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# ELECTRON CORRELATION AND THE REALITY OF XeF2

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Electron Correlation and the Reality of  $\operatorname{XeF}_2$ 

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#### **ABSTRACT**

The electronic structure of xenon difluoride has been studied using ab initio theoretical methods. The primary goal was to determine whether current theoretical methods are capable of yielding a reasonable value of the dissociation energy of XeF<sub>2</sub>. A Slater function basis set of slightly better than "double zeta plus polarization" quality was employed. Four different types of wave functions were investigated: twoconfiguration SCF, full valence configuration interaction (CI), the first-order wave function, and a larger 1234 configuration wave function including all double excitation from the  $10\sigma_{\alpha}$ orbital. Although the TCSCF symmetric stretching potential curve has both a minimum and maximum, the minimum lies above the comparable energy of separated Xe + 2F. However, the two most complete wave functions predict dissociation energies of 1.97 and 2.14 eV, in qualitative agreement with experiment, 2.78 eV. All four wave functions provide good predictions of the Xe-F equilibrium bond distance. As was the case for KrF2, the bonding in XeF, is found to conform quite closely to Coulson's model

$$F ext{ Xe}^+ ext{ } F^- ext{ } ext{ } F^- ext{ Xe}^+ ext{ } F$$

near the equilibrium geometry. The role of the "outer orbitals"

5d and 4f appear to be a quantitative rather than qualitative one.

#### INTRODUCTION

Xenon difluoride appears to be the simplest known Xecontaining molecule, although there is still some controversy concerning the existence of the XeF radical. As such, XeF<sub>2</sub> plays a special role in the chemistry of the noble gases. XeF<sub>2</sub> was first prepared in 1962, shortly after Bartlett's discovery of XePtF<sub>6</sub>, and several relatively simple methods of preparation are now available. The dissociation energy for the process

$$XeF_2 \rightarrow Xe + 2F$$
 (1)

is  $\sim$  64 kcal/mole  $^6$  = 2.78 eV. Assuming the value  $^7$  38.8  $\pm$  2.3 kcal/mole for the dissociation energy  $D_e$  of  $F_2$ , the molecular dissociation energy for the process

$$XeF_2 \rightarrow Xe + F_2$$
 (2)

is found to be  $^{\circ}$  25 kcal/mole. For comparison, the smaller KrF $_2$  molecule is known to lie energetically <u>above</u> (by  $^{\circ}$  15 kcal/mole) the analogous dissociation limit Kr + F $_2$ . This difference between KrF $_2$  and XeF $_2$  explains the transient nature of the former as compared to the relative stability of the latter. The geometrical structure of XeF $_2$  is known from infrared and Raman studies to be linear and symmetric,  $^3$  corresonding to point group D $_{\infty h}$ . Reichman and Schreiner have determined the gas-phase Xe-F bond distance to be 1.977  $^{\pm}$  0.002  $^{\circ}$  A. In the crystalline phase, a neutron diffraction study has yielded 2.00  $^{\pm}$  0.01 for the Xe-F equilibrium separation.

Since  $XeF_2$  is a well-characterized species, many other properties  $^3$  have been experimentally determined. However, the dissociation energy and structure are particularly important and of direct relevance to the present theoretical discussion.

The discovery of the existence of noble gas compounds in the early 1960's was viewed in some quarters as an "embarrassment" to theoretical chemistry. However, this would seem an unfair generalization, since only the crudest empirical and semi-empirical theoretical methods could be applied to molecules containing xenon. More recently, Rosen and Ellis $^{10}$  have carried out relativistic Dirac-Slater computations on  $XeF_2$  using the p<sup>1/3</sup> local exchange approximation. However, to date the only study of polyatomic xenon compounds which includes exchange exactly appears to be that of Basch, Moskowitz, Hollister, and Hankins $^{11}$ on XeF2, XeF4, and XeF6. Their work, although well ahead of its time, used only a small basis set and intentionally concentrated on qualitative features of the electronic structure, wisely making no attempt to predict binding energies relative to the separated atoms and molecules. It now seems well-established 12 that reliable a priori predictions of dissociation energies require a) basis sets of at least "double zeta plus polarization" quality and b) explicit treatment of electron correlation, usually by configuration interaction. The development of ab initio theoretical methods has now proceeded to the point where a reasonable theoretical description is quite feasible. Hence the aim of the present study was to determine whether the theoretical methods used successfully in recent years to study "conventional" molecules are capable of

providing accurate predictions of the properties of xenon difluoride.

## Theoretical Approach

We should state at the outset that the present treatment of XeF<sub>2</sub> is of a nonrelativistic nature. The age-honored justification for the neglect of relativistic corrections is that they affect only the inner shells and hence presumably do not affect the chemistry, which is dictated by the valence electrons. This hypothesis has recently been given some factual support by the work of Schwenzer et al. <sup>13</sup> on the PbO molecule. Nevertheless, the assumption of a nonrelativistic model is without satisfactory theoretical justification and at present must be considered a necessary evil.

The basis set of Slater-type functions is shown in Table

I. In the accepted parlance, this basis is of slightly better
than "double zeta plus polarization" calibre. 11 The sp basis
for fluorine is the "nominal" (4s 3p) basis of Bagus and Gilbert 14
and yields a self-consistent-field (SCF) total energy of -99.4081
Hartrees, as opposed to the true Hartree-Fock energy, -99.410
Hartrees. 14 As seen in Table I, this basis set has been augmented by two 3d and one 4f polarization functions.

For Xe, our basis was modeled after that of Synek and Timmons<sup>15</sup> for Pr<sup>3+</sup>. The original set (10s 8p 5d) is of double zeta quality except for the 4d functions, of which there are three. Exponent

optimization was carried out for the  $^1$ S atomic ground of xenon using the program of Roos et al.  $^{16}$  The final atomic SCF energy obtained was -7232.1204 hartrees, which may be compared to the numerical Hartree-Fock results of Mann,  $^{17}$  -7232.14 hartrees, and Fischer,  $^{18}$  -7232.153. Recently Roetti and Clementi  $^{18}$  have reported a double zeta basis yielding energy -7232.1189 hartrees  $^{19}$  and a more extended basis yielding -7232.1302 hartrees.  $^{20}$  Thus it appears that our basis is nearly optimum considering its size and yields an SCF energy within a few hundredths of a hartree of the Hartree-Fock limit. The final xenon basis evolved through the addition of two 5d and two 4f functions, which serve as polarization functions. Thus the final basis set includes 95 Slater-type orbitals (STO's), counting  $\pi_+$ ,  $\delta_+$ , and  $\phi_+$  only once.

Four different kinds of wave functions were used in the present work:

I. The Two-Configuration SCF, TCSCF, wave function required to dissociate to the three SCF atomic wavefunctions F + Xe + F. Excluding the inner 56 electrons, the two configurations are

$$\dots 6\sigma_{\mathbf{u}}^{2} 4\pi_{\mathbf{u}}^{4} 3\pi_{\mathbf{g}}^{4} 10\sigma_{\mathbf{g}}^{2} 5\pi_{\mathbf{u}}^{4}$$
 (3)

$$\dots 6\sigma_{u}^{2} 4\pi_{u}^{4} 3\pi_{g}^{4} 7\sigma_{u}^{2} 5\pi_{u}^{4}$$
 (4)

II. The eight-configuration full valence configuration interaction (CI), which in addition to (3) and (4) includes

$$4\pi_{u}^{4} 3\pi_{g}^{4} 10\sigma_{g}^{2} 5\pi_{u}^{4} 7\sigma_{u}^{2}$$
 (5)

$$6\sigma_{u} 4\pi_{u}^{4} 3\pi_{g}^{4} 10\sigma_{g}^{2} 5\pi_{u}^{4} 7\sigma_{u}$$
 (6)

$$6\sigma_{\mathbf{u}}^{2} \ 4\pi_{\mathbf{u}}^{4} \ 3\pi_{\mathbf{g}}^{4} \ 10\sigma_{\mathbf{g}}^{2} \ 5\pi_{\mathbf{u}}^{2} \ 7\sigma_{\mathbf{u}}^{2} \tag{7}$$

$$6\sigma_{\mathbf{u}}^{2} \ 4\pi_{\mathbf{u}}^{2} \ 3\pi_{\mathbf{g}}^{4} \ 10\sigma_{\mathbf{g}}^{2} \ 5\pi_{\mathbf{u}}^{4} \ 7\sigma_{\mathbf{u}}^{2} \tag{8}$$

$$6\sigma_{u}^{2} 4\pi_{u}^{4} 3\pi_{g}^{2} 10\sigma_{g}^{2} 5\pi_{u}^{4} 7\sigma_{u}^{2}$$
 (9)

$$6\sigma_{\rm u}^2 \ 4\pi_{\rm u}^3 \ 3\pi_{\rm g}^4 \ 10\sigma_{\rm g}^2 \ 5\pi_{\rm u}^3 \ 7\sigma_{\rm u}^2 \tag{10}$$

- III. The first-order wave function,  $^{12,21}$  including only those configurations in which no more than a single electron occupies an orbital beyond the valence shell, i.e., beyond  $7\sigma_{\rm u}$ . Further restrictions invoked here are:
  - a) 56 electrons are constrained to occupy the innermost two-configuration SCF orbitals in all configurations.
  - b) The space into which the CI calculations were performed was chosen in a somewhat unusual manner. The occupied TCSCF orbitals  $(10\sigma_{\rm g},~7\sigma_{\rm u},~5\pi_{\rm u},~{\rm and}~3\pi_{\rm g})$  were supplemented by 70 additional MO's. The added MO's were chosen to be single STO basis functions on Xe and symmetric (g or u) combinations of one basis function in each F. This MO set, although it is not orthogonal, spans the full space of the 95 STO basis set. The added MO's which correspond to basis functions describing the atomic ones, Xe 1s to 4s, 2p to 4p, and 3d and F 1S, were deleted  $(10\sigma,~4\pi,~{\rm and}~1\delta~{\rm in}~{\rm all})$ . The remaining MO's were

orthogonalized. This process allowed us to reduce the number

of configurations in the CI wave function without significant loss of accuracy.

In this way a total of 992 configurations were included in the present first-order wave functions.

IV. In addition to configurations of the type included in the first-order wave function, a further class of configurations has been studied. These configurations are double excitations of the type  $10\sigma_g^2 \to n\sigma$  mo, or

$$\dots 6\sigma_{\mathbf{u}}^{2} 4\pi_{\mathbf{u}}^{2} 3\pi_{\mathbf{g}}^{4} 5\pi_{\mathbf{u}}^{4} \text{ no mo}$$
 (11)

As discussed by Wahl and Das,  $^{22,23}$  these configurations have no contribution to the wave function as the molecule dissociates to the three atoms. However, a substantial contribution is possible near the equilibrium internuclear separation, and hence these configurations tend to increase the predicted dissociation energy. Our fifth wave function, which is labeled "first-order +  $\sigma$  doubles" here, includes configurations of this type in addition to those present in the conventional first-order wave function, III. A total of 1234 configurations are included in this final wave function.

Wave functions I - III were studied in our earlier study 21 of KrF<sub>2</sub>. In addition, it was found that the single-configuration SCF wave function yields a potential curve with its minimum 2.98 eV above the SCF energy of the three atoms F + Kr + F. For this reason, conventional SCF calculations are not reported here for XeF<sub>2</sub>. For KrF<sub>2</sub> two-configuration SCF and full valence CI treatments give essentially indistinguishable results. These wave functions dissociate properly to three SCF atom wave functions, but predict no minimum, only an interesting inflection point, in the symmetric stretching potential curve. Finally, the first-order CI did yield a potential minimum of depth 0.39 eV, as compared to experiment, 1.01 eV. Perhaps even more interesting, a potential maximum of 0.22 eV was found at a larger internuclear separation, 2.42 Å. From a theoretical viewpoint, then, it is of interest to

see whether the error in the predicted dissociation energy of  ${\rm KrF}_2$  is of an absolute ( $\sim 0.6$  eV) or relative nature. Should the latter be the case we would obtain only 40% of the dissociation energy of  ${\rm XeF}_2$  as well.

#### Potential Energy Curves

Total energy results are summarized in Table II and Figure 1, which illustrates the potential curves for the symmetric stretching of XeF<sub>2</sub>. Predicted bond distances and dissociation energies are given in Table III. Figure 1 does not include wave function II, the 8-configuration valence CI, since as for  $\mathrm{KrF}_2$  it is essentially indistinguishable from the TCSCF curve. Note that although there is a potential maximum in the TCSCF potential curve, its minimum lies 0.15 eV above the dissociation limit F + Xe + F. Thus the TCSCF wave function does not predict XeF, to be a thermodynamically stable molecule. However, if one went to the Hartree-Fock limit of a complete basis set, it is probable that XeF, would be bound (by perhaps 0.2 to 0.4 eV) in the TCSCF limit. It is also noteworthy that the TCSCF and Valence CI wave functions yield predicted Xe-F bond distances within a few thousandths of an angstrom of experiment.<sup>7</sup>

The first-order wave function appears to describe XeF<sub>2</sub> in a qualitatively acceptable manner. That is, a substantial dissociation energy is predicted, 71% of the experimental value.

This  $D_e$  value of 1.97 eV is large enough to guarantee the exothermicity of the process  $Xe + F_2 \rightarrow XeF_2$ . The absolute error of this  $D_e$ , 0.81 eV, is quite comparable to the 0.62 eV error found for  $KrF_2$ . The predicted value of  $r_e(X_e-F)$  is 0.02 Å longer than the experimental gas phase result, 7 but curiously in perfect agreement with the crystalline result from neutron diffraction. 9 Of particular interest is the fact that the potential maximum predicted by the comparable calculation on  $KrF_2$  has disappeared in  $XeF_2$ . This maximum was due to the ionic

$$F Kr^{+} F^{-} \longleftrightarrow F^{-} Kr^{+} F$$
 (11)

nature of the molecule to the left of the maximum and the covalent F Kr F nature to the right. The avoided crossing of these two descriptions results in the potential maximum. It is clear of course that the absence of a potential maximum for XeF<sub>2</sub> does not mean that a shift from covalent to ionic character does not occur. Among other possibilities, the F Xe F covalent curve might just be significantly flatter (less repulsive) than that for F Kr F.

In our most extensive CI wave function (IV), the first-order wave function is augmented by all double excitations of the type  $10\sigma_{\rm g}^2$  + mo no. Both Figure I and Tables II and III show that this "first-order + order doubles" wave functions yields a potential curve qualitatively similar to the first-order curve. However, the well is noticeably deeper near the equilibrium geometry and the dissociation energy is 2.14 eV, or 77% of the experiment.

Like the first-order prediction, the equilibrium internuclear separation is  $\sim 0.02$  Å longer than experiment. Thus we conclude that the same methods (including electron correlation)  $^{12}$  which reliably predict the dissociation energies of conventional molecules are applicable to noble gas compounds. The only real difficulty lies in the fact that the bond energies of noble gas compounds are small and hence are usually exceeded in magnitude by the extramolecular correlation energy. Thus the role of electron correlation is much larger than is usually seen in conventional molecules. Of course in other molecules (e.g.  $F_2$ ) with small dissociation energies, the same situation will arise.

#### Electronic Structure Considerations

Of course one of the most fascinating aspects of noble gas compounds is the search for a simple yet correct model of the chemical bonding. In this sense the principal achievement of the earlier work  $^{21}$  on  $\mathrm{KrF}_2$  was to unequivocally establish the validity of Coulson's model  $^{24}$  of  $\mathrm{KrF}_2$ , depicted by (11). In addition to the maximum in the symmetric stretching potential curve, it was found  $^{21}$  that the electric field gradient changes very rapidly as a function of internuclear separation near the position of the  $\mathrm{KrF}_2$  potential maximum. The field gradient shifts from a value at larger r(Kr-F) separation appropriate to the Kr atom to one at smaller separations (including  $\mathrm{r}_{\mathrm{e}}$ ) appropriate to the  $\mathrm{Kr}^+$  ion.

In the present work on  $XeF_2$  a much simpler route has been followed. Mulliken populations have been obtained from the TCSCF wave functions and are summarized in Table IV. These show that between r(Xe-F) values of 4.0 and 5.0 bohrs, a switch from the covalent

does not occur as rapidly as in  $KrF_2$ , but is nevertheless a very real change. This change in electronic structure is also seen of course in the TCSCF potential curve in Figure 1. Thus the present  $\frac{ab\ initio}{ab\ initio}$  calculations give strong support to Coulson's model of the bonding in  $XeF_2$ . It is also interesting to note that the difference between the dissociation energies of  $XeF_2$  and  $KrF_2$ ,  $2.78-1.01=1.77\ eV$ , is very close to the difference between the ionization potentials  $^{26}$  of  $Xe\ and\ Kr\ (IP(Kr)=14.00\ eV,\ IP(Xe)=12.13\ eV$ , and  $\Delta IP=1.87\ eV$ ). In other words the increase in  $D_e\ for\ XeF_2\ correlates\ very\ closely\ with the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with\ the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with\ the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with\ the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with\ the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ with\ the decrease in IP for <math>XeF_2\ correlates\ very\ closely\ closely$ 

Orbital energies and a detailed Mulliken population analysis  $^{25}$  are given for r(Xe-F)=3.8 bohrs in Table V. There we see that most of the molecular orbitals are primarily distorted linear combinations of atomic orbitals. The small populations indicated for the  $6\sigma_u$  orbital are due to the fact that this orbital is occupied only in configuration (4). In the TCSCF model neither the  $10\sigma_g$  nor the  $6\sigma_u$  orbital energies should be interpreted as ionization

potentials in the sense of Koopmans' theorem. Note that the  $7\sigma_{\rm u}$  orbital is doubly occupied in both configurations (3) and (4). Incidentally, the coefficients at this geometry of the two configurations in the TCSCF wave function are 0.9835 and -0.1811. At larger separations of course, both coefficients approach  $1/\sqrt{2}$ .

A topic of considerable debate in the literature  $^{27-30}$  is the importance or unimportance of "higher" or "outer" orbitals in the bonding of XeF<sub>2</sub>. Specifically, the unconventional nature of noble gas compounds has led some researchers  $^{29,30}$  to suggest that 5d and 4f orbitals might in some sense be responsible for the existence of molecules such as XeF<sub>2</sub>. Note that our basis set (Table I) does include two functions of each of these types. The present Mulliken populations suggest that 20.262 "electrons" reside in d functions, while 0.037 electrons may be assigned to f functions. Since 20.00 electrons are assigned to d functions in the  $^{50}$ g,  $^{17}$ g,  $^{10}$ g,  $^{70}$ g,  $^{27}$ g, and  $^{20}$ g orbitals, only 0.262 can be identified with 5d orbitals. Thus it appears that the importance of 5d and 4f functions is of a quantitative nature as polarization functions.  $^{12}$  We find little evidence of a qualitative role for these outer orbitals in the bonding.

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TABLE I. Basis Set of Slater Functions,  $r^{n-1}e^{-\zeta r}$ , for Calculations of Xenon Fluorides

Atom	Type	Orbital ζ
Xe	1s	55.110
	1.s	36.545
•	2s	26.283
	2s	22.451
	3s	14.881
	3s	12.067
	4s	7.620
	4s -	5.566
	5s	3.518
	5s	2.173
	2p	30.678
	2p	21.424
٠.	3p	13.721
	3p	10.709
	4p	7.422
	4p	5.036
•	5p	3.516
	5p	2.016
	3d	20.469
	3d	11.964
	4d	7.727
	4d	5.233
	4d	3.379
	5d	2.0
	5d	1.2
	4f	3.5
	4f	2.5
	41	2.3
Atom	Type	Orbital ζ
F	1s	11.011
	1s	7.917
•	2s	3.096
	2s	1.946
	2p	6.165
	2p	3.176
	2p	1.612
	3 <b>d</b>	4.0
	3d	2.0
	4f	3.0
		<del></del> -

TABLE II. Total energies (in hartrees) of the  $\text{XeF}_2$  molecule for  $\text{D}_{\infty h}$  geometries. The five types of wave functions used are described in the text.

	ī	II	III	IV
R(Xe-F), bohrs	TCSCF	Valence CI	First-Order	First-Order + σ Doubles
3.4	-7430.9065	-7430.9094	-7430.9843	-7430.9907
3.6	-7430.9293	-7430.9329	-7431.0133	-7431.0197
3.8	-7430.9300	-7430.9342	-7431.0187	-7431.0249
4.0	-7430.9201	-7430.9244	-7431.0108	-7431.0167
4.5	-7430.9053	-7430.9059	-7430.9735	-7430.9764
5.0	-7430.9215	-7430.9216	-7430.9498	-7430.9504
6.0	-7430.9378	-7430.9378	-7430.9458	-7430.9458
8.0	-7430.9405	-7430.9405	-7430.9464	-7430.9464
10.0	-7430.9404	-7430.9404	-7430.9462	-7430.9462

TABLE III. Summary of structural and energetic predictions for  ${\tt XeF}_2$ . The different types of wave functions are described in the text.

Wave Function	I TCSCF	II Valence CI	III First-Order	IV First-Order + o Doubles	Experiment
Property					•
r <sub>e</sub> (Xe-F)					
bohrs	3.713	3.724	3.781	3.777	
<b>Å</b>	1.965	1.971	2.001	1.999	1.977 ± 0.002 <sup>a</sup>
Energy, hartrees	-7430.93100	-7430.93504	-7431.01875	-7431.2502	
Dissociation energy					
eV	-0.26	-0.15	1.97	2.14	2.78 <sup>b</sup>
kcal/mole	-5.9	-3.4	45.5	49.4	64 <sup>b</sup>

a Reference 7

b Reference 6

TABLE V. Orbital energies (in hartrees) and Mulliken populations for  $XeF_2$  at an Xe-F separation of 3.8 bohrs. These results were obtained from a two-configuration SCF wave function.

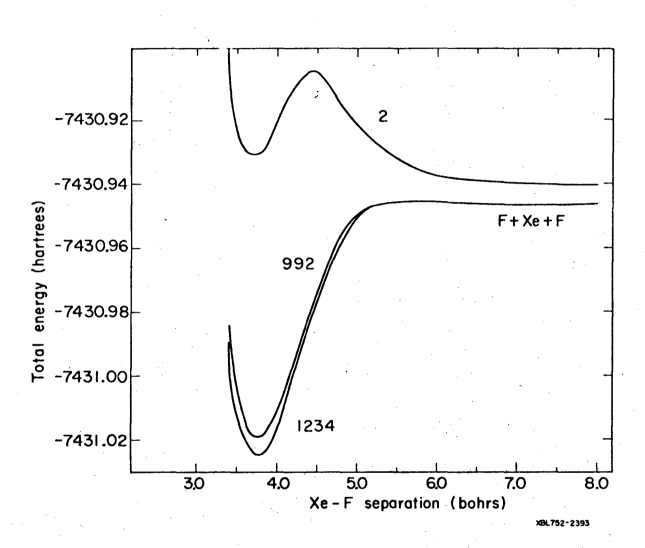
	SCF wave function.								
Orbital	Energy	s	P	Ke d	f		s	2F p	d
10 g	-1224.5139	2.00	-	-	-		-	-	<b>-</b> .
20 g	- 189.4574	2.00	- -	-			-	· <del>-</del>	- <del>-</del>
lσ <sub>u</sub>	- 177.8993	-	2.00	-	<u> </u>		-	_	-
$1\pi_{\mathbf{u}}$	- 177.8972	-	4.00	<del>-</del>	<u>.</u> .	•	· -	-	-
$3\sigma_{\mathbf{g}}$	- 40.2907	2.00	-		-	•	-	-	-
$2\sigma_{\mathbf{u}}$	- 35.3412	-	2.00	-	-	:	-	-	-
$2\pi_{\mathbf{u}}$	- 35.3333	· -	4.00	-	-			-	<b>-</b> .
4σ <sub>g</sub>	- 26.2759	-	-	-	-		2.00	-	. <del>-</del>
$3\sigma_{\mathbf{u}}$	- 26.2759	-	*	-			2.00	<b>-</b> .	-
5σ <sub>g</sub>	- 26.2402		-	2.00	-		-	<b>-</b> .	
$1\pi_{g}$	- 26.2370	-	-	4.00	<b>-</b>		-	_	-
$1\delta_{\mathbf{g}}$	- 26.2292	-	-	4.00	<b>-</b> .		-	-	-
<b>6</b> σ <sub>g</sub>	- 7.9698	2.00	-	-	-		-	-	· <b>-</b> .
4 o u	- 6.1302	-	2.00	-	- '			-	· <u>-</u> .
3 π <sub>u</sub>	- 6.1155	-	4.00	-	-		-	<b>-</b> .	-
7 σ <sub>g</sub>	- 2.9050	-	-	2.00			, <b>-</b> ,	_	
2 π g	- 2.8969	-	-	4.00	-		-	-	-
2 δ <sub>g</sub>	- 2.8785	_	-	4.00	-			-	-
5 o	- 1.5444	-	0.07	<b>-</b> ·	0.01		1.92	0.01	-
8 o	- 1.5270	0.13	-	-	-		1.86	-	<del>.</del>
6 o	- 1.0277	-	0.04	-	-		. –	0.03	
9 g	- 1.0197	1.69	_	0.01			0.08	0.20	0.01
7 o	- 0.7081		0.62	-	0.01		0.05	1.30	0.02
4π <sub>u</sub>	- 0.6629	-	0.79	-	0.02		-	3.18	0.01
3π <sub>g</sub>	- 0.6395	-	-	0.09	-		-	3.90	-
100g	- 0.5762	0.14	-	0.16			0.02	1.62	. <b>-</b>
5π <sub>u</sub>	- 0.4988	-	3.22	-	0.01		-	0.79	-0.01

TABLE IV. Mulliken populations for two-configuration SCF wave functions for xenon difluoride.

R(bohrs)	Xe	F		
3.4	52.77	9.62		
3.6	52.86	9.57		
3.8	52.98	9.51		
4.0	53.13	9.44		
4.5	53.71	9.15		
5.0	53.93	9.04		
6.0	53.99	9.01		
8.0	54.00	9.00		

# Figure Caption

Figure 1. Potential energy curves for the symmetric dissociation of XeF<sub>2</sub> to Xe + 2F. The labels 2, 992, and 1234 refer to the number of configurations included in the different wave functions under study. These wave functions are described in the text.



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