The gradient and Hessian of E are

$$H_{\varepsilon}(P) = 2 \left(P - \begin{bmatrix} 1 & \varepsilon \\ \varepsilon & 0 \end{bmatrix} \right),$$
$$\nabla^{2} E_{\varepsilon}(P) = 2.$$

Simple computations show that the set of rank-1 projectors on \mathbb{R}^2 can be parameterized as

$$\mathcal{M}_1 := \left\{ P(a, b) = \begin{bmatrix} 1 - a & b \\ b & a \end{bmatrix} \mid a \in [0, 1], \ b = \pm \sqrt{a(1 - a)} \right\}.$$

The eigenvalues of H_{ε} at $P(a,b) \in \mathcal{M}_1$ are $\pm 2\sqrt{a^2 + (b-\varepsilon)^2}$. The gap is thus $\nu(a,b) := 4\sqrt{a^2 + (b-\varepsilon)^2}$.

The case $\varepsilon = 0$. Here, the unique minimum is clearly

$$P(0,0) = \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix} \in \mathcal{M}_1$$

and the gap is zero. Since $\nabla^2 E=2$, this minimum satisfies Assumption 2.3 with $\eta=2$.

The case $\varepsilon > 0$. We compute

$$E_{\varepsilon}(P(a,b)) = 2(a + \varepsilon^2 - 2\varepsilon b),$$

and therefore

$$E_{\varepsilon}\left(P(a,\sqrt{a(1-a)})\right)\leqslant E_{\varepsilon}\left(P(a,-\sqrt{a(1-a)})\right).$$

Hence, to compute the minimizer of the energy, we can restrict ourselves to the 1D manifold

$$P(a) = \begin{bmatrix} 1 - a & \sqrt{a - a^2} \\ \sqrt{a - a^2} & a \end{bmatrix}$$

with $a \in [0,1]$. Then, the energy is

(4.2)
$$E_{\varepsilon}(P(a)) = 2\left(a + \varepsilon^2 - 2\varepsilon\sqrt{a(a-1)}\right).$$

The first-order condition yields

$$a = \frac{1 \pm \sqrt{1 - \frac{4\varepsilon^2}{1 + 4\varepsilon^2}}}{2},$$

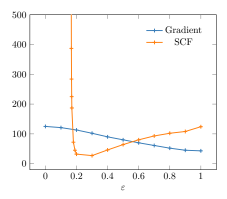
with the lowest energy achieved at

$$a(\varepsilon) \coloneqq \frac{1 - \sqrt{1 - \frac{4\varepsilon^2}{1 + 4\varepsilon^2}}}{2}.$$

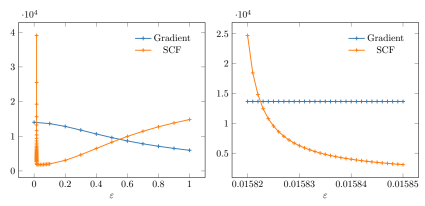
The gap $\nu(\varepsilon) := 4\sqrt{a(\varepsilon)^2 + (\sqrt{a(\varepsilon)(1-a(\varepsilon))} - \varepsilon)^2}$ goes to 0 monotonically when $\varepsilon \to 0$. In particular, for $\varepsilon \approx 0$ we have $a(\varepsilon) \approx \varepsilon^2$ and $\nu(\varepsilon) \approx 4\varepsilon^2$. This model can thus be used to study the influence of the gap on the convergence of the two algorithms.

Influence of ε on the convergence. We run Algorithms 3.1 and 3.2 with fixed β on this system. We start from a random point on the manifold \mathcal{M} . We take as a convergence criterion $\|P^k - P(a(\varepsilon))\|_{\mathcal{F}} \leq 10^{-12}$ and consider the algorithm has failed if convergence was not achieved after 50,000 iterations.

In Figure 4, we plotted the number of iterations to achieve convergence for each algorithm as a function of ε (without changing the starting point), for two different values of β : 10^{-1} and 10^{-3} . The results confirm the theory we developed in subsection 3.3: the gap has a strong influence on the convergence behavior of the SCF algorithm. Indeed, as the gap decreases, smaller and smaller damping parameters must be used, and the number of iterations increases. In fact for this system, $1 + \Omega_*^{-1} K_*$ has a single eigenvalue equal to $1 + \frac{2}{\nu(\varepsilon)} \approx 1 + \frac{1}{2\varepsilon^2}$ for ε small. Thus we expect convergence for $\beta < 4\varepsilon^2$, and therefore a critical ε_c of ≈ 0.158 for $\beta = 10^{-1}$ and 0.0158 for $\beta = 10^{-3}$, with a number of iterations proportional to $\frac{1}{\varepsilon}$ when $\varepsilon > \varepsilon_c$. The numerical results are in perfect agreement with this prediction. By contrast, the gradient algorithm is much less affected by the smallness of the gap and converges in an essentially constant number of iterations: our prediction for the convergence rate of that method is $r = 1 - \beta(\nu(\varepsilon) + 2) \approx 1 - 2\beta$ for ε small, and therefore a number of iterations for convergence to 10^{-12} of 124 for $\beta = 10^{-1}$ and 1.3×10^4 for $\beta = 10^{-3}$, again in perfect agreement with the numerical results.



(a) Number of iterations to reach convergence for both algorithms as a function of ε for $\beta = 10^{-1}$.



(b) Number of iterations to reach convergence for both algorithms as a function of ε for $\beta=10^{-3}$. On the left is a global view of the convergence for $\varepsilon\in[0,1]$ and on the right, we zoom in on the neighborhood of ε_c .

Fig. 4. Comparison of the convergence of both algorithms depending on ε for two different values of the step β .

4.3. Chaos in SCF iterations. We consider in this section another toy model, a model inspired by the one proposed in [41, section 2.1]. Let $h \in \mathbb{R}^{3\times 3}_{\text{sym}}$ and $J \in \mathbb{R}^{3\times 3}$ be the matrices defined by

$$h \coloneqq \begin{bmatrix} 1.4299 & -0.2839 & -0.4056 \\ -0.2839 & 1.1874 & 0.2678 \\ -0.4056 & 0.2678 & 2.3826 \end{bmatrix} \quad \text{and} \quad J \coloneqq h^{-1}$$

and the energy functional $E_{c_1,c_2}: \mathbb{R}^{N_b \times N_b} \to \mathbb{R}$ defined by

(4.3)
$$E_{c_1,c_2}(P) = \text{Tr}(hP) + \frac{c_1}{2} \rho_P^* J \rho_P - c_2 \sum_{j=1}^3 \rho_{P,j}^{4/3},$$

where $c_1, c_2 \in \mathbb{R}_+$ are nonnegative real parameter and where the density $\rho_P \in \mathbb{R}^3$ associated with the density matrix P is given by $\rho_{P,j} = P_{jj}$ for j = 1, 2, 3. This model is reminiscent of a Kohn–Sham LDA model, with h playing the role of the

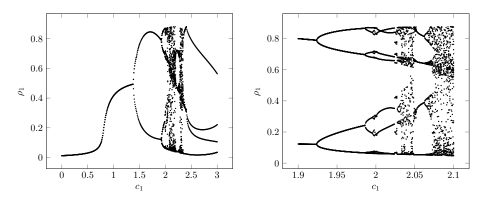


Fig. 5. Chaotic behavior of the simple SCF map for the energy functional $E_{c_1,1}$ defined by (4.3) and N=1 as a function of c_1 . On the left is a global view of the bifurcation and the right is a zoom in on part of the interval on which we observe a chaotic behavior.

core Hamiltonian, J of the Hartree operator, and the third term in E_{c_1,c_2} of the exchange-correlation energy. We seek the minimizers of E_{c_1,c_2} on \mathcal{M}_1 .

We study the behavior of the simple SCF (Roothaan) iterations $P^k = \Phi(P^k)$ with initial guess $P^0 = \phi_0 \phi_0^*$ with ϕ_0 a random vector of norm 1.

The case $c_2 = 0$. Here, the energy functional is the sum of a linear and a quadratic term. In this case, either $(P^k)_{k \in \mathbb{N}}$ converges to a critical point of the problem (in practice a local minimizer) or it has two different accumulation points P_{odd} and P_{even} , none of them being a critical point, and the iterates oscillates between the two, in the sense that $P^{2k+1} \to P_{\text{odd}}$ and $P^{2k} \to P_{\text{even}}$ when $k \to \infty$ [14, 36]. This is due to the fact that we have

$$\begin{split} P^{2k+1} &= \operatorname{argmin} \left\{ \widetilde{E}_{c_1,0}(P^{2k},P), \ P \in \mathcal{M}_1 \right\}, \\ P^{2k+2} &= \operatorname{argmin} \left\{ \widetilde{E}_{c_1,0}(P,P^{2k+1}), \ P \in \mathcal{M}_1 \right\} \end{split}$$

with

$$\widetilde{E}_{c_1,0}(P,P')\coloneqq \frac{1}{2}\operatorname{Tr}(hP)+\frac{1}{2}\operatorname{Tr}(hP')+\frac{c_1}{2}\rho_P^*J\rho_P',$$

so that (P^{2k}, P^{2k+1}) converges to a minimizer of $\widetilde{E}_{c_1,0}$ on $\mathcal{M}_1 \times \mathcal{M}_1$. When c_1 is small, the simple SCF algorithm converges: for $c_1 = 0$, the matrix h has a nondegenerate lowest eigenvalue and the algorithm converges in one iteration. When the value of c_1 increases, we observe numerically a bifurcation at a critical value $c_{1,*} \simeq 0.28$ after which the algorithm oscillates between two states.

The case $c_2 = 1$. Here the energy is not quadratic, and the previous theory does not apply. In Figure 5, we vary c_1 and plot the value of ρ_1 for the last 40 out of 1,500 SCF iterations. For this case, we still observe that the algorithm converges for c_1 small enough ($0 \le c_1 < c_{1,*} \simeq 1.38$) and oscillates between two states for c_1 slightly larger than $c_{1,*}$. However, in contrast with the $c_2 = 0$ case, this is followed by a cascade of cycles of increasing periods, transitioning to a chaotic region, following the "period-doubling route to chaos" observed for other types of chaotic systems such as the logistic map [55].

4.4. Local convergence for a 1D nonlinear Schrödinger equation. In this section, we present a simple 1D numerical experiment to validate on a more physically

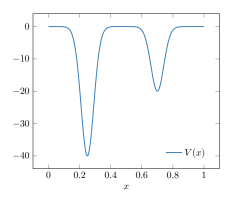


Fig. 6. V for c = 30 and C = 20.

relevant system the sharpness of the convergence rates we derived in the previous section. As the goal here is to analyze the behavior of the simplest representative of each class of algorithms when physical parameters (such as the gap) vary, we chose unpreconditioned algorithms. We consider a discretized 1D Gross-Pitaevskii model (N=1) on the torus, and its (nonphysical) fermionic counterpart for N=2. At the continuous level, the minimization set is

$$\left\{ \gamma \in \mathcal{L}(L_{\mathrm{per}}^2), \ \gamma^2 = \gamma = \gamma^*, \ \mathrm{Tr}(\gamma) = N \right\},$$

where $\mathcal{L}(L_{\mathrm{per}}^2)$ is the space of bounded operators on $L_{\mathrm{per}}^2 \coloneqq \{u \in L_{\mathrm{loc}}^2(\mathbb{R}) \mid u(\cdot - 1) = u(\cdot)\}$, and the energy functional is defined as

$$\mathcal{E}_{\alpha}(\gamma) = \operatorname{Tr}_{L_{\mathrm{per}}^2} \left(-\frac{1}{2} \Delta \gamma \right) + \int_0^1 V \rho_{\gamma} + \frac{\alpha}{2} \int_0^1 \rho_{\gamma}^2,$$

where ρ_{γ} is the density of the density matrix γ , $\alpha \in \mathbb{R}_{+}$, and V is an asymmetric double-well external potential chosen equal to

(4.4)
$$V(x) := -C\left(\exp\left(-c\cos\left(\pi(x - 0.20)\right)^2\right) + 2\exp\left(-c\cos\left(\pi(x + 0.25)\right)^2\right)\right)$$

with c = 30 and C = 20 (Figure 6).

The Euler–Lagrange equations of this minimization problem are

$$\gamma_* = \sum_{i=1}^N (\phi_i, \cdot)_{L_{\mathrm{per}}^2} \phi_i, \quad \rho_* = \sum_{i=1}^N |\phi_i|^2, \quad -\frac{1}{2} \Delta \phi_i + V \phi_i + \alpha \rho_* \phi_i = \varepsilon_i \phi_i, \quad \int_0^1 \phi_i \phi_j = \delta_{ij}.$$

We discretize this model using the finite difference method with a uniform grid of step size $\delta = 1/N_b$, which leads to the finite-dimensional model

$$\inf \left\{ E_{\alpha}(P), \ P \in \mathcal{M}_{N} \right\},\,$$

where

(4.7)
$$\forall P \in \mathcal{H}, \quad E_{\alpha}(P) := \text{Tr}(hP) + \frac{\alpha}{2} \delta \sum_{i=1}^{N_b} \left(\frac{P_{ii}}{\delta}\right)^2,$$

the nonzero entries of the matrix $h \in \mathbb{R}^{N_b \times N_b}$ being given by

$$\forall \ 1 \leq i \leq N_b, \quad h_{ii} = \frac{1}{\delta^2} + V(i\delta), \quad h_{i,i+1} = h_{i,i-1} = -\frac{1}{2\delta^2},$$

where we identify the sites 0 and N_b on the one hand and N_b+1 and 1 on the other. With this discretization, the discrete density is then given by $\rho(i\delta) \approx \rho_i := P_{ii}/\delta$. We compare the fixed-step gradient descent and damped SCF algorithms (Algorithms 3.1 and 3.2) on this problem for various values of α , using as a starting point the ground state for $\alpha=0$. The functional E is smooth. To check Assumption 2.3 we notice that E is a convex functional of P, so that $\nabla^2 E(P) \geqslant 0$. Therefore, at any local minimizer satisfying the strong Aufbau principle, $\Omega_* + K_* \geqslant \Omega_* \geqslant \eta > 0$, and therefore Assumption 2.3 is satisfied. In the case where the Aufbau principle is not satisfied, Assumption 2.3 is not a priori always satisfied, so we check it a posteriori by computing the lowest eigenvalue of $\Omega_* + K_*$.

We prove in the appendix the following lemma, which collects some of the mathematical properties of the discretized model under consideration. The proof of this lemma, given in the appendix, strongly relies on the properties of our particular model (1D difference equation with periodic boundary conditions and a specific nonlinearity).

LEMMA 4.1 (mathematical properties of (4.6)). Let $\alpha \in \mathbb{R}_+$.

- 1. For N=1, the optimization problem (4.6) has a unique minimizer P_* . In addition, P_* can be written as $P_*=\phi_*\phi_*^*$ with $\phi_*\in\mathbb{R}^{N_b}$, $\phi_*^*\phi_*=1$, and ϕ_* positive componentwise, and P_* satisfies the strong Aufbau principle.
- 2. For $2 \leqslant N \leqslant N_b$ with $N_b \neq 2N$ if $N_b \in 4\mathbb{N}^*$, the relaxed constrained optimization problem

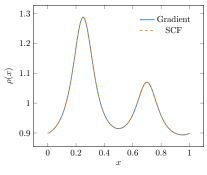
(4.8)
$$\inf \{ E_{\alpha}(P), P \in CH(\mathcal{M}_N) \},$$

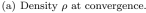
where $\operatorname{CH}(\mathcal{M}_N) = \{P \in \mathcal{H}, P = P^*, 0 \leq P \leq 1, \operatorname{Tr}(P) = N\}$ is the convex hull of \mathcal{M}_N , has a unique minimizer P_* . Either $P_* \in \mathcal{M}_N$, in which case P_* is the unique minimizer of (4.6) and satisfies the Aufbau principle, or $P_* \notin \mathcal{M}_N$, in which case the eigenvalues $\varepsilon_1 \leq \cdots \leq \varepsilon_{N_b}$ of the mean field Hamiltonian matrix $H_* = \nabla E_{\alpha}(P_*)$ satisfy $\varepsilon_{N-1} < \varepsilon_N = \varepsilon_{N+1} < \varepsilon_{N+2}$ and none of the local minimizers to (4.6) satisfy the Aufbau principle.

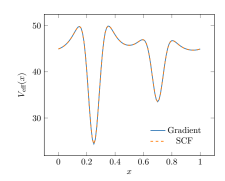
Note that the unique minimizer P_* to the relaxed constraint problem (4.8) can be computed using the ODA introduced in [13]. As shown in the proof of Lemma 4.1, the only case when the minimizer P_* is not unique is the very particular case when $N_b \in 4\mathbb{N}^*$, $N_b = 2N$, and all the entries $[V_{\text{eff}}]_i := V(i\delta) + \alpha \delta^{-1}[P_*]_{ii}$ of the effective potential are equal. In the rest of this section, we consider the cases N=1 and N=2.

The case N=1. It follows from Lemma 4.1 and Theorems 3.3 and 3.4 that Algorithms 3.1 and 3.2 locally converge to the unique minimizer P_* as long as β is chosen small enough. The resulting densities, effective potentials, and convergence behavior of both algorithms are plotted in Figure 7 for $N_b=100$. The SCF algorithm converges faster in terms of the number of iterations, as a smaller β , hence more steps, are required for the gradient to converge. This is expected from the large spectral radius of the matrix h in the absence of preconditioning.

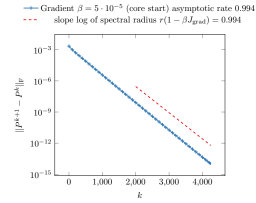
For the gradient algorithm, the convergence rate is consistent with the spectral radius of the Jacobian matrix $1 - \beta J_{\text{grad}}$. For the damped SCF algorithm with the

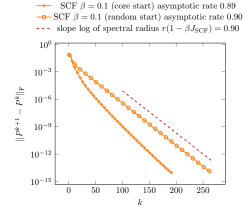






(b) Effective potential $V_{\rm eff} = V + \alpha \rho$ at convergence.



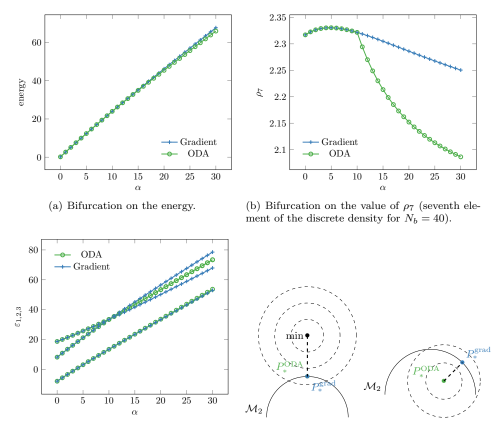


(c) Error decay for SCF and gradient descent algorithms (every few marks only are plotted for the sake of visibility). Marked lines are the evolution of the error $\|P^{k+1} - P^k\|_F$ and dashed lines represent the slope computed with the spectral radius of the Jacobian matrix, computed by finite differences.

Fig. 7. Convergence of Algorithms 3.1 and 3.2 for $N=1, \alpha=50,$ and $N_b=100.$

ground state of the core Hamiltonian as a starting point, surprisingly, we observe an asymptotic convergence rate slightly faster than that expected from the spectral radius of the Jacobian matrix $1 - \beta J_{\rm SCF}$. Using a random perturbation of the ground state of the core Hamiltonian as a starting point again gives a convergence rate consistent with the spectral radius.

The explanation of this effect is to be found in the repartition of the error among the eigenvectors of J_{SCF} . Since both Ω_* and K_* are positive semidefinite operators, the eigenvalues of $J_{\text{SCF}} = 1 + {\Omega_*}^{-1} K_*$ are greater than 1, and the convergence for β small is limited by the modes associated with eigenvalues of J_{SCF} close to 1. These eigenvalues correspond to high eigenvalues of Ω_* and therefore to highly oscillatory modes. When the initial guess is chosen as the ground state of the core Hamiltonian, these modes are only weakly excited and do not contribute to the observed convergence rate before convergence is achieved. When the initial guess is randomly perturbed, this effect is not present and the convergence rate is consistent with the spectral radius. For the gradient algorithm, the rate-limiting modes are associated with small eigenvalues of $\Omega_* + K_*$, which are not oscillatory, and this effect is not present either.



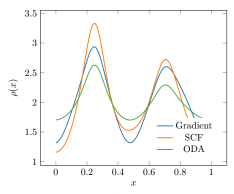
(c) Bifurcation on the first three eigenvalues (d) ODA minimum before (left) and after (right) the of the Hamiltonians $H(P_*^{\mathrm{ODA}})$ and $H(P_*^{\mathrm{grad}})$. bifurcation. Dashed lines are the energy level sets For ODA, after the collision, the second and around the minimum. third eigenvalues are degenerate.

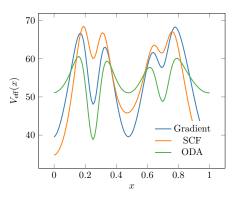
Fig. 8. Bifurcation on the energy, the density, and the eigenvalues as a function of α for $N=2,\ N_b=40.$

The case N=2. Since the second smallest eigenvalue of the matrix h is strictly lower than the third one, for α small enough, the unique minimizer P_* of (4.8) is on \mathcal{M}_2 and satisfies the strong Aufbau principle, and both the gradient descent and SCF algorithm locally converge to P_* . For larger values of α , the two alternatives of Lemma 4.1 appear. We plot the energy, the density at an arbitrarily chosen point, and the eigenvalues of the solutions P^{grad} and P^{ODA} obtained by the gradient and the ODA algorithm as a function of α in Figure 8, evidencing a bifurcation for a critical value of $\alpha_c \simeq 10$, after which these two solutions are different.

For α lower than the critical value $\alpha_c \simeq 10$, P_*^{ODA} is on the manifold \mathcal{M}_2 and satisfies the strong Aufbau principle. The algorithms all converge to this solution: $P_*^{\text{grad}} = P_*^{\text{SCF}} = P_*^{\text{ODA}} = P_*$. However, for $\alpha > \alpha_c$ in the range tested, $P_*^{\text{ODA}} \notin \mathcal{M}_2$. A geometrical interpretation of the bifurcation is sketched in Figure 8(d): the level sets of the function E_{α} are degenerate ellipsoids. Below α_c , the intersection of the nonempty closed convex set $\text{CH}(\mathcal{M}_2)$ with the level set of E_{α} of lowest energy belongs to \mathcal{M}_2 , while this is no longer the case beyond α_c .

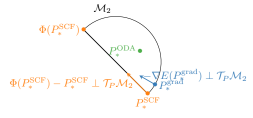
For $\alpha > \alpha_c$, the solutions obtained by the ODA, gradient, and SCF algorithm differ, as shown in Figure 9:





(a) Densities of P_*^{grad} , P_*^{SCF} , and P_*^{ODA} .

(b) Effective potential $V_{\rm eff} = V + \alpha \rho_P$ for $P = P_*^{\rm grad}$, $P_*^{\rm SCF}$, and $P_*^{\rm ODA}$.



(c) Geometrical interpretation of the limiting points
of the gradient descent, SCF and ODA algorithms.

	Gradient	SCF	ODA
ε_1	52.9	51.3	53.6
ε_2	67.9	67.8	73.4
ε_3	78.5	79.6	73.4

(d) Lowest three eigenvalues of H(P) for $P = P_*^{\text{grad}}$, $P = P_*^{\text{SCF}}$, and $P = P_*^{\text{DDA}}$.

FIG. 9. Results obtained with the gradient descent, damped SCF, and ODA algorithm for N=2, $\alpha=30$ and $N_b=100$: the limiting points $P_*^{\rm grad}$ and $P_*^{\rm SCF}$ lay by construction on the manifold \mathcal{M}_2 , while $P_*^{\rm ODA}$ does not (it only belongs to its convex hull $\mathrm{CH}(\mathcal{M}_2)$).

• The lowest second and third eigenvalues of $H(P_*^{\text{ODA}})$ are degenerate $(\varepsilon_2 = \varepsilon_3)$ and $P_*^{\text{ODA}} \notin \mathcal{M}_2$. More precisely, we have

$$P_*^{\text{ODA}} = \phi_1 \phi_1^* + (1 - f) \phi_2 \phi_2^* + f \phi_3 \phi_3^* \quad \text{with} \quad H(P_*^{\text{ODA}}) \phi_i = \varepsilon_i \phi_i, \quad \phi_i^* \phi_j = \delta_{ij}$$

with a fractional occupation 0 < f < 1.

- Algorithms 3.1 and 3.2 converge to two different limits P_*^{grad} and P_*^{SCF} , none of them satisfying the Aufbau principle:
 - P_*^{grad} is a local minimizer of E_{α} on \mathcal{M}_2 , which does not satisfy the Aufbau principle. More precisely, P_*^{grad} is the orthogonal projector on the space generated by the eigenvectors associated with the lowest first and third eigenvalues of $H(P_*^{\text{grad}})$.

 P_*^{SCF} satisfies $\Pi_{P_*^{\text{SCF}}}(\Phi(P_*^{\text{SCF}}) P_*^{\text{SCF}}) = 0$, but $\Phi(P_*^{\text{SCF}}) P_*^{\text{SCF}} \neq 0$
 - $-P_*^{\text{SCF}}$ satisfies $\Pi_{P_*^{\text{SCF}}}\left(\Phi(P_*^{\text{SCF}}) P_*^{\text{SCF}}\right) = 0$, but $\Phi(P_*^{\text{SCF}}) P_*^{\text{SCF}} \neq 0$ and $[H(P_*^{\text{SCF}}), P_*^{\text{SCF}}] \neq 0$. The iterates are trapped as the search direction is orthogonal to the tangent space (Figure 9(c)). The limit point P_*^{SCF} is a spurious stationary state of the SCF iteration which is not physically relevant, not being a critical point of E_α .
- 4.5. Kohn–Sham density functional theory. We now investigate a more realistic computation: the electronic structure of a silicon crystal using KS-DFT. We used the DFTK.jl code [28], which solves the equations of KS-DFT in a plane-wave

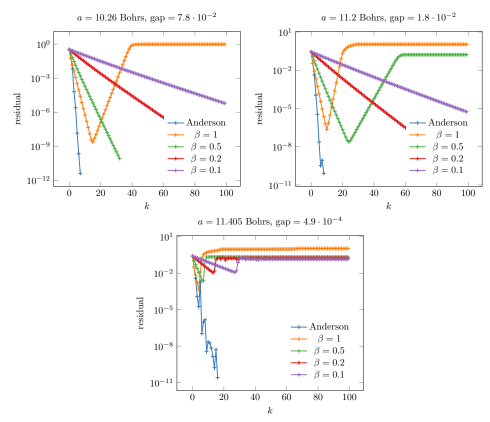


Fig. 10. Convergence curves of the density residual as a function of the number of iterations k for silicon with different lattice constants a.

basis under a pseudopotential approximation. All computations below use the LDA of the exchange-correlation energy [33, 43], Goedecker–Teter–Hutter pseudopotentials [22], a cutoff energy of 30 Hartrees, and Γ -only Brillouin zone sampling for simplicity, although the same behavior was observed with different exchange-correlation functionals and fine Brillouin zone discretizations. In all cases, the initial guess for the algorithms is a superposition of atom-centered densities. The DFTK code as well as the script used to produce these results are available at https://dftk.org/.

We consider the case of silicon in its standard face-centered cubic phase. With the chosen pseudopotentials and without spin polarization, silicon has four occupied orbitals: N=4. We examine the convergence of algorithms as a function of the lattice constant a (the size of the computational domain). In the equilibrium state of silicon $(a\approx 10.26 \text{ Bohrs})$, there is a gap of about 0.08 Hartree between the occupied and virtual states. As the lattice constant a is increased, this gap decreases, until it closes at $a\approx 11.408$ Bohrs. We examine the convergence of self-consistent iterations, using both fixed-step damped density mixing with four values of the mixing parameter β ($\beta=1$ (no damping), $\beta=0.5$, $\beta=0.2$, $\beta=0.1$), as well as the self-consistent iteration accelerated with Anderson acceleration (also known as Pulay's DIIS method [16, 47, 48]). In Figure 10, we plot the convergence of the density residual $\|\rho_{\Phi(P_n)}-\rho_{P_n}\|_2$ as a function of the iterations for three values of a, with decreasing gaps.

In the first case, with a=10.26 Bohrs, the simple (undamped) SCF method appears to be converging for almost 20 iterations but then diverges, until the density

residual stabilizes at a positive value, as predicted in [14] for the Hartree–Fock model. The damped methods appear to converge. For a=11.2 Bohrs, the damping method with $\beta=0.5$ does not converge. When the lattice constant is further increased to 11.405 Bohrs, with a small gap of 4.9×10^{-4} Hartree, the fixed-step damped SCF iterations do not converge for the tested values of the damping parameter ($\beta=1,0.5,0.2,0.1$).

When it occurs, the transient behavior of apparent-convergence before eventual divergence is unusually long. For instance, for $\beta=1$ at a=10.26 Bohrs, the method appears to be converging for almost 20 iterations, up to a reduction in residual of a factor 10^{-8} . In fact, it is consistent with an initial error of the order of machine precision (about 10^{-16} here) being amplified at a constant rate. The cause of this effect appears to be that the divergent modes break the natural inversion symmetry of the crystal in this particular case: we have checked that the divergence occurs much sooner if we break this symmetry by perturbing the positions of the atoms around their symmetric positions (at 9 iterations by perturbing the position of one atom by 10%). In practice, in the symmetric case, one way to overcome this issue is to ensure during the algorithm that, at each step, we have a symmetric solution. Note that this phenomenon is reminiscent of that observed in Figure 7, where all the modes were not fully excited, making the convergence faster than expected.

It is remarkable that the convergent methods (and even the divergent ones before their divergence) appear to have the exact same slope as with a=10.26 Bohrs. This is consistent with our result: assuming the main effect of increasing the lattice constant is to decrease the gap, while keeping the lowest eigenvalues of $\Omega_*^{-1}K_*$ constant, then for β small enough the convergence is limited by the lowest, not the highest, eigenvalues of this operator.

In all these cases, Anderson acceleration was able to converge to a solution, even in the presence of a very small gap, albeit in an irregular fashion. We attribute this to the well-known fact that, in the linear regime, Anderson acceleration is equivalent to the GMRES algorithm. Since the GMRES algorithm is a Krylov method, it is robust to the presence of a large eigenvalue and achieves convergence even though the underlying iteration is strongly divergent. This shows a limitation of our theoretical convergence rates, which do not capture the reduced sensitivity of accelerated methods to a small gap.

5. Conclusion. In this paper, we examined the convergence of two simple representatives in the class of direct minimization and SCF algorithms. We showed that both algorithms converge locally when the damping parameter is chosen small enough. We derived their convergence rates; we showed that the damped SCF algorithm is sensitive to the gap, while the gradient method is sensitive to the spectral radius of the Hamiltonian. We confirmed these results with numerical experiments. The goal here was not to propose efficient algorithms but to analyze the behavior of the simplest representatives of each class. However, accelerated algorithms are generally found to follow the trend suggested by our theoretical results, although we showed that the Anderson-accelerated SCF algorithm was able to converge quickly even in the presence of a single very small gap.

In practice, should the SCF or direct minimization class of algorithms be preferred? The answer depends not only on the convergence rate studied in this paper but also on the cost of each step and the robustness of the algorithm. We examine two prototypical situations.

In quantum chemistry using Gaussian basis sets to solve the Hartree–Fock model or KS-DFT using hybrid functionals, the rate-limiting step is often the computation of the Fock matrix H(P). In this case, an iteration of a gradient descent and a damped SCF algorithm are of roughly equal cost. In most cases, solutions for isolated molecules satisfy the Aufbau principle, and the damped SCF algorithm, suitably robustified (for instance, using the ODA algorithm) and accelerated (for instance, with the DIIS algorithm), converges reliably and efficiently toward a solution. Direct minimization algorithms are then only useful in the cases where local or semilocal functionals are used [52] and the Aufbau principle is violated, or when SCF algorithms tend to converge to saddle points (for instance, for computations involving spin).

In condensed-matter physics using plane-wave basis sets to solve the KD-DFT with local or semilocal functionals, the matrices P and H are not stored explicitly. Solving the linear eigenproblem is then done using iterative block eigensolvers, which can be understood as specialized direct minimization algorithms in the case of a linear energy functional $E(P) = \text{Tr}(H_0P)$. In this case, direct minimization algorithms effectively merge the two loops of the SCF and linear eigensolver and should therefore be more efficient. Another interest of direct minimization algorithms is their robustness, as the choice of a step size can be made in order to minimize the energy, unlike the damped SCF algorithm, where choosing an appropriate damping parameter is often done empirically.

Despite this, direct minimization algorithms are rarely used in condensed-matter physics. The main reason seems to be that challenging problems are often metallic in character and require the introduction of a finite temperature. Direct minimization algorithms then need to optimize over the occupations as well as the orbitals, a significantly more complex task [9, 21, 44]. A thorough comparison of the performance and robustness of direct minimization and a self-consistent approach for these systems would be an interesting topic of inquiry. A number of implementation "tricks" commonly used to accelerate the convergence of iterative eigensolvers (for instance, using a block size larger than the number of eigenvectors sought) might also play a large role in performance comparison for the two classes of algorithms: understanding how to generalize these to direct minimization would be interesting.

We discussed in Remark 3.7 preconditioning for both direct minimization and SCF algorithms. The concept of preconditioning for Riemannian optimization problems seems not to have been explored much in the mathematical literature, except in some specific models and preconditioners (see, for instance, [7, 61] for the Gross–Pitaevskii model), and a deeper analysis of this would be interesting. In particular, this is necessary to extend the convergence theory presented in this paper to infinite-dimensional settings.

Appendix: Proof of Lemma 4.1. For any $\alpha \in \mathbb{R}_+$ the energy functional E_{α} is smooth and convex on the nonempty compact convex set $\mathrm{CH}(\mathcal{M}_N)$. The set of minimizers to (4.8) is therefore nonempty, compact, and convex. Let P_* and P'_* be two minimizers of E_{α} and let ρ_* and ρ'_* be their densities: $\rho_{*,i} = \delta^{-1}(P_*)_{ii}$ and $\rho'_{*,i} = \delta^{-1}(P'_*)_{ii}$. For all $\theta \in [0,1]$, we have

$$I_{\alpha} = E_{\alpha}(\theta P_* + (1 - \theta)P'_*) = I_{\alpha} + \frac{\alpha}{2\delta} \sum_{i=1}^{N_b} \left((\theta \rho_{*,i} + (1 - \theta)\rho'_{*,i})^2 - (\theta \rho_{*,i}^2 + (1 - \theta)\rho'_{*,i}^2) \right),$$

where $I_{\alpha} = E_{\alpha}(P_*) = E_{\alpha}(P'_*)$ is the minimum of (4.8). Since the function $\mathbb{R} \ni x \mapsto x^2 \in \mathbb{R}$ is strictly convex, we obtain that $\rho_* = \rho'_*$. Therefore, all the minimizers of (4.8) share the same density, hence the same mean-field Hamiltonian matrix H_* . If P_* is a minimizer of (4.8), it satisfies the first-order optimality condition (Euler

inequality)

$$\forall P \in CH(\mathcal{M}_N), Tr(H_*(P - P_*)) \geqslant 0,$$

from which we infer by a classical argument that (5.1)

$$P_* = \mathbf{1}_{(-\infty,\mu)}(H_*) + Q_*$$
 with $0 \leqslant Q_* \leqslant 1$, $\operatorname{Ran}(Q_*) \subset \operatorname{Ker}(H_* - \mu)$, $\operatorname{Tr}(P_*) = N$,

for some Fermi level $\mu \in \mathbb{R}$ (the Lagrange multiplier of the constraint $\operatorname{Tr}(P) = N$). Let $\varepsilon_1 \leqslant \cdots \leqslant \varepsilon_{N_b}$ be the eigenvalues of H_* , counting multiplicities. If $\varepsilon_N < \varepsilon_{N+1}$, then we necessarily have $P_* = \mathbf{1}_{(-\infty,\varepsilon_N]}(H_*)$, so that (4.8) has a unique minimizer, P_* is on \mathcal{M}_N and therefore is also the unique minimizer of (4.6), and it satisfies the strong Aufbau principle.

Let us now consider the case when $\varepsilon_N = \varepsilon_{N+1} =: \mu$. Since the eigenvalue problem $H_*\psi = \mu\psi$ is a second-order difference equation,

(5.2)
$$\frac{-\psi_{i+1} + 2\psi_i - \psi_{i-1}}{2\delta^2} + V_{\text{eff},i}\psi_i = \mu\psi_i, \quad 1 \leqslant i \leqslant N_b$$

(here and in what follows we use the convention that $\psi_0 = \psi_{N_b}$ and $\psi_{N_b+1} = \psi_1$) with $V_{\text{eff}} = V + \alpha \rho_*$, the eigenspace $\text{Ker}(H_* - \mu)$ is at most of dimension 2. We therefore have $\varepsilon_{N-1} < \varepsilon_N = \varepsilon_{N+1} < \varepsilon_{N+2}$.

Using the variational characterization of the ground state eigenvalue, we have

(5.3)
$$\varepsilon_1 = \min_{\psi \in \mathbb{R}^{N_b}, \, \psi^* \psi = 1} \psi^* H_* \psi$$
 with $\psi^* H_* \psi = \sum_{i=1}^{N_b} \left| \frac{\psi_{i+1} - \psi_i}{\delta} \right|^2 + \sum_{i=1}^{N_b} V_{\text{eff}, i} |\psi_i|^2$.

Since $||x| - |y|| \le |x - y|$ for all $x, y \in \mathbb{R}$ with equality if and only if x and y have the same sign, we infer from (5.3) that all the entries of a ground state eigenvector of H_* have the same sign. In particular, two normalized ground state eigenvectors of H_* cannot be orthogonal. This implies that the ground state eigenvalue of H_* is simple, i.e., $\varepsilon_1 < \varepsilon_2$. The first statement of Lemma 4.1 staightforwardly follows from the results established so far.

To prove the second statement, assume that $N \ge 2$ and that (4.8) has two distinct minimizers P_* and P'_* sharing the same density. In view of (5.1), this can only occur if $\varepsilon_N = \varepsilon_{N+1} =: \mu$. Using an orthonormal basis (ϕ, ψ) of $\operatorname{Ker}(H_* - \mu)$ consisting of eigenvectors of P_* , we can assume without loss of generality that

$$P_* = \mathbf{1}_{(-\infty,\mu)}(H_*) + (1-f)\phi\phi^* + f\psi\psi^*,$$

$$P_*' = \mathbf{1}_{(-\infty,\mu)}(H_*) + (1-a)\phi\phi^* + a\psi\psi^* + b(\phi\psi^* + \psi\phi^*)$$

with $0 \leqslant f \leqslant 1$, $0 \leqslant a \leqslant 1$, and $b^2 \leqslant a(1-a)$. Since P_* and P_*' have the same density, we have for all $1 \leqslant i \leqslant N_b$, $(1-f)\phi_i^2 + f\psi_i^2 = (1-a)\phi_i^2 + a\psi_i^2 + 2b\phi_i\psi_i$, that is, $(a-f)\phi_i^2 - 2b\phi_i\psi_i - (a-f)\psi_i^2 = 0$.

If a=f, then $b\neq 0$ since $P_*\neq P'_*$ by assumption, so that $\phi_i\psi_i=0$ for all $1\leqslant i\leqslant N_b$. From (5.2), we see that it is not possible to have $\psi_i=\psi_{i+1}=0$ (otherwise, ψ would be identically equal to zero), and the same holds true for ϕ . Therefore, N_b must be even, and either all the odd entries of ϕ and all the even entries ψ must vanish, or the other way round. We then infer from (5.2) that this implies that $\phi_{i+2}+\phi_i=\psi_{i+2}+\psi_i=0$ for all $1\leqslant i\leqslant N_b$ and that all the entries

of $V_{\rm eff}$ are equal to $\mu - \delta^{-2}$. This implies that $N_b \in 4\mathbb{N}^*$ and that the states ϕ and ψ are given by $\phi_{2i} = c(-1)^i$, $\phi_{2i+1} = 0$, $\psi_{2i} = 0$, $\psi_{2i+1} = c'(-1)^i$ for all $1 \leq i \leq N_b$, where c and c' are normalization constants. By explicit diagonalization of the matrix $H_* = H(0) + (\mu - \delta^{-2})I_{N_b}$ one can check that the states ϕ and ψ are therefore those spanning the 2D space associated to the twofold degenerate eigenvalues $\varepsilon_{N_b/2} = \varepsilon_{1+N_b/2}$ of H_* . This is only possible if $N = N_b/2$. The case $\alpha = f$ can thus be excluded for $2 \leq N \leq N_b$ with $N_b \neq 2N$ if $N_b \in 4\mathbb{N}^*$.

If $a \neq f$, we have for all $1 \leqslant i \leqslant N_b$, $\phi_i^2 - 2\gamma\phi_i\psi_i - \psi_i^2 = 0$, for $\gamma = \frac{b}{a-f}$, and up to replacing ψ with $-\psi$, that we can assume without loss of generality that $\gamma \geqslant 0$. Denoting by $C_{\pm} := \gamma \pm \sqrt{1 + \gamma^2}$ the roots of the polynomial $x^2 - 2\gamma x + 1$ with $C_+C_- = -1$, we obtain that for each $1 \leqslant i \leqslant N_b$, either $\phi_i = C_+\psi_i$ or $\phi_i = C_-\psi_i$. Using the discrete Schrödinger equation (5.2) satisfied by both ϕ and ψ , we see that if $\phi_i = C_+\psi_i$ and $\phi_{i+1} = C_+\psi_{i+1}$ for some $1 \leqslant i \leqslant N_b$, then $\phi = C_+\psi$, and likewise if C_+ is replaced by C_- . This is impossible since ϕ and ψ are orthonormal. Therefore, we must have $\phi_{2i} = C_+\psi_{2i}$ and $\phi_{2i+1} = C_-\psi_{2i+1}$ (or the other way around), and N_b must be even. Using again (5.2), this leads to $\phi_{i+2} + \phi_i = 0$ and $\psi_{i+2} + \psi_i = 0$ for all $1 \leqslant i \leqslant N_b$ and therefore, as in the previous case, that $N_b \in 4\mathbb{N}^*$, $N_b = 2N$ that all the entries of V_{eff} are equal, and that ϕ and ψ span the 2D space associated to the twofold degenerate eigenvalues $\varepsilon_N = \varepsilon_{N+1}$ of H_* .

This proves that for $2 \leq N \leq N_b$ with $N_b \neq 2N$ if $N_b \in 4\mathbb{N}^*$, (4.8) has a unique minimizer P_* . If $P_* \in \mathcal{M}_N$, it is of course also the unique minimizer of (4.6), and P_* satisfies the Aufbau principle. Conversely, if $P'_* \in \mathcal{M}_N$ is a local minimizer of (4.6) satisfying the Aufbau principle, we have

$$\forall P \in CH(\mathcal{M}_N), \quad Tr(H(P'_*)(P - P'_*)) \geqslant 0,$$

which means that P'_* is a solution to the Euler inequality for (4.8), and therefore a global minimizer of this convex problem. Since the minimizer P_* of (4.8) is unique, we finally obtain that if $P_* \notin \mathcal{M}_N$, then none of the local minimizers of (4.6) satisfies the Aufbau principle.

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REFERENCES

- P.-A. ABSIL, R. MAHONY, AND R. SEPULCHRE, Optimization Algorithms on Matrix Manifolds, Princeton University Press, Princeton, NJ, 2008.
- [2] F. Alouges and C. Audouze, Preconditioned gradient flows for nonlinear eigenvalue problems and application to the Hartree-Fock functional, Numer. Methods Partial Differential Equations, 25 (2009), pp. 380–400.
- [3] X. Antoine, A. Levitt, and Q. Tang, Efficient spectral computation of the stationary states of rotating Bose-Einstein condensates by preconditioned nonlinear conjugate gradient methods, J. Comput. Phys., 343 (2017), pp. 92–109.
- [4] V. Bach, E. Lieb, L. M., and J. Solovej, There are no unfilled shells in unrestricted Hartree-Fock theory, Phys. Rev. Lett., 72 (1994), pp. 2981–2983.
- [5] G. BACSKAY, A quadratically convergent Hartree-Fock (QC-SCF) method. Application to closed shell systems, Chem. Phys., 61 (1981), pp. 385–404.
- [6] W. BAO AND Y. CAI, Mathematical theory and numerical methods for Bose-Einstein condensation, Kinet. Relat. Models, 6 (2013), pp. 1–135.
- [7] W. BAO AND Q. Du, Computing the ground state solution of Bose-Einstein condensates by a normalized gradient flow, SIAM J. Sci. Comput., 25 (2004), pp. 1674–1697.
- [8] J. W. MILNOR, Topology from the Differentiable Viewpoint, Princeton University Press, Princeton, NJ, 1997.

- [9] E. CANCÈS, Self-consistent field algorithms for Kohn-Sham models with fractional occupation numbers, J. Chem. Phys., 114 (2001), pp. 10616-10622.
- [10] E. CANCÈS, R. CHAKIR, AND Y. MADAY, Numerical analysis of nonlinear eigenvalue problems, J. Sci. Comput., 45 (2010), pp. 90–117.
- [11] E. CANCÈS, R. CHAKIR, AND Y. MADAY, Numerical analysis of the planewave discretization of some orbital-free and Kohn-Sham models, ESAIM Math. Model. Numer. Anal., 46 (2012), pp. 341–388.
- [12] E. CANCÈS, M. DEFRANCESCHI, W. KUTZELNIGG, C. LE BRIS, AND Y. MADAY, Computational Quantum Chemistry: A Primer, Handb. Numer. Anal., North-Holland, Amsterdam, 2003.
- [13] E. CANCÈS AND C. LE BRIS, Can we outperform the DIIS approach for electronic structure calculations?, Int. J. Quantum Chem., 79 (2000), pp. 82–90.
- [14] E. CANCÈS AND C. LE BRIS, On the convergence of SCF algorithms for the Hartree-Fock equations, ESAIM Math. Model. Numer. Anal., 34 (2000), pp. 749–774.
- [15] G. CHABAN, M. SCHMIDT, AND M. GORDON, Approximate second order method for orbital optimization of SCF and MCSCF wavefunctions, Theoret. Chem. Accounts, 97 (1997), pp. 88-95.
- [16] M. CHUPIN, M.-S. DUPUY, G. LEGENDRE, AND E. SÉRÉ, Convergence Analysis of Adaptive DIIS Algorithms with Application to Electronic Ground State Calculations, 2020, https://arxiv.org/abs/2002.12850.
- [17] X. Dai, Z. Liu, L. Zhang, and A. Zhou, A conjugate gradient method for electronic structure calculations, SIAM J. Sci. Comput., 39 (2017), pp. A2702–A2740.
- [18] I. Danaila and B. Protas, Computation of ground states of the Gross-Pitaevskii functional via Riemannian optimization, SIAM J. Sci. Comput., 39 (2017), pp. B1102-B1129.
- [19] P. Dederichs and R. Zeller, Self-consistency iterations in electronic-structure calculations, Phys. Rev. B, 28 (1983), 5462.
- [20] A. EDELMAN, T. ARIAS, AND S. SMITH, The geometry of algorithms with orthonormality constraints, SIAM J. Matrix Anal. Appl., 20 (1998), pp. 303-353.
- [21] C. FREYSOLDT, S. BOECK, AND J. NEUGEBAUER, Direct minimization technique for metals in density functional theory, Phys. Rev. B, 79 (2009), 241103.
- [22] S. GOEDECKER, M. TETER, AND J. HUTTER, Separable dual-space Gaussian pseudopotentials, Phys. Rev. B, 54 (1996), 1703.
- [23] X. Gonze, Towards a potential-based conjugate gradient algorithm for order-N self-consistent total energy calculations, Phys. Rev. B, 54 (1996), 4383.
- [24] D. R. HARTREE, The wave mechanics of an atom with a non-Coulomb central field. Part II. Some results and discussion, Math. Proc. Cambridge Philos. Soc., 24 (1928), pp. 111–132.
- [25] P. Heid, B. Stamm, and T. P. Wihler, Gradient Flow Finite Element Discretizations with Energy-Based Adaptivity for the Gross-Pitaevskii Equation, 2019, https://arxiv.org/abs/ 1996/96954
- [26] T. Helgaker, P. Jørgensen, and J. Olsen, Molecular Electronic-Structure Theory, John Wiley & Sons, New York, 2014.
- [27] P. Henning and D. Peterseim, Sobolev Gradient Flow for the Gross-Pitaevskii Eigenvalue Problem: Global Convergence and Computational Efficiency, 2018, https://arxiv.org/abs/ 1812.00835.
- [28] M. F. HERBST AND A. LEVITT, The Density Functional ToolKit, DFT software, https://dftk.org, 2020, https://doi.org/10.5281/zenodo.
- [29] N. J. Higham, Functions of Matrices, Other Titles Appl. Math., SIAM, Philadelphia, 2008.
- [30] R. B. HOLMES, A formula for the spectral radius of an operator, Amer. Math. Monthly, 75 (1968), pp. 163–166.
- [31] D. Johnson, Modified Broyden's method for accelerating convergence in self-consistent calculations, Phys. Rev. B, 38 (1988), pp. 12807–12813.
- [32] T. Kato, Perturbation Theory for Linear Operators, 2nd ed., Class. Math., Springer-Verlag, Berlin, 1995.
- [33] W. Kohn and L. J. Sham, Self-consistent equations including exchange and correlation effects, Phys. Rev., 140 (1965), pp. A1133–A1138.
- [34] G. Kresse and J. Furthmüller, Efficient iterative schemes for ab initio total-energy calculations using a plane-wave basis set, Phys. Rev. B, 54 (1996), pp. 11169-11186.
- [35] K. Kudin, G. Scuseria, and E. Cancès, A black-box self-consistent field convergence algorithm: One step closer, J. Chem. Phys., 116 (2002), pp. 8255–8261.
- [36] A. LEVITT, Convergence of gradient-based algorithms for the Hartree-Fock equations, ESAIM Math. Model. Numer. Anal., 46 (2012), pp. 1321–1336.
- [37] L. LIN AND J. LU, A Mathematical Introduction to Electronic Structure Theory, SIAM, Philadelphia, 2019.

- [38] L. LIN, J. LU, AND L. YING, Numerical methods for Kohn-Sham density functional theory, Acta Numer., 28 (2019), pp. 405-539.
- [39] X. LIU, X. WANG, Z. WEN, M. ULBRICH, AND Y. YUAN, On the analysis of the discretized Kohn-Sham density functional theory, SIAM J. Numer. Anal., 53 (2015), pp. 1758–1785.
- [40] X. LIU, X. WANG, Z. WEN, AND Y.-X. YUAN, On the convergence of the self-consistent field iteration in Kohn-Sham Density Functional Theory, SIAM J. Matrix Anal. Appl., 35 (2013), pp. 546-558.
- [41] X. LIU, Z. WEN, X. WANG, M. ULBRICH, AND Y. YUAN, On the analysis of the discretized Kohn-Sham density functional theory, SIAM J. Numer. Anal., 53 (2015), pp. 1758–1785.
- [42] L. Marks and D. Luke, Robust mixing for ab initio quantum mechanical calculations, Phys. Rev. B, 78 (2008), 075114.
- [43] R. M. MARTIN, Electronic Structure: Basic Theory and Practical Methods, Cambridge University Press, Cambridge, 2004.
- [44] N. MARZARI, D. VANDERBILT, AND M. C. PAYNE, Ensemble density-functional theory for ab initio molecular dynamics of metals and finite-temperature insulators, Phys. Rev. Lett., 79 (1997), 1337.
- [45] R. McWeeny, The density matrix in self-consistent field theory. I. Iterative construction of the density matrix, Proc. A, 235 (1956), pp. 496–509.
- [46] A. Mostofi, P. Haynes, C.-K. Skylaris, and M. Payne, Preconditioned iterative minimization for linear-scaling electronic structure calculations, J. Chem. Phys., 119 (2003), pp. 8842–8848.
- [47] P. Pulay, Convergence acceleration of iterative sequences. The case of SCF iteration, Chem. Phys. Lett., 73 (1980), pp. 393–398.
- [48] P. Pulay, Improved SCF convergence acceleration, J. Comput. Chem., 3 (1982), pp. 556–560.
- [49] D. RACZKOWSKI, A. CANNING, AND L. W. WANG, Thomas-Fermi charge mixing for obtaining self-consistency in density functional calculations, Phys. Rev. B, 64 (2001), 121101.
- [50] T. ROHWEDDER AND R. SCHNEIDER, An analysis for the DIIS acceleration method used in quantum chemistry calculations, J. Math. Chem., 49 (2011), 1889.
- [51] C. ROOTHAAN, New developments in molecular orbital theory, Rev. Modern Phys., 23 (1951), pp. 69–89.
- [52] E. Rudberg, Difficulties in applying pure Kohn-Sham density functional theory electronic structure methods to protein molecules, J. Phys. Condensed Matter, 24 (2012), 072202.
- [53] Y. SAAD, J. R. CHELIKOWSKY, AND S. M. SHONTZ, Numerical methods for electronic structure calculations of materials, SIAM Rev., 52 (2010), pp. 3–54.
- [54] G. Srivastava, Broyden's method for self-consistent field convergence acceleration, J. Phys. A, 17 (1984), pp. L317–L321.
- [55] S. STROGATZ, Nonlinear Dynamics and Chaos: With Applications to Physics, Biology, Chemistry, and Engineering, CRC Press, Boca Raton, FL, 2001.
- [56] P. UPADHYAYA, E. JARLEBRING, AND E. RUBENSSON, A density matrix approach to the convergence of the self-consistent field iteration, Numer. Algebra Control Optim., 11 (2021), pp. 99–115.
- [57] E. VECHARYNSKI, C. YANG, AND J. E. PASK, A projected preconditioned conjugate gradient algorithm for computing a large invariant subspace of a Hermitian matrix, J. Comput. Phys., 290 (2015), pp. 73–89.
- [58] N. WOODS, M. PAYNE, AND P. HASNIP, Computing the self-consistent field in Kohn-Sham density functional theory, J. Phys. Condensed Matter, 31 (2019), 453001.
- [59] C. Yang, W. Gao, and J. C. Meza, On the convergence of the self-consistent field iteration for a class of nonlinear eigenvalue problems, SIAM J. Matrix Anal. Appl., 30 (2009), pp. 1773–1788.
- [60] X. ZHANG, J. ZHU, Z. WEN, AND A. ZHOU, Gradient type optimization methods for electronic structure calculations, SIAM J. Sci. Comput., 36 (2014), pp. 265–289.
- [61] Z. Zhang, Exponential Convergence of Sobolev Gradient Descent for a Class of Nonlinear Eigenproblems, arXiv:1912.02135, 2019.
- [62] Z. Zhao, Z. Bai, and X. Jin, A Riemannian Newton algorithm for nonlinear eigenvalue problems, SIAM J. Matrix Anal. Appl., 36 (2015), pp. 752–774.