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PtZnH$_5^-$, A σ-Aromatic Cluster

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ABSTRACT: We report a joint photoelectron spectroscopic and theoretical study of the PtZnH$_5^-$ cluster anion. This cluster exhibited an unprecedented planar pentagonal coordination for Pt and an unusual stability and high intensity in the mass spectrum. Both are due to the σ-aromaticity found in the H$_5^-$ cycle supported by the 5d orbitals on the Pt atom. σ-Aromaticity in all-H systems has been predicted in the past but never found in experimentally observed species. Besides fundamental importance, mixed transition-metal hydrides can be found as intermediates in catalytic processes, and thus, the unexpected stability facilitated by σ-aromaticity can be appreciated also in practical applications.

SECTION: Molecular Structure, Quantum Chemistry, and General Theory

Aromaticity, one of the most significant concepts in modern chemistry, continues to gain appreciation as it is one of the stabilizing chemical bonding effects that leads to remarkable symmetries and stabilities of clusters and molecules.$^{1-16}$ Many all-metal clusters were reported to exhibit multiple types of aromaticity, including σ, π, and δ-type or a combination of these three types. σ-Aromaticity in particular has been a concept first subjected to debates and prosecution and then accepted and widely explored. The simplest systems in which σ-aromaticity can be found are clusters of H, where the bonding overlap of the 1s atomic orbitals (AOs) on atoms gives rise to aromaticity, including σ- aromaticity, π-aromaticity, and also of PtZnH$_3^-$ and PtZnH$_4^-$. These anionic hydrides are generated in the gas phase and investigated using negative ion photoelectron spectroscopy (PES). The experimental photoelectron spectra are compared with theoretical predictions for the found global minimum structures. PtZnH$_5^-$ is found to have an unusual structure, high stability manifested in the abundance in the mass spectrum, and σ-aromaticity.

Experimental Results. The mass spectrum of the PtZnH$_5^-$ cluster anion is presented in Figure 1A. One can observe several major isotopic peaks ranging from 261 to 271. In order to make sure that the spectrum has no impurities from other hydrides, we plotted the simulated isotopic distribution however, their experimental evidence is still absent. It would be exciting to see if the simplest possible type of aromaticity and antiaromaticity (σ-type) could be present in metal hydrides and govern their stability, in particular, if one day this could be explored in catalytic processes.

In this work, we study mixed Pt–Zn hydrides at the subnano scale. The Pt–Zn bimetallic systems are used as catalysts for various reactions, such as hydrogenation/dehydrogenation and fuel cell oxidation.$^{21-26}$ Doping precious metal catalysts in general is an amenable strategy for manipulating and tuning the electronic structure and thus properties of the catalyst and potentially reducing the cost. However, the research is still within the nanoregime, and subnano bimetallic Pt–Zn clusters containing only several atoms have not been studied but pose a great interest in view of the observation that small clusters are actually the sites where the majority of catalytic processes occur.$^{27-29}$ Hydrides of these small clusters are thus of interest.

We present a combined experimental and theoretical study of a σ-aromatic cluster, PtZnH$_5^-$, and also of PtZnH$_3^-$ and PtZnH$_4^-$. These anionic hydrides are generated in the gas phase and investigated using negative ion photoelectron spectroscopy (PES). The experimental photoelectron spectra are compared with theoretical predictions for the found global minimum structures. PtZnH$_5^-$ is found to have an unusual structure, high stability manifested in the abundance in the mass spectrum, and σ-aromaticity.

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the triumph of PtZnH$_5^-$ from the PtZnH$_n^-$ series indicates its special stability. Figure 2 presents the photoelectron spectrum of PtZnH$_5^-$ taken with a 355 nm (3.496 eV) laser. We took spectra from all the isotopic peaks, and all of them were identical, which is another confirmation that there is no contamination in the mass spectrum. One observes a major EBE band ranging from 2.4 to 3.2 eV and peaked at 2.75 eV, with the resolution being better than 50 meV. The electron binding energy (EBE) value corresponding to the intensity maximum in the observed band is its vertical detachment energy (VDE), the transition energy at which the Franck–Condon overlap between the wave functions of the anion and its neutral counterpart is maximal. The adiabatic electron affinity (EA) is the energy difference between the lowest energy vibronic state of the anion and the lowest energy vibronic state of its neutral counterpart. We have estimated the EA of PtZnH$_3^-$ to be 2.5 eV by extrapolating the low EBE side, with the corresponding EBE value there being taken as the EA.

Cluster Structures. Figure 3 shows the found global minima of considered clusters, PtZnH$_3^-$, PtZnH$_4^-$, and PtZnH$_5^-$; for PtZnH$_5^-$, we show the global minimum and the second competing isomer. All clusters bind atomic H, which indicates that they would likely dissociate H$_2$, should it bind to them as a molecule, for example, in a catalytic process. Indeed, Pt is a typical catalyst for dehydrogenation, in the form of extended surfaces, nanoparticles, and nanoislands, even though catalysis of really small clusters was hardly ever explored. For PtZnH$_3^-$, the global minimum is a C$_1$ (‘A) structure, with all three H atoms bound to the Pt atom. For PtZnH$_4^-$, it is the C$_{2v}$ (‘A$^+$) structure, with two of the four H atoms bridging between Pt and Zn and the remaining two bound to Pt. For PtZnH$_5^-$, there are two isomers relatively close in energy. Most theoretical methods predict isomer II (C$_{2v}$ (‘A$^+$)) to be more stable by $\sim$4 kcal/mol. One exception is the CASSCF(12,12) method, which predicts isomer I (C$_5$ (‘A$_1$)) to be more stable. Considering the expected error in the performance of the ab initio method in use, it is fair to say that these two isomers are isoenergetic. Calculated properties of both isomers are shown in the Supporting Information.

Theoretical Interpretation of the Photoelectron Spectrum of PtZnH$_5^-$.

In this work, we focus on PtZnH$_5^-$ because of its unusually high abundance in the mass spectrum as compared to the two lighter hydrides. We expect it to exhibit unusual structure and bonding characteristics. Table 1 shows calculated photoelectron spectra of the two isomers of PtZnH$_5^-$.

The simulated spectrum of isomer II does not agree with the experiment, with VDE$_{II}$ being $\sim$1 eV off, and that rules out the possibility of the isomer II being observed in the experiment. On the other hand, the theoretical spectrum of isomer I is in close agreement with the experiment. All experimental features are successfully assigned to single-electron processes of electron detachment from the valence molecular orbitals (MOs) shown in Figure 4A. The fact that isomer I is predicted to be less stable but is the one observed experimentally indicates either the inadequate performance of theoretical methods or the kinetic stabilization of the metastable isomer I in the beam. The predicted adiabatic detachment energy (ADE) is close to the VDE (Table 1), and in the experiment, the observed EA, 2.5 eV, and the theoretical ADE are effectively the same quantities in this case. The electron detachment from the completely symmetric HOMO of the cluster leads to a small geometric
HOMO contribute to the bonding in the cluster. The HOMO and Zn (HOMO diagram for the valence MOs; the MOs outlined in red in (A) are highlighted. (C) A qualitative scheme for the formation of the cluster obey the (4 AOs on H atoms in a pentagonal arrangement; the three lowest-energy MOs are analogous to those populated in the cluster; their population makes clearly pure d lone pairs of Pt (HOMO presented in Figure 4B. Some of the MOs of the cluster are Figure 4A, and a sketch of how they are formed from AOs is Figure 4. Chemical bonding in PtZnH5. (A) Valence MOs, with the ones outlined in red corresponding to the σ-aromatic unit. (B) Correlation diagram for the valence MOs; the MOs outlined in red in (A) are highlighted. (C) A qualitative scheme for the formation of σ-MOs from five 1s AOs on H atoms in a pentagonal arrangement; the three lowest-energy MOs are analogous to those populated in the cluster; their population makes the cluster obey the (4n + 2) Hückel’s rule for aromatic compounds, with n = 1.

Table 1. Calculated Photoelectron Spectra of PtZnH5− Isomers I and II, in eV

<table>
<thead>
<tr>
<th>Isomer I</th>
<th>peak expl. VDE/EA</th>
<th>MO</th>
<th>resultant e-configuration</th>
<th>TD-UPBEPE, a VDE/ADE</th>
<th>TD-UPBEPE, b VDE/ADE</th>
<th>CCSD(T)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Isomer I</td>
<td>X 2.75/2.5</td>
<td>HOMO</td>
<td>...2a1,2b1,3a1,2e1,3e1,4a1</td>
<td>2.71/2.58</td>
<td>2.70/2.51</td>
<td>2.50</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HOMO−1</td>
<td>...2a1,2b1,3a1,2e1,3e1,4a1</td>
<td>4.48d</td>
<td>4.50d</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>HOMO−2</td>
<td>...2a1,2b1,3a1,2e1,3e1,4a1</td>
<td>4.93d</td>
<td>4.92d</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>HOMO−3</td>
<td>...2a1,2b1,3a1,2e1,3e1,4a1</td>
<td>5.83</td>
<td>5.91</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>HOMO−4</td>
<td>...2a1,2b1,3a1,2e1,3e1,4a1</td>
<td>7.12d</td>
<td>7.13d</td>
<td></td>
</tr>
<tr>
<td>Isomer II</td>
<td>X 2.75/2.5</td>
<td>HOMO</td>
<td>...5a1,6a1,2b1,3b1,2a1,3a1</td>
<td>3.68</td>
<td>3.65</td>
<td>3.84</td>
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<tr>
<td></td>
<td></td>
<td>HOMO−1</td>
<td>...5a1,6a1,2b1,3b1,2a1,3a1</td>
<td>3.97</td>
<td>4.02</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>HOMO−2</td>
<td>...5a1,6a1,2b1,3b1,2a1,3a1</td>
<td>4.17</td>
<td>4.19</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td>HOMO−3</td>
<td>...5a1,6a1,2b1,3b1,2a1,3a1</td>
<td>4.46</td>
<td>4.67</td>
<td></td>
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<td></td>
<td></td>
<td>HOMO−4</td>
<td>...5a1,6a1,2b1,3b1,2a1,3a1</td>
<td>4.70</td>
<td>5.17</td>
<td></td>
</tr>
</tbody>
</table>

The aug-cc-pVTZ-pp basis set was used. bThe 6-311++G** basis was used for Zn and H, and LANL2DZ(+f) was used for Pt. cAt the geometry of the anion optimized using MP2/aug-cc-pvTZ-pp. dIndicates doubly degenerate features.

change upon relaxation, hence resulting in a small difference between the VDE and ADE.

Aromaticity of PtZnH5−. Figure 4 addresses the chemical bonding in the C5v (1A1) isomer I of PtZnH5−. First of all, its structure is highly unusual due to its high symmetry and an anomalous coordination of the Pt atom, pentagonal pyramidal. Pt is often found to be square planar, but here, we have a five-coordinated arrangement in-plane and an additional Zn atom on the axial position. To the best of our knowledge, such coordination for Pt was never observed before. What leads to the stabilization of this unusual structure and the high abundance of this species in the mass spectrum of PtZn hydride anions? Valence MOs of the cluster are shown in Figure 4A, and a sketch of how they are formed from AOs is presented in Figure 4B. Some of the MOs of the cluster are clearly pure d lone pairs of Pt (HOMO−1 and HOMO−4) and Zn (HOMO−6, HOMO−7, and HOMO−8) that do not contribute to the bonding in the cluster. The HOMO and HOMO−3 correspond to σ-bonding between Pt and Zn. These MOs are mainly bonding and antibonding combinations of the 4s AO on Zn and 5d, 6s AO on Pt. The Pt−Zn bond is partially ionic, with the partial charges on atoms being +0.011 on Zn and −1.355 on Pt (natural population analysis31). Importantly, removal of Zn from the cluster results in the PtH5− species, which is a planar D3h structure (at least in the local minimum form), and the bonding within this fragment is analogous to that in the PtH5− unit within the PtZnH5− hydride. Thus, Zn is merely a spectator in the system where Pt exhibits an unusual coordination.

The most interesting MOs are the doubly degenerate HOMO−2 and the HOMO−5 (Figure 4A). These are σ-MOs, mostly formed by the five 1s AOs on hydrides. They mix with the 6p and 6s AOs on Pt, as allowed by symmetry (Figure 4B). For simplicity, first consider just the isolated (hypothetical) H5− pentagon. The five 1s AOs would form five MOs that are shown schematically in Figure 4C. The three lowest-energy ones correspond nicely to the valence HOMO−2 and HOMO−5 of PtZnH5− in terms of contributions from

hydrides. Hence, they are depicted populated in the scheme in Figure 4C. Populated by six electrons, the HOMO−2 and HOMO−5 make the cluster σ-aromatic, in accord with the (4n + 2) Hückel’s electron counting rule for aromatic compounds with n = 1. Remarkably, σ-aromaticity was predicted for the isolated H$_5^−$ (D$_{3h}$) cluster, but it was found to be a saddle point on the potential energy surface. With the additional support of the 6s and 6p AOs on Pt, the aromatic all-H cycle is stabilized in the studied hydride. Pt is capable of sustaining the five-fold coordination in-plane because it is not really a five-fold coordination; it is a coordination to the H$_5^−$ pentagon (the complex can be described as [Zn−Pt(H$_5^−$)]$^{−}$), and the bonding within this unit is delocalized. Notably, transition metals can coordinate to five-membered ring structures, for example, in metallocenes, but there, the coordination is not in one plane. In PtZnH$_5^{−}$, Pt takes the central position in the ring to match the σ-MOs on H$_5$.

Besides the electron counting rule, we wished to gain an additional confirmation of aromaticity in this hydride. The calculation of the nucleus-independent chemical shifts (NICS) index at the center of the pentagonal aromatic unit produces meaningless results because the geometric center coincides with the Pt atom and the contribution of the aromatic ring to the chemical shift is convoluted by the contributions from the AOs on Pt. We also applied the AdNDP procedure, allowing for the maximal electron localization while preserving the electron pair paradigm. Two possible bonding pictures emerged from this analysis. One of them describes the H$_5^−$ arrangement σ-aromatic, with three localized AdNDP orbitals that look nearly identical to the HOMO−2 and HOMO−5 in Figure 4A. The contributions from Pt in this case were separated out to be lone pairs with fairly low population numbers of ~1.6 e. The alternative picture produced five 2c−2e Pt−H bonds with population numbers of ~1.9 e. Both pictures are valid from the AdNDP standpoint, and it is possible that there is a resonance between the two localized solutions. However, the presence of five covalent bonds in the plane mismatches the mutual orientations of the lobes of the AOs on Pt, making the picture counterintuitive. On the other hand, if one assumes the complete population of the 5d set and the 6s AOs and their overall spherical symmetry, then the H and Zn atoms would form an octahedral arrangement around Pt, not the pentagonal pyramidal arrangement. Thus, the five covalent bonds AdNDP picture is confusing. We therefore prefer the σ-aromaticity view of the studied hydride.

Aromaticity is a highly stabilizing effect, also responsible for high symmetry of the system. Therefore, we attribute the high abundance of the studied cluster in the mass spectrum and the apparent stability to σ-aromaticity. Our observation of the σ-aromatic arrangement of hydrides bound to metal clusters is important for catalysis applications. One may imagine that a cluster capable of hosting multiple H atoms that hold just the right charge may exhibit aromaticity and thus be unusually stable on the reaction profiles of the catalyzed reaction of dehydrogenation. This, however, remains to be demonstrated.

### Computational Methods
The search for the global minima of the studied cluster anions was done using the gradient-embedded genetic algorithm (GEGA), with the population size of 20, and the UPBEPBE/LANL2DZ method. Separate searches for several spin states were performed. The lowest-energy isomers were characterized at higher levels of theory, UPBEPBE/aug-cc-pVTZ+pp, UM06, single-point CCSD/UPBE, and CASSCF(m,n), with (m,n) being as large as (12,12), with the same basis set. It was found that the perturbation theory fails for the studied systems, and for this reason, MP-based methods and CCSD(T) were not used. The combination of Pople’s 6-311++G(d) basis for Zn and H and the LANL2DZ basis for Pt were also used to make sure that there is no unhealthy dependence on the choice for the basis set. For calculations of photoelectron spectra, linear response time-dependent DFT, TD-UPBEPBE, with different basis sets, and also CCSD/aug-cc-pVTZ+pp/UPBEPBE/aug-cc-pVTZ+pp were used. TD-DFT was used for calculations of the excited states of the neutral clusters, and then, the excitation energies were added to the value of the first vertical detachment energy (VDE$_0$) to produce VDE$_x$, VDE$_y$, and so forth. All calculations were done with Gaussian 09.

### Associated Content
#### Supporting Information
Calculated energy differences between isomers I and II and calculated properties of isomers I and II. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Notes**
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REFERENCES


