Geometric bonding effects in the X2A1, A2Σ+u, and B2IIg states of Li2F

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Geometric bonding effects in the X2A1, A2Σu+, and B2Πg states of Li2F
Kris W. A. Wright, Daniel E. Rogers, and Ian C. Lane

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I. INTRODUCTION

Recently in this journal, Koput\(^1\) has published a high-level calculation of the lowest rovibrational levels of the Li\(_2\)F molecule based on coupled cluster singlet doublet (triplet) [CCSD(T)] \textit{ab initio} points. In this paper, the lowest electronic states of Li\(_2\)F are investigated further and in particular attention is drawn to a second local minimum on the ground surface. The initial motive for looking at this radical was to calculate a high-level \textit{ab initio} potential for the study of ultracold collisions between a lithium atom and LiF, the simplest example of a class of alkali-alkali halide systems that have not been investigated by modern quantum scattering techniques.

The pseudopotential treatment by Roach and Child\(^2\) for the analogous K+NaCl system suggests that the KNaCl linear geometry has a rather shallow well, whereas KCINa exists as a saddle point on the ground state, but stabilized relative to the K+NaCl asymptote. These two linear geometries are both around $-25$ kJ mol$^{-1}$ relative to the reagents. The later geometry has a very shallow well, whereas KClNa exists as a saddle point on the ground state, but stabilized relative to the K+NaCl asymptote. These two linear geometries are both around $-25$ kJ mol$^{-1}$ relative to the reagents. The later geometry has a very shallow well, whereas KClNa exists as a saddle point on the ground state, but stabilized relative to the K+NaCl asymptote. These two linear geometries are both around $-25$ kJ mol$^{-1}$ relative to the reagents.

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Published \textit{ab initio} and pseudopotential calculations for the dialkali halide systems suggest that the preferred colinear geometry is for the metal to approach the metal end of the alkali halide. Here, \textit{ab initio} calculations on the Li\(_2\)F system reveal that the well depth on the halide side in this radical is much deeper and is a local saddle point associated with the ionic nonlinear global minima. Although many features of the pseudopotential surfaces are confirmed, significant differences are apparent including the existence of a linear excited $A^2\Sigma_u^+$ state instead of a triangular one, a considerably deeper global minimum some $50\%$ lower in energy and a close approach between the $X^2\Delta_1$ and the $A^2\Sigma_u^+$ states, with the $A^2\Sigma_u^+$ minimum $87$ kJ mol$^{-1}$ below the ground state asymptote. All the results can be rationalised as the avoided crossings between a long range, covalent potential dominant within the LiLiF geometry and an ionic state that forms the global minimum. Calculations on the third $^3A'$ potential indicate that even for Li+LiF collisions at ultracold temperatures the collision dynamics could involve as many as three electronic states. © 2009 American Institute of Physics. [doi:10.1063/1.3216373]

Sengupta and Chandra,\(^6\) Veličković \textit{et al.}\(^7\) and most recently Koput. The more recent work on the system has focused on the deep well present on the surface or on the calculation of the thermodynamic limits and has been inspired in part by the experimental verification of a stable Li\(_2\)F molecule by Polce and Wesdemiotis\(^8\) (after an earlier tentative claim had been made by Veljković \textit{et al.}\(^9\)) and its confirmation\(^10,11\) by Yokoyama \textit{et al.} in 2000. These theoretical studies have used primarily single-reference \textit{ab initio} techniques such as CCSD(T). Interestingly, Haketa \textit{et al.}\(^12\) has identified the $D_{oh}$ isomer as being deeper in energy than the corresponding $C_{sv}$ geometry, at odds with the Struve pseudopotential calculations, though this geometry is again unstable with respect to bending motion. The high-quality \textit{ab initio} calculations of the surface here will clarify the shape of the lowest potential energy surface.

Pearson \textit{et al.} also suggested that there would be no low lying excited state, a result in contradiction to the simple pseudopotential model. Gutowski and Simons\(^13\) investigated the lowest electronic states of both the neutral and anion and identified the presence of a number of excited states. The thesis work of Balint-Kurti\(^14\) presented an orthogonalised Moffitt (OM) calculation of the lowest electronic states of Li\(_2\)F, in both linear and $C_{2v}$ geometries. The OM method corrects for the poor calculation of electron affinity common in early \textit{ab initio} techniques. The most comprehensive \textit{ab initio} work on the alkali-alkali halide system is an early configuration interaction (CI) study by Yamashita and Morokuma\(^15\) who studied the lowest four $^2A'$ and two $^2A''$ surfaces of K+NaCl using an aug-cc-pVTZ equivalent basis set. They identify the predominant configuration of the ground state as the complex K$^+$+NaCl$^-$ and they suggest that as the K atom approaches the NaCl molecule it experiences a repulsive interaction until an electron transfer event takes place to form the ionic complex. This repulsive interaction...
FIG. 1. (a) CCSD(T) potential of the global minimum in the \( X^2A_1 \) potential of the Li,F molecule with the geometry held at \( C_2 \), in all calculations. (b) Equivalent surface calculated using a two state SA-CASSCF calculation. Energies are in kJ mol\(^{-1}\) with zero energy corresponding to the global minimum for the LiFLi molecule in the \( X^2A_1 \) state.

has also been identified in Li+FLi by Sengupta and Chandra\(^6\) using coupled cluster calculations. The first excited \( 2A^1 \) surface was predicted as having a minimum in the linear KClNa geometry though the potential was very flat between 180° and 120°. It is important to note that these potentials were based on just 120 calculated points, each involving just 4000 CSFs.

One drawback of the most recent theoretical work is that it has concentrated on the use of single reference quantum chemistry methods such as MP2, CCSD(T), etc. The switch in bonding, from covalent (neutral Li+LiF fragments) to ionic (at the global minimum), means that the problem is correctly treated by using multireference techniques such as Complete Active Space Self-Consistent Field (CASSCF) and MultiReference CI (MRCI). Gao et al. have conducted a CASSCF and MRCI study\(^{10}\) but have performed the calculation with seven electrons in ten orbitals, a CAS(7,10) calculation in their nomenclature, whereas a full valence calculation would be CAS(9,12). In this paper, the lowest three electronic states of Li,F are investigated by \textit{ab initio} quantum chemistry using all the valence electrons and orbitals. The CASSCF method is used to clarify the origin of the ionic-covalent interaction and in particular the geometric dependence of any curve crossings. Nonadiabatic effects, in particular the crossing of covalent and ionic potentials, are important in the diatomic alkali halide species\(^{17,18}\) though there is an absence of detailed \textit{ab initio} work on the corresponding triatomic systems.

Unfortunately, the CASSCF method, though relatively inexpensive and multireference in nature, does not capture the dynamic electron correlation, though in this system these effects might be expected to be relatively insignificant. To explore this more fully, CCSD(T) (single reference) and MRCI (multireference) calculations are presented to test the validity of the CASSCF method in such a system. Comparison will be made between the results here and earlier treatments, particularly the pseudopotential work.

II. \textit{AB INITIO} CALCULATIONS

Calculations were performed at the CCSD(T), CASSCF, and MRCI(Q) (MRCI with Davidson correction) levels using either GAUSSIAN 98 (Ref. 19) and GAUSSIAN 03 (Ref. 20) or PC GAMESS/FIREFLY (Ref. 21) running on a mixed cluster of Core 2 Duo (E6550 2.33, E6600 2.4, E6750 2.66, and E8200 2.66 GHz) and Quad (Q6700 2.66 GHz) PCs all with 2Gbyte of RAM. As this paper is concerned with two different molecular geometries of Li,F, two different coordinate systems are adopted. The first is a special case of Radau coordinates used to describe the LiFLi hypermetalated molecular system. The second is the three-atom Jacobi coordinates and describes the LiF+Li exchange reaction. In all calculations the augmented correlation consistent triple zeta (aug-cc-pVTZ or AVTZ) basis set for both fluorine\(^{22,23}\) and lithium\(^{24}\) atoms (denoted here as an AVTZ/AVTZ calculation) is adopted.

CCSD(T) and CASSCF calculations of the LiFLi molecule about its minimum geometry were conducted and compared with the results of Koput. This is to validate the CASSCF method as an accurate description of the ground potential surface. One significant distinction, however, is that the majority of the calculations presented (Fig. 1) in this work involve the valence electrons only, with the 1s orbitals on the atoms frozen. One advantage of the CASSCF method is the ability to calculate multiple potential surfaces of identical symmetry and therefore reveal avoided crossings between surfaces, though the resulting state-averaged CASSCF (SA-CASSCF) wave functions are inevitably less accurate than a single state CASSCF or CCSD(T) wave function. For the purposes of determining the influence of electron correlation on the ground state surface the SA-CASSCF calculation was performed over two states (two-state SA-CASSCF) and the ground state surface (Fig. 1) compared with the CCSD(T) potential.

The second group of surfaces, for the LiF+Li exchange reaction, used additional SA-CASSCF calculations over the
TABLE I. The equilibrium geometries for the ground and excited electronic states of the LiF molecule calculated at the CCSD(T), three-state SA-CASSCF and MRCI(Q) levels with the AVTZ basis set for both atoms. The angle measured is the interior angle in the LiFLi system formed from the two LiF bonds: this angle is not defined in the LiLiF geometry (the Li-Li separation is 3.352 Å). Zero energy is defined by the fragments LiF and Li in their ground states (calculated by each method in turn) with the LiF bond length set at the energy minimum (calculated by CASSCF as 2.97\(a_0\)) and the fragments separated by 30\(a_0\). The MRCI(Q) calculations used the same geometry that was optimized at the three-state SA-CASSCF level and therefore is not representative of the exact global MRCI(Q) minima.

<table>
<thead>
<tr>
<th>Method</th>
<th>State</th>
<th>(r(\text{Li}F)/\text{Å})</th>
<th>(\theta)</th>
<th>Energy/(E_h)</th>
<th>Relative energy/kJ mol(^{-1})</th>
</tr>
</thead>
<tbody>
<tr>
<td>HF</td>
<td>(X^3A_1)</td>
<td>1.669</td>
<td>101.3</td>
<td>-114.469 584 8</td>
<td>-138.41</td>
</tr>
<tr>
<td>CCSD(T)</td>
<td>(X^3A_1)</td>
<td>1.697</td>
<td>100.3</td>
<td>-114.763 191 10</td>
<td>-144.63</td>
</tr>
<tr>
<td>CASSCF</td>
<td>(A^3\Sigma^+)</td>
<td>1.666</td>
<td>180</td>
<td>-114.610 573 45</td>
<td>-76.56</td>
</tr>
<tr>
<td></td>
<td>(B^3\Pi_g)</td>
<td>1.662</td>
<td>180</td>
<td>-114.575 117 31</td>
<td>15.80</td>
</tr>
<tr>
<td></td>
<td>(X^5\Sigma^+ (\text{LiLiF}))</td>
<td>1.597</td>
<td>N/A</td>
<td>-114.608 062 98</td>
<td>-17.59</td>
</tr>
<tr>
<td>MRCI(Q)</td>
<td>(X^3A_1)</td>
<td>1.696</td>
<td>99.46</td>
<td>-114.766 752 91</td>
<td>-142.41</td>
</tr>
<tr>
<td></td>
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<td>-87.66</td>
</tr>
<tr>
<td></td>
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<td>(X^5\Sigma^+ (\text{LiLiF}))</td>
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<td>N/A</td>
<td>-114.719 822 44</td>
<td>-20.23</td>
</tr>
</tbody>
</table>

lowest three states of \(A^1\) symmetry (valence electrons only). A potential energy surface was constructed from over 600 CASSCF points. In addition, at the two linear configurations (0°,180°) MRCI(Q) points for the lowest two \(2\Sigma^+\) states were computed based on a two-state SA-CASSCF wave functions and calculated to assess the effects of dynamic electron correlation (or lack of) on the interacting potentials. MRCI(Q) calculations were also performed on the \(2\Pi\) state based on a single CASSCF wave function. With the MRCI(Q) calculations the lowest three orbitals were closed while the remaining 162 were opened allowing for single and double excitations out of the valence reference space. As a result the electronic wave function was generated from over 19 \(\times\) 10⁶ CSFs, creating a rather large and computationally expensive calculation. The potentials were fitted at long range so that they corresponded with the asymptotic limit thus enabling both the \(2\Sigma^+\) and \(2\Pi\) states to be compared. Details of all three potentials calculated are presented in Table I.

III. RESULTS AND DISCUSSION

A. LiFLi molecule

In Fig. 1(a), the calculated CCSD(T) potential in \(C_{2v}\) symmetry around the global minimum is presented. This dictates the bending motion through the LiFLi linear geometry and clearly shows the unstable linear \(D_{sh}\) isomer observed elsewhere as a local maximum (saddle point) in the bending potential. The minimum geometry here calculated with the AVTZ/AVTZ basis set (100.3° and 1.697 Å) is similar to that calculated by Koput with the larger V5Z/AVSZ basis set (101.2° and 1.686 Å). Indeed the resulting Li–F–Li angle lies in between that of the VTZ/AVTZ and VQZ/AVQZ calculations reported by Koput. Overall a trend emerges that triple zeta basis sets underestimate the bond angle while overestimating the bond length. The results obtained here are also consistent with the DFT, MP2, and CCSD(T) calculations by Ochsenfeld and Ahrlich and Sengupta et al. The three harmonic vibrational frequencies were calculated at the HF (241.397, 677.22, and 711 cm⁻¹), single state CASSCF (236.90, 649.81, and 678.54 cm⁻¹), and CCSD(T) (248.02, 637.51, and 673.49 cm⁻¹) levels using the same basis set, showing the effect of increased electron correlation is to reduce the vibrational frequencies of the highest energy modes. The saddle points are located in the \(D_{sh}\) linear geometry at 1.667 Å [CCSD(T) calculation] and 1.662 Å (two-state SA-CASSCF) with barrier heights of 1775 and 1846 cm⁻¹, respectively. The additional stabilization of the transition state thanks to dynamic electron correlation lowers the barrier height in the CCSD(T) result.

Koput also calculated the minimum at the complete basis set (CBS) limit and furthermore included relativistic and core-valence effects to determine the equilibrium geometry as 101.00° and 1.671 Å. The importance of electron correlation among the core electrons was highlighted in that work, with the major contribution being a reduction in the Li–F bond length of the order -0.018 Å. However, Iron et al. have shown that in general the core-valence correlation is small for lithium based compounds compared to the other alkali metals. To assess the influence of core-valence effects, calculations were performed on the ground state using the CASSCF, CCSD(T), and MRCI methods with the aug-pCVTZ basis set developed by Woon and Dunning. The dissociation energy of the molecule calculated by the three methods is compared with previous studies in Table II. When the calculation includes only the valence electrons, the changes in geometries and energies are less than 1%. However, when the core electrons are included, the changes are more significant: at the CCSD(T) level, the changes are -0.012 Å, +0.21° and +2.02 kJ mol⁻¹. These changes are still, however, within “chemical accuracy” and therefore the smaller AVTZ basis set is used in our subsequent calculations. Further support was provided by calculations of the SA-CASSCF equilibrium geometry for all three states using the aug-pCVTZ basis set and an active space including either the valence electrons alone or all the electrons. Comparing the two basis sets, the X-state geometry with a valence (full)...
The only noticeable difference is that the CASSCF potential is compares with the SA-CASSCF results are of comparable quality. In strikingly similar to the CCSD only adds a small contribution to the shape and depth of the correlation, although still important for absolute energies, between the two calculations reveals that the dynamic electron potentials to be computed simultaneously. The similarity between the two calculations was performed in [12] because the bond angle seems rather large in their work and the calculation by Chen28 which is not at the equilibrium geometry, the average theoretical dissociation energy $D_0$ is 146.11 ± 1.36(standard) kJ mol$^{-1}$.

**B. Li+LiF exchange**

To explore the ground state potential in further detail, a three-state SA-CASSCF calculation of the lowest $^2\Sigma^+$ potential was performed in $C_s$ symmetry using Jacobi coordinates to describe the Li+LiF system. This encompassed a larger range of molecular geometries than the CCSD(T) calculation and should more accurately describe the exchange reaction Li+LiF. Such potentials would be useful for scattering calculations and for modeling the cooling of LiF molecules by ultracold Li atoms (buffer gas cooling). The polar plot in Fig. 2 depicts the Li+LiF potential energy surfaces for the ground $^2\Sigma^+$ and first excited $^2\Pi_u$ states with the LiF diatomic bond length held at $3a_0$, close to the equilibrium bond distance calculated by CASSCF (2.99$a_0$). These calculations reveal the presence of a second minimum on the $^2\Sigma^+$ potential.
tial corresponding to the linear LiLiF geometry. This minimum is considerably shallower than the global minimum calculated earlier with a depth of just 17.59 kJ mol\(^{-1}\), in agreement with the result from Pearson et al., and about half the pseudopotential value of Struve. The ground state surface in Fig. 2 has a barrier, corresponding to the molecular rearrangement from the \(\text{X}_2^2\) configuration to the \(\text{X}_2^1\) configuration, of 2.6 kJ mol\(^{-1}\). Such a potential may support a ground vibrational state of the LiLiF molecule, though in this instance it is unlikely as the zero point energy for \(\text{X}_2^2\) is 6 kJ mol\(^{-1}\), greater than the angular barrier and hence the linear LiLiF isomer cannot be isolated.

The global minimum in the \(\text{X}_2^1\) state sits some 140 kJ mol\(^{-1}\) below the LiF+Li asymptote and the linear saddle point is clearly identified. The SA-CASSCF calculations demonstrate that the interaction between the Li atom and the LiF molecule is attractive on both surfaces for all lithium atom approaches. There is no evidence of a repulsive interaction as suggested by Chandra and co-workers and the SA-CASSCF ground state potential is broadly similar to the semiempirical potential of Roach and Child except that the \(D_{\text{sh}}\) geometry is lower in energy than \(C_{\text{sh}}\) for the Li\(_2\)F system, in agreement with Haketa et al. It has been postulated that the dissociation energy of M\(_2\)X molecule will be similar to the dissociation energy of the M\(_2^+\) cation. The calculated dissociation energy is 12 209 cm\(^{-1}\) (177.60 kJ mol\(^{-1}\)), while the experimental value for the Li\(_2^+\) transition is 14 903 cm\(^{-1}\) (178.28 kJ mol\(^{-1}\)). The value here is almost identical to that found by Pearson et al. and some 50% deeper than the pseudopotential calculation. Clearly, an isolated Li\(_2\)F radical in its ground state is very stable to fragmentation and its experimental fragility must be a consequence of its chemical reactivity. Figure 2 also emphasizes that the energy range studied by Koput is just a small part of the deep ground state potential well. Inspection of the electronic wave function indicates that this minimum is ionic in nature and Figs. 3 and 4 reveal that the lowest three \(2\) \(2\) states are the result of extensive avoided crossings between ionic and covalent potentials.

C. Excited electronic states

Figures 2 and 3 illustrate how the bottom of the \(A^2A^1\) potential lies within 64 kJ mol\(^{-1}\) of the global \(X^2A^1\) minimum and considerably below the Li+LiF asymptote. At long internuclear distances, the \(X^2A^1\) state is covalent and is bound thanks to a dipole-atom interaction potential, clearly visible at the larger Jacobi angles. As the angle decreases (the Li approaches the fluorine side of the diatomic) the co-
valent surface is crossed by an ionic state following electron transfer from the Li atom to the LiF diatomic. Consequently, the $A^2A'$ state is formed from the lowest covalent surface at short range and from the ionic potential beyond this inner crossing point. Furthermore, the minimum of the $A^2A'$ is actually a linear $A^2\Sigma_u^+$ state and not the triangular configuration suggested by Struve.

Three further observations about the lowest two $2\Sigma_u'$ surfaces should be made. First, care must be taken when comparing the ionic-covalent interaction seen here to a classic harpoon interaction. Comparison of the potentials at different Jacobi angles $\theta$ illustrates that the ionic-covalent interaction changes with the angle of the Li atom relative to the LiF molecule. Therefore, in this case there is no such thing as a “harpoon radius.” Rather it may be thought of as a harpoon arc or cone as the electron transfer process clearly demonstrates a significant stereodynamic character on the three lowest potentials. The ionic potential does not cross the covalent state in the LiLiF geometry and thus the minimum of the ionic surface lies above the Li+LiF asymptote. The ionic potential curve cannot be simulated by a simple Rittner potential, unlike other harpoon type mechanisms. On the ground state surface the avoided crossing takes place at around 4.5$a_0$ in the LiFLi arrangement, a result in broad agreement with Morokuma. Evidence for a similar crossing can be seen at $\sim 7a_0$ in the $A^2\Sigma_u^+$ state.

Second, the presence of the minimum on the second $2\Sigma_u'$ state below the Li+LiF asymptote means that collisions between these species involve both the $X^2A'$ and $A^2A'$ surfaces and consequently nonadiabatic effects cannot be ignored. The fact that the excited state lies deep within the ground state minimum is another new result here, as Struve places the minimum of the $A^2A'$ excited state of Na+NaCl well above the ground asymptote by at least 40 kJ mol$^{-1}$.

Third, the low lying $A^2A'$ state means that the vibrational and rotational levels of the ground $X^2A'$ state will suffer a great deal of perturbation by the interloping state. However, the calculations of Koput dealt with the energy region below the ground state saddle point and hence were not affected by the presence of the low lying $A^2A'$ potential minimum. Calculation of the lowest $2\Sigma_u'$ state reveals it is also the result of such an interaction, the ionic state just visible as a kink in the repulsive covalent state correlating to $Li^2P$+$LiF$ at $\theta=120^\circ$. The minimum of the $B^2A'$ state is ionic with $D_{oh}$ symmetry and lies within...
15 kJ mol$^{-1}$ of the ground state asymptote. At long-range it possesses a small barrier for Li/H$_2$O$^+$ collision at small values of $\theta$. 

D. Dynamic electron correlation

MRCI(Q) calculations were performed at the two colinear geometries on the potential surfaces in order to explore further any possible effects due to dynamical electron correlation on both the ground and excited surfaces. Again, the diatomic alkali halides have been a popular system for studying the effects of dynamic electron correlation effects, especially LiF.$^{32,33}$ Each CI point in $C_{2v}$ ($C_{\infty v}$) symmetry took around 30 h of CPU time and the Davidson correction was included in the calculations. Figure 4(a) shows the lowest three states calculated at the MRCI(Q) levels in the linear geometries LiLiF and LiFLi(2$^2\Sigma^+$ and 1$^2\Pi$). The MRCI(Q) and SA-CASSCF potentials are almost identical. In Fig. 4(b) the differences in relative energy between MRCI(Q) and SA-CASSCF points are plotted for the two linear geometries. It clearly shows that the difference in energy between the methods is greater for the LiF+Li approach geometry than for Li+LiF. This result is rather counterintuitive as one would expect that dynamic electron correlation would be greater in the LiLiF than in the LiFLi arrangement, due to the nonionic nature of the bonding. The SA-CASSCF barrier on the $A^2\Sigma^+$ state (green line) at zero degrees is 9.91 kJ mol$^{-1}$, though it is markedly reduced in height by including the extra electron correlation (to 5.43 kJ mol$^{-1}$) in the MRCI(Q) calculation. The most significant differences, however, are situated in the repulsive wall of the potential where the rate of change of energy with bond distance is the greatest. The similarity between the MRCI(Q), CCSD(T), and SA-CASSCF potentials is consistent with the remarkably successful semiempirical potentials that were calculated using a model of a single valence electron moving in a potential field$^2$ generated by the metal ion cores and the F$^-$ anion.

The covalent component of the $X^2A'$ surface is also attractive and the minimum is due to a long-range dipole-induced dipole interaction first discovered in the K+RbCl reactive scattering experiment of Miller et al.$^{34}$, which was also the first to demonstrate the formation of a long lived collision complex in an exchange reaction. The present calculations confirm that the long-range potential is indeed of a dipole-induced dipole type as the covalent-ionic crossing takes place at a relatively short range. The repulsive wall of all three potentials are pushed to longer distances in the LiLiF geometry because the van der Waals radius$^{35}$ of Li is greater than F and the separation is determined from the LiF
the ground state asymptote. As the zero point energy of LiF is 4.55 cm$^{-1}$ (5.4 kJ mol$^{-1}$), this suggests that even an ultracold collision with a ground rovibrational LiF molecule could involve as many as three potential energy surfaces (making the reasonable assumption that the well will deepen at the CBS limit). This is a rather unexpected situation as the Li($^3S$)+LiF($^2Σμ^+$) asymptote correlates with just a single (nondegenerate) potential energy surface and there is no reaction barrier. As the additional electronic states are bound, scattering resonances will feature prominently in the collision dynamics. Following the argument of Roach and Child, the effect of these nonadiabatic transitions will be to increase the reactive cross sections relative to those for inelastic scattering, a phenomenon first observed in the KBr+Na system by Moulton and Herschbach. In that particular experiment, the reactive cross section was ten times that of the nonreactive, a result that could have significant consequences for attempts to sympathetically cool alkali halides with ultracold alkali atoms.

It has been previously noted that electron correlation effects do not have a strong effect on the F+Li$_2$ surface (as well as H+Li$_2$). The present results indicate that this is not entirely true over the whole potential and it is clearly significant for the A$^2A'$ excited state potential about its potential minimum and for describing the barrier on the B$^2A'$ state. However, large parts of the ground state surface are indeed modeled well by the SA-CASSCF method. For ultracold collisions, it is possible that any differences are essentially irrelevant because of the long de Broglie wavelengths of the reagents. A comparison of quantum scattering calculations at ultracold energies on SA-CASSCF and MRCI(Q) surfaces would be instructive.

IV. CONCLUSION

High-level ab initio calculations on the Li$_2$F molecule have revealed the presence of two minima on the ground surface, one corresponding to an ionic molecule and the second, shallower well to a dipole-induced dipole interaction. At extended lithium distances, the ionic attraction is between Li$^+$ and the LiF anion. The global minimum for $X^2A'$ calculated here is consistent with the recent results by Koput and others but the relative well depths calculated are at variance with semiempirical potentials for similar alkali+alkali halide systems. The first two excited states possess very deep potential wells in the $D_{ob}$ geometry that will influence both the rovibrational levels of the ground state and the scattering dynamics, even at ultracold temperatures. It was also found that dynamic electron correlation effects were relatively small in this system, a result that is consistent with previous work on lithium containing compounds. Consequently, SA-CASSCF calculations prove to be surprisingly accurate, even for the electronically excited states.

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