Conditional probability density functional theory

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We present conditional probability (CP) density functional theory (DFT) as a formally exact theory. In essence, CP-DFT determines the ground-state energy of a system by finding the CP density from a series of independent Kohn-Sham (KS) DFT calculations. By directly calculating CP densities, we bypass the need for an approximate XC energy functional. In this work we discuss and derive several key properties of the CP density and corresponding CP-KS potential. Illustrative examples are used throughout to help guide the reader through the various concepts and theory presented. We explore a suitable CP-DFT approximation and discuss exact conditions, limitations, and results for selected examples.

1. INTRODUCTION

Over the past 30 years, density functional theory (DFT) has emerged as a widespread tool in many branches of physics and chemistry [1]. In particular, Kohn-Sham (KS) DFT is an especially popular method to find ground-state energies and electronic densities of molecules and materials. KS-DFT generally scales better with system size than wavefunction-based methods, such as coupled-cluster theory [2] or quantum Monte Carlo (QMC) [3], and is therefore more suitable for modeling realistic systems. However, KS-DFT relies on finding accurate exchange-correlation (XC) energies, which must be approximated as a functional of spin densities. Hundreds of distinct XC approximations currently exist in standard codes [4], reflecting the tremendous difficulty in finding general and accurate approximations. As these standard KS-DFT approximations usually yield accurate self-consistent densities [5], the resulting errors in energies are primarily due to the flaws in the XC energy approximation itself, i.e. the energy error would not change significantly if evaluated on the exact density. Moreover, major deficiencies still remain such as the self-interaction error and the inability to derive accurate energies in strongly-correlated systems [6].

In the exact theory, one can think of XC energies being determined by the pair density, $P^\lambda(r, r')$, which is the joint probability density for finding electrons at two points given electron-electron interactions with strength $\lambda/|r' - r|$ and fixed ground-state density $n(r)$ [8]. In standard DFT approximations, the on-top pair densities, $P^\lambda(r, r)$, are quite accurate [9] and several attempts to use pair densities or density matrices in DFT have been made [10][12]. This pair density can always be written as

$$P^\lambda(r, r') = n(r) \tilde{n}_e^\lambda(r'),$$  \hspace{1cm} (1)

where $\tilde{n}_e^\lambda(r')$ is the conditional probability (CP) density of finding an electron at $r'$, given an electron at $r$.

In standard KS-DFT, the exact KS potential, $v^\lambda_0[n](r)$, is defined to yield $n(r)$ in an effective non-interacting electron problem [13]. Analogously, a conditional probability KS potential (CP-KS), $v_0[\tilde{n}_e^\lambda](r')$, can be defined at each point $r$ in the system, so that it yields $\tilde{n}_e^\lambda(r')$ from a KS calculation with $N-1$ electrons. By the Hohenberg-Kohn (HK) theorem [14], the CP-KS potential, if it exists, is unique. The above equations are for pure density functionals, and their analogs for spin-decomposed systems will be presented in the next sections. This approach, recently presented as conditional probability DFT (CP-DFT) in Ref. [15], is formally exact. The exact system density and CP-KS potential yield exact pair densities and hence XC energies. However, just like KS-DFT, CP-DFT is only useful if good approximations can be found for a wide array of problems. A clear advantage of CP-DFT is that it naturally has no self-interaction error for one electron. Since standard KS-DFT calculations usually yield highly accurate total system densities [6], an accurate CP-KS potential approximation should yield highly accurate XC energies.

The CP-DFT approach is analogous to the Overhauser model [16][17], which was first proposed to approximate the pair correlation function of the uniform electron gas, but has also been extended to non-uniform systems through the average pair-density functional theory (APDFT) approach [18][19]. However, there are key differences: CP-DFT obtains the full pair density whereas APDFT obtains the system- and spherically-averaged pair density, and CP-DFT identifies an exact effective potential within a ground-state KS scheme, the CP-KS potential.

In the context of CP-DFT, authors in Ref. [15] present the blue electron approximation. In this rudimentary approximation the CP potential is simply generated by adding $1/|r' - r|$ to the external potential. This impurity potential represents the repulsion due to the missing electron as if it were a classical point particle at position...
2. DFT fundamentals

We consider non-relativistic purely electronic problems, and use Hartree atomic units (a.u.) throughout the paper. Begin with the variational principal for the exact N-electron ground-state energy

\[ E = \min_{\Psi} \langle \Psi | \hat{H} | \Psi \rangle, \]

where \( \hat{H} \) is the N-electron Hamiltonian and the search is over all antisymmetric, normalized many-body wavefunctions \( \Psi \). We consider Hamiltonians of the form

\[ \hat{H} = \hat{T} + \hat{V}_{\text{ee}} + V, \]

where \( \hat{T} \) is the usual total kinetic energy operator, \( \hat{V}_{\text{ee}} \) is the two-body electron-electron repulsion operator, and \( V \) is the one-body, possibly spin-dependent, total external potential for the system.

DFT replaces the central role of the one-body spin-dependent potentials, \( v(x) \), with the ground-state spin densities \( n(x) \), where

\[ n(x) = N \int dx_2 \cdots dx_N |\Psi(x_2, \ldots, x_N)|^2. \]

Here \( x_i = (r_i, \sigma_i) \) incorporates both spatial and spin coordinates and \( \int dx \) is shorthand to denote the integral over all space and sum over both spins and \( N \) is the total number of electrons in the system. The total density \( n(r) \) is given by a sum over spin densities, \( n(r) = \sum \nabla \cdot n(x) \). From the HK theorem generalized to spin DFT \[22,23,24\], there is a one-to-one correspondence between \( \{n(x)\} \) and \( \{v(x)\} \). From the variational principle, the ground-state energy of a system of N-electrons and external potentials \( v(x) \) is

\[ E = \min_n \left( F[n] + \int dx \, v(x) \, n(x) \right) \]

where \( F[n] \) is the universal part of the Hohenberg-Kohn functional, defined as

\[ F[n] = \min_{\Psi \to n} \langle \Psi | \hat{T} + \hat{V}_{\text{ee}} | \Psi \rangle, \]

where the minimization is over all antisymmetric wavefunctions that yield spin-densities \( n(x) \).

In the KS scheme, there exists an effective one-body potential, \( v_s(x) \), whose corresponding ground-state spin densities match those of the physical interacting system. The total energy of the real system is given in terms of KS quantities:

\[ E = \min_n \left( T_s[n] + \int dx \, v_s(x) \, n(x) + E_n[n] + E_{\text{XC}}[n] \right), \]

where \( T_s \) is the kinetic energy of the KS non-interacting electrons, \( E_n \) is the Hartree energy, \( n = n^\uparrow + n^\downarrow \) is the total density, and \( E_{\text{XC}} \) is the XC energy. We denote the KS wavefunction as \( \Phi_s[n] \), which we assume here is a single Slater determinant, as is typical. The KS wavefunction minimizes the kinetic energy functional,

\[ T_s[n] = \min_{\Psi \to n} \langle \Psi | \hat{T} | \Psi \rangle = T[\Phi_s[n]], \]

where

\[ T[\Psi] = -\frac{1}{2} \sum_{i=1}^N \langle \Psi | \nabla_i^2 | \Psi \rangle. \]

The Euler-Lagrange equation corresponding to Eq. (7) is

\[ \frac{\delta T_s}{\delta n(x)} + v_s(x) = 0, \]

and all potentials are undetermined up to a constant. The KS spin-dependent potential is

\[ v_s(x) = v(x) + \int d^3r \, \frac{n(r)}{|r - x|} + v_{\text{XC}}[n](x), \]
with
\[ v_{\text{xc}}[n](x) = \frac{\delta E_{\text{xc}}[n]}{\delta n(x)} . \] (12)

The KS orbitals \( \phi_{i\sigma} \) satisfy the KS eigenvalue equation
\[ \left[ -\frac{1}{2} \nabla^2 + v_{\text{xc}}(x) \right] \phi_i(x) = \epsilon_{i\sigma} \phi_i(x) . \] (13)

### 2.2. Adiabatic connection

In the adiabatic connection approach to KS-DFT, we can modulate the interaction strength \( \lambda \) of Coulomb-interacting electrons such that
\[ F_\lambda[n] = \min_{\Psi_{\lambda \rightarrow n}} \langle \Psi^\lambda | \hat{T} + \lambda \hat{V}_{\text{ee}} | \Psi^\lambda \rangle . \] (14)

For \( \lambda = 1 \), we have our real, physical system and ground-state density \( n \). For \( \lambda = 0 \), we have the KS system, because we have turned the electron-electron interaction off, but kept the same ground-state density \( n \). We can also consider the limit as \( \lambda \rightarrow \infty \), which is the strictly correlated electron (SCE) limit where the kinetic energy becomes negligible, see section 4.1. In all cases, the ground-state density remains fixed to that of the physical system \( n \). This implies that the external potential \( v^\lambda[n] \) is \( \lambda \)-dependent. The potential \( v^\lambda[n] \) corresponds to the unique one-body potential for which \( n(r) \) is the ground-state density for a system of Coulomb-interacting electrons with interaction strength \( \lambda \). As convention, if \( \lambda \) is absent from the notation then \( \lambda = 1 \) is assumed.

### 2.3. Pair densities and XC holes

It is natural to discuss the spin-decomposed pair densities \( P^\lambda(x, x') \) of the \( \lambda \)-dependent wavefunctions defined in Eq. (14):
\[ P^\lambda(x, x') = N(N - 1) \int dx_3 \cdots dx_N |\Psi^\lambda(x, x', x_3, \ldots, x_N)|^2 , \] (15)

which is the probability density of finding an electron of spin \( \sigma \) at \( r \) and a second electron of spin \( \sigma' \) at \( r' \) for interaction strength \( \lambda \). The total pair density is the spin-summed quantity:
\[ P^\lambda(r, r') = \sum_{\sigma \sigma'} P^\lambda(x, x') , \] (16)

which is the joint probability of finding an electron at \( r \) and a second electron at \( r' \) for interaction strength \( \lambda \). For \( \lambda = 0 \), \( \Psi_{\lambda = 0} = \Phi_0[n] \) and the pair density is
\[ P^{\lambda = 0}(x, x') = n(x)n(x') - |\gamma_0(x, x')|^2 , \] (17)

where \( \gamma_0 \) is the KS (first-order) density matrix:
\[ \gamma_0(x, x') = \delta_{\sigma \sigma'} \sum_{i=1}^{N_x} \phi_i^*(x) \phi_i(x') . \] (18)

The exchange hole is defined as
\[ n_{\text{xc}}(x, x') = -\frac{|\gamma_0(x, x')|^2}{n(x)} , \] (19)

such that
\[ P^{\lambda=0}(x, x') = n(x)(n(x') + n_{\text{xc}}(x, x')) . \] (20)

Checking the normalization allows us to deduce
\[ \int dx' n_{\text{xc}}(x, x') = -1 . \] (21)

For \( \lambda > 0 \), we can generalize Eq. (20) by introducing the \( \lambda \)-dependent correlation hole \( n_{\text{xc}}^\lambda(x, x') \):
\[ P^\lambda(x, x') = n(x)(n(x') + n_{\text{xc}}(x, x') + n_{\text{xc}}^\lambda(x, x')) , \] (22)

where \( n_{\text{xc}}^{\lambda=0}(x, x') = 0 \) and normalization dictates
\[ \int dx' n_{\text{xc}}^\lambda(x, x') = 0 . \] (23)

The XC hole, \( n_{\text{xc}}^\lambda(x, x') \), is simply the sum of the exchange and correlation holes,
\[ n_{\text{xc}}^\lambda(x, x') = n_{\text{xc}}(x, x') + n_{\text{xc}}^\lambda(x, x') \] (24)

where
\[ n_{\text{xc}}^\lambda(r, r') = \left( \sum_{\sigma \sigma'} n(x)n_{\text{xc}}^\lambda(x, x') \right) / n(r) . \] (25)

This quantity is especially of interest in DFT since it determines the XC energy in the adiabatic connection integral:
\[ E_{\text{xc}} = \frac{1}{2} \int_0^1 d\lambda \int d^3r \int d^3r' \frac{n(r)n_{\text{xc}}^\lambda(r, r')}{|r - r'|} . \] (26)

### 3. Theory

#### 3.1. CP-DFT

To simplify the discussion, we will first consider the case of \( \lambda = 1 \) and pure DFT, i.e. not spin decomposed. The subsequent sections will introduce the adaption for different interaction strengths \( \lambda \) and spins \( \sigma \). We define the total conditional probability (CP) density density \( \tilde{n}_r(r') \) as
\[ \tilde{n}_r(r') = \frac{P(r, r')}{n(r)} = n(r') + n_{\text{xc}}(r, r') , \] (27)
which corresponds to the conditional probability density of finding an electron at \( \mathbf{r}' \), given an electron at \( \mathbf{r} \). The subscript in \( \tilde{n}_r(\mathbf{r}') \) emphasizes the parametric dependence on the reference position \( \mathbf{r} \). In Figure 1 we provide example exact densities for a 1-dimensional (1D) model of the He and Be atoms. The 1D model we use throughout mimics reality by replacing 3D Coulomb interactions, \( 1/|\mathbf{r}' - \mathbf{r}| \), with 1D exponentials of the form \( A \exp(-\kappa|y - y'|) \), where \( y, y' \in \mathbb{R} \) and \( A, \kappa > 0 \) are parameters fitted to a soft Coulomb potential, see [20, 27–29] for additional information. The purpose of using 1D models here and throughout the paper is to provide simple and illustrative examples which are qualitatively similar to 3D reality. The 1D “exact” density results are obtained from high accuracy density matrix renormalization group (DMRG) calculations using the ITensor library [30] with an energy convergence threshold of \( 10^{-7} \) Hartree. Associated single-particle potentials were obtained using a modified version of the KS-inversion algorithm outlined in [31]. The code used to perform KS-inversion is publicly available at [32].

In Figure 1 we plot the total ground-state density \( n(y') \) for 1D He, which is spatially symmetric and analogous to a doubly occupied 1s orbital. The exact KS potential \( v_S(y') \) corresponding to this density is also plotted. We show the CP density \( \tilde{n}_y(y') \) for two different reference points, \( y = 0 \) and \( y = 0.8 \). Subtracting the ground-state density \( n(y') \) from the CP density yields the \( \lambda = 1 \) XC hole for a given reference position. We also show the exchange CP density which, for 2 electrons in a spin singlet, is simply half the total density. Thus differences of CP densities from this density are the correlation contributions which are very tiny when \( y = 0 \). For \( y = 0.8 \), correlation causes the CP density to be lower in the region of the electron and greater on the opposite side of the nucleus.

The upper right panel of the figure is the same as the left, but for the Be atom. Here the 1D Be ground-state density is analogous to a \( 1s^22s^2 \) configuration. Now the CP density integrates to 3 electrons and is not closely approximated by half the density. However the same effect occurs when the reference electron is off-center and charge is pushed to the other side of the nucleus.

The CP density follows several properties. For all reference points \( \mathbf{r} \), it must normalize to \( N - 1 \) electrons,

\[
\int d^3\mathbf{r}' \tilde{n}_r(\mathbf{r'}) = N - 1. \tag{28}
\]

In Figure 1, taking the area under the curve of any 1D He CP density yields 1 electron. Similarly, for any 1D Be CP density the integrated area yields 3 electrons. Moreover, since the pair density is a symmetrical function of \( \mathbf{r} \) and \( \mathbf{r}' \), the CP densities satisfy a complementary principle:

\[
\tilde{n}_r(\mathbf{r'}) = \frac{n(\mathbf{r'})}{n(\mathbf{r})} \tilde{n}_r(\mathbf{r}), \tag{29}
\]

which is simply Bayes’ theorem, and may be amenable to modern machine-learning methods [33]. In Fig. 1, for Be, this means the ratio of where the purple curve intersects

![FIG. 1. Exact CP densities and potentials in 1D He (left) and 1D Be (right): \( \tilde{v}_{S, y}(y') \) is the CP-KS potential with corresponding CP density \( \tilde{n}_y(y') \), \( v_S(y') \) the KS potential with corresponding ground-state density \( n(y') \). Quantities are plotted for reference positions \( y = 0 \) and \( y = 0.8 \).](image-url)
the orange vertical line to where the orange curve hits
the purple vertical line equals the ratio of densities (gray
curve) at the two points. For He, since the orange curve is
almost identical to the grey dashed line, any CP density
(purple curve) passes through (almost) the same value at
the origin, namely \( n(0)/2 \).

Next we turn to the CP potentials that generate CP
densities. For a given reference position \( r \) and CP
density \( n_{\lambda}(r') \), we define the CP potential, \( \tilde{v}_{\lambda}(r') \), as the
unique one-body potential whose total ground-state
density for \((N-1)\) Coulomb interacting electrons yields
the total CP density \( \tilde{n}_{\lambda}(r') \). Uniqueness of the CP
potential (up to a constant shift) is guaranteed by the
HK theorem. If it is known, the CP potential for this
\((N-1)\) electron auxiliary system is simply
\[
\tilde{v}_{\lambda}(r') = v[\tilde{n}_{\lambda}](r')
\]  
(30)

where \( v[n](r) \) is the unique one-body potential for which
\( n(r) \) is the Coulomb-interacting ground-state density. In
Figure 1 the exact CP potentials (and corresponding CP
densities) are plotted for He atom at reference points
\( y = 0.0 \) and \( y = 0.8 \). Notice that in the \( y = 0.8 \) case,
the potential is asymmetric and contains a kink at the
reference position. The kink in the CP potential exists
due to the 1D exponential interaction used. However,
for standard 3D Coulomb interactions, we instead obtain a
kink (or cusp) in the CP density at the reference
position due to the electron-electron cusp condition, see
section 3.2. A general feature in any dimension is that the
CP potential is less negative in the region of the reference
point, as it must push electrons away.

We can generalize this HK map to a system of \((N-1)\)
electrons with interaction strength \( \lambda \geq 0 \) and ground-
state density \( \tilde{n}_{\lambda} \). In analogy, we follow Section 2.2
but throughout we are careful to use \( \alpha \) to denote the
interaction strength of the associated \((N-1)\) electron
auxiliary system, while \( \lambda \) will be used exclusively to denote
the interaction strength of the total \( N \) electron system
with fixed ground-state density \( n \). That is, \( \alpha \) and \( \lambda \)
are independent variables in general. For a given \( \alpha \), we have
\[
\tilde{v}_{\alpha}^{\alpha}(r') = v^{\alpha}[\tilde{n}_{\alpha}](r'),
\]  
(31)

so that \( \tilde{v}_{\alpha}^{\alpha}(r') = \tilde{v}_{\alpha=0}^{\alpha}(r') \). In Figure 1
exact CP-KS potentials (and corresponding CP densities) are plotted
for 1D He and Be atoms. In the case of the 1D He atom,
clearly the CP potential is the CP-KS potential. In the
Be atom \( y = 0.8 \) case, we see a bump and kink in the CP-
KS potential at the reference position, resulting in a small
dip in the CP density in the region near the reference
position.

The CP-KS potential can be expressed as
\[
\tilde{v}_{\lambda}(r') = \tilde{v}_{\alpha=0}^{\alpha}(r') + \alpha \langle v_{\text{KS}}[\tilde{n}_{\alpha}](r') \rangle + v_{\text{exc}}^{\alpha}[\tilde{n}_{\lambda}](r'),
\]  
(32)

where \( v_{\text{KS}} \) and \( v_{\text{exc}}^{\alpha} \) are the usual KS-DFT Hartree-
exchange and correlation potentials, respectively. As
in standard KS-DFT, the CP-KS potential can be found
self-consistently. Following Eq. (19), for each reference
point \( r \) the corresponding CP-KS orbitals \( \tilde{\phi}_{\lambda}(r) \) satisfy
the following CP-KS eigenvalue equation
\[
\left[ -\frac{1}{2} \nabla^2 + \tilde{v}_{\lambda}(r') \right] \tilde{\phi}_{\lambda}(r') = \epsilon_{\lambda} \tilde{\phi}_{\lambda}(r'),
\]  
(33)

where \( \sum_{i=1}^{N-1} |\tilde{\phi}_{\lambda,i}(r')|^2 = \tilde{n}_{\lambda}(r') \).

Notice that by construction, the CP-DFT approach
is formally exact: knowledge of an exact CP potential,
\( \tilde{v}_{\alpha}^{\alpha}(r') \), and a corresponding exact XC potential,
\( \tilde{v}_{\alpha}^{\alpha,XC}(r') \), allows a self-consistent KS calculation for an exact CP
density \( \tilde{n}_{\lambda}(r') \).

By construction, CP densities are nowhere negative and
normalized to \( N - 1 \). Thus if their kinetic energy
is finite (and we know of no counterexample), they are
members of the usual set of well-behaved densities, \( L_N \)
Ref. [34]. However, this is insufficient to guarantee
non-interacting \( v \)-representability, just as in standard
DFT [34, 35]. In practice, CP-DFT calculations use
explicit approximations for both \( \alpha \) and \( \lambda \), as

\[
\tilde{n}_{\lambda}(r') \equiv \frac{\lambda P_{\alpha}(r', r')}{n(r)} = n(r') + n_{\alpha=0}^{\alpha}=0(r', r'),
\]  
(34)

for the conditional probability density of finding an
electron at \( r' \), given an electron at \( r \) and for interaction
strength \( \lambda \). The \( \lambda \)-dependent CP density also normalizes
to \( N - 1 \) electrons and satisfies the complementary
principle in Eq. (29).

In analogy with previous formalism, for a given \( r, \lambda, \alpha \)
and CP density, \( n_{\alpha}(r', r') \), we define the CP potential,
\( \tilde{v}_{\lambda}(r', r') \), as the unique one-body potential whose ground-
state density for \( \alpha \)-strength interacting \((N-1)\) electrons
yields the CP density, \( \tilde{n}_{\lambda}(r') \). If it exists, the CP
potential is
\[
\tilde{v}_{\lambda}(r') = v^{\alpha}[\tilde{n}_{\lambda}](r')
\]  
(35)

where
\[
v^{\alpha}[\tilde{n}_{\lambda}](r') = -\frac{\delta F^{\alpha}[\tilde{n}_{\lambda}]}{\delta \tilde{n}_{\lambda}}.
\]  
(36)

The CP-KS potential is \( \tilde{v}_{\alpha=0,\lambda}(r', r') \), where
\( \tilde{v}_{\alpha=0,\lambda}(r') = \tilde{v}_{\alpha=0,\lambda}(r') \) and
\[
\tilde{v}_{\alpha=0,\lambda}(r') = \tilde{v}_{\alpha=0,\lambda}(r') + \alpha \langle v_{\text{KS}}[\tilde{n}_{\alpha}](r') \rangle + v_{\text{exc}}^{\alpha}[\tilde{n}_{\lambda}](r').
\]  
(37)
3.2 Spin-adapted CP-DFT

While Section 3.1 presents a formally exact construction for any interacting electron system, it is useful in practice to have an analogous spin-decomposed formally exact framework. Below we omit $\alpha$ and $\lambda$ dependencies for clarity, but the dependence should follow straightforwardly from the Section 3.1.

We define the spin-CP density $\tilde{n}_x(r')$ as

$$\tilde{n}_x(r') \equiv \sum_{\sigma'} \tilde{n}_x(x'),$$

where

$$\tilde{n}_x(x') = \frac{P(x,x')}{n(x)} = n(x') + n_{xc}(x,x').$$

The spin-CP density $\tilde{n}_x(r')$ has a natural interpretation: given an electron of spin $\sigma$ at reference position $r$, it is the probability density of finding any electron at position $r'$. The spin-CP density normalizes to $N-1$ electrons,

$$\int d^3r' \tilde{n}_x(r') = N - 1,$$

making it a natural object to use as a starting point for spin-adapted CP-DFT. A weighted sum over spin-CP densities can be used to determine the total CP density:

$$\tilde{n}_r(r') = \sum_{\sigma} n_{\sigma}(r) \tilde{n}_{\sigma}(r').$$

In the case of a spin-unpolarized system, we recover

$$\tilde{n}_{\uparrow\uparrow}(r') = \tilde{n}_{\uparrow\downarrow}(r') = \tilde{n}_r(r')$$

as expected.

In Figure 2, we provide exact densities from a 1D model of the spin-polarized Li atom. For each plotted spin-CP and total CP density, taking the area under the curve yields 2 electrons. The total CP densities are plotted in Figure 2 using dashed curves. From Eq. (41), the total CP density can be obtained from a sum of the spin-CP densities, with each weighted by a fraction of density in that spin channel. In 1D Li, for spin-restricted KS orbitals we have the exchange-limit $(\lambda = 0)$ relation $n_{\sigma}(y') = \tilde{n}_{\sigma}(0,0)(y') = \tilde{n}_{\sigma}(y')$. In the upper left panel of Figure 2, we see that indeed $n_{\sigma}(y')$ closely approximates the spin-CP density, $\tilde{n}_{\sigma}(0,0)(y')$, with reference point at the origin, which is a high density region dominated by exchange. In the upper right panel, for the same reference position but opposite spin, we obtain a very similar spin-CP density, $\tilde{n}_{\sigma}(0,0)(y') \approx \tilde{n}_{\sigma}(0,0)(y')$, because, for this reference point, the 1s orbital, which is doubly occupied, is dominant. However, for a spin-down reference point the exchange limit spin-CP density is independent of the reference position, $\tilde{n}_{\sigma}(y') = n_{\sigma}(y')$, which is similar to the case in 1D He (Figure 1 and correspondingly we see little change in the down spin-CP density, $\tilde{n}_{\sigma}(y')$, as the reference position is changed from 0 to 0.8. This is not the case for $\sigma = \uparrow$, so we see large differences between $\tilde{n}_{\sigma}(0,0)(y')$ and $\tilde{n}_{\sigma}(0,0)(y')$.

The analog definition for the spin potential, $\tilde{v}_x(r')$, is straightforward: it is the unique one-body potential whose “accessible” ground-state density from $(N-1)$ Coulomb-interacting electrons yields the spin-CP density, $\tilde{n}_x(r')$, that is, $\tilde{v}_x(r') = v[\tilde{n}_x(x')]$. Note that this is within DFT, not spin DFT, and an “accessible” ground-state is one that conserves total spin, e.g., the total spins of the $N-$ and $(N-1)$-electron systems cannot differ by more than 1/2 unit of angular momentum [36, 37].

Practically, the main difference from Section 3.1 is that, for each reference point $r$ we solve two independent self-consistent CP-KS equations (one for each spin $\sigma$) to obtain corresponding spin-CP densities $\tilde{n}_x(r') = \sum_{\lambda=0}^{N-1} |\tilde{v}_{\lambda}(r')|^2$. Again, the ground-states should be accessible from a given spin configuration. For example, in 1D Li with $N_\uparrow = 2$, $N_\downarrow = 1$, if the reference electron is spin-down we must singly-occupy the lowest two CP-KS orbitals (with spin-up electrons) in the $(N-1)$ system to conserve spin. If the reference electron is spin-up, we doubly-occupy the lowest CP-KS orbital. This means that similar looking up and down spin-CP densities can have corresponding spin-CP-KS potentials that look quite different. This is why $\tilde{v}_{\uparrow,\downarrow}(y')$ looks so different from $\tilde{v}_{\uparrow,\downarrow}(y')$ in Figure 2.

Again, we emphasize that spin-CP densities should be a result from pure DFT with spin-restriction, not spin DFT. One may ask whether a spin-unrestricted CP-KS scheme could be used to obtain the fully spin-decomposed CP spin densities, $\tilde{n}_x(x')$, which correspond to the conditional probability density of finding an electron at $r'$ of spin $\sigma'$, given an electron at $r$ of spin $\sigma$. However, these do not spatially integrate (normalize) to a predetermined integer in general,

$$\int d^3r' \tilde{n}_x(x') = N_{\sigma'} - \delta_{\sigma\sigma'} + \int d^3r' n_{\sigma}(x,x'),$$

and so would require an ensemble of fractional particle numbers [13], presumably making approximations more complicated. Furthermore, these spin densities will contain nodes whenever $x' = x$ due to the exclusion principal. For instance, in the Li atom (with $N_\uparrow = 2$ and $N_\downarrow = 1$), the spin density $\tilde{n}_{\uparrow\downarrow}(r')$ is non-v-representable, as it is a single-particle density which is not strictly positive [35]. Therefore, in general we do not recommend fully spin-decomposed CP-DFT.
3.3 CP-DFT with averaged quantities

The spin-CP densities \( \tilde{n}_s^\lambda(r') \) are clearly high-dimensional quantities: in general there is a \( \lambda \) and \( x \) dependence. In the following we define exact lower-dimensional quantities which can equivalently be used to extract relevant exact energies, such as the electron-electron repulsion potential energy \( U_{ee} = E_u + E_{XC} \):

\[
U_{ee} = \frac{1}{2} \int_0^1 d\lambda \int dx \, n(r) \int d^3 r' \frac{\tilde{n}_s^\lambda(r')}{|r' - r|}. \tag{43}
\]

This may be rewritten in terms of purely radial quantities:

\[
U_{ee} = \frac{N}{2} \sum_{\sigma} \int du \, u \, \langle \tilde{n}_\sigma(u) \rangle \tag{44}
\]

where \( u = r' - r \), \( u = |r' - r| \), and the system and spherically-averaged CP density is

\[
\langle \tilde{n}_\sigma(u) \rangle = \frac{1}{N} \int_0^1 d\lambda \int d^3 r \, n(r) \int d\Omega_u \, \tilde{n}_s^\lambda(r + u). \tag{45}
\]

Note that this radial quantity integrates to \( N - 1 \). The corresponding radial CP-KS potential is defined as \( \tilde{v}_s^\lambda(\langle \tilde{n}_\sigma \rangle)(u) \). Knowledge of this much simpler functional also yields the XC energy, but now only a single 1D integral is required, instead of a 3D integral over \( r \). If useful approximations can be found directly for \( \tilde{v}_s^\lambda(\langle \tilde{n}_\sigma \rangle)(u) \), CP-DFT calculations would be much more efficient. This approach is analogous to average pair-density functional theory [18, 19] which utilizes an effective radial potential, such as the Overhauser model [16, 17], to determine spherically- and system-averaged pair densities.

3.4. connection to exact factorization?

In this section we present an explicit differential equation for \( \tilde{n}_s^\lambda(r') \). We begin by following Ref. [38] and partition the \( \lambda \)-dependent \( N \)-electron Hamiltonian as

\[
H^\lambda = H_s^1 + H_{N-1}^\lambda + \sum_{j=2}^{N} \frac{\lambda}{|r - r_j|}, \tag{46}
\]

where \( H_s^1 \) is the Hamiltonian for a single electron

\[
H_s^1 = -\frac{1}{2} \nabla^2 + \nu^\lambda |n(r)| \tag{47}
\]

and \( H_{N-1}^\lambda \) is the Hamiltonian for \( (N - 1) \) \( \lambda \)-strength interacting electrons

\[
H_{N-1}^\lambda = \sum_{i=2}^{N} \left[ -\frac{1}{2} \nabla_i^2 + \nu^\lambda |n_i(r_i)| \right] + \frac{1}{2} \sum_{i\neq j=2}^{N} \frac{\lambda}{|r_i - r_j|}. \tag{48}
\]
We factorize the normalized ground-state wavefunction \( \Psi^\lambda \) of \( H^\lambda \) as

\[
\Psi^\lambda(x, x_2, \ldots, x_N) = \frac{1}{\sqrt{N}} \tilde{\Psi}_x(x_2, \ldots, x_N). \tag{49}
\]

Here, \( \tilde{\Psi}_x^\lambda \) depends parametrically on \( x \) and is antisymmetric under interchange of the \( N - 1 \) electrons in its argument, but not under interchange of \( x \) with any \( x_i \), \( i \geq 2 \), in general. Note that \( \tilde{\Psi}_x^\lambda \) is not a ground-state wavefunction in general, but is uniquely defined by Eq. \( \tag{49} \), and is often referred to as the \textit{conditional amplitude} \[51\]. The ground-state wavefunction \( \Psi^\lambda \) yields the spin density \( n(x) \), so by construction we have, for all \( x \),

\[
\int dx_2 \cdots \int dx_N |\tilde{\Psi}_x^\lambda(x_2, x_3, \ldots, x_N)|^2 = 1. \tag{50}
\]

We also identify

\[
\tilde{n}_x^\lambda(r') = (N - 1) \sum_s dx_3 \cdots dx_N |\tilde{\Psi}_x^\lambda(x', x_3, \ldots, x_N)|^2. \tag{51}
\]

That is, the total density of \( \tilde{\Psi}_x^\lambda \) is the exact spin-CP density \( \tilde{n}_x^\lambda \). Using Eq. \( \tag{49} \), we generalize Ref. \[39\] and find an effective equation for \( \sqrt{n(x)} \),

\[
\left[ H_1^\lambda + v_{\text{eff}}^\lambda(x) \right] \sqrt{n(x)} = E^\lambda \sqrt{n(x)}, \tag{52}
\]

where \( E^\lambda \) is the ground-state energy of \( H^\lambda \) and the one-body effective potential is

\[
v_{\text{eff}}^\lambda(x) = \lambda \int d^3 r' \frac{\tilde{n}_x^\lambda(r')}{|r' - r|} + \left\langle \tilde{\Psi}_x^\lambda \right| H_{N-1}^\lambda \left| \tilde{\Psi}_x^\lambda \right\rangle + \frac{1}{2} \int dx_2 \cdots dx_N |\nabla_r \tilde{\Psi}_x^\lambda|^2.\tag{53}
\]

It is often partitioned this way, where the first term is referred to as the \textit{conditional potential} and the last term is the \textit{kinetic potential} \[35\]. From Eq. \( \tag{52} \) we also have

\[
\begin{align*}
\hat{H}_x^\lambda \sqrt{n(x)} \tilde{\Psi}_x^\lambda &= \left( E^\lambda - v_{\text{eff}}^\lambda(x) \right) \sqrt{n(x)} \tilde{\Psi}_x^\lambda \\
\ &= -\nabla_r \sqrt{n(x)} \cdot \nabla_r \tilde{\Psi}_x^\lambda - \frac{\sqrt{n(x)}}{2} \nabla^2_r \tilde{\Psi}_x^\lambda. \tag{54}
\end{align*}
\]

This yields an apparent Schrödinger equation for \( \tilde{\Psi}_x^\lambda \) with Hamiltonian \[40\]:

\[
\hat{H}_x^\lambda = \hat{H}_{N-1}^\lambda + \sum_{j=2}^{N} \frac{\lambda}{|r - r_j|} + v_{\text{muc.,}x}^\lambda(x_2, \ldots, x_N) \tag{55}
\]

where

\[
v_{\text{muc.,}x}^\lambda(x_2, \ldots, x_N) = -\frac{2 \nabla_r \sqrt{n(x)} \cdot \nabla_r \tilde{\Psi}_x^\lambda - \sqrt{n(x)} \nabla^2_r \tilde{\Psi}_x^\lambda}{2 \sqrt{n(x)} \tilde{\Psi}_x^\lambda} \tag{56}
\]

which is a non-multiplicative potential and includes gradients of \( \tilde{\Psi}_x^\lambda \) with respect to \( r \). The full equation reads:

\[
\hat{H}_x^\lambda \tilde{\Psi}_x^\lambda = v_{\text{eff}}^\lambda(x) \tilde{\Psi}_x^\lambda. \tag{57}
\]

This is \textit{not} a usual eigenvalue equation that you solve with given boundary conditions \[41\]. It is an inhomogenous differential equation satisfied by \( \tilde{\Psi}_x^\lambda \), defined by Eq. \( \tag{57} \). The total density of the solution \( \tilde{\Psi}_x^\lambda \) is the exact spin-CP density \( \tilde{n}_x^\lambda(r') \), see Eq. \( \tag{51} \). Eq. \( \tag{57} \) is an example of the exact factorization technique, which is typically used in studying nuclear dynamics \[42\]-\[44\], but can also be applied to the pure electronic problem \[41\] \[45\]-\[47\]. The solution \( \tilde{\Psi}_x^\lambda \) is not always the lowest eigenstate if one treats this as an inhomogenous eigenvalue problem, see Appendix B in Ref. \[41\]. That is, in general, \( \tilde{\Psi}_x^\lambda \) does not correspond to the \((N-1)\)-electron ground-state wavefunction in CP-DFT for CP potential \( \tilde{\psi}^\lambda(r') \) and interaction strength \( \lambda \). Worse, \( \psi_{\text{muc.,}x}^\lambda \) depends on \( N - 1 \) coordinates simultaneously, so the usual theorems of DFT cannot be applied. For these reasons, the general relationship between CP-DFT and exact electron factorization (EEF) is subtle, contrary to the connections made between the EEF and DFT through Eq. \( \tag{52} \) \[38\]. However, this analysis can still be useful to CP-DFT in some limits. In Section \ref{4}, we discuss the limit \( |r| \to \infty \), where \( \psi_{\text{muc.,}x}^\lambda \) vanishes everywhere, and we explore implications to CP-DFT.

4. \textbf{EXACT CONDITIONS}

Throughout, it will be convenient to define a CP correction potential, \( \Delta \tilde{\psi}^\alpha_{\text{x},\lambda}(r') \), which is simply the difference between the CP potential and the external potential:

\[
\Delta \tilde{\psi}^\alpha_{\text{x},\lambda}(r') = \tilde{\psi}^\alpha_{\text{x},\lambda}(r') - v^\lambda(n)(r'). \tag{58}
\]

4.1. \textbf{Two electron spin-singlet systems}

For \( N = 1 \), \( n_x^\lambda(r') = 0 \) since the CP density must normalize to 0 and thus there is naturally no self-interaction error in CP-DFT for single electron systems \[49\]. If \( N = 2 \), the CP density has just one electron and the \( \alpha \)-dependence and the exact \( \psi_{\text{muc.,}x} \) vanish. If the \( N = 2 \) electron system is unpolarized, i.e. a spin singlet, the CP densities are

\[
\tilde{n}_r^\lambda(r') = \frac{2 |\Psi^\lambda(r,r')|^2}{n(r)}, \tag{59}
\]

yielding

\[
\tilde{\psi}^\lambda_{\text{s},r}(r') = \frac{1}{2} \nabla^2 \Psi^\lambda(r,r') + \epsilon_r^\lambda. \tag{60}
\]
where $\varepsilon$ is the ground-state energy of the CP-KS potential, $\tilde{n}_{a\lambda}^\lambda({\bf r})$. Because the wavefunction satisfies the Schrödinger equation, we find

$$
\Delta \tilde{\Psi}_{a\lambda}^\lambda(r') + \Delta \tilde{\psi}_{a\lambda}^\lambda({\bf r}) = \frac{\lambda}{|r' - r|} - E^\lambda + \varepsilon_{x}^\lambda + \varepsilon_{r}^\lambda.
$$

(61)

In the limit $|r' - r| \to 0$ we have to leading order:

$$
\Delta \tilde{\psi}_{a\lambda}^\lambda({\bf r}) \to \frac{\lambda}{2|r' - r|}, \quad |r' - r| \to 0.
$$

(62)

In the next section, we will see that this exact condition can be more generally derived from the electron-electron cusp condition. In the exchange limit ($\lambda = 0$) for two electron spin-singlet systems we have $\tilde{n}_{a\lambda}^\lambda({\bf r}) = n({\bf r})/2$, exactly, which is independent of the reference position. From uniqueness, the corresponding CP-KS potential is then

$$
\tilde{\psi}_{a\lambda}^\lambda({\bf r}) = v_{0}[n]({\bf r}).
$$

(63)

In Figure 1 for 1D He we plot the CP density in the exchange limit (half the total density), which is quite close to the CP densities that include correlation.

### 4.2. Cusp condition

The generalized electron coalescence cusp condition requires

$$
\frac{\partial \tilde{n}_{a\lambda}^\lambda({\bf r}, u)}{\partial u} \bigg|_{u=0} = \lambda \tilde{n}_{a\lambda}^\lambda({\bf r}),
$$

(64)

where $u = {\bf r}' - {\bf r}$, $u = |r' - r|$, and the left-hand side has been spherically averaged over $r + u$ [26]. Setting $\lambda = 0$, we see that the cusp condition is a purely correlation-driven effect: the exchange hole does not contribute to the cusp. For generalized Coulomb systems the CP spin density follows

$$
\frac{\partial \tilde{n}_{a\lambda}^\lambda({\bf r})}{\partial u} \bigg|_{u=0} = \lambda \tilde{n}_{a\lambda}^\lambda({\bf r}),
$$

(65)

for $r \neq R_{i}$, where $R_{i}$ are nuclei positions. Following Kato’s theorem [50], the cusp condition is satisfied in CP-DFT with a CP correction potential that has the following condition:

$$
\Delta \tilde{\psi}_{a\lambda}^\lambda({\bf r}) \to \frac{\lambda}{2|r' - r|}, \quad |r' - r| \to 0.
$$

(66)

### 4.3. Long-range limits

When the reference position $r$ is sent to infinity, we can generalize Ref. [38] for arbitrary $\lambda$ to deduce the leading-order term of the ground-state wavefunction of the $\lambda$-interacting $N$-electron system as

$$
\lim_{{|r|} \to \infty} \Psi_{a\lambda}^{\lambda}(x, x_{2}, \ldots, x_{N}) = \sqrt{\frac{n(x)}{N}} \tilde{\psi}_{a\lambda}^{\lambda,N-1}(x_{2}, \ldots, x_{N}),
$$

(67)

where $\tilde{\psi}_{a\lambda}^{\lambda,N-1}$ is an accessible ground-state of the ionized $(N - 1)$-system and is parametrically dependent on $\hat{x} = (\hat{r}, \hat{\sigma})$, where $\hat{r} = r/|r|$ is the direction and $\hat{\sigma}$ is the spin of the electron sent to infinity. The $\hat{\sigma}$ dependence occurs when the ionized $(N - 1)$-system ground-state is otherwise degenerate [36]. The wavefunction $\tilde{\psi}_{a\lambda}^{\lambda,N-1}$ is a solution to Eq. (57) at the large $r$ limit. For a finite number of degenerate $(N - 1)$ ground-states we expect $\tilde{\psi}_{a\lambda}^{\lambda,N-1}$ to remain fixed for small changes in $r$, therefore the gradients in Eq. (56) vanish and $v_{\text{core}}^\lambda$ vanishes (see Refs. [51] and [52] for an in-depth discussion and exceptions). Taking $\alpha = \lambda$, we identify to leading order:

$$
\Delta \tilde{\psi}_{a\lambda}^{\lambda,N-1}[n](r') \to \frac{\lambda}{|r' - r|}, \quad |r| \to \infty.
$$

(68)

That is, in the large reference position limit, the CP correction potential in Eq. (68) simply represents an impurity perturbation that breaks the possible degeneracy in the ionized $(N - 1)$-system ground-state.

We continue this analysis and denote the fully spin-decomposed density of the wavefunction $\tilde{\psi}_{a\lambda}^{\lambda,N-1}$ as $\tilde{n}_{a\lambda}^{\lambda,N-1}({\bf r})$ and the total density of this wavefunction as $\tilde{n}_{a\lambda}^{\lambda,N-1}({\bf r})$. The latter corresponds to the asymptotic limit of the spin-CP density to leading order:

$$
\lim_{{|r'|} \to \infty} \tilde{n}_{a\lambda}^{\lambda}({\bf r}) = \tilde{n}_{a\lambda}^{\lambda,N-1}({\bf r}').
$$

(69)

From the complementary principal, for spin-CP densities we obtain:

$$
\lim_{{|r'|} \to \infty} \tilde{n}_{a\lambda}^{\lambda}({\bf r}) = \sum_{\sigma} n(x'(\hat{\sigma})) \frac{\tilde{n}_{a\lambda}^{\lambda,N-1}(x)}{n(x)}.
$$

(70)

For unpolarized systems or the total CP density we obtain

$$
\lim_{{|r'|} \to \infty} \frac{\tilde{n}_{a\lambda}^{\lambda}({\bf r})}{n({\bf r})} = \frac{\tilde{n}_{\text{HOMO}}^{\lambda,N-1}({\bf r})}{n({\bf r})}.
$$

(71)

That is, for a given reference position $r$, the asymptotic behavior of the total CP density is the same as the asymptotic behavior of the total density of the system, up to a multiplicative constant. This can be seen in Figure 1 for Be, the ionized $N - 1$ system has a 1s orbital very similar to that of the neutral system, so for $y = 0$ the ratio appearing in Eq. (71) is roughly unity and the CP density and total density match far from the nucleus. For He and $y = 0$ the ratio is roughly 1/2 and we see that the CP density and half the total density overlap closely far away.

At large $|r'|$ the highest occupied KS orbital dominates the density [39] and we have

$$
\left[ -\frac{1}{2} \nabla^2 + v_{\text{xc}}[n]({\bf r'}) \right] \sqrt{n({\bf r'})} = \epsilon_{\text{HOMO}} \sqrt{n({\bf r'})}.
$$

(72)

Since $\tilde{n}_{a\lambda}^{\lambda}({\bf r})$ is simply proportional to $n({\bf r'})$ at large $|r'|$, from uniqueness (up to a constant shift) we have

$$
\tilde{\psi}_{a\lambda}^{\lambda}({\bf r'}) \to v_{\text{xc}}[n]({\bf r'}) \quad \text{as} \quad |r'| \to \infty.
$$

(73)
The CP-KS potential approaches the original KS potential far from the system. This can also be seen in Figure 1 where the CP-KS potentials approach the original KS potential far from the nucleus. We can use Eq. (73) to determine a corresponding exact condition for the CP correction potential defined in Eq. (58). From Eqs. (32) and (58) we have

\[ \hat{V}_{\text{Hxc}}[\tilde{n}_r^{\lambda}](r') = \Delta \hat{V}_r^{\alpha,\lambda}(r') + v^\alpha_r[\tilde{n}_r^{\lambda}](r') - v_r^{\text{Hxc}}[\tilde{n}_r^{\lambda}](r'), \]

(74)

The asymptotic limit \(|r'| \to \infty\) of the usual KS-DFT potentials are well-known to leading order \(\text{[53]}\):

\[ v^\alpha_r[\tilde{n}_r^{\lambda}](r') \to \frac{\lambda(N - 1)}{r'}, \quad |r'| \to \infty. \]

(75)

Since the CP density \(\tilde{n}_r^{\lambda}\) integrates to \(N - 1\) electrons:

\[ v^\alpha_r[\tilde{n}_r^{\lambda}](r') \to \frac{\alpha(N - 2)}{r'}, \quad |r'| \to \infty. \]

(76)

From Eqs. (73) - (76) we obtain

\[ \Delta \hat{V}_r^{\alpha,\lambda}(r') \to \frac{N(\lambda - \alpha) + 2\alpha - \lambda}{r'}, \quad |r'| \to \infty. \]

(77)

### 4.4. Strictly correlated electron limit

When \(\lambda \to \infty\), we approach the strictly correlated electron (SCE) limit. The Hamiltonian of Eq. (46) has the expansion \(\text{[55]}\):

\[ H^{\lambda} \to \lambda \left( \hat{V}_{\text{ee}} + \hat{V}_{\text{SCE}} \right) + \mathcal{O}(\sqrt{\lambda}), \quad \lambda \to \infty, \]

(78)

where the last term is kinetic and is subleading. In this limit we have defined

\[ \hat{V}_{\text{SCE}}[n] = \lambda \sum_{i=1}^{N} v_{\text{SCE}}[n](r_i), \quad \lambda \to \infty, \]

(79)

where \(v_{\text{SCE}}[n]\) is the one-body potential that minimizes the classical potential energy operator \(\hat{V}_{\text{ee}} + \hat{V}_{\text{SCE}}[n]\) and delivers \(n(r)\) as the ground-state density. The ground-state wavefunction of such a Hamiltonian collapses into a distribution that can be expressed as \(\text{[54]}\):

\[ |\Psi_{\text{SCE}}(x_1, x_2, \ldots, x_N)|^2 = \frac{1}{N^!} \sum_P \int \frac{d^3 s}{N} \sum_{i=1}^{N} \delta^{(3)}(r_i - f_P(i)(s)), \]

(80)

where \(P\) denotes a permutation of \(1, \ldots, N\), ensuring \(|\Psi_{\text{SCE}}|^2\) is symmetric with respect to exchanging the coordinates of identical particles. The co-motion functions, \(f_P(r)\), dictate the positions of correlated electrons given an electron at position \(r\). The co-motion functions satisfy cyclic group properties, with \(f_1(r) \equiv r\), \(f_2(r) \equiv f(r)\), \(f_3(r) = f(f(r))\), and so on such that \(f_{N+1}(r) = r\). See Ref. \(\text{[55]}\) for additional properties. In the SCE limit, the pair density becomes

\[ p_{\text{SCE}}(r, r') = \frac{1}{N} \sum_{i \neq j = 1}^{N} \int \delta^{(3)}(r - f_i(s)) \delta^{(3)}(r' - f_j(s)) \]

(81)

\[ = \frac{1}{N} \sum_{i \neq j = 1}^{N} \int d^3 s \frac{n(s)}{N} \delta^{(3)}(r - f_i(s)) \delta^{(3)}(r' - f_j(s)). \]

We can neglect terms where neither \(i\) nor \(j\) are equal to 1 since the delta functions in the integral will not overlap for differing non-trivial co-motion functions. Evaluating the integral and applying cyclic properties we obtain

\[ p_{\text{SCE}}(r, r') = n(r) \sum_{i=2}^{N} \delta^{(3)}(r' - f_i(r)). \]

(82)

The CP density is then

\[ \tilde{n}_{\text{CP}}^{\lambda}(r') = \sum_{i=2}^{N} \delta^{(3)}(r' - f_i(r)). \]

(83)

Partitioning the SCE Hamiltonian in Eq. (78) like Eq. (46) and neglecting the subleading gradient terms allows us to identify a corresponding CP correction potential for \(\alpha, \lambda \to \infty\),

\[ \lambda \Delta \tilde{n}_{\text{CP}}^{\lambda}[n](r') = \frac{\lambda}{|r' - r|}. \]

(84)

This is the potential required in this semiclassical limit. In this limit, we have simply fixed the “missing” electron at position \(r\) as if it were a distinguishable particle and solve the resulting \((N-1)\) electron system in the presence of this impurity potential, Eq. (84). In the context of classical statistical mechanics, this is equivalent to the Percus test particle procedure \(\text{[56,58]}\).

### 4.5. Dissociation limit

Consider a simple example of bond dissociation, the stretched \(H_2\) bond. At the dissociation limit, the exact wavefunction has the Heitler-London \(\text{[59]}\) form \((\lambda \neq 0)\):

\[ \Psi^{\lambda}(r_1, r_2) = \frac{1}{\sqrt{2}} (\phi_A(r_1) \phi_B(r_2) + \phi_B(r_1) \phi_A(r_2)) \]

(85)

where \(\phi_A\) and \(\phi_B\) are atomic H orbitals localized on each of the two protons. This yields a CP density:

\[ \tilde{n}_{\text{CP}}^{\lambda}(r') = n_B(r'), \quad r \text{ near } A, \]

(86)

and vice versa, and vanishes elsewhere. Thus the total electron-electron repulsion potential energy from Eq. (43) vanishes due to the lack of overlap, and each atomic region correctly yields the one-electron energy of a isolated hydrogen atom.
We can also consider arbitrarily long neutral $H_N$ chains. In the spin-singlet case, where $N$ is an even number, we obtain a generalized version of Eq. (85) at the dissociation limit ($\lambda \neq 0$):

$$\Psi^\lambda(r_1, \ldots, r_N) = \frac{1}{\sqrt{N!}} \sum_{P} \prod_{i=1}^{N} \phi_i(r_{P(i)}),$$

where $\phi_i$ are atomic $H$ orbitals localized on nuclei $i$, with position $R_i$, in the enumerated $H_N$ chain. The corresponding CP density is:

$$\tilde{n}_r^\lambda(r') = \sum_{i \neq j=1}^{N} n_i(r'), \quad r \text{ near } R_j.$$

Again, the total electron-electron repulsion potential energy from Eq. (43) vanishes due to the lack of overlap, and each atomic region correctly yields a one-electron energy of a isolated hydrogen atom. The corresponding CP potential is

$$\tilde{v}_{r}^{\lambda,\alpha} (r') = -\sum_{i \neq j=1}^{N} \frac{1}{|r' - R_j|}, \quad r \text{ near } R_j.$$

Hence the CP correction potential is then

$$\Delta \tilde{v}_r^{\lambda,\alpha} (r') = \frac{1}{|r' - R_j|}, \quad r \text{ near } R_j.$$

### 5. BLUE ELECTRON APPROXIMATION

In the blue electron approximation (BEA) we make the simple approximation

$$\Delta \tilde{v}_r^{\lambda,\alpha,\text{BEA}} (r') = \frac{\lambda}{|r' - r|}$$

to the CP correction potential defined in Eq. (58). We call this the blue electron approximation because it corresponds to the classical result, Eq. (84). It is as if our reference electron was a distinguishable particle, painted blue for instance, fixed at position $r$, yielding an impurity potential, Eq. (91). Using the BEA and setting $\lambda = \alpha$ in Eq. (74) we obtain

$$\tilde{v}_{r}^{\lambda,\text{BEA}} (r') = v_s[n](r') + \frac{\lambda}{|r' - r|} + v_{\text{HXC}}^\lambda[\tilde{n}_r^\lambda](r') - v_{\text{HXC}}^\lambda[n](r').$$

While simple and semiclassical in origin, the BEA satisfies a surprising number of exact conditions discussed in this work.

In the exchange limit ($\lambda = 0$), the BEA CP correction potential and $v_{\text{HXC}}^\lambda$ terms vanish, leaving $\tilde{v}_{r}^{\lambda=0,\text{BEA}} (r') = v_s[n](r')$. For two electron spin-singlet systems, this is exact, see Eq. (63), but not otherwise. Therefore, in general we do not expect the BEA to perform well for $N > 2$, unless exchange-limit corrections are also incorporated into the CP correction potential.
Taking a reference position \( \mathbf{r} \) that is far away from the system we see that BEA matches the exact asymptotic behavior in Eq. (68) by construction. Similarly, taking \( \lambda = \alpha \) in Eq. (77), we obtain

\[
\Delta \tilde{v}_r^{\lambda=\alpha}(\mathbf{r}' \to \mathbf{r}) = \frac{\lambda}{|\mathbf{r}' - \mathbf{r}|}, \quad |\mathbf{r}'| \to \infty, \tag{93}
\]

which matches the asymptotics of BEA in Eq. (92) if the exact asymptotic decay of \( v_{\text{HXC}}^{\lambda}(\mathbf{r}') \), Eq. (75), is assumed. However, for approximate XC functionals, especially local and semilocal functionals, this condition is usually violated [60].

We can also show that BEA correctly dissociates neutral \( \text{H}_N \) chains. In the dissociation limit, the exact \( v_{\text{HXC}}^{\lambda}(\mathbf{r}') \) terms will vanish and we obtain:

\[
\tilde{v}_{s,y}^{\lambda,\text{BEA}}(\mathbf{r}') = \frac{\lambda}{|\mathbf{r}' - \mathbf{r}|} - \sum_{i=1}^{N} \frac{1}{|\mathbf{r}' - \mathbf{R}_i|}. \tag{94}
\]

In the absence of the first term, this KS potential would yield an \( N \)-fold degenerate ground-state eigenvalue which is equal to the H atom ground-state energy. For \( \lambda \neq 0 \) and when \( \mathbf{r} \) is near \( \{\mathbf{R}_i\} \), the first term breaks some degeneracy: the ground-state eigenvalue has the same value as before but is now \((N - 1)\)-fold degenerate. The associated ground-state CP-KS orbital, \( \tilde{\phi}_{0,\mathbf{r}} \), is

\[
\tilde{\phi}_{0,\mathbf{r}}(\mathbf{r}') = \frac{1}{\sqrt{N-1}} \sum_{i \neq j=1}^{N} \phi_i(\mathbf{r}'), \quad \mathbf{r} \text{ near } \mathbf{R}_j, \tag{95}
\]

where \( \phi_i \) are atomic H orbitals centered at position \( \mathbf{R}_i \). The resulting CP density is the exact one, Eq. (89). If an approximate \( v_{\text{HXC}}^{\lambda}(\mathbf{r}') \) is used, and there exists a self-interaction error for 1 electron systems, the atomic H orbitals in Eq. (95) are correspondingly approximate.

The BEA was proposed for 3D Coulomb-repelling electrons, for which it works well [15]. For 1D exponential repulsion, it is less accurate. In 1D the BEA takes the analog, \( \Delta \tilde{v}_{s,y}^{\lambda,\text{BEA}}(\mathbf{r}') = \lambda \, A \exp(-|\kappa|y - y'|) \). For \( \lambda = 1 \) we plot the BEA CP potential and density for 1D He in Figure 3. We see that the BEA is reasonably accurate, but it is clear that the BEA potential, Eq. (91), is overly repulsive and overestimates the CP potential at the reference position. For \( N > 2 \) and \( \lambda = 1 \) we use the 1D LDA approximation (parameterized in [20]) and solve the CP-KS equations self-consistently with the following CP-KS potential:

\[
v_{s,y}^{\text{BEA,LDA}}(y') = v(y') + A \exp(-|\kappa|y - y')) + v_{\text{HXC}}^{\text{LDA}}[\tilde{n}_{y'}](y'). \tag{96}
\]

We plot this potential for 1D Be in Figure 4. The results here are worse than in 1D He, because of the exchange-limit deficiency previously discussed.

We can also apply this potential in the spin-CP-DFT case to spin-polarized systems, such as 1D Li in Figure 5. For \( \sigma = \uparrow \), the BEA spin-CP density closely approximates the exact and the BEA does yield an accurate exchange-limit spin-CP density, \( n_{\uparrow,\text{LDA}}^{\text{BEA}}(y') \), which is similar to the exact, \( \tilde{n}_{(y,\uparrow)}^{\lambda=0}(y') = n_{\uparrow}(y') \).

With \( \lambda = 1 \) only we can calculate the total electron-electron repulsion energy, \( V_{ee} \), given the total ground-
FIG. 5. Spin-CP densities and potentials in 1D Li with $N_{\uparrow} = 2, N_{\downarrow} = 1$, and $\lambda = 1$: $\tilde{v}^{\text{BEA,LDA}}_{S,x}(y')$ is the 1D BEA approximation to the spin-CP-KS potential using the 1D LDA approximation, see Eq. (96), with corresponding spin-CP density $\tilde{n}^{\text{BEA,LDA}}_{S,x}(y')$. The exact CP-KS potential is $\tilde{v}_S(y')$ with corresponding exact CP density $\tilde{n}_y(y')$, $v_S(y')$ the exact KS potential with corresponding exact ground-state density $n(y')$, and $v(y')$ the external potential for our 1D Li atom.

state density $n$ and CP density $\tilde{n}_y$. In our 1D analog we have:

$$ V_{ee}[n, \tilde{n}_y] = \frac{1}{2} \int dy \, n(y) \int dy' \, \tilde{n}_y(y') \, A \exp(-\kappa |y - y'|). $$

(97)

In Table I we tabulate total electron-electron repulsion energies obtained from various methods. While we note that the BEA is an exceptionally crude approximation in 1D, for two-electron ions it yields energies that have comparable errors to Hartree-Fock (HF).

In 3D reality with Coulombic interactions, it can be seen from Eq. (66) that the BEA yields a CP density with a cusp that is too large by a factor of 2. To remedy
this, we interpolate with a local density approximation
\[
\Delta \tilde{\nu}_r^{\text{LBEA}}[n](\mathbf{r}') \equiv \frac{\lambda}{2|\mathbf{r} - \mathbf{r}'|} \left( 1 + \text{Erf} \left( \frac{|\mathbf{r} - \mathbf{r}'|}{r_s n(\mathbf{r})} \right) \right),
\]
where \( r_s = (3/(4\pi n))^{1/3} \) is the Wigner-Seitz radius at the reference point \( \mathbf{r} \). This local blue electron approximation (LBEA) will yield the exact cusp condition by Eq. (98) and satisfies the same discussed exact conditions as the BEA. In Figure 6 we plot the CP densities and potentials for the (3D) He atom in the \((x', y', z') = 0\) plane for reference position \( \mathbf{r} = (0.563, 0, 0) \). We see that despite noticeable errors in the LBEA CP potential, the corresponding CP density is rather accurate. The exact results were obtained using exact diagonalizations in the recently developed gausslet basis set [15, 61]. In Ref. [15], using the LBEA in Eq. (98) with HF-calculated total densities, it was found that \( E_{\text{LBEA}} = -1.0736 \text{ Ha} \) (error of 0.007 Ha from the exact XC energy). See Ref. [15] for results on other two-electron ions, Hooke’s atom, and H₂ dissociation. Within APDFT, the Overhauser model approximation, which represents a radial screened Coulomb interaction, is a close analog to the LBEA in CP-DFT. In fact, using the Overhauser model, the correlation energy results for two-electron ions in Ref. [19] are quite similar to the LBEA results of Ref. [15]. However, in non-radially symmetric systems or systems with more than two electrons, the Overhauser model and LBEA can produce quite different results. For example, in stretched H₂ the Overhauser model produces noticeable energy errors [62] whereas LBEA obtains the correct dissociation limit [15].

The CP-DFT approach can be used to address the uniform electron gas (UEG). Because the UEG is translationally invariant, the dependence on the reference position can be dropped. In Ref. [15] the LBEA with an added repulsive Gaussian potential term was used in CP-DFT calculations to accurately approximate the XC energy per particle of the UEG at all \( r_s \) values.

In addition, the accuracy does not deteriorate as the temperature of the UEG is raised or if a more primitive method (Thomas-Fermi, as opposed to KS-DFT) is used to calculate CP densities. At zero temperature, the added Gaussian potential is needed at high densities where exchange dominates but is not captured well with the LBEA alone. However, it was also necessary to dampen this Gaussian as \( r_s \) is increased, yielding a somewhat empirical procedure but high accuracy results [15]. More recently, Ref. [63] considers an alternative approach where a portion of the CP density is fixed. Here the parallel component of the spin-CP density, \( \tilde{n}_\sigma(\mathbf{r}', \sigma) \) in Eq. (38), is fixed to the exact exchange limit expression for the UEG (which is known analytically [64]). A spin-CP-DFT calculation is then performed for the antiparallel component. As highlighted in Section 3.2, using spin densities in spin-CP-DFT is complicated, as these components do not integrate up to integer particle numbers in general, see Eq. (12). However, in the setting of the UEG this is not a concern, as the particle number is infinite (or exceedingly large, in a practical calculation). Furthermore, the parallel component of the spin-CP density is fixed to the exchange limit, which does yield an integer particle number. By construction, this approach recovers the correct exchange energy in the high-density limit, and yields sensible accuracy for all other \( r_s \) in the UEG [63].

| \( N \) | Symbol | \( V_{ee}[n, \tilde{n}_{BEA}] \) | \( V_{ee}[n, \tilde{n}_{LDA}^{\text{BEA, LDA}}] \) | \( V_{HF}^{\text{exact}} \) |
|---|---|---|---|---|
| 2 | He | 0.659 (-0.031) | 0.650 (-0.040) | 0.722 (0.032) | 0.690 |
| 2 | Li⁺ | 0.739 (-0.016) | 0.735 (-0.020) | 0.773 (0.018) | 0.755 |
| 2 | Be⁺⁺ | 0.779 (-0.013) | 0.778 (-0.014) | 0.802 (0.010) | 0.792 |
| 3 | Li | - | 1.741 (0.094) | 1.682 (0.035) | 1.647 |
| 3 | Be⁺ | - | 1.971 (0.113) | 1.881 (0.023) | 1.858 |
| 4 | Be | - | 3.314 (0.126) | 3.360 (0.172) | 3.188 |

TABLE I. Electron-electron repulsion energies, \( V_{ee} \), for 1D systems using various methods. All energies are given in Hartree units. Errors with the exact are given in parenthesis. For \( N = 3 \) (spin-polarized systems), spin-CP-DFT calculations were performed self-consistently for each spin channel. 1D Hartree-Fock (HF) results are given as reference. Exact results are from 1D DMRG calculations [20]. The local BEA of the main text performs better for 3D Coulomb-interacting systems [15].
FIG. 6. CP densities and potentials for (3D) He atom ($\lambda = 1$): contour plots for the exact CP density $\tilde{n}_r(r')$ (top left) and corresponding exact CP-KS potential $\tilde{v}_r(r')$ (bottom left) are plotted within the ($x', y', z' = 0$) plane for reference position $r = (0.563, 0, 0)$ (displayed as red cross). The corresponding LBEA results are plotted on the right.
In this work we present CP-DFT as a formally exact theory. CP-DFT is an approach to directly calculate CP densities $\tilde{n}_r(r')$ from ground-state calculations of $N-1$ electrons in an external potential $\tilde{v}_r(r')$, known as the CP potential. In practice, the CP potential (or CP correction potential defined in Eq. (58)) used to obtain the CP density must be approximated. The XC energy can be extracted directly from the adiabatic connection formula using the ground-state density and CP densities. Notably, the CP-DFT approach bypasses the need for an XC energy functional approximation: only accurate ground-state densities and CP densities are needed to yield accurate XC energies within CP-DFT. We also introduce a formally exact spin-CP-DFT approach which can be used to obtain spin-CP densities. Similar to standard spin-DFT, we anticipate that the spin-CP-DFT approach may be more amenable to approximations that can address spin-polarized systems with more accuracy.

Throughout we present several exact conditions associated with CP densities and corresponding CP potentials, including: explicit expressions for two electron systems, long- and short-range asymptotic limits and conditions for general systems, the strictly correlated electron limit, and arbitrary neutral hydrogen chains at the dissociation limit.

We highlight an approximation, the blue electron approximation (BEA), which has semiclassical origins but, surprisingly, satisfies many exact conditions presented. An interpolated variation of BEA, the local blue electron approximation (LBEA), additionally satisfies the electron-electron cusp condition in resulting CP densities. For illustrative purposes, we provide select results for the BEA applied on 1D model systems for He, Li, and Be, as well as the LBEA applied on the (3D) He atom. Because the BEA was designed for 3D Coulomb-repelling systems, it works less well for 1D exponential repulsions, so we strongly recommend against using the 1D-mimic to study the efficacy of BEA. In previous work [15], the LBEA was shown to yield usefully accurate results in several systems where standard DFT approximations can fail, such as single electron systems, stretched $H_2$, and the hydrogen anion. In developing generalizeable CP-DFT approximations, it is clear that additional corrections are needed to obtain accurate CP and spin-CP densities in the exchange limit, which is not well captured using the BEA alone, except in two-electron singlet systems.

Exact conditions might play an important role in guiding construction or testing approximations in CP-DFT. In standard DFT, exact conditions on the energy functionals are most useful, rather than conditions on the densities or potentials. However, in CP-DFT, where an accurate CP density is the sole interest, only exact constraints on the CP densities and CP potentials are useful. Identifying further exact conditions on the CP density (or equivalently, the XC hole) and associating them to conditions on the CP potential may be helpful in the development of future CP-DFT approximations.

Overall, this work aims to provide a sound theoretical basis for CP-DFT and spin-CP-DFT. Simple examples are used throughout to demonstrate concepts and facilitate understanding. We hope that the content of this work will prompt future approximations and applications.

7. ACKNOWLEDGMENTS

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