**Oxidative Addition of Dihydrogen to Divanadium in Solid Ne: Multiple-Bonded Triplet HVVH and Singlet V₂(μ-H)₂**

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**Abstract:** Dinuclear compounds of early transition metals with a high metal–metal bond order are of fundamental interest due to their intriguing bonding situation and of practical interest because of their potential involvement in catalytic processes. In this work, two isomers of V₂H₂ have been generated in solid Ne by the reaction between V₂ and H₂ and detected by infrared spectroscopy: the linear HVVH molecule (\(\Sigma^+\) ground state), which is the product of the spin-allowed reaction between V₂ (\(\Sigma^+\) ground state) and H₂, and a lower-energy, folded V₂(μ-H)₂ isomer (\(\Sigma^+\) ground state) with two bridging hydrogen atoms. Both isomers are characterized by metal–metal bonding with a high bond order; the orbital occupations point to quadruple bonding. Irradiation with ultraviolet light induces the transformation of linear HVVH to folded V₂(μ-H)₂, whereas irradiation with visible light initiates the reverse reaction.

**Introduction**

Many dinuclear complexes of early transition metals have been synthesized in which there are M₂ units that show metal-metal multiple bonding.\[^4\] Due to their electronic properties, such complexes show a unique chemical reactivity that makes them suitable for involvement in catalysis.\[^5\] Among these complexes, there are paddle-wheel complexes like V₂(DPhF)₂, (DPhF = N,N-diphenylformamidinate), showing short V–V distances of less than 200 pm with a triple bond between the metal atoms.\[^3,4\] To stabilize complexes with a strong bonding between the metal atoms, it is advisable to reduce the number of ligands as far as possible. However, due to the increased reactivity of the low-coordinated metals in such complexes, it is necessary to use bulky ligands that shield the metal core. The prime example is the dinuclear chromium complex \(\text{RCrCrR} (\text{R} = \text{C}_6\text{H}_4\text{-}2\text{-}2\text{,6-P}r_3\text{C})_2\) with a minimum number of ligands in a trans-bent structure, which even shows quintuple bonding between the two chromium centers with a Cr–Cr distance of only 183.5 pm.\[^6\] The electronic structure of \(\text{HCrCrH}_2\) and some other small model systems for this complex have been analyzed theoretically to confirm the presence of fivefold bonding and to understand the origin of the trans-bent alignment.\[^7,8\] However, in general, it is not clear to what extent the bulky ligands determine the structure. Therefore, it is in particular desirable to characterize, if possible, experimentally, systems like the prototypical HMMH molecules with terminally bonded hydrogen atoms and multiple metal–metal bonding. The matrix-isolation technique is a versatile means to stabilize and investigate such molecules by isolating them in noble-gas matrices.

An obvious way to generate HMMH molecules in inert-gas matrices is the reaction of matrix-isolated bare metal dimers with H₂ in noble-gas matrices. A larger number of bare transition-metal dimers has been generated in noble-gas matrices,\[^9\] and some studies showed that metal dimers exhibit a special reactivity. As to matrix isolation, an outstanding example is the reactivity of the titanium dimer. Ti₂ reacts with dinitrogen by formation of cyclic Ti₂(μ-N), breaking even the strong triple bond, whereas Ti atoms do not show a corresponding reaction.\[^10\] Another example for the special reactivity of dimers is the platinum dimer cation Pt₂⁺, which was shown to be able to activate ammonia in the gas phase (Pt₂⁺ + NH₃ → Pt₃NH²⁺ + H₂), whereas Pt⁺, Pt²⁺, and Pt³⁺ were not.\[^11\] However, work concerning the reaction of neutral transition-metal dimers with dihydrogen is scarce. There are many studies in which the reaction of evaporated transition metals M with H₂ has been investigated in matrices.\[^12\] Some of them also report products with the composition M₂H₂. The late-transition-metal compounds Rh₂H₂[\(^{13}\) and Pd₂H₂[\(^{14}\) have been reported to feature rhombic structures (bridging hydrogens), while the compounds HZnZnH, HCdCdH,\[^15\] and HHgHgH\[^16\] show linear structures. Furthermore, absorptions have been tentatively assigned to Mn₂H₂[\(^{17}\) HTHtH₃[\(^{18}\) HZnZrH[\(^{19}\) and HCrCrH[\(^{20}\) and the dihydrogen complexes Pd₂Pd(μ-H)₂ and Pt₂Pt(μ-H)₂[\(^{21}\) have also been observed.

In this work, we report on the generation and characterization of matrix-isolated V₂H₂ molecules. So far, the reaction of V₂ with H₂ has not been reported, only V atoms have been reacted with dihydrogen. Thermally evaporated V atoms in Kr and Ar matrices have been found to react with H₂ only upon irradiation (320 nm < λ < 380 nm) by formation of VH₂, whereas V atoms in their electronic ground state do not react with H₂[\(^{22}\)] V atoms generated by laser ablation, however, react with H₂ to give VH₂ in Ne and Ar matrices also without specific irradiation.\[^23\]

Also, for some transition-metal-cluster cations, among them the dimer cations, the reactivity with respect to D₂ has been studied,\[^24\] and it was observed that V₂⁺₂ reacts in the gas phase with D₂ to yield V₁D³⁻.\[^25\] Interestingly, very recently, the reaction of V₂⁺₁ with CO₂ to yield V₁O₂⁻ has been investigated, and the surprising observation has been made that the intermediate V₂O²⁻ reacts considerably faster than V₂⁻.\[^26\]
In this work, it is shown by IR spectroscopy that neutral \( V_2 \) reacts with \( H_2 \) without a significant reaction barrier to linear HVVH under preservation of the spin multiplicity and to the \( V_2(\mu-H)_2 \) isomer with a nonplanar, folded structure (see Figure 1). The latter is generated, in particular, upon irradiation with light of a wavelength in the ultraviolet range between 250 and 385 nm.

![Figure 1. Structures of HVVH (left) and \( V_2(\mu-H)_2 \) (right).](image)

To assist in the assignment of the observed bands and to obtain information on the electronic structure of the products, quantum-chemical calculations are performed. Information on the energy and nature of the electronic ground states and low-lying excited states is obtained by multireference configuration-interaction (MRCI) calculations based on complete-active-space self-consistent-field orbitals. Vibrational frequencies are calculated by density-functional calculations.

**Results and Discussion**

**IR Spectra**

The IR spectra recorded after co-depositing vanadium atoms and a mixture of Ne and dihydrogen show broader absorptions at around 1580, 1571, 1569, and 1564 cm\(^{-1}\), and furthermore absorptions in the region between 331 and 346 cm\(^{-1}\) that belong to the same compound (see Figure 2). Upon annealing to 10 K, four sharper bands appear at 1570.7 (strong), 1569.8 (medium), 1569.4 (medium), and 1568.6 cm\(^{-1}\) (weak), as well as a band in the low-frequency regime at 343.4 cm\(^{-1}\) with a shoulder at 342.5 cm\(^{-1}\) at the expense of the broader bands. Upon irradiation with visible light, the broader bands re-appear with a slightly changed intensity pattern; there is more intensity at around 1580 cm\(^{-1}\) with two maxima at 1581.4 and 1580.2 cm\(^{-1}\), and in the low-frequency region, there are two pronounced absorptions at 342.4 and 340.3 cm\(^{-1}\). Upon renewed annealing to 10 K, the sharper bands that appeared in the first annealing again recover intensity at the expense of the broader bands. Upon irradiation with ultraviolet (UV) light, all those bands are extinguished, and upon further annealing to 10 K, the sharp bands only very weakly re-appear.

Another set of absorptions is observed essentially only after annealing (to 10 K), but neither directly after deposition nor directly after irradiation with visible light. These absorptions are found at 1425.1, 1405.3, 917.2, and 565.4 cm\(^{-1}\) (see Figure 3). The bands clearly appear upon annealing after deposition, they appear less strongly upon annealing after irradiation with visible light, and they appear most strongly upon annealing after irradiation with UV light. At least after the irradiation with UV light, there are weaker absorptions red-shifted with respect to the clear bands observed after annealing.

![Figure 2. IR spectra at around 1570 and 340 cm\(^{-1}\) of matrices obtained by deposition of V and H\(_2\) in Ne, a) after deposition, b) after annealing to 10 K, c) after irradiation with visible light, d) after further annealing to 10 K, e) after irradiation with UV light, f) after another annealing to 10 K.](image)

![Figure 3. IR spectra at around 1420, 920, and 560 cm\(^{-1}\) of matrices obtained by deposition of V and H\(_2\) in Ne, a) after deposition, b) after annealing to 10 K, c) after irradiation with visible light, d) after further annealing to 10 K, e) after irradiation with UV light, f) after another annealing to 10 K.](image)
For all those bands, corresponding counterparts are observed using D$_2$ or HD. For the first set of absorptions, there are D$_2$ counterparts of the four sharper signals that appeared after annealing to 10 K at 1132.6, 1131.9, 1131.8, and 1131.5(sh) cm$^{-1}$, and counterparts in the low-frequency regime at 247.0 and 246.2(sh) cm$^{-1}$ (see Figure 4). Using HD, one group of counterparts is found at 1579.0, 1678.6(sh), 1577.7, and 1577.0 cm$^{-1}$ near the absorptions found using H$_2$, and another group of counterparts is found at 1138.6, 1138.3(sh), 1137.8, and 1137.4(sh) cm$^{-1}$ near the absorptions found with D$_2$. In the low-frequency region, the HD counterparts are observed at 276.7 and 275.9(sh) cm$^{-1}$. Furthermore, using HD, there appears another weaker band at 442.8 cm$^{-1}$ not found in the experiments with H$_2$ or D$_2$.

For the second set of absorptions, the D$_2$ counterparts are found at 1035.7, 1021.4, 678.0, and 399.3 cm$^{-1}$ (see Figures 5 and 6). Using HD, absorptions are obtained at 1414.0 and 1412.5(sh) cm$^{-1}$ between the two high-frequency bands with H$_2$, another absorption is obtained at 1030.0 cm$^{-1}$ between the two high-frequency bands with D$_2$. The HD counterparts for the lower-frequency absorptions are observed at 886.8 and 495.6 cm$^{-1}$, between the corresponding H$_2$ and D$_2$ signals. In the present experiments, the signals previously assigned to VH$_2$ and VH$_3$(H$_2$) in solid Ne$^{[23]}$ have been observed weakly after irradiation.

![Figure 4](image1.png)

**Figure 4.** IR spectra at around 1570, 1140, and 300 cm$^{-1}$ of matrices obtained by deposition of V and H$_2$, D$_2$, or HD in Ne after irradiation with visible light and annealing to 10 K.

![Figure 5](image2.png)

**Figure 5.** IR spectra at around 1420, 1030, 900, and 680 cm$^{-1}$ of matrices obtained by deposition of V and H$_2$, D$_2$, or HD in Ne after irradiation with UV light and annealing to 10 K.

### Assignments

The first set of bands is assigned to the linear HVVH molecule, the second set of bands is assigned to the folded $V_5(\mu-H)_2$ molecule with bridging H atoms. The assignment is based on the number of bands observed, the positions of the bands, and their isotopic shifts, and it is confirmed by computational results (density functional calculations with the TPSS functional and the TZVP basis set).

Within the first set of bands, when using H$_2$, there is one band in the V–H stretching region at 1570.7 cm$^{-1}$. (Note: for VH$_2$ in Ne, the V–H stretching modes are found at 1492.9 and 1553.9 cm$^{-1}$.$^{[20]}$) When using D$_2$, the band shifts to 1322.2 cm$^{-1}$, confirming the presence of a V–H unit. Using HD, two bands in these regions at 1579.0 and 1138.6 cm$^{-1}$ are found, one in the V–H and one in the V–D region, blue-shifted with respect to their H$_2$ or D$_2$ counterparts. This indicates that the latter belong to pairs of bands, the higher-lying member of which is not observed for symmetry reasons, pointing to a symmetrical arrangement of two V–H groups in accordance with linear HVVH. For linear HVVH, a total of two modes out of five ($\rho^v_1$, $\rho^s$) should be observable in infrared spectra, and indeed, another mode is observed at 343.4 cm$^{-1}$, the underade combination of the bending motions. When using HD, due to symmetry reduction, yet another mode is observed at 442.8 cm$^{-1}$, the formerly gerade combination ($\rho_g$) of the bending vibrations. The density-functional results nicely confirm the assignment (see below).

Within the second set of bands, when using H$_2$, there are two signals at 1425.1 and 1405.3 cm$^{-1}$ in the region of the stretching vibrations. They have lower wavenumbers than the signals assigned to HVVH, consistent with H in a bridging configuration. The isotopic shifts to 1035.7 and 1021.4 cm$^{-1}$ when using D$_2$ confirm that these bands are related to the motion of H atoms. The observation of two bands in the V–H stretching region already with the homonuclear reactants points to a non-symmetric, likely non-planar, structure. For folded $C_4_2$-symmetric $V_5(\mu-H)_2$, five out of six modes ($3 \times a_1$, $a_2$, $b_2$) should, in principle, be observable in infrared spectra. Two additional modes are observed at 917.2 and 565.4 cm$^{-1}$, the shift of the H atoms along the direction of the V–V axis at 917.2 cm$^{-1}$, the folding motion at 565.4 cm$^{-1}$. Thus, out of the five in principle observable modes, four are found. The density-functional calculations confirm the assignment to folded $C_4_2$, $V_5(\mu-H)_2$ and indicate that the other mode which, in principle, could be observed, is very weak, a mode dominated by the V–V stretching motion.
Calculated Structures and Energies

The quantum-chemical calculations, both MRCl and density-functional calculations, yield as the lowest-lying isomer of V₃H₃ a folded V₃(µ-H)₂ isomer with a C₃ᵥ-symmetric structure, and, at an energy higher by about 0.5 eV, a linear HVVH isomer with D₁₉h symmetry (Figure 7).

According to the MRCl calculations, the electronic ground state of the folded C₃ᵥ isomer is a 1'A₁ state. The V–V distance amounts to 184.4 pm and the V–H distance to 179.6 pm. The torsional angle between the two HV₃ planes is 120.6°. The density-functional calculations yield a broken-symmetry state that corresponds to the 1'A₁ term. The values for the V–V and V–H distances of 182.9 and 180.1 pm, and the torsional angle between the HV₃ planes of 113.3° obtained by the density-functional calculations are close to the MRCl values and confirm that the density-functional calculations give, qualitatively, the correct picture.

The relative energies of low-lying excited states (by MRCl) at the structure of the C₃ᵥ 1'A₁ ground term are shown in Figure 7. From a relative energy of 0.79 eV above the ground state, there are different triplet and singlet terms with small energy separations, the lowest ones are a 1'B₃ and a 1'B₂ term at 0.79 and 0.89 eV. The lowest-lying term of the linear HVVH isomer is, according to the MRCl calculations, a 3Σ⁺ term at an energy of 0.48 eV relative to the 1'A₁ state. The V–V distance amounts to 170.7 pm and the V–H distance to 170.5 pm. The same term is also obtained by the density-functional calculations at a relative energy of 0.56 eV, in good agreement with the MRCl result. The values for the V–V and V–H distances of 171.1 and 171.6 pm obtained by the density-functional calculations are again close to the MRCl values and confirm the qualitatively correct picture of the density-functional calculations also in the case of the linear isomer. The relative energies of the lower-lying terms at the structure of the 3Σ⁺ term are also shown in Figure 7. At 0.15 and 0.17 eV with respect to the 3Σ⁺ term, there are a 1'G₁ and a 1'S₁ term, respectively. Further singlet and triplet terms follow at energies higher than 1.21 eV.

Calculated Vibrational Frequencies

The vibrational frequencies obtained from the density-functional calculations support the assignment of the observed bands to linear HVVH and folded V₃(µ-H)₂. All vibrational modes for which the calculations indicate transitions with substantial intensity have been observed with wavenumbers not too far from the calculated values. For the linear isomer, the wavenumbers of the allowed transitions are calculated to be 1613 and 322 cm⁻¹ (see Table 1), whereas the observed transitions have values of 1570 and 343 cm⁻¹. The differences amount to 43 and 21 cm⁻¹. The calculated isotopic shifts of 461 and 92 cm⁻¹ show an even better agreement with the measured shifts of 438 and 97 cm⁻¹. For the folded isomer, the calculated wavenumbers of 1522, 1502, 984, and 571 cm⁻¹ (see Table 1) deviate from the measured values of 1425, 1405, 917 and 565 cm⁻¹ by 97, 96, 66, and 5 cm⁻¹, respectively. Thus, the observed splitting between the symmetric and antisymmetric vibrations at 1425 and 1405 cm⁻¹ is almost quantitatively reproduced by the calculations. Again, the calculated isotopic shifts of 438, 431, 275, and 162 cm⁻¹ still show a better agreement with the observed shifts of 389, 384, 239, and 166 cm⁻¹. Such deviations of some 10 cm⁻¹ are in the typical range for frequencies obtained by density-functional calculations and have to be expected.

Electronic Configurations

For the 1'A₁ ground state of the folded V₃(µ-H)₂ isomer, the occupation of the valence-space natural orbitals of the MRCl wave function, (8a) 0.97 (3b) 0.36 (9a) 0.85 (10a) 1.81 (4b) 1.77 (11a) 0.79 (8b) 0.28 (3a) 0.03 (9b) 0.18 (10b) 0.11 (12a) 0.06 (4a) 0.03 (5b) 0.10 (11b) 0.01, clearly shows the multiconfigurational character. The leading configuration in the expansion of the wave function is (8a)²(3b)³(9a)²(10a)²(4b)²(11a)² with a coefficient of 0.77. The first two orbitals are orbitals with a leading contribution from the H 1s orbitals. They are V–H bonding orbitals, the first one mainly involving the V 4s orbitals, the second one involving the 3d orbitals. The following eight orbitals are essentially 3d orbitals pointing to a substantial amount of 3d–3d bonding. Four of them are bonding orbitals and four of them are the corresponding antibonding orbitals. From the occupation numbers of the natural orbitals, a bond order of 3.17 is obtained. To confirm this consideration, a natural-bond orbital analysis has also been performed. For the bridging hydrogen atoms, it yields two three-center H–V₂ bonding orbitals occupied with 1.96 electrons each and corresponding antibonding orbitals occupied with 0.03 electrons each. For the vanadium–vanadium interaction, it yields four V–V bonding orbitals occupied with 1.85, 1.81, 1.77, and 1.70 electrons, and corresponding antibonding orbitals occupied with 0.25, 0.21, 0.19 and 0.10 electrons. These values amount to 7.13 electrons in bonding orbitals and 0.75 electrons in antibonding orbitals, yielding a value of 3.19 for the bond order of the V–V bond, close to the value obtained by consideration of just the natural orbitals. Thus, the value of larger than three is indicative of the presence of fourfold bonding in V₃(µ-H)₂.

Figure 7. Relative energies of the low-lying excited electronic states of folded V₃(µ-H), and linear HVVH at the structures of their ground states by Davidson-corrected MRCl calculations with an ANO basis set.
Table 1: Wavenumbers (\(\tilde{\nu}/\text{cm}^{-1}\)) of vibrational transitions of linear HVVH and folded \(V_2(\mu-H)\), by density-functional calculations with the TPSS functional and from IR spectra of Ne matrices. The calculated values are unscalled harmonic vibrational frequencies. Experimental values for the most prominent site after annealing to 10 K. (Entries in parenthesis: calculated intensities).

| \(V_2(\mu-H)\)   | TPSS | Exp. | \(V_2(\mu-H)(\mu-D)\) | TPSS | Exp. | \(V_2(\mu-D)\) | TPSS | Exp. |
|------------------|------|------|----------------------|------|------|----------------|------|------|
| \(a_1\)          | 328.0 (5) | 326.9 (5) | \(a_1\)           | 324.7 (3) | 324.7 (3) | \(a_1\)           | 324.7 (3) | 324.7 (3) |
| \(a_1\)          | 571.1 (63) | 496.3 (51) | \(a_1\)           | 495.6 (36) | 495.6 (36) | \(a_1\)           | 495.6 (36) | 495.6 (36) |
| \(b_1\)          | 910.0 (0) | 678.4 (0) | \(b_1\)           | 656.0 (0) | 656.0 (0) | \(b_1\)           | 656.0 (0) | 656.0 (0) |
| \(b_1\)          | 983.7 (35) | 950.6 (20) | \(b_1\)           | 886.8 (17) | 886.8 (17) | \(b_1\)           | 886.8 (17) | 886.8 (17) |
| \(a_1\)          | 1502.0 (98) | 1077.4 (68) | \(a_1\)           | 1030.0 (47) | 1030.0 (47) | \(a_1\)           | 1030.0 (47) | 1030.0 (47) |
| \(a_2\)          | 1521.9 (171) | 1512.0 (134) | \(a_2\)           | 1414.0 (87) | 1414.0 (87) | \(a_2\)           | 1414.0 (87) | 1414.0 (87) |

The calculated fragmentation of HVVH, and the fragmentation of \(V_2(\mu-H)\), also the ground state, the same with an coefficient of 0.83. The first two orbitals are V–H bonding orbitals with the main contributions from H 1s and V 4s; orbitals; the remaining ones are V 3d orbitals. Hence, the \(\sigma^*, \pi, \tau, \delta\) orbitals contribute to the V–V bonding. From the occupation numbers of the natural orbitals, a bond order of 3.43 is obtained, still somewhat larger than for the isomer with the bridging hydrogen atoms. The triplet ground state results from the occupation of the two \(\sigma\) orbitals with two electrons. An NBO analysis as well as the Mayer bond order based on the density-functional results confirm the conclusions from the evaluation of the MRCI wave function (see the Supporting Information).

To obtain an estimate for the reaction energy of the formation of \(V_2(\mu-H)\), and the fragmentation of HVVH, additional density-functional calculations were performed on \(V_2(\mu-H)\), \(VH(\Delta)\), and \(H_2\). According to the calculations, the formation of \(V_2(\mu-H)\), via \(V + H_2 -> V_2(\mu-H)\), is exothermy by 1.18 eV. Hence, the formation of the linear HVVH isomer is still exothermy by 0.62 eV. The \(\Delta\) ground state for VH is in line with previous calculations. The calculated fragmentation energy of HVVH by HVVH(\(\Sigma_g^+\)) –2 VH(\(\Delta\)) amounts to 2.77 eV. This value is similar to the experimental value of 2.75 eV for the dissociation energy of VH, higher than the value of 1.53 eV for \(Cr_2\), and supports the presence of multiple bonding in HVVH.

**Discussion**

For both HVVH and \(V_2(\mu-H)\), clear, sharp signals are observed only after annealing (not after deposition or irradiation), pointing to the population of a few (well-defined) matrix sites. For HVVH, the broader signals found only about one quarter of the intensity of the strongest signal of HVVH. Thus, the detection of broad signals is expected to be more difficult for \(V_2(\mu-H)\).

The experiments show that \(H_2\) reacts with \(V_2\) by formation of both a linear HVVH isomer and a folded \(C_2\) \(V_2(\mu-H)\) isomer. Thus, it is likely that the reaction proceeds without or with only a small barrier. The folded isomer is the more stable one, nevertheless, the linear isomer is formed in substantial amount. This observation is consistent with the ground-state properties of folded \(V_2(\mu-H)\) and linear HVVH, as revealed by the calculations. HVVH has a \(\Sigma_g^+\) ground state, the same as the diatomic \(V_2\). Thus, the formation of HVVH from \(V_2\) and \(H_2\) is a spin-allowed process, whereas the formation of the folded \(V_2(\mu-H)\) isomer, which has a \(\Delta\) ground state, is spin-forbidden although energetically favorable. Irradiation with UV light leads to the destruction of HVVH and almost exclusive formation of the more stable \(V_2(\mu-H)\). Obviously, by irradiation with UV light, higher-lying excited states are populated that allow an easy isomerization to the folded \(V_2(\mu-H)\), for example, by change of the spin state due to spin-orbit coupling and subsequent isomerization or vice versa.

It is observed that upon irradiation with visible light, the signals of \(V_2(\mu-H)\) decrease, whereas those of HVVH increase, and upon irradiation with UV light, the signals of HVVH almost entirely vanish, whereas those of \(V_2(\mu-H)\) markedly increase. The experiments indicate the conversion of linear HVVH to non-planar \(V_2(\mu-H)\) by irradiation with UV light and the conversion of \(V_2(\mu-H)\) to HVVH by irradiation with visible light (Scheme 1).

The chemistry of \(V_2\) and \(H_2\) is a reminiscence of the reactivity of the gallium dimer with dihydrogen. The gallium dimer \(Ga_2\), despite spin restrictions, spontaneously reacts with dihydrogen by formation of cyclic \(Ga_2(\mu-H)\), whereas ground-state \(Ga_2\) atoms do not react with \(H_2\). \(GaH_2\) is formed only upon photoexcitation. Furthermore, \(Ga_2(\mu-H)\) features an interesting photochemistry. Irradiation (\(\lambda = 546\) nm) of \(Ga_2(\mu-H)\) leads to the formation of \(HGaGaH\) and \(H_2GaGa\), and upon irradiation of these products using...
other wavelengths ($\lambda = 365$ nm or $\lambda > 700$ nm), cyclic Ga$_2$(µ-H)$_2$ is reformulated. Furthermore, the present results for V$_2$H$_2$ are similar to the findings of a theoretical (density-functional) study on Ti$_2$H$_2$.[36] There, also a folded isomer with bridging H atoms and a linear HTiTiH isomer were found, the isomer with bridging ligands being the lowest in energy.

The V-V bond distance of 170.7 pm for HVVH is surprisingly short, even shorter than that of V$_2$($^{12}S_g$) of 180.4 pm,[37] although the leading configuration for V$_2$($6\sigma^2$($7\sigma^0$)($3\pi_u^2$($1\delta_u^2$)($1\delta_u^2$), even points to a quintuple bond (The occupation numbers of the CASSCF natural orbitals for V$_2$ yield a value for the bond order of 4.27). An explanation of this contradictory observation is likely the following: due to the formation of the V–H bonds, the 4s contribution to the electron density at the V atoms and in the V–V bonding region decreases, and therefore also the shielding of the V nuclear charge, and the V$_2$ core has partial V$_2$$^{3+}$ character leading to a contraction of the V–V bonding orbitals. Although for both V$_2$(µ-H)$_2$ and HVVH, the results point to quadruple bonding, the V–V bond distance in V$_2$(µ-H)$_2$ is longer by 13.7 pm than in HVVH. The values for the V–V bond order are in qualitative agreement with this observation: the values point to quadruple bonding for both isomers, but the value for HVVH it is still somewhat larger than for V$_2$(µ-H)$_2$. However, the bonding situation differs in some respects. In V$_2$(µ-H)$_2$, there are bridging H atoms, and hence, there is associated electron density close to the V–V axis. Furthermore, HVVH has a triplet ground state. Therefore, in its leading configuration, two of the five orbitals that contribute to the bonding are occupied by one electron only, whereas in V$_2$(µ-H)$_2$, there are four V–V bonding orbitals occupied by pairs of electrons. Thus, in HVVH, for the two unpaired electrons, there is the possibility to keep a larger distance. It is likely that both factors influence the final bonding situation.

The V–V bond distances of 184.4 pm and 170.7 pm in V$_2$(µ-H)$_2$, and HVVH, respectively, are still shorter than the values in the paddlewheel complexes V$_4$(DPhF)$_4$, with a formal triple bond (197.8/197.9 pm)[38] and K[V$_2$(DPhF)$_2$] with a formal bond order of 3.5 (192.9 pm),[39] which have V$_2$$^{3+}$ and V$_2$$^{5+}$ cores, and far shorter than the value of 246.0 pm in a divanadium(IV) complex with a V–V single bond,[40] supporting the presence of a quadruple bond in V$_2$(µ-H)$_2$. HVVH and V$_2$(µ-H)$_2$ are further examples of bimetallic metal hydride compounds with a high metal–metal bond order, like HCrCrH, HMoMoH, and HWWH.[39] However, the latter ones with a trans-bent alignment have been characterized only theoretically to date and are still awaiting experimental confirmation.

### Conclusion

Multiple bonds between two transition metals are intensely studied due to their intriguing electronic structure and potential applications. The bond order is particularly high if the number of ligands is decreased to an absolute minimum. However, the synthesis of RMMR compounds (M = transition metal) by conventional techniques in solution is a difficult task and, by all means, requires bulky ligands R. In this work, we demonstrate the use of the matrix-isolation technique to experimentally characterize simple HMMH compounds with multiple MM bonds. Hence, two isomers of unprecedented V$_4$H$_2$, folded V$_2$(µ-H)$_2$, and linear HVVH, are generated, and identified by their IR spectra. UV irradiation induces the transformation of linear HVVH to folded V$_2$(µ-H)$_2$. Irradiation with visible light leads to the transformation of the folded V$_2$(µ-H)$_2$, isomer to linear HVVH. Quantum-chemical calculations (MRCI) show that the folded V$_2$(µ-H)$_2$ is the lower-lying isomer with a $^1A_g$ ground state. The linear HVVH with a $^3A_g$ electronic ground state is calculated to be at an energy of 0.48 eV with respect to the folded isomer. Both isomers are characterized by multiple V–V bonding interactions and the orbital occupations point to quadruple bonding. The calculated vibrational frequencies (density-functional calculations) support the assignment of the observed transitions. Although the folded isomer is the lower-lying one, without irradiation, the linear isomer is formed in a substantial amount. This observation is consistent with the electronic ground states of the isomers. Since the V$_2$ molecule has a $^3A_g$ ground state, the formation of the folded V$_2$(µ-H)$_2$ isomer ($^1A_g$) by reaction of H$_2$ and V$_2$ is a spin-forbidden process, whereas the formation of linear HVVH is spin-allowed. Thus, it is reasonable that a substantial amount is trapped as HVVH. In ongoing work, we extend our work on other multiple-bonded HMMH compounds. The experimental characterization of these small molecules is important for the detailed understanding of the bond properties and, thereby, also the reactivity of these systems.

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### Conflict of interest

The authors declare no conflict of interest.

### Keywords:
- hydrides
- matrix isolation
- MRCI calculations
- multiple bonding
- vanadium

[Scheme 1. Formation of V$_2$(µ-H)$_2$, and HVVH, and interconversion.]

\[V + H_2 \rightarrow \text{no reaction} \]
\[V_2 + H_2 \rightarrow V_2(\mu-H)_2 + HVVH\]
\[HVH \rightarrow V_2(\mu-H)_2\]
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