





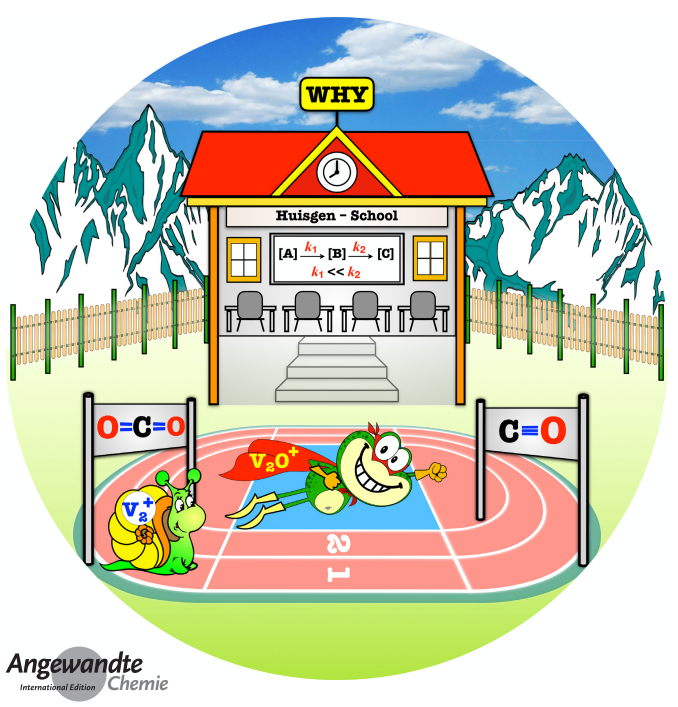
CO₂ Activation

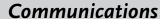
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Counter-Intuitive Gas-Phase Reactivities of [V2] and [V₂O]⁺ towards CO₂ Reduction: Insight from Electronic **Structure Calculations**

Jilai Li,* Caiyun Geng, Thomas Weiske, and Helmut Schwarz*

In memory of Rolf Huisgen









Abstract: $[V_2O]^+$ remains "invisible" in the thermal gas-phase reaction of bare $[V_2]^+$ with CO_2 giving rise to $[V_2O_2]^+$; this is because the $[V_2O]^+$ intermediate is being consumed more than 230 times faster than it is generated. However, the fleeting existence of $[V_2O]^+$ and its involvement in the $[V_2]^+ \rightarrow [V_2O_2]^+$ chemistry are demonstrated by a cross-over labeling experiment with a 1:1 mixture of $C^{16}O_2/C^{18}O_2$, generating the product ions $[V_2^{16}O_2]^+$, $[V_2^{16}O^{18}O]^+$, and $[V_2^{18}O_2]^+$ in a 1:2:1 ratio. Density functional theory (DFT) calculations help to understand the remarkable and unexpected reactivity differences of $[V_2]^+$ versus $[V_2O]^+$ towards CO_2 .

Kolf Huisgen has shaped physical organic chemistry in a nearly unparalleled fashion.^[1] His extensive, seminal work on, for example, 1,3-dipolar cycloadditions $^{[1b]}$ forms a pillar of contemporary mechanistic organic chemistry, not to speak of their role in the synthesis of heterocyclic compounds. By using the then available experimental repertoire, the Huisgen school unraveled the finest mechanistic details of numerous chemical processes, and their work gives testament for chemist generations to come.[2]

A familiar kinetic scheme of a multi-step process is depicted in Equation (1), and one problem here concerns the identification of "intermediate" [B]. This holds true in particular when the rate constants k_1 and k_2 differ significantly, that is, $k_2 \gg k_1$.

$$[A] \xrightarrow{k_1} [B] \xrightarrow{k_2} [C]$$

A typical example is provided by the gas-phase chemistry of the [Pd]⁺/CH₃I couple which gives rise to the formation of [Pd(CH₂I)]⁺.^[3] In this case, the product ion is not formed by the entropically favored direct cleavage of the C-H bond of CH₃I.^[3a] Rather, in the first step a short lived [Pd(CH₃)]⁺ intermediate is generated, which reacts further with CH3I in a very fast metathesis process to form [Pd(CH₂I)]⁺ and CH₄.[3b] Recently, in the context of combined experimental/ computational studies on the timely topic of CO₂ activation, [4] we came across a similar, unexpected situation in the thermal gas-phase reactions of bare diatomic $[V_2]^+$ with CO_2 .

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The cluster $[V_2]^+$ was formed by supersonic expansion of helium into a vanadium plasma generated by laser ablation/ ionization of a vanadium disk using a Nd:YAG laser, operating at 532 nm inside the external cluster source of a Fourier-transform ion cyclotron resonance (FT-ICR) mass spectrometer as described previously (for details, see the Supporting Information).^[5] A fraction of the ion population was then guided by a static ion optical system into the ICR cell. Next, in a sequence of pulses, argon was admitted to the ICR cell such that the ions collide on average about 1×10^5 times with argon. This procedure ensures thermalization of hot ions and quenching of excited electronic states. After mass-selection of the thermalized $[V_2]^+$ ions, they were reacted with CO2 or C18O2 at a constant pressure low enough to ensure single collision conditions. The elementary compositions of the charged species have been confirmed by exact mass measurements using high resolution mass spectrometry.

The result of the reaction of $[V_2]^+$ with CO_2 at a partial pressure of 1.0×10^{-7} mbar and a reaction time of 10 s inside the ion trap of the FT-ICR machine is shown in Figure 1 b. In addition to the signal of the starting material $[V_2]^+$ (signal A, Figure 1), a signal **D** appears, which has been assigned the formula [V₂O₂]⁺ by exact mass measurements; also, very small amounts of atomic $[V]^+$ are present (signal **B**). Formally, the formation of $[V_2O_2]^+$ corresponds to the loss of one carbon atom. If $C^{18}O_2$ acts as a reaction partner, signal **D** from Figure 1b is shifted by four mass units (Figure 1c; signal **E**). Clearly, oxygen atom transfer (OAT) from CO2 to the vanadium cluster takes place. However, since a carbon atom represents an extremely poor leaving group, the suggestion was raised as to whether the formation of $[V_2O_2]^+$ from the [V₂]⁺/CO₂ couple is the result of a multi-step process, although the spectra in Figure 1b,c do not exhibit any clue for the involvement of conceivable intermediates. The assumption for the operation of a multi-step process is corroborated by a crossover-labeling experiment using a 1:1 mixture of $C^{16}O_2$ and $C^{18}O_2$: The appearance of $[V_2^{16}O^{18}O]^+$ (signal F in Figure 1 d) together with the other two isotopomers in a 1:2:1 ratio verifies this hypothesis. According to Equations (2) and (3) and in line with Figure 1 d, one has to conclude that $[V_2]^+$ reacts consecutively with two CO_2 molecules via an OAT by the release of one CO molecule each time; the monoxide $[V_2O]^+$ serves as a short-lived intermediate (Scheme 1). Further oxidation processes according to Equation (4) in Scheme 1 do not occur at ambient temperature.

In the $[V_2]^+/CO_2$ couple, the supposed intermediate $[V_2O]^+$ can only be detected experimentally as an extremely weak signal under highly optimized conditions of both pressure and reaction time (signal G in Figure 1e). However and fortunately, $[V_2O]^+$ can be synthesized independently and in sufficient quantities in the external ion source by adding traces of O₂ to the helium buffer gas (Figure 1 f). As shown in Figure 1 g, the so-formed $[V_2^{16}O]^+$ indeed reacts with $C^{18}O_2$ to produce $[V_2^{16}O^{18}O]^+$ by the expulsion of $C^{18}O$. The collision efficiency ϕ of that reaction is close to 100%. The high reactivity of $[V_2O]^+$ explains why, $[V_2O]^+$ from the reaction of $[V_2]^+$ with CO_2 , remains elusive. As soon as the monoxide has



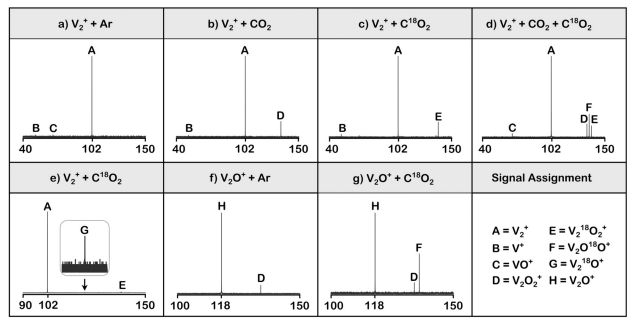


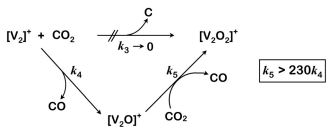
Figure 1. Representative mass spectra: a)–e) are for the thermal reactions of $[V_2]^+$ with a) Ar at 1.0×10^{-7} mbar, b) $C^{16}O_2$ at 1.0×10^{-7} mbar, c) $C^{18}O_2$ at 1.0×10^{-7} mbar, and d) a 1:1 mixture of $C^{16}O_2$ and $C^{18}O_2$ at 2.0×10^{-7} mbar after a reaction time of 10 s, respectively; e) $C^{18}O_2$ at 1.0×10^{-7} mbar and a reaction time of 1s; f) and g) represent the thermal reactions of $[V_2O]^+$ with f) Ar at 4.0×10^{-9} mbar, g) $C^{18}O_2$ at 4.0×10^{-9} mbar, after a reaction time of 2s, respectively. All x-axes are scaled in m/z, and the y-axes are normalized relative ion abundances.

formed within the ICR cell, it immediately reacts with another CO_2 molecule. The high reactivity of the $[V_2\mathrm{O}]^+$ intermediate, generated from the $[V_2]^+/\mathrm{CO}_2$ couple, is further enhanced due to the substantial exothermicity of the process (see below) and the fact that, owing to the absence of a heat bath, part if not most of the excess energy is stored in the ion by partitioning the energy released between $[V_2\mathrm{O}]^+$ and CO . To distinguish possible reactions of the charged particles with omnipresent background gases from those of the selected substrate, spectra with argon were recorded as well. As shown in Figure 1 a,f side reactions with background gases like trace amounts of oxygen are limited to the generation of small quantities of products typical for reactions of $[V_2]^+$ with O_2 , $^{[9e]}$ which are more pronounced in $[V_2\mathrm{O}]^+$, since O_2 has been added to the He buffer gas.

$$[V_2]^+ + CO_2 \longrightarrow [V_2O]^+ + CO$$
 (2)

$$[V_2O]^+ + CO_2 \longrightarrow [V_2O_2]^+ + CO$$
 (3)

$$[V_2O_2]^+ + nCO_2 \longrightarrow [V_2O_{2+n}]^+ + nCO$$
 (4)

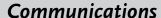


Scheme 1. Multi-step activation of CO_2 by $[V_2]^+$ with $[V_2O]^+$ serving as a fleeting intermediate.

The rate constants $k_4([V_2]^+/CO_2)$ and $k_5([V_2O]^+/CO_2)$ for reactions shown in Equations (2) and (3) in Scheme 1 were measured and they amount to 3.8×10^{-12} (efficiency $\phi = 0.5\%$) and 8.8×10^{-10} cm³ molecule⁻¹ s⁻¹ ($\phi = 1.15$), respectively; these numbers translate to a ratio of $k_5/k_4 > 230$. Owing to an uncertainty in the determination of the absolute pressure of CO_2 , an error of $\pm 30\%$ is associated with the rate measurements.^[5b] For the relative rate constants, however, the error is much smaller, typically around $\pm 5\%$.

The considerable reactivity difference of $[V_2]^+$ versus $[V_2O]^+$ is unexpected and counter-intuitive on the ground that the reactions in Equations (2) and (3) constitute redox processes in which electrons are transferred from the metal-containing cluster ions to the CO_2 molecule. Therefore, one would expect that $[V_2]^+$ reduces CO_2 to CO more readily than the already oxidized and electron-depleted $[V_2O]^+$. Consequently, k_4 should be larger than k_5 ; this is not the case. Why does the reactivity of $[V_2]^+$ and $[V_2O]^+$ not reflect their respective oxidation states? Further, does $[V_2O]^+$ react mechanistically the same way with CO_2 as $[V_2]^+$? These questions have been addressed by density functional theory (DFT) calculations.

Earlier theoretical work had already shown that a description of vanadium dimers constitutes a notoriously difficult case to be dealt with by DFT calculations. This dilemma holds true as well for $[V_2O_2]^+$. While for $[V_2O_2]^+$ the gasphase IR spectrum suggests a planar, four-membered V-O-V-O ring structure of C_s symmetry, an unambiguous assignment of the electronic ground state $(^2A' \text{ or } ^6A')$ remained unresolved at the DFT levels employed. The problems are compounded by the observation that the gas-phase chemistry of vanadium ions of the married by a two- or multi-state







reactivity scenario. [10] As to the bare $[V_2]^+$ cluster, due to its multireference character^[7,11] we applied the ZORA-NEVPT2(9e,12o)/ZORA-def2-QZVPP Scheme (Table S1 in the Supporting Information).^[12] According to these high-level calculations, the ground state of $[V_2]^+$ corresponds to ${}^4\Sigma_{\sigma}^-$. This assignment agrees well with experimental data and other, less sophisticated calculations.^[13] The lowest excited state of $[V_2]^+$ is a doublet $(^2\Delta)$ and exceeds the ground state by only about 3 kJ mol⁻¹ (Table S2). Therefore, it is very likely that under the given experimental conditions a mixture of two electronic states, ${}^{2,4}[V_2]^+$, is present, in which the quartet may dominate. We mention in passing that low-energy excited states of the neutral V2 dimer were identified by Hübner and Himmel using matrix isolation spectroscopy.^[11] To construct a simplified potential energy surface (PES) for the [V₂]⁺/[V₂O]⁺/CO₂ systems, extensive quantum chemical calculations were performed at the ZORA-M06L/ZORA-def2-QZVPP//M06L/def2-TZVP level of theory. Electronic structure calculations using the multi-reference ZORA-NEVPT2 scheme are not feasible technically. Rather, the M06L functional was chosen based on the exhaustive testing carried out by Truhlar and co-workers.^[7] This functional performed best also among all the density functionals we benchmarked (see Tables S2, S6 and S7). In addition, M06L delivers fairly good results for the electronic states of $[V_2O]^+$ and $[V_2O_2]^+$ (see Tables S3 and S4).

Figure 2 displays the PESs of the most favorable pathways for the reduction of CO₂ with vanadium ions having doublet spin multiplicity as well as selected structural parameters of stationary points along the reaction coordinate. This particular spin state was chosen on the ground that along the whole transformation $[V_2]^+ \rightarrow [V_2O_2]^+$ the spin multiplicity was conserved. It should be mentioned at this point, however, that because of the strongly correlated electrons, the energetic and geometric properties of some stationary points with different spin multiplicities are located fairly close to each other. Therefore, it is quite possible that there is a superposition of the doublet with the quartet state and vice versa, facilitating a multi-state reactivity scenario pending on the efficiency of spin-orbit coupling (see below for a further comment).^[10] However, our aim is not to provide a quantitatively correct description of the whole PESs; rather, we aim at offering a qualitative understanding of the amazing reactivity behavior in the $[V_2]^+/[V_2O]^+/CO_2$ systems. As the reactivity pattern proved to be quite complex, herein we limit ourselves to the main results. For further details, see Figures S2 and S3.

As displayed in Figure 2a, at the doublet surface the exothermic OAT^[14] in the system $[V_2]^+/CO_2$ commences with the barrier-free formation of an encounter complex ${\bf 1a}$ by coordinating the incoming ligand to only one of the two vanadium atoms in an end-on (η^1) mode. Subsequently, intermediate ${\bf 2a}$ is generated by surmounting transition state ${\bf TS1a}$, which is accompanied by a substantial elongation of the V–V bond from about 1.6 Å to 2.5 Å; this bond length does not change that much further in the remaining steps. In contrast, significant structural reorganization happens for CO_2 at these stages. The initially linear substrate gets bent with an O-C-O angle of 141° in ${\bf 2a}$, and the C–O bond length increases from 1.17 to 1.27 Å; clearly, CO_2 is already markedly

activated in this initial step. Next, the complete scission of one C–O bond of the CO_2 moiety occurs along the route ${\bf 2a} \rightarrow {\bf TS2a} \rightarrow {\bf 3a}$. The reaction is completed by the release of CO to generate $^2[V_2O]^+$. The energetically submerged transition state ${\bf TS1a}$, as part of the reduction sequence of the first CO_2 molecule, is with $29~{\rm kJ\,mol}^{-1}$ located below the entrance asymptote and represents the rate-limiting step of the entire reaction.

As to the exothermic reaction of $[V_2O]^+$ with CO_2 and as displayed in Figure 2b, $^2[V_2O]^+$ is approached by carbon dioxide as well, to then also form an η^1 encounter complex ${\bf 1b}$; subsequently, this complex has sufficient internal energy to easily surmount transition state ${\bf TS1b}$ and generate ${\bf 2b}$. Next, the complete cleavage of the activated C–O bond takes place along the sequence ${\bf 2b} \to {\bf TS2b} \to {\bf 3b}$, and release of CO from ${\bf 3b}$ provides $^2[V_2O_2]^+$. In contrast to the $[V_2]^+/CO_2$ couple, here it is transition state ${\bf TS2b}$ ($-48~{\rm kJ}~{\rm mol}^{-1}$) in which the C–O bond of the CO_2 molecule is broken and which represents the rate-limiting step in the reduction of CO_2 to CO_2 .

At first glance, the reductions of the two CO₂ molecules by $[V_2]^+$ and $[V_2O]^+$ do not seem to differ too much in terms of structural features of the intermediates. So, why do they exhibit quite pronounced reactivity differences? As will be shown, these features can be attributed to both intrinsic structural and electronic properties. As suggested by a reviewer, one reason may be due to the spin situation. For the [V₂O]⁺/CO₂ couple the OAT is both exothermic and spinallowed (Figure 2b); the whole reaction is confined to the doublet surface. This must not necessarily be the case for the $[V_2]^+/CO_2$ system. While for electronically excited ${}^2[V_2]^+$ the reaction with CO₂ is also both exothermic and spin-allowed (Figure 2a), for the ground state ${}^{4}[V_{2}]^{+}$ a quartet \rightarrow doublet spin change has to occur along the overall exothermic reaction coordinate (Table S5). If this electronic reorganization is inefficient, as has been reported for the analogous exothermic [V]+/CO₂ conversion, [9f] as well as for the [V]+/ N_2O couple, [9a] the reaction of $[V_2]^+$ with CO_2 will be slowed down as well. Further, while thermochemical arguments on the endo-versus exothermicity of a reaction, [7] based only on DFT calculations, may be considered with caution, the insensitivity of the [V2]+/CO2 reactivity to intentional excitation of the $[V_2]^+$ projectile, strongly suggests that process (2) is exothermic and that its much reduced reactivity is not due to a thermodynamic impediment.

Indeed, there are other factors to be considered to explain the much higher reactivity of the $[V_2O]^+/CO_2$ couple. Let us begin by looking at the charge distribution in the ionic actors. According to Figure 3a, the reactive vanadium atoms in ${}^2[V_2O]^+$ carry more positive charge as compared to ${}^2[V_2]^+$. It is therefore expected that $[V_2O]^+$ will act as a worse electron donor for the reduction of CO_2 compared to $[V_2]^+$. $[^{1.5]}$ However, this is not the whole truth. In fact, the presence of the bridging oxygen atom in $[V_2O]^+$ actually plays a supportive role in the CO_2 activation which overcompensates the unfavorable charge situation.

First, the presence of the oxygen atom induces a structural rearrangement of the active site, thus, providing a "preorganized" structure, which reduces substantially the energy

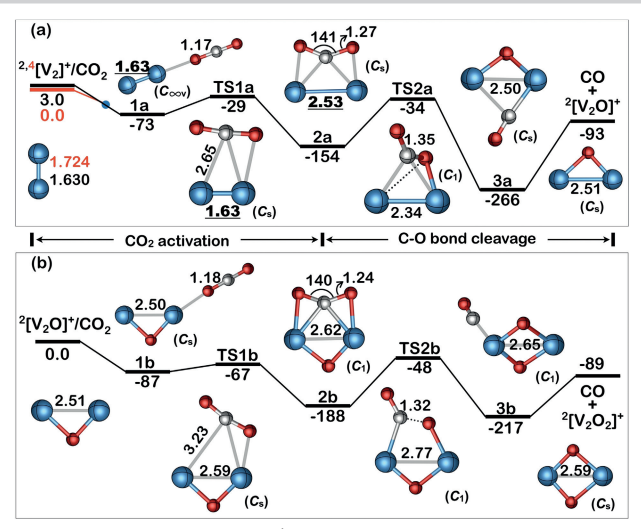


Figure 2. Schematic potential energy surfaces $(\Delta H_{298K} \text{ in kJ mol}^{-1})$ obtained at the ZORA-M06L/ZORA-def2-QZVPP//M06L/def2-TZVP level of theory for the reactions of a) $^2[V_2]^+$ and b) $^2[V_2Q]^+$ with CO₂, respectively. Key structures with selected geometric parameters and symmetries are also provided. Bond lengths [Å]; angles [°]. The relative energies of $^{2,4}[V_2]^+$ were obtained at the ZORA-NEVPT2(9e,12o)/ZORA-def2-QZVPP level of theory. For details, see the Supporting Information.

demand to reach transition state **TS1b**; this effect is well-known from enzymatic processes. [16] The calculated Wiberg Bond Index (WBI)[17] of 4.5 indicates a rather strong and short V–V bond for $[V_2]^+$ (1.63 Å); a much smaller value of the WBI of 1.2 and an elongated V–V bond of 2.53 Å holds true

for 2a (Figure 2 and Figure 3b). As a consequence, stretching of the V-V bond in $[V_2]^+$ in the step $1a \rightarrow 2a$ is quite energy-demanding (Figure 2a). In contrast, the corresponding transformation 1b to 2b in $[V_2O]^+$ requires much less energy as the V-V bond is already sufficiently elongated due to the presence of the oxygen bridge. Thus, $[V_2O]^+$ possesses a "prepared" structure which greatly enhances the reduction of CO_2 . In the remaining stages of the reaction there is no further dramatic change in the properties of the V-V bond, as indicated by the rather small oscillations of the WBI's around a value of 1. The moderate movements of the two vanadium atoms relative to each other do not require that much energy; rather, it is

this structural change of the "active" site which assists in lowering **TS1b** as compared with **TS1a**.

There is yet another aspect to be considered. In $^2[V_2O]^+$ the presence of an oxygen atom raises the energy level of the electron-donating d-orbitals. As shown in Figure 4, in **TS1 a** of

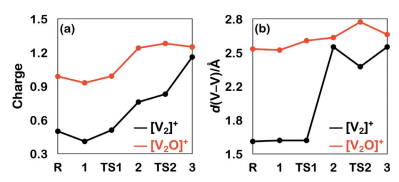


Figure 3. a) NBO-calculated charges at the vanadium atoms coordinated to CO_2 and b) development of the V–V bond length [Å] in the reactions of $^2[V_2]^+$ and $^2[V_2O]^+$ with CO_2 .





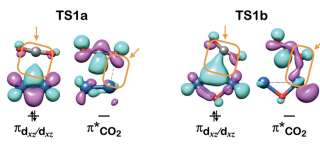


Figure 4. The frontier orbitals of the transition states of TS1 a,b from the doublet surface of the CO_2 activation. Arrows and rounded rectangles indicate the overlap regions between the doubly occupied π-electron-donating orbital of the metals and the lowest empty orbital of CO_2 .

 $[V_2]^+$ the overlap between the lobes of the doubly-occupied $\pi d_{xz}/d_{xz}$ -orbital of the metal center and the vacant π^* -orbital of the carbon atom of CO_2 has yet to be established. In contrast, for the early transition state $\mathbf{TS1b}$, the doubly-occupied $\pi d_{xz}/d_{xz}$ -orbital is already delocalized quite extensively to the carbon atom of CO_2 , and the root cause for this difference can be traced back to orbital energy differences. The introduction of an oxygen bridge raises the $\pi d_{xz}/d_{xz}$ -orbital energy from -10.5 eV in $[V_2]^+$ to -9.5 eV in $[V_2O]^+$; this greatly eases the transfer of the d-electrons into the empty π^* -orbital of CO_2 thus favoring the reduction of $[V_2O]^+$.

As shown repeatedly,^[16,18] the presence or absence of "prepared states" are crucial in controlling the structural, electronic and energetic features of transition states. While there exist quite a number of examples for this scenario in the area of gas-phase C–H bond activation,^[19] to our knowledge the present study describes for the first time this effect in the context of CO₂ activation.

In conclusion, in the present experimental/computational work we report and explain quite a few unexpected observations:

- 1) A fleeting $[V_2O]^+$ intermediate is involved in the multistep oxidation of $[V_2]^+$ by CO_2 .
- 2) The counter-intuitive oxidation behavior of [V₂]⁺ versus [V₂O]⁺ and the quite enormous reactivity differences of the two cluster ions have been traced back to the supportive role of the oxygen bridge in [V₂O]⁺. This cluster oxide, in contrast to bare [V₂]⁺, exhibits structural and electronic features which aid in the activation of CO₂. While thermochemical aspects are not likely to cause the relative "inertness" of [V₂]⁺, an inefficient quartet → doublet spin change in the [V₂]⁺/CO₂ couple may well matter in slowing down its reactivity.

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Conflict of interest

The authors declare no conflict of interest.

Keywords: CO_2 activation \cdot gas-phase reactions \cdot oxidation state \cdot quantum chemical calculations

- a) R. Huisgen, The Adventure Playground of Mechanisms and Novel Reactions (Profiles, Pathways, and Dreams), American Chemical Society, Washington DC, 1994; b) R. Huisgen, Angew. Chem. Int. Ed. Engl. 1968, 7, 321-328; Angew. Chem. 1968, 80, 329-337; c) R. Huisgen, J. Org. Chem. 1968, 33, 2291-2297; d) R. Huisgen, Angew. Chem. Int. Ed. Engl. 1963, 2, 633-645; Angew. Chem. 1963, 75, 742-754.
- [2] For an Essay on Huisgen's legacy, see: a) K. N. Houk, H.-U. Reissig, Chem 2019, 5, 2499-2505; Also, see: b) V. V. Rostovtsev, L. G. Green, V. V. Fokin, K. B. Sharpless, Angew. Chem. Int. Ed. 2002, 41, 2596-2599; Angew. Chem. 2002, 114, 2708-2711.
- [3] a) J. Schwarz, C. Heinemann, D. Schröder, H. Schwarz, J. Hrušák, Helv. Chim. Acta 1996, 79, 1-5; b) J. Schwarz, D. Schröder, H. Schwarz, C. Heinemann, J. Hrušák, Helv. Chim. Acta 1996, 79, 1110-1120.
- [4] For a Review on gas-phase aspects, see: H. Schwarz, Coord. Chem. Rev. 2017, 334, 112–123.
- [5] a) M. Engeser, T. Weiske, D. Schröder, H. Schwarz, J. Phys. Chem. A 2003, 107, 2855-2859; b) D. Schröder, H. Schwarz, D. E. Clemmer, Y. Chen, P. B. Armentrout, V. I. Baranov, D. K. Böhme, Int. J. Mass Spectrom. 1997, 161, 175-191; c) K. Eller, H. Schwarz, Int. J. Mass Spectrom. 1989, 93, 243-257.
- [6] a) G. Kummerlöwe, M. K. Beyer, Int. J. Mass Spectrom. 2005, 244, 84-90; b) T. Su, M. T. Bowers, J. Chem. Phys. 1973, 58, 3027-3037; c) M. T. Bowers, J. B. Laudenslager, J. Chem. Phys. 1972, 56, 4711-4712.
- [7] W. Zhang, D. G. Truhlar, M. Tang, J. Chem. Theory Comput. 2014, 10, 2399 – 2409.
- [8] a) R.-Z. Li, H.-G. Xu, G.-J. Cao, Y.-C. Zhao, W.-J. Zheng, *Chin. J. Chem. Phys.* 2011, 24, 572-579; b) K. R. Asmis, G. Meijer, M. Brümmer, C. Kaposta, G. Santambrogio, L. Wöste, J. Sauer, *J. Chem. Phys.* 2004, 120, 6461-6470; c) M. Calatayud, J. Andrés, A. Beltrán, *J. Phys. Chem. A* 2001, 105, 9760-9775.
- [9] For some typical examples, see: a) S. G. Ard, B. C. Sweeny, D. C. McDonald, A. A. Viggiano, N. S. Shuman, J. Phys. Chem. A 2020, 124, 30-38; b) Y. C. Chang, Y. Xu, C.-Y. Ng, Phys. Chem. Chem. Phys. 2019, 21, 6868-6877; c) H. Zhang, H. Wu, L. Geng, Y. Jia, M. Yang, Z. Luo, Phys. Chem. Chem. Phys. 2019, 21, 11234-11241, and references therein; d) G. K. Koyanagi, D. K. Bohme, J. Phys. Chem. A 2006, 110, 1232-1241; e) C. Rue, P. B. Armentrout, I. Kretzschmar, D. Schröder, J. N. Harvey, H. Schwarz, J. Chem. Phys. 1999, 110, 7858-7870; f) J. Xu, M. T. Rodgers, J. B. Griffin, P. B. Armentrout, J. Chem. Phys. 1998, 108, 9339-9350; g) M. R. Sievers, P. B. Armentrout, J. Chem. Phys. 1995, 102, 754-762; h) M. M. Kappes, R. H. Staley, J. Phys. Chem. 1981, 85, 942-944.
- [10] For a selection of Reviews, see: a) S. Shaik, *Isr. J. Chem.* 2020, DOI: 10.1002/ijch.202000002; b) D. Ricciarelli, L. Belpassi, J. N. Harvey, P. Belanzoni, *Chem. Eur. J.* 2020, 26, 3080-3089; c) D. Mandal, D. Mallick, S. Shaik, *Acc. Chem. Res.* 2018, 51, 107-117; d) T. Takayanagi, T. Nakatomi, *J. Comput. Chem.* 2018, 39, 1319-1326; e) J. N. Harvey, *Wiley Interdiscip. Rev.: Comput. Mol. Sci.* 2014, 4, 1-14; f) S. Shaik, H. Hirao, D. Kumar, *Acc.*







Chem. Res. 2007, 40, 532-542; g) F. Neese, T. Petrenko, D. Ganyushin, G. Olbrich, Coord. Chem. Rev. 2007, 251, 288-327; h) J. N. Harvey, Phys. Chem. Chem. Phys. 2007, 9, 331-343; i) S. Shaik, D. Kumar, S. P. de Visser, A. Altun, W. Thiel, Chem. Rev. 2005, 105, 2279-2328; j) H. Schwarz, Int. J. Mass Spectrom. 2004, 237, 75-105; k) R. Poli, J. N. Harvey, Chem. Soc. Rev. 2003, 32, 1-8; l) S. Shaik, S. P. de Visser, F. Ogliaro, H. Schwarz, D. Schröder, Curr. Opin. Chem. Biol. 2002, 6, 556-567; m) D. Schröder, S. S. Shaik, H. Schwarz, Acc. Chem. Res. 2000, 33, 139-145; n) S. Shaik, M. Filatov, D. Schröder, H. Schwarz, Chem. Eur. J. 1998, 4, 193-199; o) P. B. Armentrout, Science 1991, 251, 175-179; p) P. B. Armentrout, Annu. Rev. Phys. Chem. 1990, 41, 313-344.

- [11] O. Hübner, H.-J. Himmel, Phys. Chem. Chem. Phys. 2016, 18, 14667-14677.
- [12] a) C. Angeli, R. Cimiraglia, J.-P. Malrieu, J. Chem. Phys. 2002, 117, 9138-9153; b) C. Angeli, R. Cimiraglia, S. Evangelisti, T. Leininger, J. P. Malrieu, J. Chem. Phys. 2001, 114, 10252-10264; c) C. Angeli, R. Cimiraglia, J.-P. Malrieu, Chem. Phys. Lett. 2001, 350, 297-305.
- [13] a) D. S. Yang, A. M. James, D. M. Rayner, P. A. Hackett, J. Chem. Phys. 1995, 102, 3129-3134; b) A. M. James, P. Kowalczyk, E. Langlois, M. D. Campbell, A. Ogawa, B. Simard, J. Chem. Phys. 1994, 101, 4485-4495; c) L. M. Russon, S. A. Heidecke, M. K. Birke, J. Conceicao, M. D. Morse, P. B. Armentrout, J. Chem. Phys. 1994, 100, 4747-4755; d) B. Simard, A. James, P. Kowalczyk, R. Fournier, P. Hackett, Proc. SPIE-Int. Soc. Opt. Eng. 1994, 2124, 376-387.
- [14] a) Given the well-established problems with DFT-based energetics (see, for example, Ref. [7], a reviewer has raised the possibility that the reduced reactivity of [V₂]⁺ towards CO₂ is due to an unfavorable thermochemistry, that is, the formation of [V₂O]⁺ is either thermoneutral or slightly endothermic. This option has been tested by intentional kinetic excitation of [V₂]⁺ prior to its reaction with CO₂ (for technical details, see: M. Beyer, V. E. Bondybey, *Rapid Commun. Mass Spectrom.* 1997, 11, 1588–1591): Up to a time regime of 40 μs, neither the amount of the precursor ion [V₂]⁺ nor that of the product ion

 $[V_2O_2]^+$ are affected by kinetically exciting the $[V_2]^+$ projectile. At much longer excitation time, the intensity of the [V₂O₂]⁺ signal goes through a maximum while that of $[V_2]^+$ gets continuously weaker due to the ions' ejection from the ICR cell. This behavior corresponds to an exothermic reactivity scenario of the $[V_2]^+/CO_2$ couple and is further experimentally supported by the fact that at room temperature no formation of $[V_2]^+$ is observed in the reaction of $[V_2O]^+$ with CO, which represents the reversal of [Eq. (2)]. Since there is no oxygen exchange when $[V_2O]^+$ is exposed to $^{13}C^{18}O$, the energy of **TS2a** in Figure 2a must exceed that of the product pair [V2O]+/CO, confirming the results of the calculations. b) According to previous experimental work, [9d,g,h] the reaction of ground-state ⁵[V]⁺ with CO₂ to form ³[VO]⁺ and CO is exothermic and spinforbidden. For this process our M06L/DEF2-TZVP calculations predict a barrier of 74 kJ mol⁻¹ above the entrance channel; this explains why this redox process has not been observed using SIFT or ICR. $^{[9d,h]}$

- [15] T. Fan, X. Chen, Z. Lin, Chem. Commun. 2012, 48, 10808-10828.
- [16] A. J. T. Smith, R. Müller, M. D. Toscano, P. Kast, H. W. Hellinga, D. Hilvert, K. N. Houk, J. Am. Chem. Soc. 2008, 130, 15361– 15373.
- [17] K. B. Wiberg, Tetrahedron 1968, 24, 1083-1096.
- [18] a) W. Lai, C. Li, H. Chen, S. Shaik, Angew. Chem. Int. Ed. 2012, 51, 5556-5578; Angew. Chem. 2012, 124, 5652-5676; b) S. Ye, F. Neese, Proc. Natl. Acad. Sci. USA 2011, 108, 1228-1233; c) S. S. Shaik, J. Am. Chem. Soc. 1981, 103, 3692-3701.
- [19] a) C. Geng, J. Li, T. Weiske, H. Schwarz, Chem. Eur. J. 2019, 25, 12940–12945; b) J. Li, X.-N. Wu, M. Schlangen, S. Zhou, P. González-Navarrete, S. Tang, H. Schwarz, Angew. Chem. Int. Ed. 2015, 54, 5074–5078; Angew. Chem. 2015, 127, 5163–5167; c) H. Schwarz, Isr. J. Chem. 2014, 54, 1413–1431.

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