brought to you by 🗓 CORE

Low-lying electronic states and molecular structure of Fe₂O₂

Zexing Cao,^a Miquel Duran^b and Miquel Solà^{*b}

^a Department of Chemistry and Institute of Physical Chemistry, Xiamen University, Xiamen 361005, People's Republic of China

^b Institute of Computational Chemistry and Department of Chemistry, Universitat de Girona, 17071 Girona, Catalonia, Spain. E-mail: miquel@iqc.udg.es

Received 18th May 1998, Accepted 3rd August 1998

The structure, bonding and relative stabilities of the ground and low-lying excited states of Fe_2O_2 have been studied by the hybrid B3LYP density-functional and coupled-cluster molecular orbital methods. Calculations indicate that the $(\mu$ -O)₂ rhombic ⁷B_{2u} state is the ground state for Fe_2O_2 . Stable molecular diiron oxo Fe_2 -O₂ complexes in distorted tetrahedral and planar side-on modes have been also located on the potential-energy hypersurfaces of Fe_2O_2 . The calculated IR-active frequencies corresponding to two in-plane deformations of the rhombic ring agree well with the observed values. The bonding features of the $(\mu$ -O)₂ rhombic Fe_2O_2 have been discussed based on natural bond orbital and Bader topological analyses. These analyses show that an effective Fe—Fe bonding across the ring exists in the ⁷B_{2u} ground state.

Introduction

The interaction of iron with molecular oxygen has achieved a considerable interest in current experimental and theoretical studies because it is involved in a wide range of processes, from corrosion of materials to oxygen transportation in biological systems. Many studies have been devoted to monoiron oxides. Among them, the diatomic FeO molecule has been the most thoroughly investigated.^{1–9} Experimental and theoretical investigations of the triatomic FeO₂, FeO₂⁻ and FeO₂⁺ have been also carried out.^{1,10–18} The higher monoiron oxides FeO₄ and FeO₄⁻ have been observed experimentally,^{18,19} and their structures and molecular bonding have been studied theoretically.²⁰ However, to date, only a few studies have been performed on the structure and bonding of diiron oxides.

Diiron oxo complexes are a common structural motif in a class of metalloproteins that are important as oxygen carriers.^{21,22} In a recent study,²³ it has been demonstrated that a high-valent Fe_2O_2 diamond core can constitute the common structural unit for high-valent intermediates in the oxygen activation mechanisms of non-heme iron enzymes.²⁴ A highvalent non-heme diiron intermediate and the first bis(µ-oxo) diiron(III) complex have been recently synthesized.^{2,26} Further, extended X-ray absorption fine structure (EXAFS) and Mössbauer spectroscopy studies have provided direct experimental evidence for the participation of a $(\mu$ -O)₂ rhombic Fe₂O₂ diamond core (structure I in Fig. 1) in the catalytic cycle of methane monoxygenase.²⁷ On the other hand, a recent anion photoelectron spectroscopy study has suggested the existence of $(\mu$ -O)₂ rhombic Fe₂O₂ and possible isomers containing an unbroken O–O bond.¹⁹ IR spectra, complemented with density functional calculations of several electronic states of different iron oxide product molecules in the reactions of laser-ablated iron atoms with molecular oxygen, have indicated that the $(\mu$ -O)₂ rhombic Fe₂O₂ absorbs at 517.4 and 660.6 cm⁻¹.¹⁸ Finally, Fe_2O_2 species have also been produced through reaction of free iron clusters with molecular oxygen.²⁸

Despite all these important contributions, the question of how many Fe_2O_2 isomers exist and which are the most stable

remains still open. In this work, we have carried out a theoretical investigation on Fe_2O_2 with the B3LYP hybrid DFT functional and *ab initio* molecular methods. Eight possible structures have been analyzed in our calculations (Fig. 1).

Computational details

Density functional theory (DFT) calculations have been performed with the Becke's three-parameters hybrid exchange functional²⁹ and the non-local correlation functional of Lee, Yang and Parr³⁰ (B3LYP) to obtain the optimized geometries, energies, vibrational frequencies and corresponding intensities



Fig. 1 The eight possible structures for $\mathrm{Fe_2O_2}$ considered in this study.

of the ground and low-lying excited states of Fe_2O_2 . The computational efficiency and accuracy of the B3LYP method have been very well documented for transition metal complexes.^{31–34} In particular, previous studies on FeO_2 ^{10.18} showed that the hybrid B3LYP approach works reasonably well. Single and double substitutions with non-iterative triple excitations coupled cluster [CCSD(T)]^{35,36} energy calculations for several selected states have been carried out at the optimized B3LYP geometries (CCSD(T)//B3LYP) to confirm the validity of this approach.

The initial guesses of electron configurations for the lowlying states of Fe_2O_2 are obtained by altering molecular orbitals from converged SCF calculations at different geometries. All calculations have been done within the unrestricted methodology, except for singlets, which have been calculated within the restricted formalism. The basis set used for all atoms has been the 6-311G*,^{37,38} which, for iron, is the allelectron basis set of Wachters and Hay^{39,40} with the scaling factors of Raghavachari and Trucks.⁴¹ The polarization functions for iron and oxygen atoms are an *f* function with an exponent of 1.05 and a *d* function with an exponent of 1.292, respectively. All calculations have been performed with Gaussian 94,⁴² except for Bader topological analyses that were carried out by means of the Electra program.⁴³

Results and discussion

The electron configurations, optimized geometries, harmonic vibrational frequencies and relative energies for some selected low-lying states of the $(\mu$ -O)₂ rhombic, distorted tetrahedral, and side-on Fe₂O₂ are presented in Table 1. Only the most stable states obtained for each multiplicity are included. Information about other states can be found in the supplementary material[†]

† Supplementary material (SUP 57424, 9 pp.) deposited with the British Library. Details are available from the Editorial Office.

Energetics

The results in Table 1 show that the ground state of Fe_2O_2 is the ${}^{7}B_{2u}$ state of the (µ-O)₂ rhombic structure D_{2h} I. At the CCSD(T)//B3LYP level, the ${}^{5}A_{g}$, ${}^{3}B_{1u}$ and the ${}^{1}A_{g}$ states of the rhombic structure are higher in energy than the lowest ${}^{7}B_{2u}$ state by 117, 150 and 160 kJ mol⁻¹, respectively. The energy ordering predicted by the B3LYP method is in agreement with the CCSD(T)//B3LYP calculation for these states, although the larger energy differences between the low-spin and high-spin states indicate that the B3LYP method tends to overestimate the stability of the latter. The higher stability of the state with high spin was also found for Fe₂⁴⁴ and for rhombic (CoO)₂.⁴⁵ All attempts to find a distorted (μ -O)₂ rhombic state ³A₂, which was the most stable among the several states considered in a previous study,45 have failed. It is noteworthy that the same authors⁴⁵ indicated that this state was probably an artifact of their calculation. Furthermore, the single-point calculation of the ${}^{3}A_{2}$ state at the geometry reported⁴⁵ has been found to be higher in energy than other triplet states of the $(\mu$ -O)₂ rhombic structure. In addition to the most stable $(\mu$ -O)₂ rhombic isomer, much less stable states corresponding to C_{2v} II and C_{2v} VII structures were located. These can be considered as molecular diiron oxo Fe₂-O₂ complexes and they might account for the low electron affinity isomers detected in a previous photoelectron spectroscopy experiment.19

Very few stable states with the other possible structures depicted in Fig. 1 were found in the present study. The singlet state ${}^{1}A_{1}$ in C_{2v} IV structure has a large Fe—Fe separation of 3.107 Å and can be considered a van der Waals complex between Fe and FeO₂. Similar behaviour has been found for the states of higher spin in the C_{2v} IV structure. This implies that the interaction of the d orbitals of Fe and the bonding $1\pi_{u}$ and antibonding $1\pi_{g}$ of O₂ is strong and breaks the Fe—Fe bond. No stable complexes with C_{2v} V structure have been found. During the optimization of the states of lower spin for the (μ - η^{2} : η^{2} -O₂) D_{2h} VI isomer the O—O bond is

Table 1 Electronic configurations, geometries, energies and vibrational frequencies obtained with the B3LYP method for some selected low-lying states of Fe_2O_2

State	Configuration ^a	Geometry ^b	ΔE^{c}	ZPE^{d}	Frequencies ^e
D _{2h} I					
¹ A.	$4a^{2}2b_{2}^{2}$ $3b_{1}^{2}$ b_{2}^{2} b_{2}^{2} b_{1}^{2} $a^{2}b_{2}^{2}$	2.476, 1.714, 2.372	423.9(159.9)	21.8	285(74), 379(0), 629(2), 725(126), 801(0), 758(0)
${}^{3}B_{1}^{g}$	$4a_{a}^{2}2b_{2u}^{2}3b_{1u}^{2}b_{3a}^{2}b_{1a}^{2}b_{3u}a_{u}^{2}b_{2a}b_{3u}$	2.364, 1.731, 2.530	231.4(149.8)	19.5	279(72), 353(0), 523(9), 617(0), 692(223), 804(0)
${}^{5}A_{a}^{1u}$	$4a_{a}^{5}2b_{2u}^{2}3b_{1u}^{2}b_{3a}^{2}b_{1a}^{2}b_{3u}a_{u}b_{2a}b_{1a}$	2.447, 1.770, 2.558	79.5(116.9)	18.4	180(94), 321(0), 563(0), 574(218), 686(142), 750(0)
${}^7\mathrm{B_{2u}^{s}}$	$4a_{g}^{2}2b_{2u}^{2}2b_{1u}^{2}b_{3u}^{2}b_{3u}^{2}b_{3g}^{2}b_{1g}^{2}a_{u}b_{2g}b_{3g}^{1}$ $b_{1u}b_{1g}b_{3u}$	2.203, 1.790, 2.821	0.0(0.0)	19.2	203(94), 401(0), 484(0), 657(87), 737(0), 738(109)
C_{2v} II	C C				
$^{1}A_{1}$	$6a_1^2 3b_2^2 3b_1^2 2a_2^2$	1.885, 1.466, 76.7	702.6	17.7	245(0), 340(0), 437(16), 503(9), 568(32), 874(43)
${}^{5}A_{1}$	$6a_1^2 3b_2^2 2b_1^2 a_2^2 2b_1 2a_2$	1.551, 2.377, 73.6	564.1	16.2	220(8), 338(257), 395(0), 489(33), 500(0), 767(30)
C_{2v} VII		, ,			
¹ A.	$6a_{2}^{2}3b_{2}^{2}3b_{2}^{2}2a_{2}^{2}$	1.913, 1.437, 82.2	751.1	18.5	230(0), 297(20), 423(7), 640(2), 686(22), 816(105)
${}^{3}B_{1}$	$6a_1^2 3b_2^2 3b_1^2 a_2^2 a_3 b_3$	1.942, 1.424, 81.5	665.3	20.3	345(0), 375(6), 492(0), 656(2), 728(4), 804(158)
${}^{5}A_{2}^{1}$	$6a_1^2 2b_2^2 3b_1^2 a_2^2 a_3^2 2b_3 a_1$	1.847, 1.303, 82.2	585.3	17.5	209(16), 254(0), 324(9), 427(12), 502(3), 1213(131)
${}^{7}A_{1}^{2}$	$6a_{1}^{2}2b_{2}^{2}2b_{1}^{2}a_{2}^{2}a_{2}^{2}3b_{2}^{1}b_{1}^{1}a_{1}$	2.011, 1.317, 80.5	448.9	15.9	162(6), 181(0), 329(26), 373(9), 418(0), 1194(90)
Fe ₂					
⁵ Δ.,	$2\sigma_a^2\pi_a^4\delta_a^3\delta_a^2\pi_a^2\sigma_a^{B}$	1.876	75.4	2.9	483(0)
$^{7}\Delta_{u}^{u}$	$2\sigma_{a}^{2}\pi_{u}^{4}\delta_{a}^{5}\delta_{u}^{2}\pi_{a}^{2}\sigma_{u}^{1}$	1.906	0.0	2.8	469(0)
FeO	5 4 5 4 5 4				
$^{3}\Sigma$	$2\sigma^2\pi^4\delta^2\pi^2$	1.614	207.1	5.7	951(70)
⁵ Δ	$\sigma^2 \pi^4 \delta^3 \pi^2 \sigma^1$	1.606	0.0	5.4	906(97)
0,					
- 1Λ	$2\sigma^2\sigma^2\pi^4\pi^2$	1.207	163.3	9.7	1626(0)
${}^{3}\overline{\Sigma}_{g}^{g}$	$2\sigma_{g}^{2}\sigma_{u}^{2}\pi_{u}^{4}\pi_{g}^{2}$	1.206	0.0	9.8	1641(0)

^{*a*} Core electrons are omitted, and numbers in front of molecular orbitals indicate the number of orbitals with the same symmetry in the wavefunction. ^{*b*} The bond lengths in Å, and bond angles in degrees. $D_{2h}I$: Fe–Fe, Fe–O, O–O, respectively; $C_{2v}II$: Fe–O, O–O, Fe–O–Fe, respectively; $C_{2v}VII$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}VII$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O–O, Fe–O–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}II$: Fe–Fe, O–O, O–Fe–Fe, respectively; $C_{2v}I$

broken, thus yielding the $(\mu$ -O)₂ rhombic D_{2h} I species. The low stability of Fe₂O₂ complexes with the $(\mu$ - η^2 : η^2 -O₂) rhombic structure VI is in accordance with the fact that no Fe₂O₂ complex with this structure has been synthesized yet.⁴⁶ The states of high spin for the linear $D_{\infty h}$ III structure are not stable. The same has been found for the linear Fe–Fe– -O–O (structure VIII in Fig. 1), and the states of high spin in $C_{\infty v}$ symmetry dissociate to Fe + FeOO. Calculated results for these less stable isomers and higher-order saddle points are not presented here.

Note that the results obtained for the diiron oxo $Fe_2 - O_2$ complexes are different from those reported for the Fe_2-N_2 species.⁴⁷ Among the Fe₂-N₂ species, the linear Fe-Fe-N-N septet state was found to be the most stable, and stable states of lower spin in linear symmetry were also located. The difference in the most stable molecular structure between $Fe_2 - O_2$ and $Fe_2 - N_2$ can be attributed to the different highest occupied molecular orbitals (HOMO) of N2 and O_2 . The HOMO of N_2 is a $3\sigma_g$ orbital, while the HOMOs of O_2 are singly occupied (for the triplet) and doubly occupied (for the singlet) $1\pi_{g}$ orbitals. The linear end-on mode is thus favoured in $Fe_2 - N_2$, through interaction between the lowest unoccupied molecular orbital (LUMO) $7\sigma_u$ and singly occupied HOMO $6\sigma_u$ of Fe₂ in its ground state and the $3\sigma_g$ of N_2 . For O_2 the same $3\sigma_g$ orbital is much lower in energy and, therefore, less suitable for the linear interaction than the $3\sigma_{\alpha}$ of N_2 . The Fe_2-O_2 complexes prefer the side-on and tetrahedral modes through the interaction between singly occupied lower in energy $1\delta_g$ and $3\pi_g$ orbitals of Fe₂ and the HOMO $1\pi_{\sigma}$ of O₂. Comparison between the results obtained for $Fe_2 - O_2$ and $Fe_2 - N_2$ species shows that the interaction of Fe_2 with O_2 is stronger than that of Fe_2 and N_2 .

Table 2 lists the dissociation energies computed at the B3LYP level corresponding to different dissociation products for some selected states of the rhombic, tetrahedral and side-on structures. The states of the rhombic isomer in Table 2 are lower in energy than the products of the six dissociation channels considered here, except for the ¹A_g state relative to the ground state of FeO. Dissociation of the $(\mu$ -O)₂ rhombic $^{7}B_{2u}$ state to two $^{5}\Delta$ FeO is endothermic by 364 kJ mol⁻¹. This value can be compared with the 301 kJ mol⁻¹ for the dissociation of the ⁷A_u (CoO)₂ system to 2CoO molecules.⁴⁵ The tetrahedral and side-on diiron oxo Fe_2-O_2 complexes are more stable than the products $Fe_2 + O_2$, except for the singlet states relative to the ground states $Fe_2(^7\Delta_u) + O_2(^3\Sigma_g^-)$. Considering the spin multiplicity of the initial and final states, the dissociation of the singlet states of $Fe_2 - O_2$ into the septet Fe_2 and the triplet O_2 is not likely, and the singlet molecular



Fig. 2 The three IR-active vibrational modes for rhombic $(\mu\text{-}O)_2$ Fe_2O_2 .

diiron oxo $Fe_2 - O_2$ complexes should be correlated to certain excited states of Fe_2 with lower spin.

Vibrational frequencies

For the $(\mu$ -O)₂ rhombic isomer, there are only three IR-active vibrational modes corresponding to two deformations in plane and one out of plane, which are depicted in Fig. 2 The 517.4 and 660.6 cm⁻¹ frequencies with 16/18 isotopic ratios of 1.0457 and 1.0454, respectively, observed in reactions of laserablated iron atoms with oxygen molecules have been assigned to Fe_2O_2 species.^{10,18,45} The B3LYP calculated frequencies in Table 1 agree with these observed values for certain stable states of the $(\mu$ -O)₂ rhombic structure. For the ³B_{1µ} state, the strong band at 692(223) cm⁻¹ is also in reasonable agreement with the observed 660.6 cm^{-1} . Further, its calculated harmonic 16/18 isotopic ratio of 1.0453 agrees with the observed 1.0454. The lower ${}^{5}A_{g}$ quintet state has strong absorptions at 574(218) and 686(142) cm⁻¹, which are only slightly higher than the observed values. The ${}^{5}A_{g}$ state has calculated 16/18 isotopic ratios of 1.0475 for the 574(218) cm⁻¹ and 1.0457 for the 686(142) cm⁻¹, to be compared to the observed 1.0457 and 1.0454, respectively. Finally, the ground-state ${}^{7}B_{2u}$ has a band at 657(87) cm⁻¹ and a corresponding 16/18 isotopic ratio of 1.0462. Unfortunately, states of different symmetry and spin multiplicity can account for the observed values and, therefore, a definite assignment is not possible. However, it is worth noting that all vibrational modes with frequencies close to the observed bands at 517.4 and 660.6 cm^{-1} are in plane b_{1u} and b_{2u} deformations of the rhombic ring.

Bonding features

We have carried out a natural population and natural bond orbital (NBO) analysis in order to get a further insight into the bonding features of the $(\mu$ -O)₂ rhombic Fe₂O₂. Tables 3 and 4 present the results of this natural population and NBO analysis. Note that the natural charges, as well as the Mulliken charges in Table 3, increase as the spin multiplicity

 Table 2
 Dissociation energies^a corresponding to different dissociation limits for the lower stable states in the rhombic, tetrahedral and side-on isomers computed at the B3LYP level

State	Dissociation limits						
	$2 \text{FeO}(^5\Delta)$	$2 \text{FeO}(^{3}\Sigma)$	$\mathrm{Fe}_2(^7\Delta_u) + \mathrm{O}_2(^3\Sigma_g^-)$	$\mathrm{Fe}_2(^7\Delta_u) + \mathrm{O}_2(^1\Delta_g)$	$\mathrm{Fe}_2({}^5\Delta_u) + \mathrm{O}_2({}^3\Sigma_g^-)$	$\mathrm{Fe}_2({}^5\Delta_u) + \mathrm{O}_2({}^1\Delta_g)$	
D_{2h} I							
${}^{1}A_{a}$	-62.8	351.9	258.1	1226.7	333.5	496.6	
${}^{3}B_{1}$	136.4	546.4	452.7	615.9	528.4	691.6	
⁵ A _g	284.9	699.6	606.3	769.4	681.2	844.3	
${}^{7}B_{2u}^{*}$	363.6	777.8	684.1	847.3	759.4	922.6	
C_{2v} II							
${}^{1}A_{1}$	-337.2	77.4	-16.7	146.4	59.0	222.2	
${}^{5}A_{1}^{1}$	-197.1	217.1	123.4	286.6	198.7	361.9	
C_{2v} VII							
${}^{1}A_{1}$	-386.2	28.0	-65.7	97.5	9.6	172.8	
${}^{3}B_{1}$	-302.5	112.1	18.0	181.2	93.7	256.9	
${}^{5}A_{2}$	-219.7	194.6	100.8	264.0	176.6	339.7	
${}^{7}A_{1}^{2}$	-81.6	332.6	209.6	372.8	314.2	477.4	

^{*a*} The dissociation energies are defined by the difference: E (dissociation limits) -E (Fe₂O₂), in kJ mol⁻¹. Zero-point vibrational energies have been included in the calculation.

Table 3 Natural population analysis of the lower states of the rhombic Fe_2O_2 at the B3LYP level, showing natural charge q (with corrsponding Mulliken charges in parentheses) and effective valence-shell natural electron configuration (NEC)

State	Atom	q^a	NEC
¹ A _g ³ B _{1u} ⁵ A _g ⁷ B _{2u}	Fe O Fe O Fe O Fe O	$\begin{array}{c} 0.729(0.690) \\ -0.729(-0.690) \\ 0.811(0.745) \\ -0.811(-0.745) \\ 0.919(0.825) \\ -0.919(-0.825) \\ 0.972(0.860) \\ -0.972(-0.860) \end{array}$	$\begin{array}{c} 4s^{0.68}3d^{6.59}\\ 2s^{1.90}2p^{4.82}\\ 4s^{0.67}3d^{6.51}\\ 2s^{1.91}2p^{4.89}\\ 4s^{0.66}3d^{6.41}\\ 2s^{1.92}2p^{4.99}\\ 4s^{0.61}3d^{6.40}\\ 2s^{1.90}2p^{4.82} \end{array}$

^a In atomic units (electron charge).

increases, showing that more electronic charge is transferred from Fe to O, and implying that the bonding interaction becomes stronger with the increase in spin multiplicity. This is consistent with the relative stabilities predicted by the B3LYP and CCDS(T)//B3LYP methods. Results of the Bader analysis⁴⁸ collected in Table 5, indicate that the Fe—O bond has ionic character, as can be deduced from the small density, positive laplacian, and small ratio between the perpendicular and the parallel curvatures at the bond critical point ($|\lambda_1| \lambda_3$).

It is interesting to discuss whether an Fe–Fe bond exists in the $(\mu$ -O)₂ rhombic Fe₂O₂. From the NBO results in Table 4 one can see that an effective Fe–Fe antibonding interaction exists for the ¹A_g and ³B_{1u} states, while an effective bonding for the Fe–Fe bond is present in the lowest ⁷B_{2u} state. Finally, no Fe–Fe bonding across the ring is observed for the

Table 4 Principal NBOs for the lower states of the $(\mu$ -O)₂ rhombic Fe₂O₂ species calculated at the B3LYP level, with orbital energies, occupancies and compositions $(c_Ah_A + c_Bh_B)$ in terms of hybrids (h) on A and B with c_A and c_B being the corresponding expansion coefficients (A,B = Fe,O)

			Composition			
NBO ^a						
A—B	Energy [®]	Occupancy	c _A	h _A	$c_{\mathbf{B}}$	h _B
¹ A _a						
Fe-Fe*(2)	-0.1518	1.9999	0.7071	d	-0.7071	d
Fe-O(4)	-0.5004	1.9862	0.4463	sd ^{1.5}	0.8949	sp ^{7.9}
Fe-Fe	-0.1384	0.4279	0.7071	d	0.7071	d
Fe-Fe	-0.1289	0.7624	0.7071	d	0.7071	d
³ B ₁						
Alpha electro	ons					
Fe-Fe*	-0.1982	0.9999	0.7071	d	-0.7071	d
Fe-O(4)	-0.4763	0.9907	0.4686	sd1.33	0.8834	sp ^{11.39}
Fe-Fe	-0.1635	0.2879	0.7071	d	0.7071	d
Beta electron	18					
Fe-Fe*	-0.1687	0.9998	0.7071	d	-0.7071	d
Fe-O(4)	-0.4918	0.9935	0.4423	sd ^{1.62}	0.8969	sp ^{9.15}
Fe-Fe	-0.1330	0.2523	0.7071	d	0.7071	d
⁵ A _e						
Alpha electro	ons					
Fe-O(4)	-0.4236	0.9863	0.5117	sd ^{1.25}	0.8591	sp ^{17.4}
Beta electron	15					
Fe-O(4)	-0.4539	0.9946	0.4449	sd ^{1.5}	0.8956	sp ⁹
Fe-O(2)	-0.2602	0.9317	0.2703	d	0.9628	р
${}^{7}B_{2u}$						
Alpha electro	ons					
Fe-Fe*	-0.2708	0.9972	0.7071	d	-0.7071	d
Fe-Fe	-0.2749	0.5943	0.7071	d	0.7071	d
0-0	-0.3444	0.8512	0.7071	sp ^{38.77}	0.7071	sp ^{38.77}
0-0*	-0.2432	0.4102	0.7071	sp ^{38.77}	-0.7071	sp ^{38.77}
Beta electrons						
Fe-Fe	-0.2081	0.8820	0.7071	sd ^{3.04}	0.7071	sd ^{3.04}
Fe-O(4)	-0.4417	0.9895	0.4298	sd ^{2.49}	0.9029	sp ^{11.28}
Fe-O(2)	-0.2575	0.9269	0.2805	d	0.9599	р
4 Tho num	have in m	anonthagan in	adiaata	the mum	han of ag	

^a The numbers in parentheses indicate the number of equivalent NBOs. ^b In atomic units.

Table 5 Bader analysis for the lower states of the $(\mu$ -O)₂ rhombic Fe_2O_2 species computed at the B3LYP level

State	Critical point	r_{c}^{a}	$ ho(r_{\rm c})^b$	$\nabla^2 \rho(r_{\rm c})^b$	$ \lambda_1 /\lambda_3$
¹ A ₂	(3, 1)	1.238	0.061	0.272	0.385
g	Fe - O(3, -1)	0.850	0.180	0.849	0.187
³ B ₁	(3, 1)	1.182	0.069	0.241	0.329
Iu	Fe - O(3, -1)	0.868	0.179	0.739	0.236
⁵ A.	(3, 1)	1.223	0.060	0.205	0.351
g	Fe - O(3, -1)	0.891	0.166	0.592	0.267
${}^{7}B_{2}$	Fe-Fe(3, -1)	1.102	0.097	0.187	0.343
- 2u	Fe = O(3, -1)	0.912	0.162	0.572	0.258
					. ? h.

^{*a*} Distance from the critical point to the closest Fe atom in Å. ^{*b*} In atomic units (e a_0^{-3} and e a_0^{-5}).

 ${}^{5}A_{g}$ state. This is in agreement with the results of Table 5, which show that a bond critical point between Fe—Fe exists only for the most stable ${}^{7}B_{2u}$ state. Surprisingly, there are NBOs for the alpha electrons between the two O atoms in the lowest septet state, despite the large O—O separation of 2.821 Å and the fact that no bond critical point has been found connecting the two oxygen atoms. This suggests a strong electron delocalization in this state.

Conclusions

We have investigated the low-lying states of Fe₂O₂ with the B3LYP and CCSD(T) methodologies. The molecular structure and harmonic vibrational frequencies for these states have been reported. The ${}^{7}B_{2u}$ (µ-O)₂ rhombic state has been found to be the ground state for Fe_2O_2 . The existence of an effective Fe-Fe bonding for this state is confirmed by the NBO and Bader analyses. At the CCSD(T) level, the ${}^{5}A_{g}$, ${}^{3}B_{1u}$ and ${}^{1}A_{g}$ states are less stable than the ${}^{7}B_{2u}$ state by 117, 150 and 160 kJ mol⁻¹, respectively. Interestingly, the energy ordering for the lower states of the $(\mu$ -O)₂ rhombic Fe₂-O₂ structure given by the B3LYP approach is in agreement with the CCSD(T) calculations, although the energy differences between low-spin and high-spin states are overestimated by the B3LYP approach. The observed IR bands at 660.6 and 514.7 cm⁻¹ have been assigned to two in-plane deformation modes of the rhombic ring. Finally, the distorted tetrahedral and planar side-on modes are the most stable structures for the diiron oxo $Fe_2 - O_2$ complexes.

This work has been supported by the Spanish DGICYT Project No. PB95-0762. ZC gratefully acknowledges the Catalan Comissionat per a Universitats i Recerca for a fellow-ship to work in Girona. This work is also part of the China NSF Project No. 29773036.

References

- 1 J. Fan and L. S. Wang, J. Chem. Phys., 1995, 102, 8714.
- 2 A. J. Merer, Annu. Rev. Phys. Chem., 1989, 40, 407.
- 3 M. Dolg, U. Wedig, H. Stoll and H. Preuss, J. Chem. Phys., 1987, 86, 2123.
- 4 T. C. Steimle, D. F. Nachman and J. E. Shirley, J. Chem. Phys., 1989, 90, 5360.
- 5 T. Andersen, K. R. Lykke, D. M. Neumark and W. C. Lineberger, J. Chem. Phys., 1987, 6, 1858.
- 6 T. Kroeckertskothen, H. Knockel and E. Tiemann, Chem. Phys., 1986, 103, 335; Mol. Phys., 1987, 62, 1031.
- 7 A. W. Taylor, A. S.-C. Cheung and A. J. Merer, J. Mol. Spectrosc., 1985, 113, 487.
- 8 D. W. Green and G. T. Reedy, J. Mol. Spectrosc., 1979, 78, 257 and references therein.
- 9 P. C. Engelking and W. C. Lineberger, J. Chem. Phys., 1977, 62, 2566.
- 10 L. Andrews, G. V. Chertihin, A. Ricca and C. W. Bauschlicher, J. Am. Chem. Soc., 1996, 118, 467.
- 11 Z. Cao, M. Duran and M. Solà, Chem. Phys. Lett., 1997, 274, 411.

- 12 D. Schröder, A. Fiedler, J. Schwarz and H. Schwarz, *Inorg. Chem.*, 1994, 33, 5094.
- 13 M. Fanfarillo, H. E. Cribb, A. J. Downs, T. M. Greene and M. J. Almond, *Inorg. Chem.*, 1992, **31**, 2962.
- 14 P. D. Lyne, D. M. P. Mingos, T. Ziegler and A. J. Downs, *Inorg. Chem.*, 1993, **32**, 4785.
- 15 S. A. Mitchell and P. A. Hackett, J. Chem. Phys., 1990, 93, 7822.
- 16 S. Abramowitz, N. Acquista and I. W. Levin, Chem. Phys. Lett., 1977, 50, 423.
- (a) S. Chang, G. Blyholder and J. Fernandez, *Inorg. Chem.*, 1981,
 20, 2813; (b) G. Blyholder, J. Head and F. Ruette, *Inorg. Chem.*,
 1982, 21, 1539; (c) M. Helmer and J. M. C. Plane, *J. Chem. Soc.*, *Faraday Trans.*, 1994, 90, 395.
- 18 G. V. Chertihin, W. Saffel, J. T. Yustein, L. Andrews, M. Neurock, A. Ricca and C. W. Bauschlicher, J. Phys. Chem., 1996, 100, 5261.
- 19 H. Wu, S. R. Desai and L. S. Wang, J. Am. Chem. Soc., 1996, 118, 5296.
- 20 Z. Cao, W. Wu and Q. Zhang, submitted for publication.
- 21 B. P. Murch, P. D. Boyle and L. Que, Jr., J. Am. Chem. Soc., 1985, 107, 6728.
- 22 A. C. Rosenzweig and S. Lippard, Acc. Chem. Res., 1994, 27, 229.
- 23 C. Kim, Y. Dong and L. Que, Jr., J. Am. Chem. Soc., 1997, 119, 3635.
- 24 L. Que, Jr. and Y. Dong, Acc. Chem. Res., 1996, 29, 190.
- 25 Y. Dong, H. Fujii, M. P. Hendrich, R. A. Leising, G. Pan, C. R. Randall, E. C. Wilkinson, Y. Zang, L. Que, Jr., B. G. Fox, K. Kauffmann and E. Münck, J. Am. Chem. Soc., 1995, 117, 2778.
- 26 (a) Y. Zang, Y. Dong, L. Que, Jr., K. Kauffmann and E. Münch, J. Am. Chem. Soc., 1995, 117, 1169; (b) E. C. Wilkinson, Y. Dong, Y. Zang, H. Fujii, R. Fraczkiewicz, G. Fraczkiewicz, R. S. Czernuszewicz and L. Que, Jr., J. Am. Chem. Soc., 1998, 120, 955.
- 27 L. Shu, J. C. Nesheim, K. Kauffmann, E. Münck, J. Lipscomb and L. Que, Jr., *Science*, 1997, **275**, 515.
- 28 R. L. Whetten, D. M. Cox, D. J. Trevor and A. Kaldor, J. Phys. Chem., 1985, 89, 566.
- 29 (a) A. D. Becke, J. Chem. Phys., 1993, 98, 5648; (b) J. Chem. Phys., 1992, 97, 9173; (c) J. Chem. Phys., 1992, 96, 2155.
- 30 C. Lee, W. Yang and R. G. Parr, Phys. Rev. B, 1988, 37, 785.

- 31 C. W. Bauschlicher Jr., Chem. Phys. Lett., 1995, 246, 40. View Online
- 32 T. Wagener and G. Frenking, Inorg. Chem., 1998, 37, 1805.
- 33 F. A. Cotton and X. Feng, J. Am. Chem. Soc., 1997, 119, 7514; 1998, 120, 3387.
- 34 B. S. Jursic, Int. J. Quantum Chem., 1997, 64, 255.
- 35 R. J. Bartlett and G. D. Purvis, Int. J. Quantum Chem., 1978, 14, 516.
- 36 G. E. Scuseria and H. F. Schaefer III, J. Chem. Phys., 1989, 90, 3700.
- 37 A. D. McLean and G. S. Chandler, J. Chem. Phys., 1980, 72, 5639.
- 38 R. Krishnan, J. S. Binkley, R. Seeger and J. A. Pople, J. Chem. Phys., 1980, 72, 650.
- 39 A. J. H. Wachters, J. Chem. Phys., 1970, 52, 1033.
- 40 P. J. Hay, J. Chem. Phys., 1977, 66, 4377.
- 41 K. Raghavachari and G. W. Trucks, J. Chem. Phys., 1989, 91, 1062.
- 42 M. J. Frisch, G. W. Trucks, H. B. Schlegel, P. M. W. Gill, B. G. Johnson, M. A. Robb, J. R. Cheesman, T. Keith, G. A. Petersson, J. A. Montgomery, K. Raghavachari, M. A. Al-Laham, V. G. Zakrzewski, J. V. Ortiz, J. B. Foresman, C. Y. Peng, P. Y. Ayala, W. Chen, M. W. Wong, J. L. Andres, E. S. Replogle, R. Gomerts, R. L. Martin, D. J. Fox, J. S. Binkley, D. J. Defrees, J. Baker, J. J. P. Stewart, M. Head-Gordon, C. Gonzalez and J. A. Pople, Gaussian 94 Revision E.2, Gaussian, Inc., Pittsburgh PA, 1995.
- 43 J. Mestres, Electra, Girona, CAT, 1994.
- 44 T. Noro, C. Ballard, M. H. Palmer and H. Tatewaki, J. Chem. Phys., 1994, 100, 452.
- 45 G. V. Chertihin, A. Citra, L. Andrews and C. W. Bauschlicher, J. Phys. Chem., 1997, 101, 8793.
- 46 K. Yoshizawa, T. Ohta, T. Yamabe and R. Hoffmann, J. Am. Chem. Soc., 1997, **119**, 12311.
- 47 A. Zacarias and M. Castro, Int. J. Quantum Chem.: Quantum Chem. Symp., 1996, 30, 207.
- 48 R. F. W. Bader, Atoms in Molecules: A Quantum Theory, Clarendon, Oxford, 1990.

Paper 8/03725F