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# Fe-V sulfur clusters studied through photoelectron spectroscopy and density functional theory†

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Iron-vanadium sulfur cluster anions are studied by photoelectron spectroscopy (PES) at 3.492 eV (355 nm) and 4.661 eV (266 nm) photon energies, and by density functional theory (DFT) calculations. The structural properties, relative energies of different structural isomers, and the calculated first vertical detachment energies (VDEs) of different structural isomers for cluster anions  $FeVS_{1-3}^-$  and  $Fe_mV_nS_{m+n}^-$ (m + n = 3, 4; m > 0, n > 0) are investigated at a BPW91/TZVP theory level. The experimental first VDEs for these Fe-V sulfur clusters are reported. The most probable ground state structures and spin multiplicities for these clusters are tentatively assigned by comparing their theoretical and experiment first VDE values. For  $FeVS_{1-3}^-$  clusters, their first VDEs are generally observed to increase with the number of sulfur atoms from 1.45 eV to 2.86 eV. The NBO/HOMOs of the ground state of FeVS<sub>1-3</sub> clusters are localized in a p orbital on a S atom; the partial charge distribution on the NBO/HOMO localized site of each cluster anion is responsible for the trend of their first VDEs. A less negative localized charge distribution is correlated with a higher first VDE. Structure and steric effect differences for  $Fe_mV_nS_{m+n}$ (m + n = 3, m > 0, n > 0) clusters are suggested to be responsible for their different first VDEs and properties. Two types of structural isomers are identified for  $Fe_mV_nS_{m+n}^-$  (m+n=4, m>0, n>0) clusters: a tower structure isomer and a cubic structure isomer. The first VDEs for tower like isomers are generally higher than those for cubic like isomers of  $Fe_m V_n S_{m+n}^-$  (m + n = 4, m > 0, n > 0) clusters. Their first VDEs are can be understood through: (1) NBO/HOMO distributions, (2) structures (steric effects), and (3) partial charge numbers on the NBO/HOMO's localized sites. EBEs for excited state transitions for all Fe-V sulfur clusters are calculated employing OVGF and TDDFT approaches at the TZVP level. The OVGF approach for these Fe/V/S cluster anions is better for the higher transition energies than the TDDTF approach. The experimental and theoretical results for these Fe/V/S cluster anions are compared with their related pure iron sulfur cluster anions. Properties of the NBO/HOMO are essential for understanding and estimating the different first VDEs for Fe/V/S, and comparing them to those of the pure Fe/S cluster anions.

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# Introduction

Iron sulfur clusters are prevalent in both biological and industrial systems, 1-7 as has been recognized for many decades.

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 $\dagger$  Electronic supplementary information (ESI) available: The following results are supplied as additional detailed information for these studies: (1) the mass spectrum of Fe<sub>m</sub>V<sub>n</sub>S<sub>x</sub>C<sub>y</sub><sup>-</sup> cluster anions generated by laser ablation of a mixed Fe:V = 1:1 target in the presence of 0.1% CS2 in He carrier gas is presented in Fig. S1; (2) DFT optimized structures of the FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> cluster at the BPW91/TZVP level are displayed in Fig. S2; (3) NBO plots showing HOMO to HOMO-4 orbitals of  $\text{FeVS}_{1-3}$  and  $\text{Fe}_m V_n S_{m+n}$  (m+n=3,4;m>0,n>0) clusters are displayed in Fig. S3-S5; (4) a brief description of broken symmetry; and (5) spin orbit coupling (SOC) corrected calculational results for (FeS)<sub>1,2</sub> clusters. See DOI: 10.1039/ c8cp03157f

Investigations of iron-sulfur systems, ranging from bare Fe-S clusters to analogue complexes and proteins, are common throughout bioinorganic chemistry. Synthesis and characterization of iron sulfur clusters and complexes comprise a large sub-field of organometallic chemistry.8 A number of studies have been performed on gas phase cationic, 9,10 neutral, 11,12 and anionic13-20 iron sulfur clusters for investigation of their composition, stability, structure, and reactivity. Extensive theoretical efforts devoted to the structural evolution of electronic properties of iron-sulfur complexes have also appeared.<sup>21-25</sup> For example, the structure of the Fe<sub>2</sub>S<sub>2</sub><sup>-</sup> cluster is assigned to be a planar rhomboid, the structure of the Fe<sub>3</sub>S<sub>3</sub> cluster is assigned to be a planar six-member ring, and a cubic structure is assigned for the  $\mathrm{Fe_4S_4}^-$  cluster.  $^{26}$ 

Trace elements, such as vanadium, are found to be essential for both biological and general catalytic systems.<sup>27-34</sup> The bacterial enzyme nitrogenase can catalyze the reduction of atmospheric  $N_2$  to  $NH_3$  and is responsible for cycling about 108 tons of N per year from the atmosphere to the soil. <sup>29</sup> Modeling the enzymatic  $N_2$  fixation process remains one of the great challenges for bioinorganic chemists. Many studies find that Fe–V mixed metal sulfur clusters in enzymes can be characterized as an active catalytic site; for example, an Fe/V/S anionic cluster is suggested to be responsible for the conversion of  $N_2$  to  $NH_3$ . <sup>33</sup> Therefore, investigations of iron–vanadium mixed metal sulfur systems, ranging from small Fe–V sulfur clusters to analogous complexes and proteins, are common throughout bioinorganic chemistry.

Photoelectron spectroscopy (PES) has been proven to be a successful approach for study of electronic and geometric structures of atomic and molecular clusters,35 as it combines size selectivity with spectral sensitivity and can generate information on both ionic and neutral species. These gas phase experimental results can then be directly and accurately compared to the calculated ones for the clusters of interest.<sup>13</sup> Computational chemistry has a very important role to play in helping to predict and rationalize the nature of the electronic ground state of transition metal compounds.<sup>36</sup> PES experimental results for cluster anions are also essential as tests for the performance of appropriate computational and theoretical methods. Electron binding energies of Fe-V sulfur clusters obtained from theory can be compared with those obtained from experiment to justify the theoretical method. Theoretical results obtained by provably reliable calculations can then be employed to analyze further PES spectra and finally generate geometric and electronic structures for those Fe-V sulfur clusters that are not directly observable experimentally. Thus, theoretical calculations for both anionic and neutral systems can be employed to explore cationic species, as well.

This report presents a PES study of a series of Fe–V mixed metal sulfur cluster anions, employing a magnetic-bottle time-of-flight (MBTOF) photoelectron spectroscopy (PES) apparatus. The PES spectra of these cluster anions at 355 nm and 266 nm photon energies are reported, and the structural and electronic properties of these cluster anions are investigated by density functional theory (DFT). The most probable ground state structures and spin multiplicities of this small, neutral, and anionic cluster series are thereby assigned by comparing the theoretical first vertical detachment energies (VDEs) with their experiment values.

#### **Methods**

#### A. Experimental

The MBTOF-PES experimental setup, consisting of a laser vaporization cluster/molecular source, an orthogonal acceleration/extraction reflectron time of flight (oaRETOF) mass spectrometer (MS), a mass gate, a momentum decelerator, and a MBTOF electron analyzer, employed in this work has been described previously in detail.<sup>37,38</sup> Only a brief outline of the apparatus is given below. In this work, Fe/V/S clusters are generated by laser ablation of a mixed iron/vanadium target [made by pressing a

mixture of iron (99.9%, Sigma Aldrich) and vanadium (99.9%, Sigma Aldrich) powders with a ratio of Fe:V = 1:1] in the presence of 0.1% CS2 in helium carrier gas. A 10 Hz, focused, 532 nm Nd<sup>3+</sup>:YAG (Nd<sup>3+</sup>:yttrium aluminum garnet) laser with  $\sim 5$  mJ per pulse energy is used for the laser ablation. The expansion gas is pulsed into the vacuum by a supersonic nozzle (R. M. Jordan, Co.) with a backing pressure of typically 100 psi. The mass spectrum obtained from the above generation methods is given as Fig. S1 in the ESI.† The ablation laser energy is adjusted from 1 to 10 mJ per pulse to seek the best generation condition for Fe-V sulfur clusters. The intensity of mass spectral features for  $Fe_m V_n S_x^-$  cluster anions is observed to be related to the ablation laser energy. The intensity of the mass spectrum increases when the ablation laser energy is adjusted from 1 to  $\sim$  5 mJ per pulse, and then decreases if the ablation laser energy keeps increasing from 5 to 10 mJ per pulse. This behavior suggests that the obtained MS and PES profiles are probably affected by cluster source conditions. The mass selected and decelerated anions are exposed to different laser wavelengths (355 nm, 266 nm) at the photo-detachment region. The photodetached electrons are energy analyzed by the MBTOF-PES spectrometer. PES spectra are collected and calibrated at this resolution with known spectra of Cu<sup>-</sup>.<sup>39</sup>

#### B. Theoretical

All calculations are performed using the Gaussian 09 program package.40 The structures of Fe/V/S clusters are optimized for different isomers and spin multiplicities using DFT without constraints. For each cluster, different initial structures are employed as the input in the optimization procedure. There are many possible structures, especially for the larger clusters. Unfortunately, we are probably not able to calculate all possibilities, but an extensive search for the global minimum structure has been pursued for each cluster: for example, see calculated structures of the FeV<sub>3</sub>S<sub>4</sub> cluster in Fig. S2 in the ESI.† The magnetic (spin) properties of iron containing clusters stand as their most fundamental characteristic and thereby provide an indispensable and essential means for their characterization. Therefore, spin-dependent delocalization, depicting ferromagnetic and antiferromagnetic spin alignments, is one of the most interesting, essential, and challenging topics for iron containing cluster studies. 24,41,42 In this work, the relative energies for each cluster with different spin multiplicities (M = (2S + 1)) from low to high are investigated at the BPW91/ TZVP level. Broken-symmetry is employed for low spin calculations (see details in the ESI†). Generally, the lowest relative energy spin state is selected and discussed for each structure isomer of small  $Fe_m V_n S_x^-$  (m + n = 2, 3) clusters. For larger  $\operatorname{Fe}_{m} \operatorname{V}_{n} \operatorname{S}_{x}^{-} (m+n=4)$  clusters, several spin states are found to be relative energy degenerate and lowest, so these spin states are all discussed in Section C below. In our previous studies of iron sulfur clusters, 26 the spin-orbit coupling (SOC) corrections are found to be insignificant for the calculations of relative energies of the spin states and the first VDE of small clusters, such as (FeS)<sub>1,2</sub> (details are given in the ESI†). Thus, SOC corrections are neglected for calculations of relative energies of the spin

states and the first VDEs of Fe-V sulfur clusters studied in this work at the present level of theory. Nonetheless, SOC may be important for larger clusters, however, which present a symmetrical environment for iron centers. 43 All relative energies are zero point energy corrected. Vibrational frequency calculations are further performed to confirm global minima, which have zero imaginary frequency. The Perdew-Wang<sup>44</sup> correlation functional (BPW91) and the triple-ζ valence plus polarization (TZVP)<sup>45</sup> basis set, which were proved to have good performance in previous studies of iron sulfur 12,37 and vanadium sulfur clusters, 46 are employed to explore these Fe/V/S clusters. In our previous studies of iron sulfur and vanadium sulfur clusters, different reasonable functionals (B3LYP, 47,48 B3PW91, 49,50 and APFD<sup>51</sup>) and basis sets (6-311+G(d)<sup>52-54</sup>) are employed to calculate the relative energy and the first VDE of the studied clusters (such as (FeS)<sub>4</sub><sup>-26</sup>) with different spin multiplicities. The basis sets aug-cc-PV5Z<sup>55</sup> for sulfur atoms and TZVP for iron atoms are also employed and tested to seek a more secure assignment. All the above calculation methods are selected for comparison with BPW91/TZVP level calculations to ensure that the latter method is sufficient to explore the description of iron sulfur anion clusters studied by us previously. Calculated results for the relative energy ( $\Delta E$ ) and the first VDE of studied clusters with different spin multiplicity employing different functionals and basis sets are indistinguishable from one another and consistent with the experimental results (VDEs).26 Such comparisons make clear that the performance of BPW91/TZVP is more than sufficient for investigation of Fe-V sulfur clusters, so the method BPW91/TZVP is adopted for the present studies. All calculations are treated in a spin-unrestricted manner.

In this approach, for each spin state of the Fe-V sulfur cluster anion, the first vertical detachment energy (VDE =  $E_{\text{neutral at optimized anion geometry}} - E_{\text{optimized anion}}$ ) is calculated as the lowest transition from the spin state (M) of the anion into the spin state (M + 1 or M - 1, M = 2S + 1) of the related neutral species at the geometry optimized for the anion. The M + 1 spin state of the related neutral species is considered for the process in which the photo detached electron comes from paired electrons, and the M-1 spin state of the related neutral species is considered for that in which the photo detached electron comes from an unpaired electron. The other spin states of the related neutral species are not considered, because

a second electron transition or spin conversion process must be considered for such a transition. Compared with the transition of the M spin state ion to M + 1 or M - 1 spin states of the related neutral species, a second electron transition or spin conversion process in the related neutral species should have significantly lower probability (intensity). The optimized anion geometries are used for the further calculations of the photoelectron spectra employing time-dependent density functional theory (TDDFT).56 Vertical excitation energies of the neutral species are added to the first VDE of the corresponding Fe-V sulfur anion clusters to obtain their second and higher EBEs. The outer valence Green function method (OVGF/TZVP)<sup>57</sup> is also used to calculate the second and higher VDEs of these Fe-V sulfur anion clusters.

NBO analysis is an often employed orbital (wave function) localization and population analysis method to help understand the electron distribution in a molecule or cluster around particular sites or moieties of interest. Within this method, natural atomic orbitals (NAOs), determined for the particular species under consideration, are evaluated and employed: NAOs are the effective orbitals of an atom in the particular molecular environment (rather than for isolated atoms). NAOs are also the maximum occupancy orbitals. Information obtained from an NBO analysis, such as partial charges and HOMO-LUMO orbitals, is reported to explain, for example, a number of experimental phenomena of gas phase 1-butyl-3-methylimidazolium chloride ion pairs.58 The NBO calculations in this work are performed using the NBO 3.1 program as implemented in the Gaussian 09 package. Partial charge distributions of cluster anions studied in this work are calculated using NBO analysis.

#### Results and discussion

#### A. Studies of the FeVS<sub>1-3</sub> cluster anions

Photoelectron spectra of  $FeVS_{1-3}^{-}$ . The obtained PE spectra for the FeVS<sub>1-3</sub> cluster anions at different photon energies are shown in Fig. 1. Photo detachment transitions occur between the ground state of an anion and the ground and excited states of its neutral counterpart, at the structure of the anion. The profile of the transition is governed by the Franck-Condon overlap between the two species, the anion and the neutral.

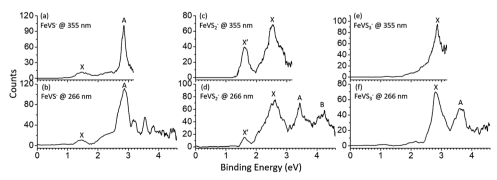


Fig. 1 Photoelectron spectra of FeVS $_{1-3}^-$  cluster anions at 355 nm and 266 nm. X and X' label the ground state transition peaks, and A and B label the first and second low-lying transition peaks at high VDE (see assignment details presented in Table 1).

The electron binding energy (EBE) value at the intensity maximum in the Franck-Condon profile is the vertical detachment energy (VDE). The first VDE, proving important in establishing a cluster's electronic and geometric structure, is derived from the energy of the first peak maxima in the photoelectron spectra. For the PE spectra of the FeVS cluster shown in Fig. 1a and b, two features are observed at both 355 nm and 266 nm, and their measured VDEs are 1.45 (X) and 2.85 (A) eV, respectively. In Fig. 1c, two features are observed in the PE spectrum of the FeVS<sub>2</sub><sup>-</sup> cluster at 355 nm, and their measured VDEs are 1.63 (X') and 2.55 (X) eV. Two peaks are observed for the higher transition labeled feature A (3.45 eV) and B (4.20 eV) at 266 nm photon energy (Fig. 1d). For the PE spectrum obtained for the FeVS<sub>3</sub> cluster (Fig. 1e), only one peak (2.86 eV, X) is observed at 355 nm. The second peak is observed for the higher transition labeled feature A (3.65 eV) at 266 nm photon energy (Fig. 1f).

**DFT calculations for FeVS**<sub>1-3</sub><sup>-</sup>. Determination of the geometrical structures of the clusters is important, since this cluster property is the basis for the description of all other cluster characteristics (e.g., electronic structure, electron density, charge and spin densities, etc.). Various structural isomers of Fe-V sulfur clusters discussed in this work are investigated, and different spin multiplicities from low to high are considered for each isomer. The relative energy differences ( $\Delta E$ ,  $\Delta G$ ) between different structural isomers with different spin multiplicities are calculated and compared to evaluate their relative stability. The structure and spin multiplicity for the ground state of a cluster anion are assigned mainly based on agreement of the calculated first VDEs compared to the experimental values. The calculated VDEs (in eV) for FeVS<sub>1-3</sub> clusters at the BPW91/ TZVP level, as well as the experimental results for comparison, are shown in Table 1.

Three types of structural isomers are obtained from calculations for the FeVS<sup>-</sup> cluster anion. Structural details of the three isomers are displayed in Fig. 2a. Comparing their calculated first VDEs with the experimental values obtained in Fig. 1a, b and Table 1, the calculated VDEs of isomer I (1.82 eV) and isomer III (1.61 eV) of FeVS are both in reasonable agreement with the experimental value of the X labeled feature (1.45 eV). These results suggest both structural isomers I and III of FeVS probably exist under the experimental conditions and contribute to the PE spectrum for the FeVS cluster, but structural isomer II of FeVS<sup>-</sup> probably does not exist in the anion beam. Comparison of PES experimental VDEs to calculated ones is the appropriate method by which to study and assign the actual structures and properties of the clusters present.

Two types of structural isomers are obtained theoretically for the FeVS<sub>2</sub> cluster anion: the geometric details of isomers I and II are displayed in Fig. 2b. Isomer I of FeVS<sub>2</sub><sup>-</sup> has one terminal sulfur bonded to a vanadium site, and isomer II of FeVS<sub>2</sub> has a four-member ring structure. Comparing their calculated first VDEs with the experimental values obtained in Fig. 1c, d and Table 1, the calculated VDE of isomer I (2.46 eV) of FeVS<sub>2</sub><sup>-</sup> is in good agreement with the experimental value of the X labeled feature (2.55 eV), and the calculated VDE of isomer II (1.76 eV)

**Table 1** The calculated VDEs (in eV) for  $FeVS_{1-3}^-$  clusters at the BPW91/ TZVP level, as well as the experimental results for comparison. Following EBE values after the first VDE are calculated employing TDDFT at the BPW91/TZVP level and the OVGF/TZVP method. The relative energies ( $\Delta E$ ) of different isomers with their various spin multiplicities are presented for each cluster ( $\Delta G$ s are given in the brackets)

	Structural isomer	Spin $(M = 2S + 1)$	BPW91/TZ	Exp.		
Cluster				Calculated VDE	Observed feature	VDE
FeVS <sup>-</sup>	Isomer I	1	0.00 (0.00)	1.82 <sup>a</sup>		1.45
				$3.22^b$ $2.17^c$	A	2.85
	Isomer II	3	0.06 (0.03)	$0.61^{a}$	Not obser	ved
	Isomer III	1	0.49(0.50)	1.61 <sup>a</sup>		1.45
			, ,	$2.52^b$ $2.45^c$	A	2.85
FeVS <sub>2</sub>	Isomer I	3	0.00 (0.00)	2.46 <sup>a</sup>	X	2.55
			, ,	3.46 <sup>b</sup> 3.13 <sup>c</sup>	A	3.45
				3.94 <sup>b</sup> —	В	4.20
	Isomer II	3	0.32 (0.39)	1.76 <sup>a</sup>	X'	1.63
			,	$3.42^b 2.08^c$		3.45
				$4.07^{b}$ —	В	4.20
FeVS <sub>2</sub> -	Isomer I	3	0.00 (0.00)	$2.46^{a}$	X	2.86
3			( )	$4.43^b$ $2.86^c$		3.65
	Isomer II	3	0.31 (0.33)	2.87 <sup>a</sup>	X	2.86
			( )	$3.81^b$ $3.88^c$	Α	3.65
	Isomer III	3	0.93 (0.89)	$2.76^{a}$		2.86
		-	(0.03)	$3.91^b$ $3.16^c$	A	3.65
				3.51 3.10	11	5.05

<sup>a</sup> The calculated first VDE. <sup>b</sup> The calculated following EBE value after the first VDE employing the OVGF/TZVP method. <sup>c</sup> The calculated following EBE value after the first VDE employing TDDFT at the BPW91/TZVP level.

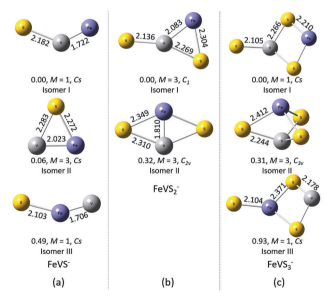


Fig. 2 DFT optimized structures of (a) FeVS- (b) FeVS2- and (c) FeVS3clusters at the BPW91/TZVP level. The lowest relative energy spin state geometry of each isomer is displayed in this figure. Geometries of other spin states for each cluster are generally similar to the one shown but with slightly different bond lengths and angles. Bond lengths (in angstroms), relative energy (in eV), spin multiplicity M, and point group symmetry are indicated on the structures. [Grey = V, blue = Fe, yellow = S.]

of FeVS<sub>2</sub><sup>-</sup> is also in good agreement with the experimental value of the X' labeled feature (1.63 eV).

For the FeVS<sub>3</sub><sup>-</sup> cluster anion, three types of structural isomer are obtained theoretically and their geometric details are displayed in Fig. 2c. Comparing their calculated first VDEs with the experimental values obtained in Fig. 1e, f and Table 1, the calculated first VDEs of isomers I (2.46 eV), II (2.87 eV), and III (2.76 eV) of FeVS<sub>3</sub> are close (within  $\sim 0.4$  eV), and are all in good agreement with the experimental value of the X labeled feature (2.86 eV). These results suggest structural isomers I, II, and III of FeVS<sub>3</sub> probably all exist under the experimental conditions and contribute to the PE spectrum for the FeVS<sub>3</sub><sup>-</sup> cluster.

In sum, the experimental first VDEs are generally observed to increase with the number of sulfur atoms from 1.45 eV to 2.85 eV for these FeVS<sub>1-3</sub> clusters. Diverse types of structural isomers are found for each  $FeVS_x^-$  (x = 1-3) cluster. One type of structural isomer with a terminal sulfur bonded to a vanadium site (isomer I) is found for all FeVS<sub>1-3</sub> clusters, and its calculated relative energy ( $\Delta E$ ) is obtained to be the lowest among all structural isomers for each species  $FeVS_x^-$  (x = 1 - 3).

Understanding the first VDEs for FeVS<sub>1-3</sub> through structures, NBO/HOMO distributions, and partial charge distributions. To estimate the first VDEs of cluster anions, one electron is removed from the highest occupied molecular orbital (HOMO) of the cluster maintaining the cluster optimized geometry. Therefore, studies of HOMO properties of these FeVS<sub>1-3</sub> clusters are helpful to understand their different first VDEs. Furthermore, partial atomic charges are suggested to play a decisive role in determining core electron binding energy in small molecules.<sup>59</sup> The partial charge at the HOMO localized site in these Fe-V sulfur cluster anions may to some extent affect the energy (first VDE) required to remove an electron from such clusters through a "charge effect". A small negative charge number at the site means less electron density distribution on that site (i.e., the site is more positive than anticipated), and therefore removal of an electron from that site may require more energy than otherwise estimated based simply on the NBO/HOMO distribution for that site. In order to investigate and understand the above interesting physical behavior for FeVS<sub>1-3</sub> clusters, NBOs and NBO charges for each atom are calculated for all assigned ground state isomers: isomers I of FeVS<sub>1.3</sub> cluster anions, and isomers I and II of FeVS<sub>2</sub> clusters anions. For the latter case, both features X' and X are identified.

Electron density distribution plots for FeVS<sub>1-3</sub> cluster NBO/ HOMOs are shown in Fig. 3. The experimental first VDEs of  $\text{FeVS}_{1-3}^{-}$  clusters increase with the number of sulfur atoms: from 1.45 eV of FeVS-, to 2.55 eV of FeVS2-, and to 2.86 eV of FeVS<sub>3</sub><sup>-</sup>. The NBO/HOMOs of all isomers I of FeVS<sub>1-3</sub><sup>-</sup> present a major electron distribution similar to that of localized p orbitals on the terminal S atom. The NBO partial charge numbers of the S sites, on which the NBO/HOMO is localized, are -0.871 for isomer I of FeVS<sup>-</sup>, -0.560 for isomer I of FeVS<sub>2</sub><sup>-</sup>, and -0.309 for isomer I of FeVS<sub>3</sub> (see Fig. 3). Since the S atom is more electronegative than iron or vanadium atoms, the negative charge number at the S site in these Fe-V sulfur clusters is not unreasonable. The observed increase of the first VDE for these FeVS<sub>1-3</sub> clusters with the number of sulfur atoms is consistent with the decrease of the negative charge number of their

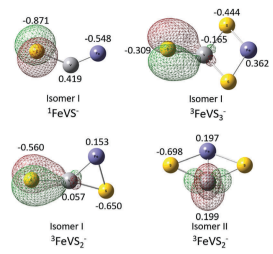


Fig. 3 NBO plots showing the highest occupied molecular orbital (HOMO) of  $FeVS_{1-3}^-$  cluster anions. The spin multiplicity (M) is listed as  $^{M}$ FeVS<sub>1-3</sub> $^{-}$ . The NBO charges for important atoms are given in the figure. [Grey = V, blue = Fe, yellow = S.]

NBO/HOMO localized S site. These results suggest that the above proposed "charge effect" is probably an important factor responsible for the trend of the first VDEs of these ground state (isomer I) FeVS<sub>1-3</sub> cluster anions.

#### B. Studies of the $\operatorname{Fe}_m V_n S_{m+n}^-$ (m+n=3, m>0, n>0) cluster anions

Photoelectron spectra of  $Fe_m V_n S_{m+n}^-$  (m+n=3, m>0, n>0). The obtained PE spectra for the  $Fe_m V_n S_{m+n}^-$  (m + n = 3, m > 0, n > 0) cluster anions at different photon energies are shown in Fig. 4. For the PE spectrum of the FeV<sub>2</sub>S<sub>3</sub> cluster shown in Fig. 4a, one broad feature (X) is observed at 355 nm, and its measured first VDE is 2.00 eV. Another broad peak is observed for the higher transition labeled feature A (3.17 eV) at 266 nm photon energy (Fig. 4b). In the PE spectra of the Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> cluster, one peak (3.18 eV, X) is partly observed at 355 nm (Fig. 4c), and the second peak is observed for the higher transition labeled feature A (~4.0 eV) at 266 nm photon energy (Fig. 4d).

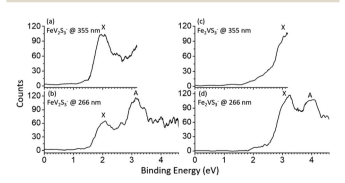


Fig. 4 Photoelectron spectra of  $\text{FeV}_2\text{S}_3^-$  and  $\text{Fe}_2\text{VS}_3^-$  cluster anions at 355 nm and 266 nm. X labels the ground state transition peak, and A labels the first low-lying transition peak at high VDE (see assignment details presented in Table 2)

**Table 2** The calculated VDEs (in eV) for  $FeV_2S_3^-$  and  $Fe_2VS_3^-$  clusters at the BPW91/TZVP level, as well as the experimental results for comparison. Following EBE values after the first VDE are calculated employing TDDFT at the BPW91/TZVP level and the OVGF/TZVP method. The relative energies  $(\Delta E)$  of different isomers with their various spin multiplicities are presented for each cluster ( $\Delta G$ s are given in the brackets)

			BPW91/TZVP (eV)		Exp.	
Cluster	Structural isomer	. *	$\Delta E$	Calculated VDE	Observed feature	VDE
FeV <sub>2</sub> S <sub>3</sub>	Isomer I	4	0.00 (0.00)	$2.55^a$ $4.13^b$ $2.90^c$		2.00 3.17
	Isomer II	4	0.12 (0.16)	$2.22^a$ $3.77^b$ $3.02^c$		2.00 3.17
Fe <sub>2</sub> VS <sub>3</sub>	Isomer I	5	0.00 (0.00)	$2.58^a$ $4.17^b$ $3.15^c$	X A	3.18 ~ 4.0
	Isomer II	5	2.55 (2.60)	1.31 <sup>a</sup>	Not obser	ved

<sup>&</sup>lt;sup>a</sup> The calculated first VDE. <sup>b</sup> The calculated following EBE value after the first VDE employing the OVGF/TZVP method. The calculated following EBE value after the first VDE employing TDDFT at the BPW91/TZVP level.

DFT calculations for  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-}$  (m+n=3, m>0, n>0). Calculated VDEs (in eV) for FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> clusters at the BPW91/TZVP level, as well as the experimental results for comparison, are shown in Table 2. Two types of structural isomers are obtained calculationally for the FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> cluster anion; the geometric details are displayed in Fig. 5a. Comparing their calculated first VDEs with the experimental values obtained in Fig. 4a, b and Table 2, the calculated VDE for isomer II (2.22 eV) of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> is in slightly better agreement with the experimental value of the X labeled feature (2.00 eV) than that for isomer I (2.55 eV) of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup>.

For the Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> cluster anion, two types of structural isomer are also obtained theoretically: geometric details for isomers I and II are displayed in Fig. 5b. As presented in Table 2, the calculated VDE of isomer I of Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> is 2.58 eV, which is similar to the experimental value 3.18 eV obtained from the X labeled feature in Fig. 4c and d.

Understanding the first VDEs for  $Fe_m V_n S_{m+n}^-$  (m + n = 3, m > 0, n > 0) through structures, NBO/HOMO distributions, and partial charge distributions. Due to comparison of calculated VDEs with experimental values as discussed above, the ground state of the Fe<sub>2</sub>VS<sub>3</sub> cluster anion is assigned to be isomer I, whose structure is a planar six-member ring as shown in Fig. 5b. The ground state of the FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> cluster anion is assigned to be isomer II, whose structure is 3-dimensional as shown in Fig. 5b. The experimental first VDEs for the above three metal, three sulfur FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> clusters, are about one eV different. The experimental first VDE of Fe<sub>2</sub>VS<sub>3</sub> (3.18 eV) is  $\sim 1.2$  eV higher than that of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> (2.00 eV) as displayed in Table 2. With the same analysis strategy used above in Section A for two metal atom containing FeVS<sub>1-3</sub> clusters, NBOs and NBO partial charges for each atom are calculated for both assigned ground state isomers (II) of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and (I) of Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> clusters anions to understand their different properties.

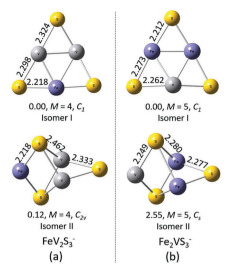


Fig. 5 DFT optimized structures of (a)  $FeV_2S_3^-$  and (b)  $Fe_2VS_3^-$  at the BPW91/TZVP level. The lowest relative energy spin state geometry of each isomer is displayed in this figure. Geometries of other spin states for each cluster are generally similar to the one shown but with slightly different bond lengths and angles. Bond lengths (in angstroms), relative energy (in eV), spin multiplicity M, and point group symmetry are indicated on the structures. [Grey = V, blue = Fe, yellow = S.]

In plots of distributions for their NBO/HOMOs show in Fig. 6, NBO/HOMOs of both isomers present a major electron distribution similar to that of localized p orbitals on a S atom. The NBO partial charge numbers of the S sites, on which the NBO/HOMO is localized, are -0.468 for isomer II of FeV<sub>2</sub>S<sub>3</sub> $^-$ , and -0.638 for isomer I of Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup>. Note that the charge number for the NBO/HOMO localized site of isomer II of Fe<sub>2</sub>VS<sub>3</sub> is more negative than that of isomer I of FeV<sub>2</sub>S<sub>3</sub>, but the first VDE for  $Fe_2VS_3^-$  (3.18 eV) is higher than that of  $FeV_2S_3^-$ (2.00 eV). These results suggest the "charge effect" proposed in Section A is not the only factor to affect the first VDEs of these large three metal containing Fe-V sulfur cluster anions. For the 3-dimensional structural isomer II of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup>, the NBO/HOMO is localized on sulfur atom bonds with two vanadium atoms, and that  $\angle$  VSV angle is 47.78° (Fig. 5a and 6). For the planar structural isomer I of Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup>, the NBO/HOMO is localized on sulfur atom bonds with two iron atoms, and that ∠ FeSFe angle

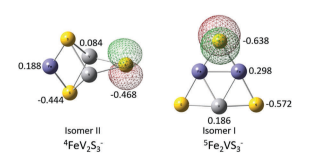


Fig. 6 NBO plots showing the highest occupied molecular orbital (HOMO) of  ${\rm FeV_2S_3}^-$  and  ${\rm Fe_2VS_3}^-$  cluster anions. The spin multiplicity (M) is listed as  ${}^{M}\text{Fe}_{m}\text{V}_{n}\text{S}_{m+n}^{-}$ . The NBO charges for important atoms are given in the figure. [Grey = V, blue = Fe, yellow = S.]

is 62.77° (see Fig. 5b and 6). Their structures are different, and the bigger metal-sulfur (NBO/HOMO localized)-metal angle (62.77°) for isomer I of Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> means a larger steric effect for the electron detachment than that for isomer II of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup>. This structure and steric effect difference between FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and Fe<sub>2</sub>VS<sub>3</sub> clusters probably is responsible for their different first VDEs.

In summary, three metal atom containing Fe/V/S anions. FeV<sub>2</sub>S<sub>3</sub> and Fe<sub>2</sub>VS<sub>3</sub> clusters, are discussed in this section through PES and DFT. The first VDE for Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> is observed  $\sim 1.2$  eV higher than that of FeV<sub>2</sub>S<sub>3</sub>. Their NBOs are found to be the same and localize on S atom p orbitals. The "charge effect" for the NBO/HOMO localized site is not suggested to be a major factor affecting their first VDEs. With different Fe/V ratios, their ground state structures are assigned to be different, and their structure and steric effect differences are suggested to be responsible for their different first VDEs and properties. Note that the structures of ground state FeVS<sub>1-3</sub> clusters discussed in Section A are simpler (all close to planar) than these three metal atom containing Fe-V sulfur (FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup>) clusters: the trend of the first VDEs of FeVS<sub>1-3</sub><sup>-</sup> cluster anions can be rationalized by a "charge effect" at their comparable NBO/HOMO localized sites. Therefore, structure apparently plays a more important role for the properties of these Fe-V mixed sulfur cluster as their size increases: this correlation is especially important for understanding and estimating their first VDEs.

#### C. Studies of $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-}$ (m+n=4, m>0, n>0) cluster anions

Photoelectron spectra of  $\operatorname{Fe}_m V_n S_{m+n}^-$  (m+n=4, m>0, n>0). In this section, four metal atom containing Fe-V sulfur clusters  $\operatorname{Fe}_{m} \operatorname{V}_{n} \operatorname{S}_{m+n}^{-} (m+n=4, m>0, n>0)$  are discussed. Their PE spectra are shown in Fig. 7. The PE spectrum of the FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> cluster shown in Fig. 7a evidences one broad feature at 355 nm excitation: its measured first VDE is  $\sim 2.7$  (X) eV. Another broad peak is observed for the higher transition labeled feature A ( $\sim$ 4.0 eV) at 266 nm photon energy (Fig. 7b). In Fig. 7c, two features are observed in the PE spectrum of the Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> cluster at 355 nm: their measured VDEs are 1.50 (X') and 2.17 (X) eV. One broad peak is observed for the higher transition labeled

feature A ( $\sim$  3.5 eV) obtained at 266 nm photon energy (Fig. 7d). For the Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> cluster anion, only broad peaks are observed (Fig. 7f). The first peak (X'') is around 1.60 eV, and the second peak (X') is around 2.3 eV. Another feature labeled X is observed at 3.20 eV. At higher binding energy, a broad peak (A) is detected from 3.5 eV to 4.4 eV.

DFT calculations for  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-} (m+n=4, m>0, n>0)$ . Calculated VDEs (in eV) for  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-} (m+n=4, m>0, n>0)$ cluster anions at the BPW91/TZVP level, as well as experimental results for comparison, are presented in Table 3. Two types of structural isomers are obtained from these calculations for each FeV<sub>3</sub>S<sub>4</sub><sup>-</sup>, Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup>, and Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> cluster anion. Isomer I of each  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-}$  (m+n=4, m>0, n>0) has a tower like structure and isomer II of each  $Fe_m V_n S_{m+n}^-$  (m + n = 4, m > 0,n > 0) has a cubic like structure.

As displayed in Table 3, the calculated first VDE for isomer I (2.64 eV) of FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> is in very good agreement with the experimental value of the X labeled feature ( $\sim 2.7$  eV).

For Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> cluster anions, the two lowest relative energy  $(\Delta E)$  spin states of each structural isomer are found to be degenerate (Table 3). Note that the intensity of peak X' is lower than that of peak X in Fig. 7c and d.  $\Delta E$  for isomer II of  $\text{Fe}_2\text{V}_2\text{S}_4^-$  is  $\sim 0.4$  eV higher than that for isomer I, which agrees with that intensity difference observation for X and X' peaks, considering that a higher  $\Delta E$  implies a less stable state with a lower population. These results imply that both structural isomer I and II of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> exist under the experimental conditions and contribute to the PE spectrum for Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub> cluster anions, and that their multiple low  $\Delta E$  spin states should also be considered when evaluating their properties and behavior in real chemical and biological systems.

The first VDEs and EBEs for both tower like structure isomer I and cubic like structure isomer II (see details in Fig. 8c) of Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> are also calculated and presented in Table 3. From Fig. 7f, the intensity of peaks decreases in the order X, X', X''. This trend for Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> clusters can be understood based on the increase of  $\Delta Es$  for nonet isomer I (the X feature) to 11-et isomer I (the X' feature), and to nonet and 11-et isomer II (the X" feature). These observations and comparisons illustrate that not only do the lowest relative energy spin state and isomer exist in the experimental system, but also that the relative

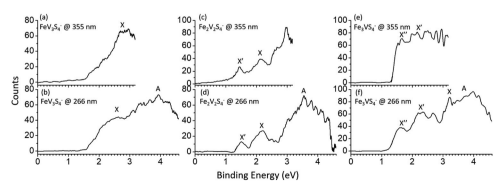


Fig. 7 Photoelectron spectra of  $\operatorname{Fe}_m \operatorname{V}_n \operatorname{S}_{m+n}^-$  (m+n=4, m>0, n>0) cluster anions at 355 nm and 266 nm. X, X' and X" label the ground state transition peaks, and A and B label the first and second low-lying transition peaks at high VDE (see assignment details presented in Table 3).

**Table 3** The calculated VDEs (in eV) for  $Fe_m V_n S_{m+n}^-$  (m + n = 4, m > 0, n > 0) clusters at the BPW91/TZVP level, as well as the experimental results for comparison. Following EBE values after the first VDE are calculated employing TDDFT at the BPW91/TZVP level and the OVGF/TZVP method. The relative energies ( $\Delta E$ ) of different isomers with their various spin multiplicities are presented for each cluster ( $\Delta G$ s are given in the brackets)

•			BPW91/TZVP (eV)		Exp.		
Cluster	Structural isomer	Spin $(M = 2S + 1)$	$\Delta E$	Calcula VDE	ated	Observed feature	VDE
FeV <sub>3</sub> S <sub>4</sub>	Isomer I	3		2.64 <sup>a</sup>			~2.7
	Isomer II	3	(0.00) 0.87 (0.88)			A ~4.0 Not observed	
$\mathrm{Fe_2V_2S_4}^-$	Isomer I	4		$2.39^{a}$		X	2.17
		-		4.47 <sup>b</sup>			~3.5
		6		$2.31^a$ $3.82^b$		X A	$2.17$ $\sim 3.5$
	Isomer II	6		$1.46^a$		X'	1.50
			(0.48)	3.93 <sup>b</sup>	$1.86^{c}$		~3.5
		8		$1.49^{a}$		X'	1.50
			(0.44)	3.77 <sup>b</sup>	1.91 <sup>c</sup>	A	~3.5
$\mathrm{Fe_3VS_4}^-$	Isomer I	9		$3.19^{a}$ $4.76^{b}$		X A	3.20 3.5-4.4
		11		$2.32^{a}$		X'	2.30
				4.53 <sup>b</sup>			3.5-4.4
	Isomer II	9	0.97	$1.70^{a}$		$X^{\prime\prime}$	1.60
				$4.26^{b}$			3.5-4.4
		11	0.87			X"	1.60
			(0.88)	$4.00^{b}$	2./9°	A	3.5-4.4

<sup>&</sup>lt;sup>a</sup> The calculated first VDE. <sup>b</sup> The calculated following EBE value after the first VDE employing the OVGF/TZVP method. c The calculated following EBE value after the first VDE employing TDDFT at the BPW91/TZVP level.

populations of the diverse co-existing spin states and isomers may relate to their relative energies.

Important details for whether or not such relative energy populations and spectroscopic intensities are appropriately compared and related to one another are typically contained in the particular characteristics and cross over points associated with the potential energy surfaces (stationary points, barrier heights, and intermediates states) for the various isomers of each cluster anion.

Understanding the first VDEs for  $Fe_m V_n S_{m+n}^-$  (m + n = 4,m > 0, n > 0) through structures, NBO/HOMO distributions, and partial charges. From the above results and discussion for four metal atom containing Fe-V sulfur clusters  $Fe_m V_n S_{m+n}^{-}$ (m + n = 4, m > 0, n > 0), we find that the experimental VDEs for tower like structure isomers are generally higher than those for cubic like structure isomers of these  $Fe_m V_n S_{m+n}^-$  (m + n = 4, m > 0, n > 0) clusters. To understand these results, NBOs and NBO partial charges for each atom are calculated for observed structural isomers of  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-} (m+n=4, m>0, n>0)$ clusters (Fig. 9).

Regarding  $Fe_m V_n S_{m+n}$  (m + n = 4, m > 0, n > 0) cluster anion experimental VDEs, the following comparison consequences are obtained. First, the experimental VDE for tower like structure isomer I of Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> is the highest, 3.20 eV. Its NBO/HOMO presents an electron distribution similar to that of

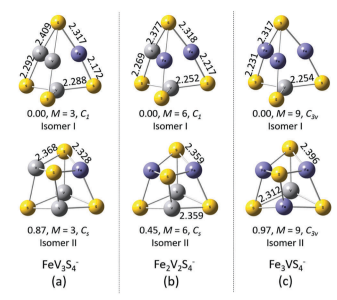


Fig. 8 DFT optimized structures of (a)  $FeV_3S_4^-$  (b)  $Fe_2V_2S_4^-$  and (c) Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> clusters at the BPW91/TZVP level. The lowest relative energy spin state geometry of each isomer is displayed in this figure. Geometries of other spin states for each cluster are generally similar to the one shown but with slightly different bond lengths and angles. Bond lengths (in angstroms), relative energy (in eV), spin multiplicity M, and point group symmetry are indicated on the structures. [Grey = V, blue = Fe, yellow = S.]

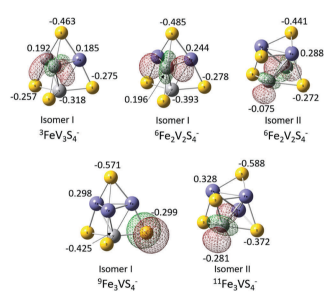


Fig. 9 NBO plots showing the highest occupied molecular orbital (HOMO) of  $Fe_mV_nS_{m+n}^-$  (m+n=4, m>0, n>0) cluster anions. The spin multiplicity (M) is listed as  ${}^{M}\text{Fe}_{m}\text{V}_{n}\text{S}_{m+n}^{-}$ . The NBO charges for important atoms are given in the figure. [Grey = V, blue = Fe, yellow = S.]

localized p orbitals on an S atom. The NBO partial charge number on its NBO/HOMO localized S site is -0.299. Second, for the other two tower like structure isomers, the experimental VDE for isomer I of FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> is 2.70 eV, and for isomer I of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub> is 2.17 eV. Their NBO/HOMOs both present an electron distribution similar to that of localized d orbitals on V atoms. The NBO partial charge numbers on their NBO/HOMO

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localized V sites are positive, 0.192 (isomer I of FeV<sub>3</sub>S<sub>4</sub><sup>-</sup>) and 0.196 (isomer I of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup>). Third, for cubic like structure isomers, the experimental VDE for isomer II of Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> is 1.60 eV, and for isomer II of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub> is 1.50 eV. Their NBO/HOMOs also present an electron distribution similar to that of localized d orbitals on V atoms. The NBO partial charge numbers on their NBO/HOMO localized V sites are negative, -0.281 (isomer II of Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup>) and -0.075 (isomer II of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup>).

The above three comparisons suggest that more complex factors affect the electronic properties (the first VDEs) simultaneously for these four metal atom containing Fe-V sulfur clusters  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-}$  (m+n=4, m>0, n>0), instead of only one major factor. The structure with regard to steric effects, NBO/HOMO distribution, and partial charge on the NBO/HOMO localized site (see the "charge effect" proposed in Section A) should all be considered to understand and estimate the changes of their first VDEs. The following three general rules can be derived based on the above results. First, NBO/HOMO distributions on S atoms compared to those on V atoms generate a higher first VDE for the cluster, due to the higher electronegativity of S than that of V (the Pauling electronegativity of S is 2.58, and of V is 1.63). This may be the reason that the highest first VDE (3.20 eV) observed for all  $\text{Fe}_{m}V_{n}S_{m+n}^{-}$  (m + n = 4, m > 0, n > 0)cluster anions is for isomer I of Fe<sub>3</sub>VS<sub>4</sub> (the NBO/HOMO distribution on the S atom). Second, the tower structure tends to have a higher first VDE than the cubic structure due to its relatively strong steric effect. For example, NBO/HOMOs for both isomer I (tower like structure) and isomer II (cubic like structure) of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> are localized on d orbital V atoms. The larger S-V (NBO/HOMO localized)-S angle for isomer I (165.58°) than that for isomer II ( $\sim 110.30^{\circ}$ ) implies a stronger steric effect for the tower structural isomer I of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub>-. This structural effect is perhaps responsible for the higher first VDE for isomer I (2.17 eV) than that for isomer II (1.50 eV) of Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup>. Third, the proposed "charge effect" still applies for these four metal atom Fe-V sulfur clusters: the positive NBO partial charge numbers on the NBO/HOMO localized V sites of FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> isomer I and Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> isomer I may play a role in their higher first VDEs, as compared to the negative NBO partial charge numbers for the NBO/HOMO localized V sites of Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> isomer II and Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> isomer II.

In summary, four metal atom containing Fe-V sulfur clusters,  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-} (m+n=4, m>0, n>0)$  cluster anions, are analyzed in this section through PES and DFT. Two types of structural isomers (tower like and cubic like) are observed for Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> and Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> clusters; only tower like structural isomers are observed for the FeV<sub>3</sub>S<sub>4</sub><sup>-</sup> cluster anions in the PES experiments. This suggests that the ratio of Fe and V atoms can affect the structure properties for these Fe-V sulfur clusters. The first VDEs of  $\operatorname{Fe}_m V_n S_{m+n}^-$  (m+n=4, m>0, n>0) clusters are reported and suggested to be understood through three electronic properties: (1) NBO/HOMO distributions, (2) structures (steric effects), and (3) partial charge numbers on the NBO/HOMOs localized sites (the "charge effect"). Note that comparisons of Tables 1-3 emphasize two distinct behaviors: (1) addition of metal atoms to these  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-} (m+n=2-4, m>0, n>0)$ 

clusters does not affect their first VDEs systematically; and (2) addition of V or Fe atoms to these clusters also does not generate systematic changes for their first VDEs.

#### D. EBE values of low-lying transition peaks for $FeVS_{1-3}$ and $\text{Fe}_{m}V_{n}S_{m+n}^{-}$ (m+n=3, 4; m>0, n>0) cluster anions

The electron binding energy (EBE) values, following the first VDE, are calculated employing TDDFT (BPW91/TZVP level) and OVGF/TZVP methods. The calculated results for FeVS<sub>1-3</sub> and  $\text{Fe}_{m}\text{V}_{n}\text{S}_{m+n}^{-}$  (m + n = 3, 4; m > 0, n > 0) cluster anions are summarized and compared with their experimentally measured values in Tables 1 to 3, respectively.

EBEs regarding excited state transitions for related Fe/V/S are calculated employing OVGF and TDDFT approaches at a TZVP basis set level. The OVGF approach is found to be generally better for higher transition energy theoretical studies of FeVS<sub>1-3</sub>-, Fe<sub>2</sub>VS<sub>3</sub>-, and FeV<sub>3</sub>S<sub>4</sub>- cluster anions, but not for FeV<sub>2</sub>S<sub>3</sub><sup>-</sup>. For Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> and Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> clusters, both approaches seem acceptable, mostly due to the breadth of the feature associated with their higher energy transition peaks. Therefore, both OVGF and TDDFT approaches are acceptable for these latter two cluster EBE calculations of Fe-V sulfur clusters regarding excited state transitions to obtain cautiously tentative assignments. The OVGF method may be preferred generally, however, due to its better agreement for most Fe-V sulfur clusters studied in this work.

### E. Comparison of $\text{Fe}_{m}V_{n}S_{m+n}^{-}$ (m+n=2-4, m>0, n>0)with pure (FeS)2-4 cluster anions

Since pure iron sulfur clusters are also found to be important in biological and industrial systems, comparing properties of the Fe-V sulfur clusters discussed above with properties of their related pure iron sulfur clusters becomes a useful endeavor.

As we reported previously, <sup>26</sup> Fe<sub>2</sub>S<sub>2</sub> - cluster anions have a fourmember ring structure, which is similar to isomer II of FeVS<sub>2</sub><sup>-</sup>. The NBO/HOMO of the Fe<sub>2</sub>S<sub>2</sub><sup>-</sup> cluster is localized on a p orbital on a S atom, however, which is similar to the electronic NBO/ HOMO of isomer I of FeVS<sub>2</sub><sup>-</sup>. Interestingly, the experimental first VDE of the Fe<sub>2</sub>S<sub>2</sub><sup>-</sup> cluster is 2.34 eV, which is close to that of isomer I of FeVS<sub>2</sub><sup>-</sup> (2.55 eV), but not close to that of structures like isomer II of FeVS<sub>2</sub><sup>-</sup> (1.63 eV). This result suggests the electron distribution property of the NBO/HOMO is a principal factor to be considered with regard to the first VDE of these small two metal containing metal sulfide clusters (Fe<sub>2</sub>S<sub>2</sub><sup>-</sup> and FeVS<sub>2</sub><sup>-</sup>).

Our previously reported experimental first VDE of the Fe<sub>3</sub>S<sub>3</sub><sup>-</sup> cluster anion  $(3.57 \text{ eV})^{26}$  is about  $\sim 0.4$  and  $\sim 1.5 \text{ eV}$  higher than those of the  $Fe_2VS_3^-$  (3.18 eV) and  $FeV_2S_3^-$  (2.00 eV) cluster anions. The NBO/HOMO of Fe<sub>3</sub>S<sub>3</sub><sup>-</sup> clusters is delocalized as a Fe-Fe bonding orbital. Interestingly, the NBO/HOMOs of FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> (isomer II) and Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> (isomer I) clusters both appear to be localized p orbitals on the S site; they are not observed localized on an Fe site. As proposed and discussed in the studies of  $(FeS)_m H_{0.1}$  (m = 2-4) cluster anions, the change of cluster VDE from low to high can be associated with the change in the nature of their NBO/HOMO from a valence p orbital on S to an

Fe–Fe delocalized valence bonding orbital.<sup>60</sup> Therefore, the distinctly different first VDEs of pure iron sulfur  $Fe_3S_3^-$  and Fe–V sulfur  $Fe_mV_{3-m}S_3^-$  (m=1,2) cluster anions can be related to the different NBO/HOMO electron distribution (wave function) properties of each cluster anion.

The experimental first VDE for the  $Fe_4S_4^-$  cluster previously reported<sup>26</sup> is 2.71 eV. The structure of this cluster anion is that of a distorted cube and its NBO/HOMO is that of a localized d orbital on an Fe site. Cubic like structures are also observed for  $Fe_2V_2S_4^-$  and  $Fe_3VS_4^-$  cluster anions. Their experimental first VDEs are 1.50 eV for  $Fe_2V_2S_4^-$ , and 1.60 eV for  $Fe_3VS_4^-$ ; their NBO/HOMOs appear to be localized d orbitals on the V sites. The lower first VDEs for these  $Fe_2V_2S_4^-$  and  $Fe_3VS_4^-$ ) than those for  $Fe_4S_4^-$  cluster anions may be related the smaller Pauling electronegativity for V (1.63) than that for Fe (1.83). These results also suggest that the electron distribution of the the NBO/HOMO is an essential characteristic through which one can understand and estimate the first VDEs of these big four metal containing metal sulfide cluster anions  $(Fe_4S_4^-, Fe_2V_2S_4^-)$  and  $Fe_3VS_4^-)$ .

In summary, the structure evolution from the Fe<sub>2</sub>S<sub>2</sub> to the Fe<sub>4</sub>S<sub>4</sub> cluster is from planar ring to three-dimensional cubic structures.26 In this work, a similar trend is observed for Fe-V sulfur clusters  $\operatorname{Fe}_{m} V_{n} S_{m+n}$  (m+n=2-4, m>0, n>0) with increased numbers of Fe and V atoms. A planar rhomboid structure is observed for the FeVS<sub>2</sub> (isomer II) cluster, planar six-member ring structures are found for both Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup> (isomer I) and FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> (isomer I) clusters, and cubic structures are assigned for four metal center containing clusters Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> (isomer II) and Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> (isomer II). Other structural isomers are also observed/assigned for these V atom involving clusters: for example, a planar structure containing a terminal S on V is also assigned for FeVS<sub>2</sub><sup>-</sup> (isomer I), a three-dimensional structure is only observed for FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> (isomer II), and tower like structures are found for all four metal center containing  $Fe_m V_n S_{m+n}$ (m + n = 4, m > 0, n > 0) clusters (isomer I). Note that the FeV<sub>3</sub>S<sub>4</sub> cluster is only assigned as a tower like structure (isomer I). Compared to pure iron sulfur clusters, the Fe-V sulfur clusters have a more diverse set of structural isomers, probably due to the varied oxidation state properties of V compared to those of Fe. The relative energy differences ( $\Delta E$ ) between different structural isomers with different spin multiplicities are calculated and compared to evaluate the relative stability of these Fe-V sulfur clusters. The structure and spin multiplicity for the ground state of a cluster anion is assigned mainly based on agreement of the calculated first VDEs compared to the experimental values. Interestingly, the calculated first VDEs of all lowest  $\Delta E$  structural isomers are in good agreement with the experimental value of studied Fe-V sulfur clusters, so the lowest  $\Delta E$  structural isomer is generally assigned to be the ground state structure. Nonetheless, some structural isomers with high  $\Delta E$  (less than 1 eV) are also observed in experiments, such as isomer II of  ${}^{3}\text{FeVS}_{2}^{-}$  ( $\Delta E = 0.32$ ) and isomer II of  ${}^{6}\text{Fe}_{2}\text{V}_{2}\text{S}_{4}^{-}$  ( $\Delta E = 0.45$ ). Some very low  $\Delta E$  structural isomers (for example, for isomer II of  ${}^{3}\text{FeVS}^{-}$   $\Delta E = 0.06$ , and first VDE = 0.61 eV) are not observed, although their calculated

first VDEs are not overlapped with those of other isomers. These results suggest that the distributions/populations of structural isomers of Fe–V sulfur cluster anions can be generated either thermodynamically (through  $\Delta E, \Delta G$ ) or kinetically (through transition state barriers). The obtained PES profile can provide an experimental method to assign coexisting structural isomers. Furthermore, electron distribution properties of the NBO/HOMO must also be an essential factor through which one can understand the first VDEs for Fe–V sulfur clusters compared to pure iron sulfur clusters. More comparison studies between larger Fe–V sulfur clusters and related pure iron sulfur clusters, with regard to how NBO/HOMO distributions affect their first VDEs, should certainly prove informative.

# Conclusions

Iron-vanadium sulfur cluster anions are studied by PES at 3.492 eV (355 nm) and 4.661 eV (266 nm) photon energies, and by DFT calculations. The structural properties, relative energies of different structural isomers, and calculated first VDEs of different structural isomers for these cluster anions are investigated at BPW91/TZVP theory levels. The most probable structures and ground state spin multiplicities for these clusters are tentatively assigned by comparing their theoretical and experiment first VDE values.

The first VDEs for FeVS<sub>1-3</sub> clusters are generally observed to increase with the number of sulfur atoms from 1.5 eV to 2.8 eV. Diverse types of structural isomers are found for each FeVS<sub>x</sub> (x = 1-3) cluster. One type of structural isomer with a terminal sulfur bonded to a vanadium site (isomer I) is found for all FeVS<sub>1-3</sub> clusters: its calculated relative energy ( $\Delta E$ ) is obtained to be the lowest among all structural isomers for each species FeVS<sub>x</sub><sup>-</sup> (x = 1-3). The NBO/HOMOs of ground states (isomers I) of  $FeVS_{1-3}^{-}$  clusters are localized in a p orbital on a S atom. The partial charge distribution on the NBO/HOMO localized sites of each cluster anion (considering the "charge effect") is probably responsible for the trend of their first VDEs. The "charge effect" on the NBO/HOMO localized site is suggested not to be an essential factor that affects the first VDEs for three metal atom containing Fe/V/S (FeV<sub>2</sub>S<sub>3</sub><sup>-</sup> and Fe<sub>2</sub>VS<sub>3</sub><sup>-</sup>) clusters. With different Fe/V ratios for these clusters, the ground state structures are different: structure and steric effect differences are responsible for their different first VDEs and properties. Two types of structural isomers are found for  $Fe_m V_n S_{m+n}^-$  (m + n = 4, m > 0, n > 0) clusters: tower structure and cubic structure isomers. For the FeV<sub>3</sub>S<sub>4</sub> cluster anion, only the tower structure isomer is observed. Both tower and cubic structure isomers are observed for Fe<sub>2</sub>V<sub>2</sub>S<sub>4</sub><sup>-</sup> and Fe<sub>3</sub>VS<sub>4</sub><sup>-</sup> clusters. The metal ratio in these four metal atom containing Fe/V/S cluster anions is probably the main factor that affects the structure properties for these Fe/V/S clusters. The first VDEs for tower like isomers are generally higher than those for cubic like isomers of  $\operatorname{Fe}_{m} V_{n} S_{m+n}^{-}$  (m+n=4, m>0, n>0) clusters. Their first VDEs are reported and suggested to be understood through: (1) NBO/HOMO distributions, (2) structures (steric effects), and

(3) partial charge numbers on the NBO/HOMOs localized sites (the "charge effect").

EBEs regarding excited state transitions for related Fe/V/S are calculated employing OVGF and TDDFT approaches at a TZVP basis set level. The OVGF approach is generally better for higher transition energy theoretical studies of Fe/V/S cluster anions. The experimental and theoretical results of these Fe/V/S cluster anions are compared with their related pure Fe/S cluster anions. The electron distribution (wave function) properties of the NBO/HOMO are suggested to be an essential factor for understanding and comparing the different first VDEs of Fe/V/S cluster anions to those of pure Fe/S cluster anions.

# Conflicts of interest

**PCCP** 

There are no conflicts to declare.

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