

Nonuniqueness in spin-density-functional theory on lattices

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(Received 19 May 2005; published 2 August 2005)

In electronic many-particle systems, the mapping between densities and spin magnetizations, $\{n(\mathbf{r}), \mathbf{m}(\mathbf{r})\}$, and potentials and magnetic fields, $\{v(\mathbf{r}), \mathbf{B}(\mathbf{r})\}$, is known to be nonunique, which has fundamental and practical implications for spin-density-functional theory (SDFT). This paper studies the nonuniqueness (NU) in SDFT on arbitrary lattices. Two new, nontrivial cases are discovered, here called *local saturation* and *global noncollinear* NU, and their properties are discussed and illustrated. In the continuum limit, only some well-known special cases of NU survive.

DOI: 10.1103/PhysRevB.72.073102

PACS number(s): 71.15.Mb, 75.10.Lp, 31.15.Ew

Density-functional theory (DFT)¹⁻³ is a widely used approach for calculating the electronic structure of atoms, molecules, and many types of materials. The fundamental theorem of Hohenberg and Kohn¹ establishes that the ground-state density $n(\mathbf{r})$ of a nonmagnetic, nondegenerate electronic system uniquely determines the scalar potential $v(\mathbf{r})$, apart from an arbitrary additive constant C . However, many applications of interest involve electronic ground states that have a spin magnetization, $\mathbf{m}(\mathbf{r})$, and/or are exposed to external magnetic fields, $\mathbf{B}(\mathbf{r})$. Such situations can be handled with spin-density-functional theory (SDFT),⁴⁻⁹ where the fundamental variable is the four-density $\{n(\mathbf{r}), \mathbf{m}(\mathbf{r})\}$, which couples to the four-potential $\{v(\mathbf{r}), \mathbf{B}(\mathbf{r})\}$.

Using the Rayleigh-Ritz variational principle, one can prove that a given physical four-density uniquely determines the ground-state wave function Ψ_0 (apart from trivial phase factors). This guarantees that any property of the system expressible in terms of Ψ_0 is a functional of the four-density. Following the Hohenberg-Kohn theorem of DFT, one might expect that there is also a unique map from ground-state wave functions to four-potentials. However, it was recognized long ago by von Barth and Hedin,⁴ and more recently by Capelle and Vignale⁵ and Eschrig and Pickett⁶ that such a unique correspondence does not exist. These authors showed that the extent of nonuniqueness (NU) in SDFT is much greater than the addition of a mere constant C to $v(\mathbf{r})$ for the nonmagnetic systems considered in DFT. As a consequence, some applications of SDFT such as the calculation of excitation energies or of one-electron spin gaps in half-metallic ferromagnets must be critically reexamined.

NU in SDFT means that an N -particle ground state Ψ_0 remains unchanged under addition of a four-potential $\{\Delta v(\mathbf{r}), \Delta \mathbf{B}(\mathbf{r})\}$. This happens if and only if Ψ_0 is an eigenstate of the operator⁵

$$\Delta H = \sum_{j=1}^N [\Delta v(\mathbf{r}_j) - \Delta \mathbf{B}(\mathbf{r}_j) \cdot \vec{\sigma}_j], \quad (1)$$

where $\vec{\sigma}_j$ is the vector of Pauli matrices acting on the spin of the j th electron, and we use units where the Bohr magneton $\mu_B = 1$. The entire spectrum $\{\Psi\}$ is invariant if ΔH is related

to a constant of motion (*systematic* NU); all other cases are called *accidental* NU.⁵ Similar conditions for NU can be given for other multicomponent generalizations of DFT, e.g., for current-carrying or superconducting systems.¹⁰ Thus, NU appears to be a characteristic feature of generalized DFTs, and its fundamental and practical consequences need to be explored in detail. In the first place, it is important to know what types of NU can occur in practice. The following classes of examples have been identified in SDFT:

(a) $\{\Delta v = C, \Delta \mathbf{B} = B \hat{e}_z\}$, with constant B , for systems with an energy gap and a collinear spin arrangement along the z axis, where the $\{\Psi\}$ are eigenstates of \hat{S}_z (*systematic* NU). B should be sufficiently small to avoid level crossings in order for Ψ_0 to remain the ground state. As shown by Gidopoulos,⁸ the mapping in the collinear case is unique in the broader sense that spin-potentials $\{v_\uparrow(\mathbf{r}), v_\downarrow(\mathbf{r})\}$ which differ by more than a spin-dependent constant always have different ground states.

(b) In a fully spin polarized Kohn-Sham system with $n_\uparrow(\mathbf{r}) = n(\mathbf{r})$ and $n_\downarrow = 0$ and an energy gap, there is an infinite number of spin-down Kohn-Sham potentials that produce the same ground state (*accidental* NU).

(c) $\{\Delta v = \lambda u(\mathbf{r}), \Delta \mathbf{B} = \lambda u(\mathbf{r}) \mathbf{m}(\mathbf{r}) / m(\mathbf{r})\}$ for one-electron systems only, where $u(\mathbf{r})$ is an arbitrary function and λ is sufficiently small (*accidental* NU).⁴

In Ref. 6, a general condition for NU in N -electron systems is given: If Ψ_0 is invariant under the addition of a four-potential, then this four-potential must have the form $\{\Delta v(\mathbf{r}) = C, \Delta \mathbf{B}(\mathbf{r}) = B \hat{e}(\mathbf{r})\}$, i.e., $\Delta \mathbf{B}$ may possibly be noncollinear but must be constant in magnitude. However, Argaman and Makov⁷ have raised doubts whether such noncollinear four-potentials can really be found.

It thus appears that NU for $N > 1$ electrons, while interesting and of potential practical relevance, is limited to rather simple situations of collinear spin arrangement or full, ferromagnetic spin polarization. The purpose of this paper is to examine NU in SDFT for arbitrary lattice systems. This has technical advantages over dealing with continuum systems since one can use linear algebra methods in finite vector spaces. We discover two new, nontrivial classes of NU for

N -electron systems with noncollinear spins, both of the “accidental” type. These examples require the ground state four-density to satisfy certain constraints on the lattice. In the continuum limit, we show that only some well-known special cases survive, and we discuss consequences for practical applications.

We consider a noninteracting N -electron system on a finite-size lattice with P lattice points whose specific geometry is not important for the following. We assume that the kinetic-energy operator \hat{T} has been suitably discretized on this lattice (in practice, we use a finite-difference approach). The single-particle wave functions ψ_j obey the following Schrödinger equation:

$$[\hat{T} + \hat{V} - \hat{\mathbf{B}} \cdot \vec{\sigma}] \psi_j = E_j \psi_j, \quad j = 1, \dots, 2P. \quad (2)$$

For the spatial part of the wave functions we use a localized basis, $\varphi_{ik} = \delta_{ik}$, $i, k = 1, \dots, P$. The j th eigenstate on lattice site k can then be written as

$$\psi_{jk} = \sum_{i=1}^P \varphi_{ik} (c_{ji} \alpha + d_{ji} \beta) = c_{jk} \alpha + d_{jk} \beta, \quad (3)$$

where α, β are the usual two-component spinors. The coefficients c_{jk}, d_{jk} follow from diagonalizing the Hamiltonian matrix associated with the lattice-specific \hat{T} and the given four-potential $\{v, \mathbf{B}\}$. The resulting four-density on lattice site k is

$$\begin{pmatrix} n_k \\ m_k^x \\ m_k^y \\ m_k^z \end{pmatrix} = \sum_{j=1}^N \begin{pmatrix} |c_{jk}|^2 + |d_{jk}|^2 \\ c_{jk} d_{jk}^* + c_{jk}^* d_{jk} \\ ic_{jk} d_{jk}^* - ic_{jk}^* d_{jk} \\ |c_{jk}|^2 - |d_{jk}|^2 \end{pmatrix}. \quad (4)$$

A trivial case of NU arises when the lattice holds the maximum number of electrons allowed by the Pauli principle, $N = 2P$, for then $n=2$ and $\mathbf{m}=0$ for all external four-potentials. We now formulate the first new nontrivial example for NU on lattice systems, which we call *local saturation* NU.

Theorem I. A noninteracting N -electron ground state on a P -point lattice is invariant under a perturbation with four-potential $\{v'_k, \mathbf{B}'_k = v'_k \mathbf{m}_k / m_k\}$, with arbitrary v'_k , that acts locally only on those lattice sites where $n_k = m_k$.

Proof. Notice that for $N=1$ this reduces to the example (c) above, since in that case $n=m$ on all lattice sites. For $N>1$,

it is not hard to show that the condition $n_k = m_k$ on a specific site k is satisfied if and only if

$$\psi_{jk} = f_j (C_k \alpha + D_k \beta), \quad j = 1, \dots, N, \quad (5)$$

i.e., the lowest N ψ_j must have the same spin part, and therefore parallel magnetizations, on point k . Let us now act on these states with a four-potential that is nonzero on site k only and vanishes on all other sites $i \neq k$:

$$\begin{aligned} (v'_k - \mathbf{B}'_k \cdot \vec{\sigma}) f_j (C_k \alpha + D_k \beta) \\ = f_j [C_k v'_k - D_k B'_{x,k} + i D_k B'_{y,k} - C_k B'_{z,k}] \alpha \\ + f_j [D_k v'_k - C_k B'_{x,k} - i C_k B'_{y,k} + D_k B'_{z,k}] \beta. \end{aligned} \quad (6)$$

From this, it is straightforward to show that

$$v'_k [1 - (\mathbf{m}_k / n_k) \cdot \vec{\sigma}] f_j (C_k \alpha + D_k \beta) = 0 \quad (7)$$

for each j , which proves Theorem I. Local saturation NU can occur on one or more isolated lattice sites, but also on groups of sites, which includes examples associated with the formation of ferromagnetic domains on the lattice.

The second new class of examples belongs to the Eschrig-Pickett type⁶ and will be referred to in the following as *global noncollinear* NU.

Theorem II. A noninteracting two-electron ground state on a P -point lattice is invariant under a perturbation with four-potential $\{v' = 0, \mathbf{B}' = \lambda \mathbf{m} / m\}$, $|\lambda| = \text{const}$, if the ground state satisfies $\text{sign}(\lambda)(\bar{n}_1 - \bar{n}_2) / m = \text{const}$. Here, \bar{n}_1 and \bar{n}_2 denote the two occupied orbital densities, with $\bar{n}_{jk} = |c_{jk}|^2 + |d_{jk}|^2$, $j=1, 2$.

Proof. One needs to show that the magnetic field $\mathbf{B}' = \lambda \mathbf{m} / m$ causes at most an orthogonal rotation within the space spanned by the two lowest single-particle eigenstates, ψ_1 and ψ_2 , which leaves the associated two-particle Slater determinant invariant. Thus,

$$\left(\hat{H}_0 - \lambda \frac{\mathbf{m} \cdot \vec{\sigma}}{m} \right) (\gamma_{1i} \psi_1 + \gamma_{2i} \psi_2) = \varepsilon_i (\gamma_{1i} \psi_1 + \gamma_{2i} \psi_2), \quad (8)$$

for $i=1, 2$, where the γ_{ji} form an orthogonal 2×2 matrix, and \hat{H}_0 is the unperturbed single-particle Hamiltonian whose first two eigenstates and energies are $\psi_{1,2}$ and $E_{1,2}$, see Eq. (2). Now consider a lattice site k . After some straightforward algebra, using $m^\pm = m^x \pm im^y$, one arrives at the following expression:

$$\begin{pmatrix} \left[c_{1k} E_1 - \frac{\lambda_k}{m_k} (c_{1k} m_k^z + d_{1k} m_k^-) \right] \\ \left[d_{1k} E_1 - \frac{\lambda_k}{m_k} (c_{1k} m_k^+ - d_{1k} m_k^-) \right] \end{pmatrix} \begin{pmatrix} \left[c_{2k} E_2 - \frac{\lambda_k}{m_k} (c_{2k} m_k^z + d_{2k} m_k^-) \right] \\ \left[d_{2k} E_2 - \frac{\lambda_k}{m_k} (c_{2k} m_k^+ - d_{2k} m_k^-) \right] \end{pmatrix} \begin{pmatrix} \gamma_{1i} \\ \gamma_{2i} \end{pmatrix} = \varepsilon_i \begin{pmatrix} c_{1k} & c_{2k} \\ d_{1k} & d_{2k} \end{pmatrix} \begin{pmatrix} \gamma_{1i} \\ \gamma_{2i} \end{pmatrix}, \quad (9)$$

which has the form of a generalized 2×2 eigenvalue problem. Equation (9) can be easily transformed into a standard eigenvalue problem by multiplying with the inverse of the right-hand coefficient matrix. Using relations (4), one finds after some manipulation

$$\left(\begin{array}{c} \left[\begin{array}{c} -\frac{\lambda_k}{m_k}(\bar{n}_{1k} - \bar{n}_{2k}) + E_1 \\ -2\frac{\lambda_k}{m_k}(c_{1k}c_{2k}^* + d_{1k}d_{2k}^*) \end{array} \right] \left[\begin{array}{c} -2\frac{\lambda_k}{m_k}(c_{1k}^*c_{2k} + d_{1k}^*d_{2k}) \\ \frac{\lambda_k}{m_k}(\bar{n}_{1k} - \bar{n}_{2k}) + E_2 \end{array} \right] \\ \times \begin{pmatrix} \gamma_{1i} \\ \gamma_{2i} \end{pmatrix} = \varepsilon_i \begin{pmatrix} \gamma_{1i} \\ \gamma_{2i} \end{pmatrix}, \end{array} \right) \quad (10)$$

which leads to a characteristic second-degree polynomial with solution

$$\varepsilon_{1,2} = \frac{E_1 + E_2}{2} \pm \sqrt{\frac{(\Delta E)^2}{4} + \lambda_k^2 - \lambda_k \Delta E \frac{\bar{n}_{1k} - \bar{n}_{2k}}{m_k}}, \quad (11)$$

where $\Delta E = E_1 - E_2$. We see immediately that for $\lambda=0$ this reduces to $\varepsilon_{1,2} = E_{1,2}$.

So far, the derivation was for a specific lattice site k . To ensure that the solution $\varepsilon_{1,2}$ and the associated orthogonal eigenvectors γ_{ji} are the same for all P lattice sites, we need to impose the constraints

$$\text{sign}(\lambda_1) \frac{\bar{n}_{11} - \bar{n}_{21}}{m_1} = \dots = \text{sign}(\lambda_p) \frac{\bar{n}_{1p} - \bar{n}_{2p}}{m_p}, \quad (12)$$

which completes the proof of Theorem II, and determines $\text{sign}(\lambda_k)$. Global noncollinear NU thus requires two-electron ground states whose orbital densities and total magnetization are related according to Eq. (12). An explicit example for this will be given below.

Combining Eq. (12) and the normalization of the orbital densities, $\sum_{i=1}^P (\bar{n}_{1i} - \bar{n}_{2i}) = 0$, one obtains another characteristic signature of global noncollinear NU:

$$\text{sign}(\lambda_1)m_1 + \text{sign}(\lambda_2)m_2 + \dots + \text{sign}(\lambda_p)m_p = 0, \quad (13)$$

which shows that this type of NU, although not ‘‘systematic’’ in the sense of Ref. 5, is nevertheless associated with a simple global invariance.

Some additional remarks are in order.

One-electron case. For a single electron, Eq. (9) reduces to

$$\varepsilon = E_1 - \lambda_k n_k / m_k. \quad (14)$$

Again, we require this to be the same on all lattice sites. But, of course, $n=m$ everywhere for a single electron, so that we end up with the condition $\lambda = \text{const}$, i.e., $\varepsilon = E_1 - \lambda$. This leads to the statement that any one-electron ground state is unchanged under the influence of a magnetic field $\mathbf{B}' = \lambda \mathbf{m} / m$ (provided λ is sufficiently small such that the order of the lowest levels is not changed). This is a special case already contained in the one-electron limit of Theorem I.

N-electron case. Global noncollinear NU cannot occur for systems with more than two electrons, which can be seen as follows. The two-electron derivation is easily generalized up to the point where one arrives at a generalized eigenvalue problem similar to Eq. (9), but of the type $R\vec{\gamma} = \varepsilon S\vec{\gamma}$ where R, S are $2 \times N$ rectangular matrices, and $\vec{\gamma}$ is an N -component column vector. Such underdetermined problems are singular, that is, one can find at most two eigenvalues, all remaining $N-2$ eigenvalues are undefined.¹¹ Often one finds no eigenvalues at all. This means that, except for

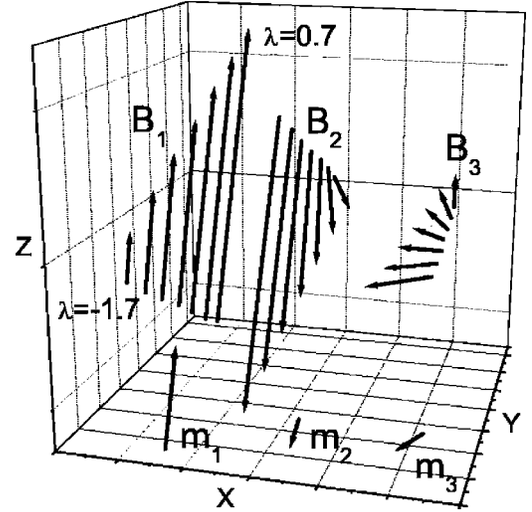


FIG. 1. Global noncollinear NU for a linear three-point lattice with $m_1 = m_2 + m_3$. All magnetic fields shown here, where $\mathbf{B}'_1 = \mathbf{B}_1 + \lambda \mathbf{m}_1 / m_1$ and $\mathbf{B}'_{2,3} = \mathbf{B}_{2,3} - \lambda \mathbf{m}_{2,3} / m_{2,3}$ (see Table I), produce the same four-density (keeping v fixed).

trivial situations or by accident, there is no noncollinear field \mathbf{B}' that results only in a rotation within the single-particle ground-eigenspace. The N -particle ground-state Slater determinant is thus not invariant for $N > 2$.

2-point lattices. For the special case $P=2$ one can show that *all* well-behaved four-potentials produce two-electron ground states whose magnetization has the *same* magnitude on the two lattice sites, $m_1 = m_2$. This result is independent of electron interactions. Global noncollinear NU is thus *always* present on two-point lattices.

An interesting implication of this is that magnetizations with $m_1 \neq m_2$ can arise on a two-point lattice only as ensemble four-densities of degenerate ground states. Similar consequences of NU are expected for lattices with more than two points, i.e., certain classes of four-densities can only come from ensembles of degenerate ground states. For the case of nonmagnetic DFT, the topology of the v and n spaces on lattices was recently clarified,¹³ with the result that pure- and ensemble- v -representable densities have the same mathematical measure. In SDFT, this general statement no longer holds due to the much richer NU, as is evident from the two-point lattice example.

Figure 1 illustrates an example for global noncollinear NU on a linear three-point lattice with lattice constant a and sites 1, 2, 3.¹⁴ To discover this and many other examples, the four-potential parameter space was numerically searched with a multidimensional simplex algorithm¹² until a two-electron ground state was found to satisfy conditions (12) and (13), with $m_1 = m_2 + m_3$, to within an accuracy of 10^{-14} (similar numerical techniques yield examples for local saturation NU). Measuring energies in units of $\hbar^2 / 2m_e a^2$, with electron mass m_e , we give the four-potential and resulting four-density in Table I. All magnetic fields $\mathbf{B}' = \mathbf{B} \pm \lambda \mathbf{m} / m$ (+ on site 1, - on sites 2, 3) produce the *same* two-electron ground state four-density (keeping v fixed), for $-1.7 < \lambda < 0.7$. Values of λ outside that range result in level crossings and thus different ground states.

TABLE I. The four-potential and four-density used in Fig. 1, on lattice sites 1, 2, and 3. Potentials and magnetic fields are measured in units of $\hbar^2/2m_e a^2$ (setting $\mu_B=1$). The lattice four-density is dimensionless. The constraints (12) and (13) are satisfied, with $|\bar{n}_1 - \bar{n}_2|/m=0.99756$ on each lattice site.

	1	2	3
v	-1.62192	1.55381	0.0
B^x	0.87156	-0.14000	-0.50808
B^y	0.15523	0.23990	-0.69702
B^z	1.76179	-0.77994	0.11107
\bar{n}_1, \bar{n}_2	0.96869, 0.07157	0.02926, 0.20498	0.00205, 0.72345
m^x	0.40883	-0.10282	-0.46681
m^y	0.08899	-0.05986	-0.54760
m^z	0.79605	-0.12989	-0.07209
m	0.89931	0.17614	0.72317

We now turn to the continuum limits of our lattice examples.

1. *Local saturation NU.* It is possible that the local condition $n(\mathbf{r})=m(\mathbf{r})$ is satisfied in lower-dimensional subspaces (e.g., points or lines) for a continuum system. But four-potentials that are confined to the same local subspaces and vanish everywhere else are highly pathological (involving delta or step functions). Thus only the well-known special cases of local saturation NU survive in the continuum limit, namely, the one-electron and the completely polarized, ferromagnetic case.

2. *Global noncollinear NU.* The number of constraints, see Eq. (12), that need to be imposed on ground states to exhibit this type of NU grows with the number of lattice sites. Therefore, global noncollinear NU becomes increasingly rare for larger lattices, and is thus ruled out in the continuum limit, in agreement with Ref. 7. Again, only the one-electron special case survives.

These findings are reassuring for the practical application of SDFT to electronic structure calculations in atoms, molecules, and solids. In the collinear case, all that is required in a spin-dependent Kohn-Sham calculation is to fix two constants in the spin-up and spin-down channel, for example through the asymptotic behavior of the potentials. In the noncollinear case, a single constant appears to be sufficient. In situations with full spin polarization, such as in half-metallic ferromagnets, the NU in SDFT is likely to result in the occurrence of discontinuities in the exchange-correlation potential, which will require further study.^{5,6}

On the other hand, spin systems on small lattices are of great interest in the field of spintronics and quantum computation. From a basic point of view, for example, a quantum dot molecule constitutes a two-point lattice. The results presented in this paper will be relevant for the manipulation of electronic charges and spins on such small lattice systems, for instance in performing qubit operations using external magnetic fields.

This work was supported in part by DOE Grant No. DE-FG02-04ER46151. The author thanks Walter Kohn for suggesting this problem, and Giovanni Vignale and Klaus Capelle for valuable discussions.

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¹⁴The spatial orientation of the lattice with respect to the applied magnetic fields is irrelevant in the absence of spin-orbit coupling.