



Cite this: *Phys. Chem. Chem. Phys.*,
2022, 24, 16576

Rearrangement and decomposition pathways of bare and hydrogenated molybdenum oxysulfides in the gas phase†

Marco Pritzi,^a Tobias F. Pascher,^a Marie-Luise Grutza,^b Philipp Kurz,^b
Milan Ončák^{*a} and Martin K. Beyer^{*a}

Molybdenum sulfides and molybdenum oxysulfides are considered a promising and cheap alternative to platinum as a catalyst for the hydrogen evolution reaction (HER). To better understand possible rearrangements during catalyst activation, we perform collision induced dissociation experiments in the gas phase with eight different molybdenum oxysulfides, namely $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$, $[\text{Mo}_2\text{O}_2\text{S}_6]^-$, $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$, $[\text{Mo}_2\text{O}_2\text{S}_5]^-$, $[\text{Mo}_2\text{O}_2\text{S}_4]^-$, $[\text{HMo}_2\text{O}_2\text{S}_6]^-$, $[\text{HMo}_2\text{O}_2\text{S}_5]^-$ and $[\text{HMo}_2\text{O}_2\text{S}_4]^-$, on a Fourier transform ion cyclotron resonance (FT-ICR) mass spectrometer. We identify fragmentation channels of the molybdenum oxysulfides and their interconnections. Together with quantum chemical calculations, the results show that $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is a particularly stable species against further dissociation, which is reached from all starting species with relatively low collision energies. Most interestingly, H atom loss is the only fragmentation channel observed for $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ at low collision energies, which relates to potential HER activity, since two such H atom binding sites on a surface may act together to release H_2 . The calculations reveal that multiple isomers are often very close in energy, especially for the hydrogenated species, *i.e.*, atomic hydrogen can bind at various sites of the clusters. S_2 groups play a decisive role in hydrogen adsorption. These are further features with potential relevance for HER catalysis.

Received 11th March 2022,
Accepted 23rd June 2022

DOI: 10.1039/d2cp01189a

rscl.li/pccp

Introduction

Hydrogen plays a major role in sustainable energy economy schemes,¹ *e.g.* for the storage of renewable electricity from solar or wind farms or for transportation, either directly or *via* power-to-gas processes.² Water electrolysis is crucial for these plans, and both half-cell reactions, the oxygen evolution reaction (OER) and the hydrogen evolution reaction (HER), must be optimized to increase efficiency and reduce costs.^{3,4} The standard HER catalyst under acidic conditions, where the highest efficiencies are obtained, is platinum.⁵ Given the limited availability of this element in the Earth's crust, large-scale electrolyzers often rely on more abundant catalyst materials which only work under alkaline conditions,⁶ which reduces the energy efficiency of the process.

Molybdenum sulfides⁷ are extensively investigated as a promising alternative for platinum.⁸ After electrochemical

reduction, they catalyze the HER in acidic solution.⁷ $[\text{Mo}_3\text{S}_{13}]^{2-}$ clusters have been synthesized to optimize the number of edge sites in an effort to increase the number of active sites for HER catalysis,⁹ which are also suitable for visible light driven HER.¹⁰ Also amorphous molybdenum sulfides MoS_{2+x} , $x < 1$, show high catalytic activity.^{11–14} Replacing terminal disulfides by halogenides results in $[\text{Mo}_3\text{S}_7\text{Cl}_6]^{2-}$ and $[\text{Mo}_3\text{S}_7\text{Br}_6]^{2-}$, which have significantly reduced catalytic activity, illustrating the importance of terminal S_2 units in HER catalysis.¹⁰

Gas phase ion chemistry offers unique ways to study elementary steps of homogeneous as well as heterogeneous catalysis at a molecular level.^{15–19} Reactions of small cationic iron sulfide clusters with small alkanes elucidated the oxidative power of these compounds.²⁰ Also molybdenum chalcogenide cations were investigated with gas-phase techniques.²¹ Jarrold, Raghavachari and co-workers have focused on molybdenum oxide and sulphide cluster reactivity,^{22–26} with studies involving H_2 and H_2O bearing relevance for HER catalysis.^{24,27–31} We have shown that cationic molybdenum clusters readily react with dimethyl disulfide, which reflects the high sulfur affinity of molybdenum.³²

$[\text{Mo}_3\text{S}_{13}]^{2-}$ species are particularly attractive for mass spectrometric studies since the ions and their protonated counterparts are readily transferred to the gas phase by electrospray

^a Institut für Ionenphysik und Angewandte Physik, Universität Innsbruck, Technikerstrasse 25, 6020 Innsbruck, Austria. E-mail: milan.oncak@uibk.ac.at, martin.beyer@uibk.ac.at

^b Institut für Anorganische und Analytische Chemie, Albert-Ludwigs-Universität Freiburg, Albertstrasse 21, 79104 Freiburg, Germany

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d2cp01189a>



ionization.³³ Their stability, decomposition pathways and chemical rearrangements were studied extensively by collision induced dissociation in combination with quantum chemistry.³³ Terminal disulfide was identified as the protonation site by infrared multiple photon dissociation spectroscopy of $[\text{HMo}_3\text{S}_{13}]^-$, again guided by quantum chemistry.³⁴

Since the synthesis of molybdenum sulfide cluster compounds starts from the heptamolybdate, it is not surprising that a variety of oxysulfides can be obtained by slightly modified synthesis routes. These also exhibit HER catalytic activity^{35,36} and have been shown to be capable of catalyst self-healing.³⁷

In the gas phase, we recently probed protonation of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ in reactions with organic acids, which proceeds *via* Coulomb explosion into the protonated species and the deprotonated acid anion.³⁸ The fragmentation pathways of molybdenum oxide clusters in the gas phase provide important additional information.³⁹ To obtain insight into catalytic properties of molybdenum oxysulfides, we investigate here their structure, flexibility and chemical reactivity as well as hydrogen adsorption/desorption through collision-induced dissociation of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$, $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ and their fragments, as well as quantum chemical calculations. The results suggest that these species represent potential reactive intermediates in HER catalysis.

Experimental and computational details

The experiments were performed on a Bruker APEX Qe FT-ICR mass spectrometer, equipped with a Nanobay Console, a 9.4 T superconducting magnet and an Apollo Dual Source II.⁴⁰ The ions are brought to the gas phase by electrospray ionization (ESI⁺) of a ~ 0.3 mM solution of the tetramethylammonium salt of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ in acetonitrile (ACN) and transferred into the ICR cell. Isotopically enriched $[(\text{CH}_3)_4\text{N}]_2[\text{Mo}_2\text{O}_2\text{S}_6]$ was prepared from $^{92}\text{Mo}_2\text{O}_3$ (STB Isotope Germany GmbH), following the procedure established by Müller *et al.*,^{41–43} with the modifications described in detail before.³⁸

The ultra-high vacuum region of the ICR cell has a base pressure in the low 10^{-9} mbar range. For sustained off-resonance irradiation (SORI)⁴⁴ collision induced dissociation (CID), the collision gas argon is introduced *via* a leak valve at a constant pressure in the 10^{-8} mbar range. SORI-CID was performed with a fixed frequency offset of 500 Hz and an irradiation pulse length of 1 s. The amplitude of the SORI excitation is controlled *via* the parameter “SORI power” in % values in the control software Compass 3.0.0 of the instrument. The acquisition of multiple mass spectra yields fragmentation curves as a function of gradually increased SORI power, which provide qualitative information on the relative stability of the different ions.

The high mass resolving power afforded by the 9.4 T magnet proved advantageous for the study of oxysulfides, as the mass of O_2 and S differs by only 0.01776 amu. We were able to distinguish between ions where one S atom is replaced by O_2 ,

for example $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ and $[\text{Mo}_2\text{O}_2\text{S}_6]^-$, even with the collision gas background which reduces the coherence time of the transient ion signal during detection and thus the mass resolution.

The mass spectrum obtained from ESI⁺ of the sample solution contains numerous peaks, as shown in Fig. S1 (ESI⁺). Based on their exact mass, we identified different molybdenum oxysulfides of interest in the mass spectrum, namely $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$, $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$, $[\text{Mo}_2\text{O}_2\text{S}_6]^-$, $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ and $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. To obtain protonated species, we added 5% formic acid to the sample solution, which yielded $[\text{H}^{92}\text{Mo}_2\text{O}_2\text{S}_6]^-$, $[\text{H}^{92}\text{Mo}_2\text{O}_2\text{S}_5]^-$ and $[\text{H}^{92}\text{Mo}_2\text{O}_2\text{S}_4]^-$. SORI-CID experiments were performed on these eight molybdenum oxysulfide species. Prior to SORI excitation, the ions are mass-selected by ejecting all unwanted ions from the cell *via* broadband frequency sweep and single-frequency shot excitation.

The geometries of all precursor and fragment ions were optimized at the B3LYP/def2TZVP level of theory using the D3 dispersion as suggested by Grimme *et al.*,⁴⁵ which performed well for molybdenum sulfides.³³ Candidate structures were derived from known crystal structure data, modified by educated guess. From our previous studies, we know that molybdenum oxysulfides tend to be low-spin compounds, therefore only the two lowest spin multiplicities, namely singlet and triplet or doublet and quartet, were considered; other than singlet and doublet spin multiplicities are given explicitly in the respective tables and figures; all spin multiplicities are given in the ESI⁺. Spin contamination in open-shell species was mostly below 4% before spin projection. Among ten structures with higher spin contamination, there are $[\text{MoOS}_3]^-$ and the least stable $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ isomer found, two structures of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ with a loosely bound S_2 and six structures of $[\text{H}_2\text{Mo}_2\text{O}_2\text{S}_6]$ or $[\text{Mo}_2\text{O}_2\text{S}_6]$. To estimate the induced error, we performed benchmark calculations for S_2 dissociation from $[\text{Mo}_2\text{O}_2\text{S}_6]^-$ using the BLYP functional without Hartree–Fock exchange (Table S2, ESI⁺). Reaction energies are calculated from electronic energies corrected for zero-point energy, without thermal corrections.

For evaluation of electron affinities of doubly charged species, a larger basis set is required. Therefore, we performed further calculations using the aug-cc-pVDZ basis set for sulfur and oxygen, and aug-cc-pVDZ-PP for Mo, as found on the Basis Set Exchange Website,⁴⁶ for these species. The stability of the wave function was verified in all calculations. The connection of the transition state (with a single imaginary frequency) with the local minima was verified, small structural changes (*e.g.*, rotation of an OH group) were ignored. However, intrinsic reaction coordinate (IRC) calculations failed to start in many cases because of the shallow shape of the potential energy surface. In those cases, a small offset of the transition state geometry was generated in both directions along the displacement vector of the vibrational mode carrying the imaginary frequency, followed by optimization. The Gaussian 16 program package was used for all calculations.⁴⁷ Cartesian coordinates and electronic energies of the optimized structures are available in the ESI⁺.



Results and discussion

$[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$

The mass-selected spectrum of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ that serves as starting point of the SORI-CID experiment is given as an example in Fig. S2 (ESI†) while Fig. S3 (ESI†) shows a typical decomposition mass spectrum at 0.5% SORI power. The full fragmentation curve is illustrated in Fig. 1(a). The fragmentation curve shows significant intensity of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ at 0% SORI power, which indicates that thermal collisions and residual kinetic excitation of some ions from the mass selection procedure is sufficient to induce electron detachment. Subsequent S_2 loss leads to $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. Direct S_2^- loss is not likely, as S_2^- was not observed in the mass spectrum. In a potential Coulomb explosion of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ into $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ and S_2^- , the latter would carry most of the kinetic energy release due to momentum conservation, and we cannot rule out that all S_2^- escapes from the cell. However, since $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ only becomes the dominant fragment above a SORI power of about 0.3%, where the $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ curve drops in intensity, the sequence of electron detachment followed by S_2 loss at increasing energies is more plausible.

$[\text{Mo}_2\text{O}_2\text{S}_4]^-$ undergoes splitting into $[\text{MoOS}_2]^-$ and, most likely, neutral $[\text{MoOS}_2]$. The isobaric $[\text{Mo}_2\text{O}_2\text{S}_4]^{2-}$ structure can be ruled out, based on the late appearance of the peak in the SORI-CID fragmentation curve. It should be noted that secondary fragments are mostly formed in the same collision event as the primary fragment. At high enough collision energies, the first neutral fragment is lost, taking away a small amount of kinetic and internal energy in addition to its binding energy. The major part of the excess energy, however, remains as internal energy of the fragment with the higher density of states, in this case the charged molybdenum oxysulfide fragment. Once dissociation has taken place, however, the SORI

irradiation is too far off resonance to yield appreciable collision energies for the fragment ions. Of course, the kinetic energy of the larger fragment remaining after the first collision and dissociation event can be converted into internal energy in subsequent collisions with argon, but this process is likely playing a minor role.

In Fig. 1(a), at around 0.5% SORI power a decrease of the $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ intensity is observed, while three other fragments rise. This suggests that the three fragments are formed by fragmentation of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. To verify this hypothesis, SORI-CID of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ was performed.

$[\text{Mo}_2\text{O}_2\text{S}_6]^-$

The fragmentation curve of the singly charged ion $[\text{Mo}_2\text{O}_2\text{S}_6]^-$, shown in Fig. 1(b), documents a sharp increase in the $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ signal starting at 0% SORI power, consistent with the sequential electron detachment and S_2 loss mechanism discussed above. At 0.8% SORI power, two other fragments arise, most likely secondary fragments of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. The $[\text{MoS}_3]^-$ ion observed before is missing here, probably due to the higher noise level. Comparison of threshold energies in Fig. 1(a) and (b) is difficult since the kinetic energy of ions excited in an FT-ICR instrument depends on the square of the charge, *i.e.* significantly higher collision energies are reached for doubly charged ions compared to singly charged at nominally identical SORI power.

$[\text{Mo}_2\text{O}_2\text{S}_4]^-$

$[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is obtained in high yields from ESI,† yet no fragment is present at 0% as shown in Fig. 2(a). This underlines the considerable stability of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. The fragments $[\text{MoOS}_2]^-$, $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ and $[\text{MoS}_3]^-$ emerge at similar thresholds, confirming that $[\text{MoS}_3]^-$ is a direct fragment of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. At higher SORI power more fragments arise, namely $[\text{MoO}_2\text{S}]^-$, $[\text{MoOS}]^-$ and $[\text{MoS}_2]^-$ in the order of their appearance. It is interesting to note that all but one fragment contain only one molybdenum atom.

$[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ and $[\text{Mo}_2\text{O}_2\text{S}_5]^-$

The intensity of $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ was very weak and accompanied by two other peaks at the same nominal mass. Therefore, proper isolation and a complete SORI-CID experiment was not possible with $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$. Nevertheless, we were able to identify electron detachment as the first fragmentation step. Fig. S4 (ESI†) shows the mass spectrum of mass-selected $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ exposed to collisions with argon at 0% SORI power. Electron detachment is proceeding efficiently, leading to significant intensity of $[\text{Mo}_2\text{O}_2\text{S}_5]^-$. This suggests that electron detachment in this case is even easier than for $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$. A 1000-fold zoom of the m/z axis reveals that the peak at m/z 187.83 actually consists of three separated peaks. The ion of interest $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ is assigned to the center peak, found at m/z 187.831. The peak at m/z 187.823 corresponds to either $[\text{Mo}_2\text{S}_6]^{2-}$ or $[\text{MoS}_3]^-$, and the peak at m/z 187.840 is due to $[\text{Mo}_2\text{O}_4\text{S}_4]^{2-}$ or $[\text{MoO}_2\text{S}_2]^-$. At a SORI power of 0.2% the $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ peak is already significantly higher than the

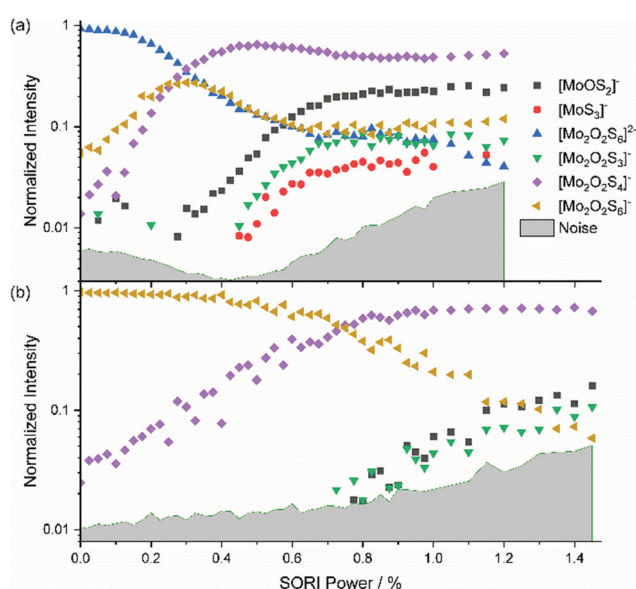


Fig. 1 Fragmentation curve obtained from the SORI-CID experiment of (a) $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and (b) $[\text{Mo}_2\text{O}_2\text{S}_6]^-$.



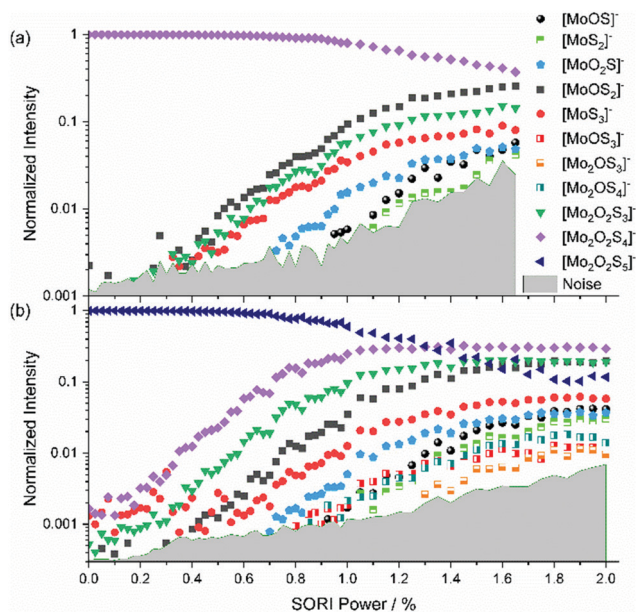


Fig. 2 SORI-CID fragmentation curve of (a) $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ and (b) $[\text{Mo}_2\text{O}_2\text{S}_5]^-$.

precursor, Fig. S5 (ESI[†]). We did not observe the corresponding side peaks for $[\text{Mo}_2\text{O}_2\text{S}_5]^-$, suggesting that the side peaks of $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ are singly charged ions, *i.e.*, $[\text{MoS}_3]^-$ and $[\text{MoO}_2\text{S}_2]^-$. Also $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is observed in small amounts.

We were able to isolate the singly charged species with considerably higher intensity than the doubly charged one, with no side peak issues. This made SORI-CID feasible, see Fig. 2(b). The first fragment observed in the fragmentation curve of $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ is again $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. The fact that this structure has been seen in every fragmentation curve so far further hints at its high stability against further dissociation. Next in the fragmentation curve, the six fragments of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ can be seen again, namely $[\text{Mo}_2\text{O}_2\text{S}_3]^-$, $[\text{MoOS}_2]^-$, $[\text{MoS}_3]^-$, $[\text{MoO}_2\text{S}]^-$, $[\text{MoOS}]^-$ and $[\text{MoS}_2]^-$. However, the order of appearance of $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ and $[\text{MoOS}_2]^-$ is reversed, with the former arising first. This suggests that the precursor $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ also undergoes direct fragmentation into $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ by loss of S_2 . Around 1.0% SORI power, three additional fragments appear which were not seen in any other experiment, namely $[\text{MoOS}_3]^-$, $[\text{MoOS}_4]^-$ and $[\text{MoOS}_5]^-$. These ions are most likely direct fragments of the precursor $[\text{Mo}_2\text{O}_2\text{S}_5]^-$, implying loss of MoOS_2 , SO and S_2O , respectively.

Protonated species $[\text{HMo}_2\text{O}_2\text{S}_6]^-$, $[\text{HMo}_2\text{O}_2\text{S}_5]^-$ and $[\text{HMo}_2\text{O}_2\text{S}_4]^-$

Addition of 5% formic acid to the sample solution led to the appearance of protonated species in the mass spectrum. The first experiment discussed here is the SORI-CID of $[\text{HMo}_2\text{O}_2\text{S}_6]^-$, with the fragmentation curve shown in Fig. 3(a). Similar to $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$, S_2 loss has the lowest threshold, leading to $[\text{HMo}_2\text{O}_2\text{S}_4]^-$. The next fragment is $[\text{Mo}_2\text{O}_2\text{S}_4]^-$, which appears at around 0.3% SORI power in the mass spectrum. It is presumably not a direct fragment of the precursor

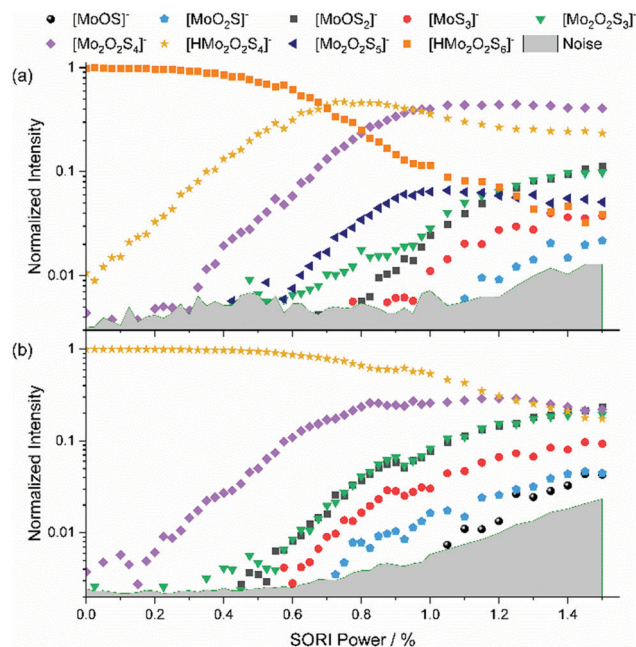


Fig. 3 (a) SORI-CID fragmentation curve of (a) $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ and (b) $[\text{HMo}_2\text{O}_2\text{S}_4]^-$.

but a fragment of $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ as the intensity of the latter decreases above 0.6% SORI power. This suggests that a hydrogen atom can be removed from $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ without destroying the $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ structure. SORI-CID of $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ is discussed below to validate this interpretation. Interestingly, beginning at a SORI power of about 0.6%, $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ is observed. This suggests that here the loss of HS is more favorable than the loss of S or H. After that, four fragments of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ appear, where $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ is formed as a secondary fragment of both $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ and $[\text{Mo}_2\text{O}_2\text{S}_4]^-$, as discussed previously.

We were also able to perform a SORI-CID experiment with $[\text{HMo}_2\text{O}_2\text{S}_5]^-$, shown in Fig. S6 (ESI[†]). Here, $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is formed *via* loss of HS. Fragments of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ again follow at higher SORI power. SORI-CID of $[\text{HMo}_2\text{O}_2\text{S}_4]^-$, Fig. 3(b), confirms the interpretation of the $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ experiment. As expected, hydrogen atom loss is the first, and over a wide range dominant, fragmentation channel. Again, the well-known fragments of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ arise above 0.4% SORI power, namely $[\text{Mo}_2\text{O}_2\text{S}_3]^-$, $[\text{MoOS}_2]^-$, $[\text{MoS}_3]^-$, $[\text{MoO}_2\text{S}]^-$, and $[\text{MoOS}]^-$.

Quantum chemical calculations

Optimized structures and relative energies of low-lying isomers for the most important species investigated are summarized in Fig. 4. In most cases, several isomers are found within 0.2 eV of the lowest energy structure, especially for protonated species. For $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$, the lowest-lying structure found in our calculations corresponds closely to the crystal structure.^{36,48} The two molybdenum atoms are connected by two bridging sulfur atoms. Each molybdenum center is additionally bound to an

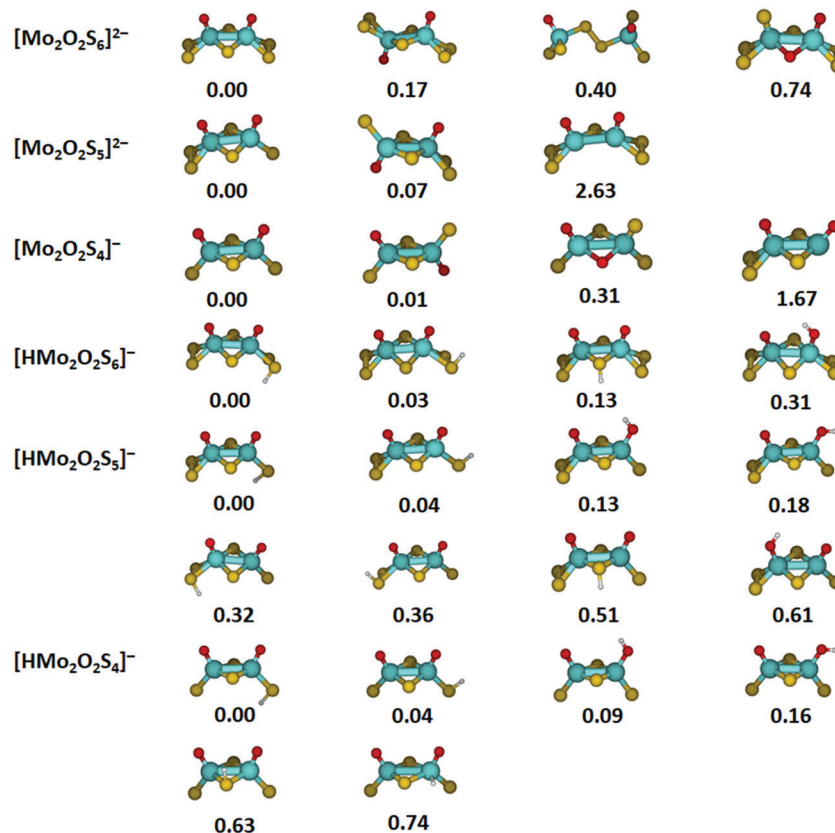


Fig. 4 Low-lying isomers for six of the eight studied molybdenum oxysulfides optimized at the B3LYP+D3/def2TZVP level. Zero-point corrected energies are given in eV. Color code: Mo – cyan, S – yellow, O – red, H – white.

oxygen atom and a terminal disulfide unit, overall in C_{2v} symmetry. However, turning one half of the molecule by 180 degrees results in a C_{2h} structure which lies only 0.17 eV higher. Isomers with a bridging disulfide unit or bridging oxygen atoms lie significantly higher in energy.

The energetically most favorable structure of $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$ is reached by removing a terminal sulfur atom from $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$. The loss of a bridging sulfur atom is significantly more demanding and can be ruled out. Removing a terminal sulfur atom from the second molybdenum center together with an electron leads to the lowest-energy structures of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$. For this ion, the energy difference between the C_{2v} and C_{2h} structure is reduced to 0.01 eV. Structures with two non-equivalent molybdenum centers are energetically not competitive.

Fig. 5(a) and (b) shows the isomerization pathway between the C_{2v} and C_{2h} structures of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and $[\text{Mo}_2\text{O}_2\text{S}_4]^-$, respectively. The predicted pathway for $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ includes an exchange of a bridging sulfur atom with an oxygen atom followed by a transfer of a terminal sulfur atom and returning of the bridging oxygen atom to the terminal position. For $[\text{Mo}_2\text{O}_2\text{S}_4]^-$, a similar pathway is obtained, without the need of sulfur exchange on the terminal groups as no S_2 unit is present. Still, the oxygen transfer reactions in both clusters are energetically demanding, requiring more than 1.6 eV.

For the protonated species, the protonation site and the orientation of the S–H or O–H bond have a small effect on the

energetics, corresponding to the dispersed distribution of the negative charge across all sulfur and oxygen atoms. In all cases, the energetically favorable protonation site is a terminal sulfur atom with the hydrogen atom pointing away from the nearest oxygen atom. However, changing the orientation of the S–H bond towards the oxygen atom is only 0.03–0.04 eV higher in energy for the three ions.

Protonation of a bridging sulfur atom lies by 0.13 eV higher than terminal disulfide for $[\text{HMo}_2\text{O}_2\text{S}_6]^-$, but the energy difference increases significantly if single terminal sulfur atoms are available. In this case, also the neighboring oxygen atoms are good proton acceptors, only about 0.1 eV above the preferred structures for $[\text{HMo}_2\text{O}_2\text{S}_5]^-$ and $[\text{HMo}_2\text{O}_2\text{S}_4]^-$. The affinity of the oxygen atoms towards a proton depends critically on the coordination of the molybdenum center with single sulfur atoms vs. terminal disulfide units. Protonation of a molybdenum atom seems generally unfeasible, lying 0.74 eV above the minimum for $[\text{HMo}_2\text{O}_2\text{S}_4]^-$.

As mentioned above, the fragmentation of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ to $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ presumably happens in two steps, with electron detachment followed by loss of S_2 . To reach a favorable $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ structure, however, the two sulfur atoms must come from the two terminal disulfide units. The lowest energy pathway for S_2 elimination from $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ that was found is depicted in Fig. 5(c). First, a terminal disulfide group rotates, reaching LM_9 at 0.68 eV. Then, the other terminal disulfide also



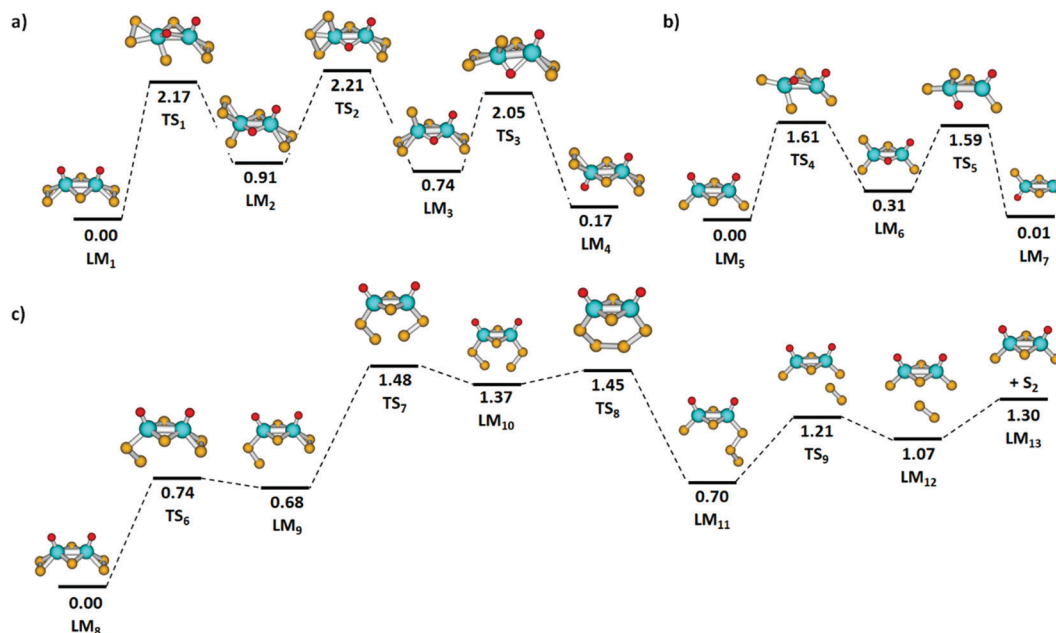


Fig. 5 Reaction pathways for isomerization reactions in (a) $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and (b) $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ clusters. (c) Simplified reaction profile from $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ to $[\text{Mo}_2\text{O}_2\text{S}_4]^- + \text{S}_2$. Calculated at the B3LYP+D3/def2TZVP level; relative energies are given in eV, local minima (LM) and transition states (TS) are denoted. Color code: Mo – cyan, S – yellow, O – red, H – white.

rotates so that two sulfur atoms of the disulfide units can form a bond, LM₁₀. The strain involved in this rearrangement leads to the highest barrier TS₇ along the reaction path, 1.48 eV. Passing through TS₈, the newly formed trisulfur unit rotates away from the molybdenum atoms to reach LM₁₁ and dissociates S₂ via TS₉. For comparison, the direct dissociation of a terminal disulfide unit would require at least 2.97 eV in energy without accounting for any transition state or rearrangement.

The fragmentation pathway shown in Fig. 5(c) requires considerably less energy.

Additional quantum chemical calculations were performed for all charged fragments observed and the corresponding neutrals. This results in a concise picture of the decomposition of the studied species, summarized in Fig. 6 for major pathways along with Table S1 and Fig. S7 (ESI†) that include also minor fragmentation channels. The energies reported in the figure

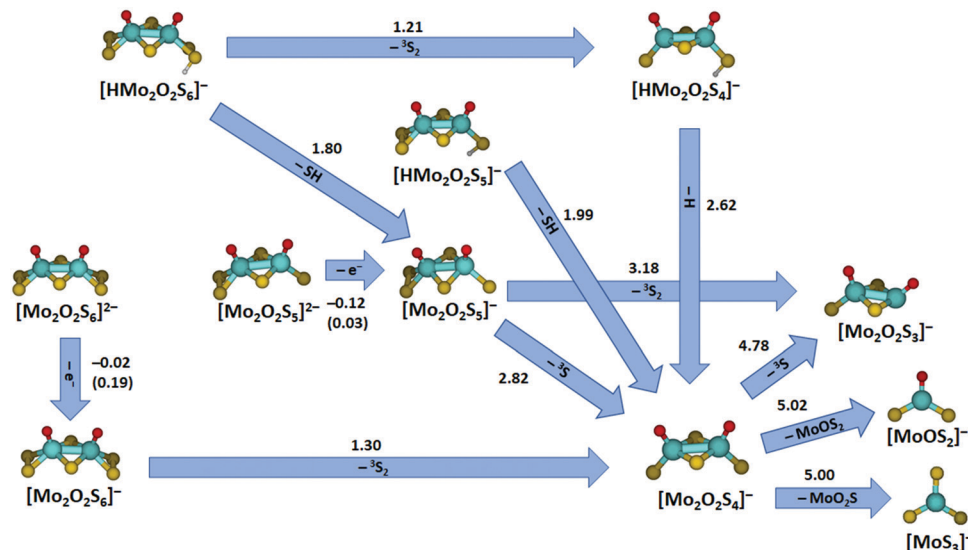


Fig. 6 Major fragmentation pathways of the molybdenum oxysulfides studied by SORI-CID along with reaction energies at the B3LYP+D3/def2TZVP level; for electron detachment, values calculated with aug-cc-pVDZ(O,S), aug-cc-pVDZ-PP(Mo) basis sets are given in parentheses. Reaction energies are given in eV. For all minor fragmentation paths, see Fig. S7 and Table S1 (ESI†). Color code: Mo – cyan, S – yellow, O – red, H – white. Other than singlet or doublet spin multiplicities are given explicitly.



compare reactant and product energies without considering any higher-lying transition states involved in the fragmentation. However, the reactions represent mostly direct dissociation reactions or require rearrangement of S₂ or O groups (similar to Fig. 5) or hydrogen atom transfer (see below), which involve transition states that most likely lie below or close to the respective dissociation energies. The doubly charged species [Mo₂O₂S₆]^{2−} and [Mo₂O₂S₅]^{2−} undergo electron detachment

with low energetic thresholds in the experiment. With the basis set used for most calculations, electron detachment is slightly exothermic in both cases, −0.02 eV for [Mo₂O₂S₆]^{2−} and −0.12 eV for [Mo₂O₂S₅]^{2−}. Using a larger basis set, aug-cc-pVDZ on O, S and aug-cc-pVDZ-PP on Mo, the electron detachment energies increase to 0.19 eV and 0.03 eV, respectively. In addition to the binding energy, however, the doubly charged structures are stabilized by a Coulomb barrier against electron

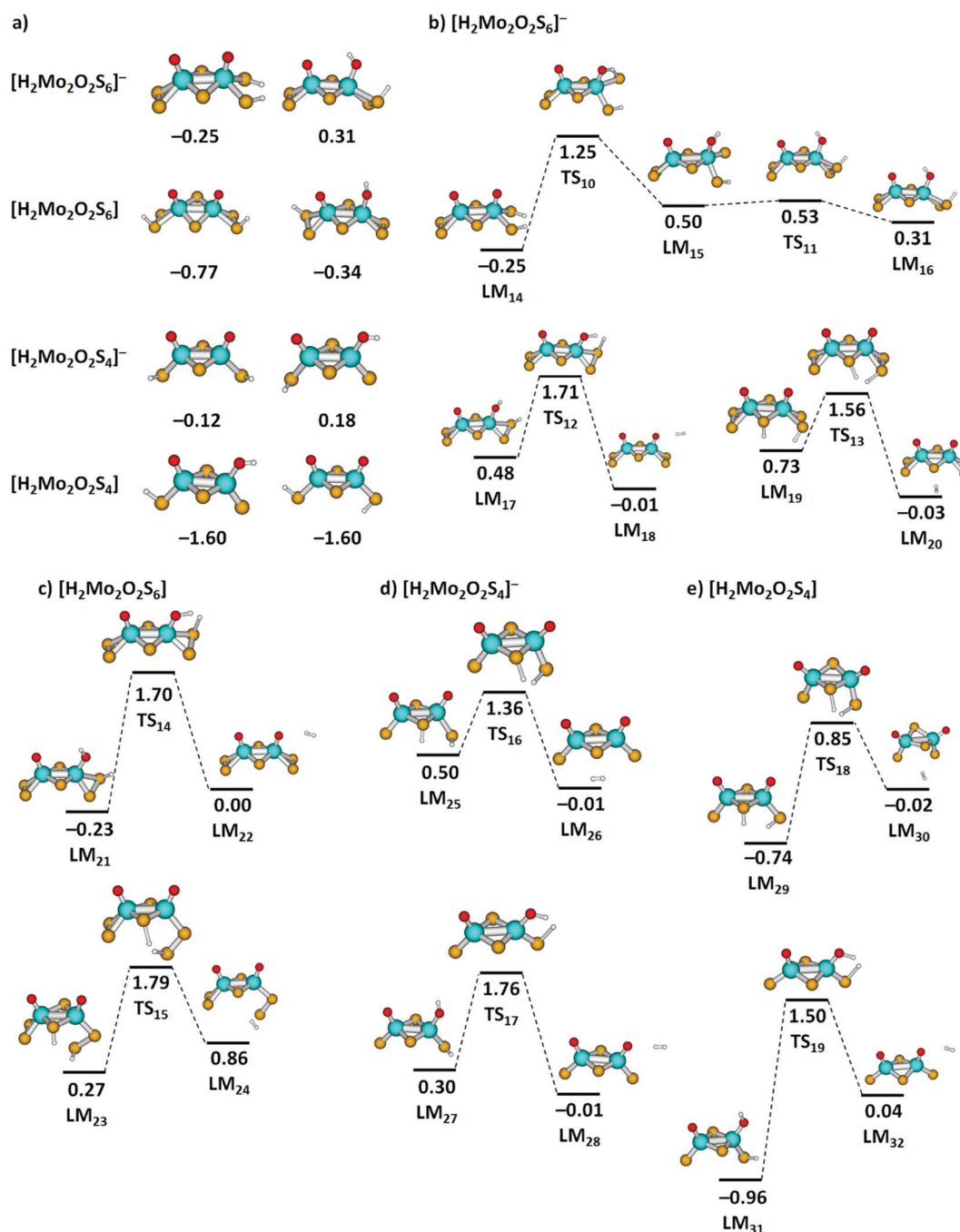


Fig. 7 (a) Two most stable located minima of [H₂Mo₂O₂S₆]^{2−}, [H₂Mo₂O₂S₆], [H₂Mo₂O₂S₄]^{2−}, and [H₂Mo₂O₂S₄] clusters, further structures are shown in Fig. S8 (ESI†). (b)–(e) Reaction pathways for H₂ dissociation from [H₂Mo₂O₂S₆]^{2−}, [H₂Mo₂O₂S₆], [H₂Mo₂O₂S₄]^{2−}, and [H₂Mo₂O₂S₄] clusters. For [H₂Mo₂O₂S₆]^{2−}, a simplified reaction pathway for hydrogen transfer is also shown. Calculated at the B3LYP+D3/def2TZVP level; relative energies are given in eV with respect to the limit of [cluster + H₂]; local minima (LM) and transition states (TS) are denoted. Color code: Mo – cyan, S – yellow, O – red, H – white.

detachment.⁴⁹ The calculations also explain why we were not able to obtain any signal for the doubly charged $[\text{Mo}_2\text{O}_2\text{S}_4]^{2-}$. Electron detachment for this ion is with -0.39 eV even more exothermic at the B3LYP/def2TZVP theory level, and still -0.26 eV with the bigger basis set defined above. This also shifts the Coulomb barrier down so that it can be overcome in low energy collisions, assisted by tunneling.

Turning to the singly charged species, many different fragments were seen in SORI-CID of $[\text{Mo}_2\text{O}_2\text{S}_5]^-$, five of which have been assigned as direct fragments. The energies needed for the fragmentation correspond to the order of appearance in the fragmentation curve, consistent with this interpretation. The first and energetically least demanding fragment is $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ with 2.82 eV, followed by $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ with 3.18 eV. Other fragments observed in minor intensity are $[\text{Mo}_2\text{OS}_4]^-$ with 3.94 eV, $[\text{MoOS}_3]^-$ with 4.23 eV, and $[\text{Mo}_2\text{OS}_3]^-$ with 5.79 eV.

Loss of S_2 from $[\text{Mo}_2\text{O}_2\text{S}_6]^-$ to form $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is endothermic by 1.30 eV. In agreement with experiment, fragmentation of $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ requires higher energies, see Table S1 (ESI†). The first three fragments appear at very similar SORI power, perfectly reflected in the calculations: formation of $[\text{Mo}_2\text{O}_2\text{S}_3]^-$ is endothermic with 4.78 eV, $[\text{MoOS}_2]^-$ with 5.02 eV and $[\text{MoS}_3]^-$ with 5.00 eV. The next fragments appear at significantly higher collision energies, corresponding to the increased energy demand forming $[\text{MoO}_2\text{S}]^-$ with 5.13 eV, $[\text{MoOS}]^-$ with 7.51 eV and $[\text{MoS}_2]^-$ with 7.53 eV, see Table S1 (ESI†).

The protonated species complete the picture. $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ decays by H atom loss to $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ with 2.62 eV, quite demanding compared to SH loss from $[\text{HMo}_2\text{O}_2\text{S}_5]^-$ with 1.99 eV. The SH loss from $[\text{HMo}_2\text{O}_2\text{S}_4]^-$, however, would be quite expensive with 3.77 eV. For comparison, the dissociation of a hydrogen atom from $[\text{HMo}_2\text{O}_2\text{S}_5]^-$ requires 2.81 eV while the dissociation of a terminal sulfur atom with 3.01 eV is even more expensive, in agreement with the experimental observation.

The first fragmentation channel of $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ in the experiment corresponding to S_2 elimination is calculated to be 1.21 eV endothermic. However, a barrier similar to the one shown in Fig. 5(c) may be expected. At higher SORI power, $[\text{Mo}_2\text{O}_2\text{S}_5]^-$ appeared in the mass spectrum, most likely resulting from HS loss, which is 1.80 eV endothermic according to calculations and becomes entropically favored compared to a tight transition state involved in S_2 elimination, like TS_7 in Fig. 5(b). Interestingly, hydrogen atom loss from $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ requires 2.52 eV, 0.10 eV less than from $[\text{HMo}_2\text{O}_2\text{S}_4]^-$. Since two energetically lower lying channels are available for $[\text{HMo}_2\text{O}_2\text{S}_6]^-$, H atom loss is not observed in this case. It is intriguing to note that the S–H bond dissociation energy is 55% of the bond dissociation energy of H_2 , 4.52 eV. This means that the elimination of two H atoms from such neighboring S–H groups with concomitant formation of H_2 in a catalytic environment would be 0.52 eV endothermic, which lies in the thermally accessible regime.

The fragmentation of molybdenum oxysulfides bears some similarities with $[\text{Mo}_3\text{S}_{13}]^{2-}$, $[\text{HMo}_3\text{S}_{13}]^-$ and $[\text{H}_3\text{Mo}_3\text{S}_{13}]^+$ studied previously.³³ In both cases the energetically favorable protonation/hydrogenation site was found to be a terminal sulfur atom or a terminal disulfide unit. Also, in both cases,

H_xS_y elimination is primarily seen, and all molybdenum atoms are retained in the ionic fragment at low collision energies. A completely new fragmentation channel in the present study is the loss of a single hydrogen atom for $[\text{HMo}_2\text{O}_2\text{S}_4]^-$. While $[\text{Mo}_3\text{S}_{13}]^{2-}$ do not undergo electron detachment during CID, this pathway was readily observed for the doubly charged molybdenum oxysulfides.

Parallels can also be drawn to our proton transfer reactivity study of $[\text{Mo}_2\text{O}_2\text{S}_6]^{2-}$ and $[\text{Mo}_2\text{O}_2\text{S}_5]^{2-}$.³⁸ A terminal sulfur atom was identified as the protonation site, and a Coulomb barrier hindered charge separation reactions.^{38,50–54}

The thermochemistry of H atom loss prompted us to calculate H_2 elimination pathways from clusters containing two H atoms. Although we do not have any experimental data for these species, we found it worthwhile to gain more insight into potential final steps of HER. We calculated a series of $[\text{H}_2\text{Mo}_2\text{O}_2\text{S}_6]^-$, $[\text{H}_2\text{Mo}_2\text{O}_2\text{S}_4]^-$, $[\text{H}_2\text{Mo}_2\text{O}_2\text{S}_6]$, and $[\text{H}_2\text{Mo}_2\text{O}_2\text{S}_4]$ clusters, see Fig. 7(a) for low-lying isomers. For all investigated species, formation of two S–H groups leads to a very stable cluster, while clusters with one S–H and one O–H groups are close in energy (see Fig. S8, ESI† for further structures), with S_2 groups playing a prominent role in H_2 adsorption. For the anionic clusters, H_2 adsorption is only mildly exothermic, with a very low adsorption energy. In neutral clusters, H_2 is bound much stronger. This indicates that in an electrochemical environment, addition of an electron will promote H_2 elimination from these clusters.

For selected H_2 dissociation pathways from the clusters starting either from two S–H groups or one S–H and one O–H group, we see consistently lower barriers for H_2 dissociation for the anionic species. Particularly low barriers are obtained for anionic species with H atoms attached to a bridging and a terminal sulfur unit, with 0.83 eV and 0.86 eV from LM_{19} via TS_{13} and LM_{25} via TS_{16} , respectively.

Conclusion

SORI-CID experiments on eight different molybdenum oxysulfide species showed that $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is a common fragment of all larger molybdenum oxysulfides. While $[\text{Mo}_2\text{O}_2\text{S}_6]^-$ and $[\text{HMo}_2\text{O}_2\text{S}_6]^-$ readily lose an S_2 unit after some rearrangement at low SORI power, $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ shows fragmentation only at relatively high collision energies. Overall, the OMoS_2MoO backbone is retained up to high collision energies, while quantum chemical calculations confirm that all decompositions of the $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ structure require at least 5 eV, indicating a particular stability of this species with respect to further dissociation reactions. $[\text{Mo}_2\text{O}_2\text{S}_4]^-$ is the smallest unit with two fully saturated Mo centers, and may be representative of the structures formed during catalyst activation in electrocatalytic environments. Electron detachment is almost thermoneutral across the doubly charged species, consistent with the catalytically relevant redox properties of these species. Electrons can readily be accepted or donated. An interesting fragmentation channel of $[\text{HMo}_2\text{O}_2\text{S}_4]^-$ is the dissociation of an H atom, leaving the



[Mo₂O₂S₄][−] structure intact. Calculations reveal that the hydrogen atom can favorably bind to a variety of almost isoenergetic sites. We also show that S₂ groups play a prominent role as a protonation site. In the calculations, it was possible to add a second H atom to the protonated species, and we investigated H₂ elimination from neutral as well as negatively charged clusters. Activation energies for H₂ formation below 1 eV were obtained in anionic species, with H atoms attached to a bridging and a terminal sulfur unit. Facile electron attachment and removal, a variety of H atom adsorption sites and low barriers for H₂ elimination are all promising properties of a HER catalyst.

Conflicts of interest

There are no conflicts of interest to declare.

Acknowledgements

This project is supported by the Klima- und Energiefonds, administered by the FFG under Project No. 853639, and put into effect in the framework of the programme “ENERGIE DER ZUKUNFT”. We thank Dr Aristeidis Baloglou for preliminary work and Reviewer #2 for providing us with several calculated structures of [Mo₂O₂S₄][−]. The computational results presented have been achieved using the HPC infrastructure LEO of the University of Innsbruck.

References

- 1 J. A. Turner, *Science*, 2004, **305**, 972.
- 2 C. Wulf, J. Linssen and P. Zapp, *Hydrogen Supply Chains*, Elsevier, 2018, pp. 309–345.
- 3 M. G. Walter, E. L. Warren, J. R. McKone, S. W. Boettcher, Q. Mi, E. A. Santori and N. S. Lewis, *Chem. Rev.*, 2010, **110**, 6446.
- 4 Z. Shi, X. Wang, J. Ge, C. Liu and W. Xing, *Nanoscale*, 2020, **12**, 13249.
- 5 J. K. Nørskov, T. Bligaard, A. Logadottir, J. R. Kitchin, J. G. Chen, S. Pandelov and U. Stimming, *J. Electrochem. Soc.*, 2005, **152**, J23.
- 6 M. Bodner, A. Hofer and V. Hacker, *Wiley Interdiscip. Rev.: Energy Environ.*, 2015, **4**, 365.
- 7 H. I. Karunadasa, E. Montalvo, Y. Sun, M. Majda, J. R. Long and C. J. Chang, *Science*, 2012, **335**, 698.
- 8 P. C. K. Vesborg, B. Seger and I. Chorkendorff, *J. Phys. Chem. Lett.*, 2015, **6**, 951.
- 9 J. Kibsgaard, T. F. Jaramillo and F. Besenbacher, *Nat. Chem.*, 2014, **6**, 248.
- 10 M. Dave, A. Rajagopal, M. Damm-Ruttensperger, B. Schwarz, F. Nägele, L. Daccache, D. Fantauzzi, T. Jacob and C. Streb, *Sustainable Energy Fuels*, 2018, **2**, 1020.
- 11 J. D. Benck, Z. Chen, L. Y. Kuritzky, A. J. Forman and T. F. Jaramillo, *ACS Catal.*, 2012, **2**, 1916.
- 12 C. G. Morales-Guio and X. Hu, *Acc. Chem. Res.*, 2014, **47**, 2671.
- 13 D. Merki, S. Fierro, H. Vrubel and X. Hu, *Chem. Sci.*, 2011, **2**, 1262.
- 14 P. D. Tran, T. V. Tran, M. Orio, S. Torelli, Q. D. Truong, K. Nayuki, Y. Sasaki, S. Y. Chiam, R. Yi, I. Honma, J. Barber and V. Artero, *Nat. Mater.*, 2016, **15**, 640–646.
- 15 D. K. Bohme and H. Schwarz, *Angew. Chem., Int. Ed.*, 2005, **44**, 2336.
- 16 S. M. Lang and T. M. Bernhardt, *Phys. Chem. Chem. Phys.*, 2012, **14**, 9255.
- 17 T. F. Pascher, M. Ončák, C. van der Linde and M. K. Beyer, *ChemPhysChem*, 2019, **20**, 1420.
- 18 T. F. Pascher, M. Ončák, C. van der Linde and M. K. Beyer, *ChemistryOpen*, 2019, **8**, 1453–1459.
- 19 E. Barwa, T. F. Pascher, M. Ončák, C. van der Linde and M. K. Beyer, *Angew. Chem., Int. Ed.*, 2020, **59**, 7467–7471.
- 20 S. M. Lang, S. Zhou and H. Schwarz, *Phys. Chem. Chem. Phys.*, 2017, **19**, 8055.
- 21 I. Kretzschmar, A. Fiedler, J. N. Harvey, D. Schröder and H. Schwarz, *J. Phys. Chem. A*, 1997, **101**, 6252.
- 22 N. J. Mayhall, E. L. Becher, A. Chowdhury and K. Raghavachari, *J. Phys. Chem. A*, 2011, **115**, 2291.
- 23 N. J. Mayhall, E. L. Becher III., A. Chowdhury and K. Raghavachari, *J. Phys. Chem. A*, 2011, **115**, 2291.
- 24 A. K. Gupta, J. E. Topolski, K. A. Nickson, C. C. Jarrold and K. Raghavachari, *J. Phys. Chem. A*, 2019, **123**, 7261.
- 25 J. L. Mason, A. K. Gupta, A. J. McMahon, C. N. Folluo, K. Raghavachari and C. C. Jarrold, *J. Chem. Phys.*, 2020, **152**, 54301.
- 26 S. E. Waller and C. C. Jarrold, *J. Phys. Chem. A*, 2014, **118**, 8493.
- 27 J. E. Topolski, A. K. Gupta, K. A. Nickson, K. Raghavachari and C. C. Jarrold, *Int. J. Mass Spectrom.*, 2018, **434**, 193.
- 28 A. Saha and K. Raghavachari, *J. Chem. Phys.*, 2013, **139**, 204301.
- 29 M. Ray, S. E. Waller, A. Saha, K. Raghavachari and C. C. Jarrold, *J. Chem. Phys.*, 2014, **141**, 104310.
- 30 M. Ray, A. Saha and K. Raghavachari, *Phys. Chem. Chem. Phys.*, 2016, **18**, 25687.
- 31 M. Ray, R. N. Schauggaard, J. E. Topolski, J. O. Kafader, K. Raghavachari and C. C. Jarrold, *J. Phys. Chem. A*, 2018, **122**, 41.
- 32 A. Baloglou, M. Ončák, C. van der Linde and M. K. Beyer, *Top. Catal.*, 2018, **61**, 20.
- 33 A. Baloglou, M. Ončák, M.-L. Grutza, C. van der Linde, P. Kurz and M. K. Beyer, *J. Phys. Chem. C*, 2019, 8177.
- 34 A. Baloglou, M. Plattner, M. Ončák, M.-L. Grutza, P. Kurz and M. K. Beyer, *Angew. Chem., Int. Ed.*, 2021, **60**, 5074.
- 35 D. N. Chirdon, R. F. Lalisie, J. Sun, S. Zhang, B. R. Garrett, C. M. Hadad and Y. Wu, *SN Appl. Sci.*, 2020, **2**, 889.
- 36 J. McAllister, N. A. G. Bandeira, J. C. McGlynn, A. Y. Ganin, Y.-F. Song, C. Bo and H. N. Miras, *Nat. Commun.*, 2019, **10**, 370.
- 37 J. Barros Barbosa, P. L. Taberna, V. Bourdon, I. C. Gerber, R. Poteau, A. Balocchi, X. Marie, J. Esvan, P. Puech,



- A. Barnabé, L. Da Gama Fernandes Vieira, I.-T. Moraru and J. Y. Chane-Ching, *Appl. Catal., B*, 2020, **278**, 119288.
- 38 A. Baloglou, M. Pritzi, T. F. Pascher, J. C. Hartmann, M.-L. Grutza, M. Ončák, P. Kurz and M. K. Beyer, *Int. J. Mass Spectrom.*, 2021, **464**, 116558.
- 39 M. Plattner, A. Baloglou, M. Ončák, C. van der Linde and M. K. Beyer, *J. Am. Soc. Mass Spectrom.*, 2019, **30**, 1946.
- 40 A. Herburger, C. van der Linde and M. K. Beyer, *Phys. Chem. Chem. Phys.*, 2017, **19**, 10786.
- 41 A. Müller, R. G. Bhattacharyya and B. Pfefferkorn, *Chem. Ber.*, 1979, **112**, 778.
- 42 A. Müller, E. Krickemeyer and U. Reinsch, *Z. Anorg. Allg. Chem.*, 1980, **470**, 35.
- 43 A. Müller, S. Sarkar, R. G. Bhattacharyya, S. Pohl and M. Dartmann, *Angew. Chem., Int. Ed. Engl.*, 1978, **17**, 535.
- 44 J. W. Gauthier, T. R. Trautman and D. B. Jacobson, *Anal. Chim. Acta*, 1991, **246**, 211.
- 45 S. Grimme, J. Antony, S. Ehrlich and H. Krieg, *J. Chem. Phys.*, 2010, **132**, 154104.
- 46 B. P. Pritchard, D. Altarawy, B. Didier, T. D. Gibson and T. L. Windus, *J. Chem. Inf. Model.*, 2019, **59**, 4814.
- 47 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, *Gaussian 16 Revision A.03*, 2016.
- 48 W. Rittner, A. Müller, A. Neumann, W. Bätther and R. C. Sharma, *Angew. Chem., Int. Ed. Engl.*, 1979, **18**, 530.
- 49 P. Weis, O. Hampe, S. Gilb and M. M. Kappes, *Chem. Phys. Lett.*, 2000, **321**, 426.
- 50 A. Révész, B. Sztáray, D. Schröder, K. Franzreb, J. Fiser, S. D. Price and J. Roithová, *Phys. Chem. Chem. Phys.*, 2009, **11**, 6192.
- 51 M. K. Beyer and R. B. Metz, *J. Phys. Chem. A*, 2003, **107**, 1760.
- 52 M. Beyer, E. R. Williams and V. E. Bondybey, *J. Am. Chem. Soc.*, 1999, **121**, 1565.
- 53 D. Schröder and H. Schwarz, *J. Phys. Chem. A*, 1999, **103**, 7385.
- 54 B. L. J. Poad, N. D. Reed, C. S. Hansen, A. J. Trevitt, S. J. Blanksby, E. G. Mackay, M. S. Sherburn, B. Chan and L. Radom, *Chem. Sci.*, 2016, **7**, 6245.

