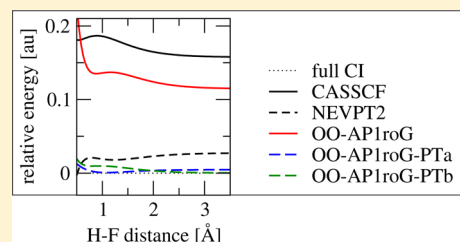


## Assessing the Accuracy of New Geminal-Based Approaches

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## Supporting Information

**ABSTRACT:** We present a systematic theoretical study on the dissociation of diatomic molecules and their spectroscopic constants using our recently presented geminal-based wave function ansätze. Specifically, the performance of the antisymmetric product of rank two geminals (APr2G), the antisymmetric product of 1-reference-orbital geminals (AP1roG) and its orbital-optimized variant (OO-AP1roG) are assessed against standard quantum chemistry methods. Our study indicates that these new geminal-based approaches provide a cheap, robust, and accurate alternative for the description of bond-breaking processes in closed-shell systems requiring only mean-field-like computational cost. In particular, the spectroscopic constants obtained from OO-AP1roG are in very good agreement with reference theoretical and experimental data.



## INTRODUCTION

Present-day quantum chemistry seeks simple, robust, and efficient algorithms for a qualitatively correct description of strongly correlated many-body problems. The difficulty in accurately modeling strongly correlated systems emerges from the fact that molecular orbitals become quasi-degenerate.<sup>1–3</sup> In such cases, the orbital model is qualitatively incorrect and the electronic wave function is no longer well-represented by a single Slater determinant. Instead, a qualitatively accurate description of electronic structures requires several Slater determinants, and the system is said to possess strong multireference character.<sup>1–6</sup> Strong electron correlation becomes important in single- and multiple-bond-breaking processes.<sup>7–10</sup> The electronic properties of transition-metal,<sup>11,12</sup> lanthanide,<sup>13</sup> and actinide compounds<sup>14,15</sup> also show the effects of strong electron correlations.

Standard, well-established multireference methods are, for instance, multireference configuration interaction,<sup>16</sup> complete active-space self-consistent field theory,<sup>17,18</sup> restricted active-space self-consistent field theory,<sup>19</sup> and the density matrix renormalization group.<sup>20–24</sup> Different techniques to model the strong electron correlation problem also emerged within multireference coupled-cluster theory,<sup>25</sup> different variants of coupled-cluster, and valence bond approaches<sup>26</sup> and the projected Hartree–Fock method.<sup>27</sup> For recent examples, we refer the reader to refs 28–42. By construction, such multireference approaches may correctly capture strong (nondynamic and static) correlation effects.<sup>2,3,6,43</sup> However, to ensure computational tractability, the number of Slater determinants needs to be restricted to handle the factorially growing dimension of the many-particle Hilbert space. This

problem is commonly addressed by introducing an active orbital space, possibly composed of different subspaces with method-dependent occupational restrictions. Note that the projected Hartree–Fock method does not require the definition of an active space.<sup>27</sup> The missing weak (dynamic) electron correlation effects, attributed to excitations to the external (i.e., inactive virtual) orbitals, can be added *a posteriori* in a perturbative way.<sup>44–50</sup>

A conceptually different group of approaches that are well-suited for strongly correlated electrons uses two-electron functions, called geminals, to build electron correlation effects into the many-particle wave function.<sup>51–55</sup> In its second-quantized form, a geminal wave function can be written as

$$|\Psi_{\text{geminal}}\rangle = \Psi_1^\dagger \Psi_2^\dagger \dots \Psi_{N/2}^\dagger |\text{vac}\rangle \quad (1)$$

where the index  $N$  is the number of electrons,  $|\text{vac}\rangle$  denotes some vacuum state with respect to the creation of geminals and  $\Psi_i^\dagger$  is a geminal, that is, a correlated two-electron function. If we confine geminals to be singlet functions, a pair-creation then bears the form

$$\Psi_i^\dagger = \sum_{p,q=1}^{M_i} C_{p,q}^i a_{p\uparrow}^\dagger a_{q\downarrow}^\dagger \quad (2)$$

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where  $M_i$  is the number of one-particle functions (orbitals) used to create geminal  $i$ ,  $C_{p,q}^i$  is a geminal matrix coefficient for subspace  $M_i$ , and  $a_{p\uparrow}^\dagger$  and  $a_{q\downarrow}^\dagger$  are the standard electron creation operators for up- and down-spin electrons ( $\uparrow$ ,  $\downarrow$ ). The geminal matrix coefficient links the geminal wave function with the underlying one-particle basis functions (orbitals). Further, the  $C_{p,q}^i$  matrix can be diagonalized for each subspace by an appropriate unitary transformation that brings the geminals into their *natural* form, constructed from the transformed set of natural orbitals,

$$\Psi_i^\dagger = \sum_{p=1}^{M_i} C_p^i a_{p\uparrow}^\dagger a_{p\downarrow}^\dagger \quad (3)$$

Specifically, the geminal matrix structure depends on the geminal wave function ansatz used.<sup>55–58</sup>

Over the past decade, various geminal-based approaches have been introduced in the literature that impose different restrictions on the geminal coefficient matrix  $\{C_p^i\}$  (or  $\{C_{p,q}^i\}$ , respectively) to ease computational cost.<sup>54,55</sup> The most popular approaches are based on the antisymmetric product of strongly orthogonal geminals<sup>51,59–63</sup> (APSG), the antisymmetrized geminal power<sup>52,64</sup> (which is a special case of projected Hartree–Fock–Bogoliubov), or the antisymmetric product of interacting geminals<sup>54–56,65–74</sup> (APIG). In the APSG approach, the subspaces  $M_i$  are disjoint, i.e., each geminal is constructed from different orbitals. Thereby, the APSG wave function has a simple and computationally tractable form, but misses a large fraction of the electron correlation energy resulting from the interaction between geminals.<sup>55</sup> The APIG ansatz alleviates this problem by relaxing the strong-orthogonality constraint. However, restoring correlation between all orbital pairs forfeits computational tractability. As a consequence, the application of geminal-based approaches to large and chemically more interesting systems has been limited.

Recently, we have presented new geminal-based wave function ansätze that approximate the APIG wave function, but require significantly less computational cost.<sup>57,58</sup> Specifically, the antisymmetric product of 1-reference orbital geminal (AP1roG)<sup>58,75</sup> provides an alternative parametrization of the doubly occupied configuration interaction (DOCI),<sup>76</sup> but demands only mean-field computational cost in contrast to the factorial scaling of traditional DOCI implementations. The AP1roG wave function ansatz can be rewritten in terms of one-particle functions as a fully general pair-coupled-cluster wave function, that is,

$$|\Psi_{\text{AP1roG}}\rangle = \exp\left(\sum_{i=1}^P \sum_{a=p+1}^K C_i^a a_a^\dagger a_{\bar{a}}^\dagger a_{\bar{a}} a_i\right) |\Phi_0\rangle \quad (4)$$

where  $a_p^\dagger$ ,  $a_{\bar{p}}^\dagger$  and  $a_p$ ,  $a_{\bar{p}}$  are the electron creation and annihilation operators, in which  $p$  and  $\bar{p}$  denote  $\alpha$  and  $\beta$  spins, respectively, and  $|\Phi_0\rangle$  is some independent-particle wave function (usually the Hartree–Fock determinant). Indices  $i$  and  $a$  correspond to virtual and occupied orbitals with respect to  $|\Phi_0\rangle$ ,  $P$  and  $K$  denote the number of electron pairs ( $P = N/2$  with  $N$  being the total number of electrons) and orbitals, respectively, and  $\{C_i^a\}$  are the geminal coefficients (cf., eqs 2 and 3). For AP1roG, the electronic Schrödinger equation can be solved very efficiently by projection onto the set of pair excitations,<sup>58</sup>

$$\langle \Phi_0 | \hat{H} | \Psi_{\text{AP1roG}} \rangle = E \langle \Phi_0 | \Psi_{\text{AP1roG}} \rangle \quad (5)$$

$$\langle \Phi_{i\bar{i}}^{a\bar{a}} | \hat{H} | \Psi_{\text{AP1roG}} \rangle = E \langle \Phi_{i\bar{i}}^{a\bar{a}} | \Psi_{\text{AP1roG}} \rangle \quad (6)$$

where  $\hat{H}$  is some Hamiltonian and  $\langle \Phi_{i\bar{i}}^{a\bar{a}} |$  are pair excitations with respect to some reference wave function  $\langle \Phi_0 |$ . We should emphasize that the exponential form of eq 4 guarantees size-extensivity.

The AP1roG approach can be combined with an orbital optimization protocol<sup>77</sup> (abbreviated by the acronym OO-AP1roG) that is analogous to the orbital-optimized coupled cluster approach.<sup>78–80</sup> The (variational) optimization of the one-particle basis functions restores size-consistency, provided that the optimized orbitals are localized,<sup>77</sup> that is, symmetry-broken. In particular, OO-AP1roG poses a mean-field-like approximation to DO-Self-Consistent-Field (DOSCF).<sup>76</sup>

In our second (variational) approach, we have employed the eigenvectors of the reduced Bardeen–Cooper–Schrieffer (BCS) Hamiltonian<sup>81</sup>

$$\hat{H}_{\text{BCS}} = \sum_{i=1}^N \varepsilon_i (a_{i\uparrow}^\dagger a_{i\uparrow} + a_{i\downarrow}^\dagger a_{i\downarrow}) - g \sum_{ij} a_{i\uparrow}^\dagger a_{i\downarrow}^\dagger a_{j\downarrow} a_{j\uparrow} \quad (7)$$

as trial wave functions. The system is defined by the set of single-particle energies  $\{\varepsilon_i\}$ , and the pairing strength  $g$ . Eigenvectors of the reduced BCS Hamiltonian have the form of a Bethe ansatz<sup>82</sup> in terms of the geminals

$$\Psi_i^\dagger = \sum_{p=1}^M \frac{1}{\lambda_i - \varepsilon_p} a_{p\uparrow}^\dagger a_{p\downarrow}^\dagger \quad (8)$$

Each pair of electrons is labeled with a *rapidity* or *pair energy*  $\lambda$ . With the use of the reduced BCS Hamiltonian on a product of these rank 2 geminals one can readily observe that an eigenvector is obtained provided that the rapidities satisfy Richardson's equations<sup>83,84</sup>

$$\frac{1}{g} + \sum_{p=1}^M \frac{1}{\lambda_i - \varepsilon_p} - 2 \sum_{j \neq i} \frac{1}{\lambda_i - \lambda_j} = 0, \quad i = 1, \dots, N/2. \quad (9)$$

To solve this set of equations numerically we have employed the algorithm proposed by De Baerdemacker et al.,<sup>85</sup> though other algorithms are prevalent.<sup>86–88</sup> Practical expressions for the 1- and 2-reduced density matrices are known.<sup>89</sup> We will refer to this trial wave function as the antisymmetric product of rank 2 geminals (APr2G).<sup>57</sup>

In this work, we assess the accuracy of our recently presented geminal-based approaches (AP1roG and APr2G) in describing the ground-state electronic structure of a test set of diatomic molecules and their spectroscopic constants (bond lengths, potential depths, and harmonic vibrational frequencies). Specifically, we focus on the LiH, Li<sub>2</sub>, HF, and C<sub>2</sub> molecules, where accurate theoretical benchmark and experimental spectroscopic data are available. In addition, we investigate the performance of simple perturbative corrections<sup>90</sup> to the OO-AP1roG wave function.

## COMPUTATIONAL DETAILS

The full configuration interaction (FCI), the DOCI, and the DOSCF energies were obtained using the MOLPRO2010 software package.<sup>91</sup> The (restricted) second-order Møller–Plesset (MP2) perturbation theory, (restricted) coupled cluster theory with singles, doubles, and perturbative triples (CCSD-(T)) and including full triples (CCSDT), as well as the

**Table 1. Spectroscopic Constants: Bond Distances ( $R_e$ ), Potential Energy Depths ( $D_e$ ) and Harmonic Vibrational Frequencies ( $\omega_e$ ) for the  $X^1\Sigma^+$  State of the LiH Molecule. Differences with Respect to FCI (in a Given Basis Set) Are Listed in Parentheses**

method	basis set	$R_e$ [Å]	$D_e$ [eV]	$\omega_e$ [cm <sup>-1</sup> ]
APr2G-A	cc-pVDZ	1.619 (+0.004)	3.569 (+1.314)	1388.7 (+35.9)
AP1roG	cc-pVDZ	1.621 (+0.006)	3.506 (+1.251)	1385.5 (+32.7)
DOCI	cc-pVDZ	1.621 (+0.006)	3.506 (+1.251)	1385.5 (+32.7)
APr2G-B	cc-pVDZ	1.616 (+0.001)	2.250 (−0.005)	1344.6 (−8.2)
OO-AP1roG	cc-pVDZ	1.616 (+0.001)	2.253 (−0.002)	1354.8 (+2.0)
DOSCF	cc-pVDZ	1.616 (+0.001)	2.253 (−0.002)	1354.8 (+2.0)
OO-AP1roG-PTa	cc-pVDZ	1.615 (+0.000)	2.254 (−0.001)	1352.4 (−0.4)
OO-AP1roG-PTb	cc-pVDZ	1.615 (+0.000)	2.254 (−0.001)	1353.1 (+0.3)
DFT/PBE	cc-pVDZ	1.623 (+0.008)	2.895 (+0.640)	1354.7 (+1.9)
MP2	cc-pVDZ	1.612 (−0.003)	3.418 (+1.163)	1379.5 (+26.7)
CCSD(T)	cc-pVDZ	1.614 (−0.001)	2.249 (−0.006)	1352.9 (+0.1)
FCI	cc-pVDZ	1.615	2.255	1352.8
AP1roG	aug-cc-pVDZ	1.595 (+0.011)	4.000 (+1.572)	1456.4 (+22.3)
DOCI	aug-cc-pVDZ	1.595 (+0.011)	4.000 (+1.572)	1456.4 (+22.3)
OO-AP1roG	aug-cc-pVDZ	1.590 (+0.006)	2.413 (−0.015)	1421.4 (−12.7)
DOSCF	aug-cc-pVDZ	1.590 (+0.006)	2.413 (−0.015)	1421.6 (−12.5)
OO-AP1roG-PTa	aug-cc-pVDZ	1.585 (+0.001)	2.425 (−0.003)	1432.1 (−2.0)
OO-AP1roG-PTb	aug-cc-pVDZ	1.586 (+0.002)	2.425 (−0.003)	1437.4 (+3.3)
DFT/PBE	aug-cc-pVDZ	1.617 (+0.033)	2.997 (+0.569)	1347.6 (−86.5)
MP2	aug-cc-pVDZ	1.587 (+0.003)	3.551 (+1.123)	1457.6 (+23.5)
CCSD(T)	aug-cc-pVDZ	1.583 (−0.001)	2.411 (−0.017)	1405.0 (−29.1)
FCI	aug-cc-pVDZ	1.584	2.428	1434.1
AP1roG	aug-cc-pVTZ	1.649 (+0.057)	3.841 (+1.311)	1359.2 (−60.0)
OO-AP1roG	aug-cc-pVTZ	1.600 (+0.008)	2.504 (−0.026)	1395.1 (−24.1)
OO-AP1roG-PTa	aug-cc-pVTZ	1.594 (+0.002)	2.527 (−0.003)	1416.0 (−3.2)
OO-AP1roG-PTb	aug-cc-pVTZ	1.594 (+0.002)	2.525 (−0.005)	1420.5 (+1.3)
DFT/PBE	aug-cc-pVTZ	1.613 (+0.021)	2.698 (+0.168)	1350.9 (−68.3)
MP2	aug-cc-pVTZ	1.589 (−0.003)	3.629 (+1.099)	1441.5 (+22.3)
CCSD(T)	aug-cc-pVTZ	1.592 (−0.000)	2.460 (−0.070)	1406.2 (−13.0)
FCI	aug-cc-pVTZ	1.592	2.530	1419.2
Exp.		1.596 <sup>111</sup>	2.515 <sup>112</sup>	1405.7 <sup>113</sup>

(unrestricted) density functional theory calculations employing the PBE<sup>92</sup> exchange–correlation functional were performed with the NWChem6.1 quantum chemistry software.<sup>93–95</sup>

**AP1roG/APr2G.** The optimization of the (restricted) AP1roG<sup>58</sup> and OO-AP1roG<sup>77</sup> wave functions was done with a local developer version of the HORTON program package.<sup>96</sup> In the OO-AP1roG approach, we used a Newton–Raphson optimizer and a diagonal approximation to the orbital Hessian to obtain the rotated set of orbital expansion coefficients. Furthermore, spatial symmetry was preserved during the orbital optimization procedure; we refer the reader to ref 97 where spectroscopic constants obtained from symmetry-broken solutions are discussed. Two types of second-order perturbative corrections, PTa and PTb, were applied on top of OO-AP1roG, to account for weak (dynamic) electron correlation effects. The main difference between these two approaches is that the former uses the Hartree–Fock determinant and the latter uses the AP1roG wave function as its dual (cf. ref 90 for more details).

The variational APr2G energies were calculated using an “in-house” FORTRAN90 code. Two types of APr2G calculations were performed: using (i) canonical Hartree–Fock orbitals (labeled as APr2G-A) and (ii) OO-AP1roG orbitals (labeled as APr2G-B). The one- and two-electron molecular orbital integrals used

for APr2G-A were generated with the DALTON2013 quantum chemistry software.<sup>98</sup> In all calculations mentioned above, we correlated all electrons. (One strength of geminal-based approaches is the removal of the need to specify an active space.)

**CASSCF/NEVPT2.** The CASSCF and NEVPT2 calculations were carried out using the DALTON2013 software package.<sup>98</sup> We employed a “full-valence” complete active space, which results in eight electrons correlated in five orbitals (CAS(8,5)) for HF and eight electrons correlated in eight orbitals (CAS(8,8)) for C<sub>2</sub>. The same active orbitals were used in subsequent NEVPT2 calculations.

**Basis Sets.** For the LiH and Li<sub>2</sub> molecules, we utilized Dunning’s cc-pVDZ (H:(4s1p) → [2s1p], Li:(9s4p1d) → [3s2p1d]), aug-cc-pVDZ (H:(5s2p) → [3s2p], Li:(10s5p2d) → [4s3p2d]) and aug-cc-pVTZ (H:(6s3p2d) → [4s3p2d], Li:(12s6p3d2f) → [5s4p3d2f]) basis sets.<sup>99–101</sup> For HF and C<sub>2</sub> the cc-pVDZ (C, F:(9s4p1d) → [3s2p1d]), aug-cc-pVDZ (C, F:(10s5p2d) → [4s3p2d]) and aug-cc-pVTZ (C, F:(11s6p3d2f) → [5s4p3d2f]) basis sets were used.<sup>99–101</sup>

**Fitting Procedure.** The potential energy curves of diatomic molecules were obtained by varying bond-lengths in a range of 1.2–10.0 Å, 2.2–10.0 Å, 0.7–5.0 Å, and 1.1–3.2 Å for the LiH, Li<sub>2</sub>, HF, and C<sub>2</sub> molecules, respectively. The points on the

resulting potential energy curve were used for a subsequent generalized Morse function<sup>102</sup> fit to obtain the equilibrium bond lengths ( $R_e$ ) and potential energy depths ( $D_e$ ). The harmonic vibrational frequencies ( $\omega_e$ ) were calculated numerically using the five-point finite difference stencil.<sup>103</sup>

## RESULTS AND DISCUSSION

**LiH.** Although LiH is the smallest stable, neutral heteroatomic molecule, it is a particularly interesting system for quantum chemists; its electronic structure changes considerably with the internuclear distance, and the resulting potential energy surface is difficult to describe with traditional (single-reference) methods.<sup>104–107</sup> While the  $X^1\Sigma^+$  state of LiH is best represented by the single  $1\sigma^22\sigma^2$  electronic configuration around the equilibrium structure, its contribution changes when the Li-H bond is stretched. For increasing bond distances, the electronic wave function becomes a mixture of four dominant electronic configurations,  $1\sigma^22\sigma^2$ ,  $1\sigma^23\sigma^2$ , and  $1\sigma^22\sigma^13\sigma^1$  (doubly degenerate). Multireference approaches are thus required to correctly model the dissociation limit of the single-bond breaking process in the LiH molecule.<sup>105,107–110</sup>

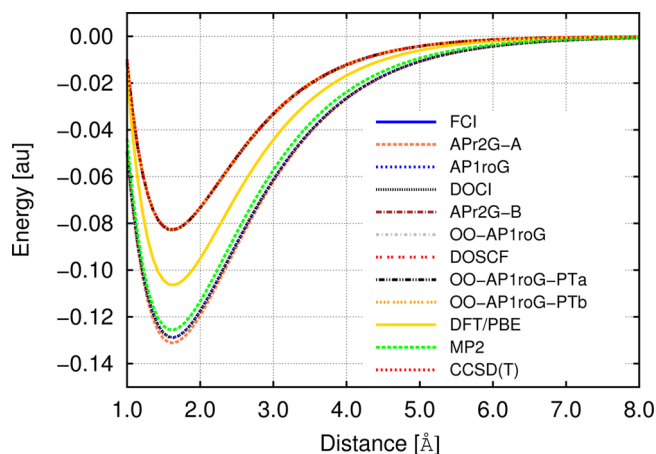
Table 1 lists all bond distances ( $R_e$ ), potential energy depths ( $D_e$ ), and harmonic vibrational frequencies ( $\omega_e$ ) for the  $X^1\Sigma^+$  state of LiH determined from different quantum chemistry methods and basis sets. For each set of spectroscopic constants, the difference with respect to the exact result in a given basis set is provided in parentheses.

In general, APr2G and AP1roG result in similar spectroscopic constants, which agree well with the DOCI reference data. While equilibrium bond lengths and harmonic vibrational frequencies are in reasonable agreement with FCI, the potential depth is overestimated by more than 1 eV. Similarly, DFT and MP2 fail to predict accurate dissociation energies.

Orbital optimization within the AP1roG approach alleviates these large discrepancies and reduces the differences in the potential well depth to only 0.005 and 0.002 eV for APr2G-B and OO-AP1roG, respectively. We should note that OO-AP1roG can reproduce DOSCF along the dissociation pathway, and thus both methods yield identical spectroscopic constants. The missing weak (dynamic) electron correlation effects can be easily captured by an *a posteriori* perturbative treatment on top of the OO-AP1roG wave function (cf. Table 1 for OO-AP1roG-PTa and OO-AP1roG-PTb results).

Potential energy surfaces for the dissociation process of LiH using the cc-pVDZ basis set are presented in Figure 1. It is important to stress that the dissociation limit has been shifted to zero a.u. in all approaches and, apart from our DFT/PBE results, all total energies are higher than the FCI reference curve for all basis sets studied (see Figures S1 and S2 of the Supporting Information for further details). A systematic improvement in the basis set size (cc-pVDZ  $\rightarrow$  aug-cc-pVDZ  $\rightarrow$  aug-cc-pVTZ) brings the theoretically determined spectroscopic constants very close to experimental data (cf. Table 1). Additional figures (S1 and S2) with potential energy surfaces for larger basis sets can be found in the Supporting Information.

**Li<sub>2</sub>.** Although being the second smallest, just after molecular hydrogen, stable homonuclear molecule, Li<sub>2</sub> has attracted a lot of attention from quantum chemists.<sup>114–116</sup> Specifically, the lithium dimer serves as a convenient test system of theoretical methods and represents the smallest prototype for a theoretical



**Figure 1.** Fitted potential energy surface of the ground state of the LiH molecule obtained from different quantum chemistry methods using the cc-pVDZ basis set. Note that the dissociation limit is shifted to zero for all approaches.

description of long-range potential energy curves in molecular photoassociative spectroscopy of cold alkali atoms.<sup>117</sup>

Spectroscopic constants for the  $^1\Sigma_g^+$  ground state of Li<sub>2</sub> are collected in Table 2. APr2G-A and AP1roG provide a qualitatively incorrect dissociation limit and overestimate equilibrium bond-lengths. The same is true for DOCI, which gives similar results to AP1roG (see Table 2). We should note that APr2G includes AP1roG as a special case only for four-electron systems.<sup>57</sup>

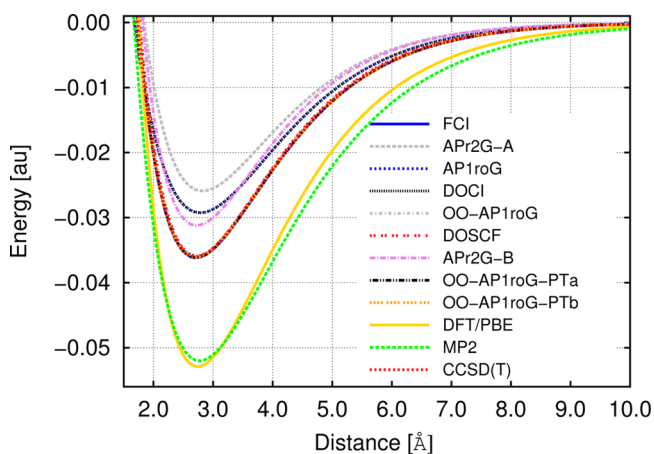
Optimizing the one-particle basis functions for geminals improves all spectroscopic constants, giving excellent agreement with FCI or CCSD(T)/CCSDT reference data for all basis sets studied (see also Figure 2). Minor differences for larger basis sets are attributed to weak (dynamic) electron correlation effects (see Figures S3 and S4 of the Supporting Information). This type of electron correlation is not directly included in our geminal-based wave function ansätze, but can easily be recovered *a posteriori* in, e.g., OO-AP1roG-PTa and OO-AP1roG-PTb. In particular, spectroscopic constants determined from OO-AP1roG-PTa or OO-AP1roG-PTb are in line with the corresponding CCSD(T) and CCSDT results. Finally, we should emphasize that only OO-AP1roG-PTa and OO-AP1roG-PTb as well as CCSD(T) and CCSDT in combination with a large basis set yield spectroscopic constants that are in agreement with experimental data (cf. Table 2), while DFT and MP2 fail in describing the potential energy curve of Li<sub>2</sub> overestimating equilibrium bond distances and dissociation energies.

**HF.** The dissociation of hydrogen fluoride represents another toy model for multireference approaches.<sup>8,119</sup> Similar to lithium hydride, the HF molecule is well represented by a single  $1\sigma^22\sigma^23\sigma^21\pi^4$  electronic configuration around the equilibrium interatomic distance, while for stretched interatomic bond lengths, two additional electronic configurations,  $1\sigma^22\sigma^21\pi^44\sigma^2$  and  $1\sigma^22\sigma^21\pi^43\sigma^14\sigma^1$ , become important.<sup>110,119</sup> The alternating interplay of weak and strong electron correlation effects along the dissociation pathway makes the HF molecule a very difficult test case for single-reference quantum chemistry methods.<sup>110</sup>

All spectroscopic constants for the  $X^1\Sigma^+$  state of the HF molecule are listed in Table 3, while the potential energy surfaces are illustrated in Figure 3. We should stress that larger deviations between the OO-AP1roG and DOSCF results arise

**Table 2. Spectroscopic Constants: Bond Distances ( $R_e$ ), Potential Energy Depths ( $D_e$ ) and Harmonic Vibrational Frequencies ( $\omega_e$ ) for the  $X^1\Sigma_g^+$  State of the  $\text{Li}_2$  Molecule. Differences with Respect to Reference Data (in a Given Basis Set) Are Listed in Parentheses**

method	basis set	$R_e$ [Å]	$D_e$ [eV]	$\omega_e$ [ $\text{cm}^{-1}$ ]
APr2G-A	cc-pVDZ	2.821 (+0.109)	0.705 (−0.278)	310.3 (−23.9)
AP1roG	cc-pVDZ	2.785 (+0.073)	0.796 (−0.187)	324.3 (−9.9)
DOCI	cc-pVDZ	2.785 (+0.073)	0.796 (−0.187)	324.3 (−9.9)
APr2G-B	cc-pVDZ	2.719 (+0.007)	0.849 (−0.134)	330.8 (−3.4)
OO-AP1roG	cc-pVDZ	2.720 (+0.008)	0.977 (−0.006)	336.7 (+2.5)
DOSCF	cc-pVDZ	2.721 (+0.009)	0.978 (−0.005)	336.6 (+2.4)
OO-AP1roG-PTa	cc-pVDZ	2.711 (−0.001)	0.984 (+0.001)	334.4 (+0.2)
OO-AP1roG-PTb	cc-pVDZ	2.713 (+0.001)	0.982 (−0.001)	334.1 (−0.1)
DFT/PBE	cc-pVDZ	2.743 (+0.031)	1.440 (+0.457)	329.6 (−4.6)
MP2	cc-pVDZ	2.773 (+0.061)	1.415 (+0.432)	329.7 (−4.5)
CCSD(T)	cc-pVDZ	2.712 (−0.000)	0.979 (−0.004)	334.5 (+0.3)
FCI	cc-pVDZ	2.712	0.983	334.2
AP1roG	aug-cc-pVDZ	2.771 (+0.065)	0.669 (−0.327)	338.8 (+7.8)
OO-AP1roG	aug-cc-pVDZ	2.718 (+0.012)	0.987 (−0.009)	334.4 (+3.4)
OO-AP1roG-PTa	aug-cc-pVDZ	2.703 (−0.002)	0.996 (+0.000)	331.0 (+0.0)
OO-AP1roG-PTb	aug-cc-pVDZ	2.703 (−0.002)	0.994 (−0.002)	329.3 (−1.7)
DFT/PBE	aug-cc-pVDZ	2.741 (+0.035)	1.439 (+0.443)	329.9 (−1.1)
MP2	aug-cc-pVDZ	2.764 (+0.058)	1.434 (+0.438)	328.3 (−2.7)
CCSD(T)	aug-cc-pVDZ	2.704 (−0.002)	0.991 (−0.005)	330.6 (−0.4)
FCI	aug-cc-pVDZ	2.706	0.996	331.0
AP1roG	aug-cc-pVTZ	2.784 (+0.123)	0.628 (−0.452)	349.1 (+2.2)
OO-AP1roG	aug-cc-pVTZ	2.690 (+0.029)	1.039 (−0.041)	340.4 (−6.5)
OO-AP1roG-PTa	aug-cc-pVTZ	2.661 (+0.000)	1.086 (+0.006)	344.9 (−2.0)
OO-AP1roG-PTb	aug-cc-pVTZ	2.666 (+0.005)	1.066 (−0.014)	342.7 (−4.2)
DFT/PBE	aug-cc-pVTZ	2.727 (+0.066)	1.451 (+0.371)	331.4 (−15.5)
MP2	aug-cc-pVTZ	2.721 (+0.060)	1.391 (+0.311)	343.2 (−3.7)
CCSD(T)	aug-cc-pVTZ	2.660 (−0.001)	1.054 (−0.026)	247.2 (+0.3)
CCSDT	aug-cc-pVTZ	2.661	1.080	346.9
Exp.		2.673 <sup>118</sup>	1.050 <sup>118</sup>	351.4 <sup>118</sup>



**Figure 2.** Fitted potential energy surface of the ground state of the  $\text{Li}_2$  molecule obtained from different quantum chemistry methods using the cc-pVDZ basis set. Note that the dissociation limit is shifted to zero for all approaches.

out of the optimization complexity in DOSCF calculations. While the latter approach has difficulties in dodging local minima, the OO-AP1roG optimization seems more robust.

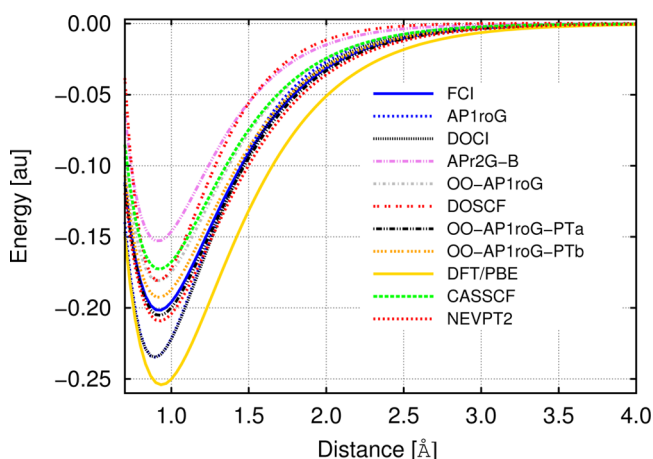
Optimization of the one-particle basis function is necessary to obtain a qualitatively correct description of spectroscopic

constants, especially potential energy depths, within AP1roG. While OO-AP1roG works well for all the spectroscopic parameters, APr2G-B provides only a qualitatively correct description of bond lengths. Perturbative corrections upon OO-AP1roG, that is, OO-AP1roG-PTa, and OO-AP1roG-PTb, further improve bond distances and potential depths. However, opposite to what we observed for LiH and  $\text{Li}_2$ , these two flavors of perturbation theory yield qualitatively different results (cf. Figure 3 and Table 3). While, OO-AP1roG-PTa systematically overestimates, OO-AP1roG-PTb systematically underestimates dissociation energies for all basis sets. Specifically, OO-AP1roG-PTb yields bond distances and vibrational frequencies that are closest to MR-CISD results for all basis sets studied (cf. Table 3). Yet, the simple PTa-correction is not able to recover dynamic electron correlation effects in a balanced way along the dissociation pathway if the basis set size increases (cc-pVDZ  $\rightarrow$  aug-cc-pVDZ  $\rightarrow$  aug-cc-pVTZ). This drawback amounts to growing discrepancies in potential depths (cf. Table 3 and Figures S5 and S6 of the Supporting Information).

Nonetheless, OO-AP1roG-PTa and OO-AP1roG-PTb outperform DFT as well as standard multireference approaches like CASSCF and NEVPT2 for all basis sets studied and are in closest agreement with reference theoretical (FCI and MR-CISD) and experimental data.

**Table 3. Spectroscopic Constants: bond Distance ( $R_e$ ), Potential Energy Depths ( $D_e$ ) and Harmonic Vibrational Frequencies ( $\omega_e$ ) for the  $X^1\Sigma^+$  State of the HF Molecule. Differences with Respect to Reference Data (in a Given Basis Set) Are Listed in Parentheses**

method	basis set	$R_e$ [Å]	$D_e$ [eV]	$\omega_e$ [cm <sup>-1</sup> ]
AP1roG	cc-pVDZ	0.893 (-0.027)	6.393 (+0.902)	4586.8 (+451.6)
DOCI	cc-pVDZ	0.893 (-0.027)	6.389 (+0.898)	4523.6 (+388.4)
APr2G-B	cc-pVDZ	0.913 (-0.007)	4.166 (-1.325)	4754.1 (+618.8)
OO-AP1roG	cc-pVDZ	0.916 (-0.004)	4.928 (-0.563)	4050.8 (-84.4)
DOSCF	cc-pVDZ	0.916 (-0.004)	4.927 (-0.564)	4139.0 (+3.8)
OO-AP1roG-PTa	cc-pVDZ	0.923 (+0.003)	5.588 (+0.097)	4064.6 (-70.6)
OO-AP1roG-PTb	cc-pVDZ	0.920 (+0.000)	5.245 (-0.246)	4126.8 (-8.4)
DFT/PBE	cc-pVDZ	0.936 (+0.016)	6.916 (+1.425)	3877.6 (-257.6)
CASSCF	cc-pVDZ	0.922 (+0.002)	4.703 (-0.788)	4030.6 (-105.2)
NEVPT2	cc-pVDZ	0.925 (+0.005)	5.695 (+0.204)	4118.5 (-16.7)
MR-CISD <sup>8</sup>	cc-pVDZ	0.920 (+0.000)	5.434 (-0.057)	4143.6 (+8.4)
FCI	cc-pVDZ	0.920	5.491	4135.2
AP1roG	aug-cc-pVDZ	0.901 (-0.022)	7.721 (+2.008)	4302.0 (+263.2)
DOCI	aug-cc-pVDZ	0.903 (-0.020)	7.659 (+1.900)	4209.6 (+115.8)
OO-AP1roG	aug-cc-pVDZ	0.916 (-0.007)	5.221 (-0.538)	4184.2 (+90.4)
OO-AP1roG-PTa	aug-cc-pVDZ	0.927 (+0.004)	5.953 (+0.194)	4006.8 (-87.0)
OO-AP1roG-PTb	aug-cc-pVDZ	0.923 (+0.000)	5.594 (-0.165)	4082.8 (-11.0)
DOSCF	aug-cc-pVDZ	0.919 (-0.004)	5.143 (-0.616)	4161.8 (+68.0)
DFT/PBE	aug-cc-pVDZ	0.935 (+0.012)	6.972 (+1.213)	3903.6 (-190.2)
CASSCF	aug-cc-pVDZ	0.919 (-0.004)	4.928 (-0.831)	4057.6 (-36.2)
NEVPT2	aug-cc-pVDZ	0.930 (+0.007)	6.043 (+0.284)	4058.5 (-35.3)
MR-CISD <sup>8</sup>	aug-cc-pVDZ	0.923	5.759	4093.8
AP1roG	aug-cc-pVTZ	0.904 (-0.015)	7.543 (+1.606)	4528.0 (+382.0)
OO-AP1roG	aug-cc-pVTZ	0.912 (-0.007)	5.336 (-0.601)	4196.7 (+50.7)
OO-AP1roG-PTa	aug-cc-pVTZ	0.921 (+0.002)	6.181 (+0.244)	4095.0 (-51.0)
OO-AP1roG-PTb	aug-cc-pVTZ	0.919 (+0.000)	5.783 (-0.154)	4159.0 (+13.0)
DFT/PBE	aug-cc-pVTZ	0.933 (+0.014)	7.510 (+1.573)	3942.8 (-203.2)
CASSCF	aug-cc-pVTZ	0.917 (-0.002)	4.993 (-0.944)	4135.6 (-10.4)
NEVPT2	aug-cc-pVTZ	0.927 (+0.008)	6.247 (+0.310)	4179.3 (+33.3)
MR-CISD <sup>8</sup>	aug-cc-pVTZ	0.919	5.937	4146.0
Exp.		0.917 <sup>118</sup>	6.119 <sup>120</sup>	4138.3 <sup>118</sup>



**Figure 3.** Fitted potential energy surface of the ground state of the HF molecule obtained from different quantum chemistry methods using the cc-pVDZ basis set. Note that the dissociation limit is shifted to zero for all approaches.

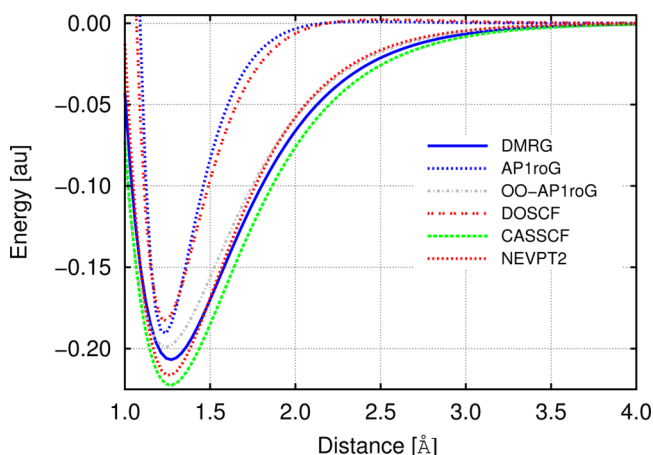
**C<sub>2</sub>.** The carbon dimer molecule is an ideal system to test the performance of new electron correlation approaches. This

diatomic molecule is known to be poorly described at the single configuration Hartree–Fock level; the ground state electronic structure of C<sub>2</sub> is characterized by two dominant configurations  $1\sigma_g^2 1\sigma_u^2 2\sigma_g^2 2\sigma_u^2 1\pi_u^4$  and  $1\sigma_g^2 1\sigma_u^2 2\sigma_g^2 1\pi_u^4 3\sigma_g^2$  as well as a number of other configurations arising from low-lying excited states.<sup>7,121–123</sup> The complicated multireference nature of C<sub>2</sub>, and the peculiar nature of its chemical bonding, has attracted a lot of attention from theoretical chemists in the past two years.<sup>42,124–127</sup>

Spectroscopic constants of the ground state of the C<sub>2</sub> molecule are summarized in Table 4, while Figure 4 illustrates the optimized potential energy surfaces along the reaction coordinate. We should note that our DOCI, OO-AP1roG-PTa, and OO-AP1roG-PTb results are not shown in Table 4. The DOCI solutions could not be converged due to convergence difficulties and local minima problems; this issue could only partially be resolved for DOSCF calculations. However, we did not make an effort to bypass the local minimum problem in DOSCF calculations due to the unfavorable scaling of the method. Besides, OO-AP1roG-PTa and OO-AP1roG-PTb calculations suffered from intruder-states. As a consequence, we could not obtain reasonable fitting parameters for DOCI, OO-AP1roG-PTa, and OO-AP1roG-PTb.

**Table 4. Spectroscopic Constants: Bond Distances ( $R_e$ ), Potential Energy Depths ( $D_e$ ) and Harmonic Vibrational Frequencies ( $\omega_e$ ) for the  $X^1\Sigma_g^+$  State of the  $C_2$  Molecule. Differences with Respect to Reference Data (in a Given Basis Set) Are Listed in Parentheses**

method	basis set	$R_e$ [Å]	$D_e$ [eV]	$\omega_e$ [cm $^{-1}$ ]
AP1roG	cc-pVDZ	1.231 (−0.042)	5.201 (−0.423)	2663.3 (+849.6)
OO-AP1roG	cc-pVDZ	1.240 (−0.033)	5.414 (−0.210)	1930.0 (+116.3)
DOSCF	cc-pVDZ	1.235 (−0.038)	4.981 (−0.643)	2122.7 (+309.0)
CASSCF	cc-pVDZ	1.267 (−0.006)	6.056 (+0.432)	1846.6 (+32.9)
NEVPT2	cc-pVDZ	1.259 (−0.014)	5.893 (+0.269)	1918.3 (+104.6)
DFT/PBE	cc-pVDZ	1.326 (+0.053)	7.261 (+1.637)	718.7 (−1095.0)
DMRG <sup>123</sup>	cc-pVDZ	1.269 (−0.004)	5.628 (+0.004)	1811.4 (−2.3)
FCI <sup>119</sup>	cc-pVDZ	1.273 (+0.000)	not provided	1812.9 (−0.8)
C-MRCI+Q <sup>9</sup>	cc-pVDZ	1.273	5.624	1813.7
AP1roG	aug-cc-pVDZ	1.221 (−0.052)	4.718 (−0.932)	2114.0 (+305.4)
OO-AP1roG	aug-cc-pVDZ	1.240 (−0.033)	5.452 (−0.198)	1938.3 (+129.7)
CASSCF	aug-cc-pVDZ	1.267 (−0.006)	6.047 (+0.397)	1841.2 (+32.6)
NEVPT2	aug-cc-pVDZ	1.259 (−0.014)	5.854 (+0.203)	1924.3 (+115.7)
DFT/PBE	aug-cc-pVDZ	1.193 (−0.080)	7.194 (+1.544)	2257.5 (+448.9)
MR-CISD <sup>7</sup>	aug-cc-pVDZ	1.273	5.650	1808.6
AP1roG	aug-cc-pVTZ	1.199 (−0.053)	4.951 (−1.168)	2862.8 (+1026.8)
OO-AP1roG	aug-cc-pVTZ	1.227 (−0.025)	5.765 (−0.354)	1853.8 (+17.8)
CASSCF	aug-cc-pVTZ	1.255 (+0.003)	6.203 (+0.084)	1841.1 (+5.1)
NEVPT2	aug-cc-pVTZ	1.244 (−0.008)	6.420 (+0.301)	1885.9 (+49.9)
DFT/PBE	aug-cc-pVTZ	1.193 (−0.059)	7.463 (+1.344)	2701.9 (+865.9)
MR-CISD <sup>7</sup>	aug-cc-pVTZ	1.252	6.119	1836.0
Exp.		1.242 <sup>118</sup>	6.409 ± 0.022 <sup>128</sup>	1854.7 <sup>118</sup>



**Figure 4.** Fitted potential energy surface of the ground state of the  $C_2$  molecule obtained from different quantum chemistry methods using the cc-pVDZ basis set. Note that the dissociation limit is shifted to zero for all approaches.

Since Hartree–Fock provides a poor zeroth-order wave function, APr2G-A and AP1roG give inaccurate spectroscopic constants and qualitatively wrong potential energy surfaces (see Figure 4). This problem can be alleviated by orbital optimization; OO-AP1roG yields spectroscopic constants that are in line with both reference theoretical (DMRG and MR-CISD) and experimental data. Yet, APr2G-B, that is, APr2G with OO-AP1roG-optimized orbitals, converged into an excited state for stretched interatomic distances. Therefore, the APr2G-B spectroscopic constants are not included in our discussion.

Remarkably, the OO-AP1roG outperforms CASSCF and NEVPT2 for  $D_e$  and, to some extent,  $\omega_e$  parameters for all basis sets studied. Larger differences in equilibrium bond lengths originate from the lack of dynamic electron correlation effects that cannot be captured completely by (OO-)AP1roG.

Our DFT/PBE calculations resulted in an unphysical double-minimum potential energy surface caused by spin-contamination in the range of 2.0–3.00 Å of interatomic distance. The fitted dissociation curve is thus not shown in Figure 4. This problem diminishes if larger basis sets (larger than cc-pVDZ) are used (see Figures S7 and S8 of the Supporting Information).

## CONCLUSIONS

In this work we have investigated the performance of APr2G and AP1roG (using canonical Hartree–Fock orbitals as well as the variationally optimized orbitals of AP1roG). We investigated the spectroscopic constants of LiH, Li<sub>2</sub>, HF, and C<sub>2</sub> by comparing the performance of our geminal-based approaches to standard quantum chemistry methods and experimental results. In general, the accuracy of our new geminal-based wave function approaches trends as APr2G-A(canonical HF orbitals) < AP1roG(canonical HF orbitals) < APr2G-B(OO-AP1roG orbitals) < OO-AP1roG. While all four methods yield qualitatively correct bond lengths, a quantitatively accurate description of potential energy depths ( $D_e$ ) and vibrational frequencies ( $\omega_e$ ) requires optimization of the one-particle basis functions that form the building blocks of geminals. Hence, APr2G-B and OO-AP1roG provide spectroscopic constants ( $R_e$ ,  $D_e$  and  $\omega_e$ ) that are in good agreement with reference theoretical and experimental data.

The simple and cheap perturbative corrections PTa and PTb on top of OO-AP1roG improve spectroscopic constants for LiH and Li<sub>2</sub> so that spectroscopic accuracy can be achieved. Yet, for strongly multireference systems, like HF and C<sub>2</sub>, the single-reference flavoured PTa and PTb approaches are not appropriate and fail in predicting dissociation energies and vibrational frequencies within spectroscopic accuracy. Different flavors of multireference perturbation theory are currently under investigation in our research group.

In summary, AP1roG and APr2G represent an accurate, cheap, and robust alternative to standard multireference quantum chemistry methods in studying single- and multiple-bond breaking processes in closed-shell systems provided the orbitals are optimized. These new geminal-based methods are size-consistent and size-extensive, and do not require the introduction of active spaces on grounds of computational efficiency. An orbital optimization protocol for APr2G is currently being developed in our laboratory.

## ■ ASSOCIATED CONTENT

### 📄 Supporting Information

Potential energy surfaces for the aug-cc-pVDZ and aug-cc-pVTZ basis sets (Figures S1–S8) and potential energy difference plots for the cc-pVDZ basis set (Figures S9–S12). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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### Notes

The authors declare no competing financial interest.

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