Aromaticity

δ Aromaticity in \([\text{Ta}_3\text{O}_3]\) **

Hua-Jin Zhai, Boris B. Averkiev, Dmitry Yu. Zubarev, Lai-Sheng Wang,* and Alexander I. Boldyrev*

The concept of aromaticity was introduced into organic chemistry to describe delocalized π bonding in planar, cyclic, and conjugate molecules possessing \((4n+2)\) π electrons.\[8\] In recent years, this concept has been advanced into main-group molecules including organometallic compounds with cyclic cores of metal atoms\[9\] and, in particular, all-metal clusters.\[10\] It has been shown that main-group clusters may exhibit multiple aromaticity (\(\delta\) and π), multiple antiaromaticity (\(\delta\) and π), and conflicting aromaticity (\(\delta\) aromaticity and π antiaromaticity or π aromaticity and \(\delta\) aromaticity).\[11,12\] Here, we report experimental and theoretical evidence of δ aromaticity, which is only possible in transition-metal systems. It is discovered in the \([\text{Ta}_3\text{O}_3]^−\) cluster through combined photoelectron spectroscopy and ab initio studies. Well-resolved low-lying electronic transitions are observed in the photoelectron spectra of \([\text{Ta}_3\text{O}_3]^−\) and are compared with ab initio calculations, which show that the \([\text{Ta}_3\text{O}_3]^−\) cluster has a planar \(D_{3h}\) triangular structure. Chemical-bonding analyses reveal that among the five valence molecular orbitals involved in the multicenter metal–metal bonding, there is a completely bonding \(\delta\) and π orbital formed from the 5d atomic orbitals of Ta. The totally delocalized multicenter \(\delta\) bond renders δ aromaticity for \([\text{Ta}_3\text{O}_3]^−\) and represents a new mode of chemical bonding. \([\text{Ta}_3\text{O}_3]^−\) is the first δ-aromatic molecule confirmed experimentally and theoretically, which suggests that δ aromaticity may exist in many multinuclear, low-oxidation-state transition-metal compounds.

In 1964, Cotton and co-workers published a milestone work on \(K_2[\text{Re}_2\text{Cl}_8]\cdot2\text{H}_2\text{O}\)[7] in which they showed the presence of a new type of chemical bond—a δ bond between the two Re atoms. Since then, a branch of inorganic chemistry has been developed that involves multiple metal–metal bonding\[9\] with bond orders higher than three, the maximum allowed for main-group systems. Power and co-workers recently reported the synthesis of a \(\text{Cr}_2\) compound with a quintuple bond (\(\delta^4\pi^2\)) between the two \(\text{Cr}\) atoms.\[9\] This work, along with recent quantum chemical studies of multiple bonds in \(\text{U}_2\) and \([\text{Re}_2\text{Cl}_8]^−\)[10] has generated renewed interest in multiple metal–metal bonding.\[11–13\] The presence of δ bonds between two transition-metal atoms suggests that multicenter transition-metal species with a completely delocalized cyclic δ bond may exist, thus raising the possibility of δ aromaticity analogous to π or \(\delta\) aromaticity in main-group systems. We have been interested in understanding the electronic structure and chemical bonding of early transition-metal oxide clusters as a function of size and composition, and in using them as potential molecular models for oxide catalysts.\[14–16\] During our investigation of tantalum oxide clusters, we found the presence of δ aromaticity in the \([\text{Ta}_3\text{O}_3]^−\) cluster, in which each Ta atom is in a low oxidation state of Ta\[\text{II}\] and still possesses three electrons for Ta–Ta bonding.

The experiment was conducted by using a magnetic-bottle-type photoelectron spectroscopy apparatus equipped with a laser vaporization cluster source.\[17\] \([\text{Ta}_3\text{O}_3]^−\) clusters with various compositions were produced by laser vaporization of a pure tantalum disk target in the presence of a helium carrier gas seeded with \(\text{O}_2\) and were size-separated by time-of-flight mass spectrometry. The \([\text{Ta}_3\text{O}_3]^−\) species was mass-selected and decelerated before photodetachment by a pulsed laser beam. Photoelectron spectra were obtained at two relatively high photon energies, 193 nm (6.424 eV) and 157 nm (7.866 eV), to guarantee access to all valence electronic transitions (Figure 1). Three well-resolved bands (X, A, and B) were observed at the lower-binding-energy side. The X band is much more intense and shows a discernible splitting at 193 nm (Figure 1a). Surprisingly, no well-defined electronic transitions were observed beyond 3.7 eV, where continuous signals were present, probably as a result of multielectron transitions. The vertical detachment energies (VDEs) of the observed transitions at the low-binding-energy side are given in Table 1, where they are compared with theoretical calculations by two different methods.
We initially performed an extensive search for the \([\text{Ta}_3\text{O}_3]\) global minimum for the singlet, triplet, and quintet states at the B3LYP/LANL2DZ level of theory, and then recalculated the global minimum structure and the three lowest isomers at three other levels of theory (see the Supporting Information for references, details of theoretical calculations, and more theoretical results). We found that the \([\text{Ta}_3\text{O}_3]\) global minimum has a perfect \(D_{\text{inh}}\) (1\(A_1\)) planar triangular structure I (Figure 2). The closest isomer II is 6.6 kcal mol\(^{-1}\) higher in energy than the \(D_{\text{inh}}\) ground state. The theoretical VDEs of the global minimum at the two highest levels of theory are compared with those calculated for the \(D_{\text{inh}}\) global minimum.

![Figure 1. Photoelectron spectra of \([\text{Ta}_3\text{O}_3]\) - a) 193 nm (6.424 eV); b) 157 nm (7.866 eV).](image)

![Figure 2. Optimized structures for the global minimum of \([\text{Ta}_3\text{O}_3]^+\) (\(D_{\text{inh}}, 1\(A_1\)) and selected low-lying isomers. The relative energies \(\Delta E_{\text{rel}}\) [kcal mol\(^{-1}\)] and interatomic distances [\(\text{Å}\)] were calculated at the B3LYP/Ta/Stuttgart + 2\(f\)g/O/aug-cc-pVTZ level of theory (\(\Delta E_{\text{rel}}\) at the B3LYP/Ta/Stuttgart + 2\(f\)g/O/aug-cc-pVTZ level is shown in brackets).](image)

<table>
<thead>
<tr>
<th>VDE (exp.)</th>
<th>Final state and configuration</th>
<th>VDE (B3LYP) ([\text{kcal mol}^{-1}])</th>
<th>VDE (B3PW91) ([\text{kcal mol}^{-1}])</th>
</tr>
</thead>
<tbody>
<tr>
<td>X 2.25 (\pm) 0.03 ([\text{eV}])</td>
<td>1(E) (3(a_2)(\bar{2})(a_1)(\bar{4})(a_2)(\bar{4})(e))</td>
<td>0.52</td>
<td>2.27</td>
</tr>
<tr>
<td>A 2.39 (\pm) 0.02</td>
<td>2(a_2) (3(a_2)(\bar{2})(a_2)(\bar{4})(a_2)(\bar{4})(e))</td>
<td>2.26</td>
<td>2.96</td>
</tr>
<tr>
<td>B 3.44 (\pm) 0.01</td>
<td>2(a_1) (3(a_2)(\bar{2})(a_2)(\bar{4})(a_2)(\bar{4})(e))</td>
<td>3.27</td>
<td>3.36</td>
</tr>
</tbody>
</table>

[a] Using the Ta/Stuttgart + 2\(f\)g/O/aug-cc-pVTZ basis set. [b] The adiabatic electron-detachment energy was measured to be (2.22 \(\pm\) 0.03) eV.

To help understand the structure and bonding in \([\text{Ta}_3\text{O}_3]^+\) we performed a detailed molecular orbital (MO) analysis. Out of the 34 valence electrons in \([\text{Ta}_3\text{O}_3]^+\), 24 belong to either pure oxygen lone pairs or those polarized towards Ta (responsible for the covalent contributions to Ta–O bonding). The remaining ten valence electrons are primarily Ta-based and are involved in direct metal–metal bonding (Figure 3). Among the five MOs, three are responsible for o bonding of the triangular Ta framework. They include the partially bonding/antibonding doubly degenerate 4e orbital HOMO and the lowest HOMO-1. The antibonding nature of the HOMO significantly reduces the \(\pi\)-bonding character according to the \((4n+2)\) Hückel rule for \(\pi\) aromaticity.[10]

The most interesting MO is HOMO-1 (4\(a_1\)), which is a completely bonding orbital that comes mainly from the overlap of the \(d_{\pi}\) orbital on each Ta atom. This orbital has the “appearance” of a \(\pi\) orbital with major overlaps above and below the molecular plane, but it is not a \(\pi\)-type MO because it is symmetric with respect to the molecular plane. However, perpendicular to the molecular \(C_3\) axis this MO has two nodal surfaces, and thus it is a \(\delta\) orbital.[29] In fact, a similar \(\delta\)-bonding MO exists in the recently synthesized quintuple-bond Cr\(_3\) complex,[30] in which it is a two-center bond formed from a \(d_{\pi}\) orbital on each Cr atom.[31] Analogous to the
Aromaticity in transition-metal systems has been discussed in the literature,[4,5,21–30] particularly since the discovery of aromaticity in all-metal clusters.[31,32] King[32] and Li[33] have considered aromaticity in transition-metal oxides as a result of metal–metal interactions through M-O-M bridges. The \([\text{Hg}_2\text{O}_2]^-\) cluster, which is a building block of the \([\text{Na}_3\text{Hg}_2]\) amalgam, has been shown by Kuznetsov et al.[34] to be aromatic and similar to the all-metal \([\text{Al}_2\text{Cl}_2]^-\) unit.[35] Tsipis et al.[36] explained the planar structure of cyclic coinage-metal hydrides on the basis of their aromatic character. Aromaticity in square-planar coinage-metal clusters was discussed by Wannere et al.[37] and Lin et al.[38] and Alexander et al.[39] suggested the presence of aromaticity in the \([\text{Cu}_2\text{Cl}_4]^+\) cluster. Datta et al.[40] used d-orbital aromaticity to explain the metal-ring structure in tiara nickel thiolates. Recently, Huang et al.[41] demonstrated the presence of d-orbital aromaticity in the 4d and 5d transition-metal-oxide clusters \([\text{Mo}_3\text{O}_9]^{2-}\) and \([\text{W}_3\text{O}_9]^{2-}\). The claim of d-orbital aromaticity in the square-planar coinage-metal clusters[27] was questioned by Lin et al.,[28] who showed that the completely filled d orbitals do not play any significant role in the bonding in these clusters. Instead, aromaticity in these systems comes primarily from \(\sigma\)-bonding interactions of the valence \(\sigma\) electrons. Thus, today the \([\text{Mo}_3\text{O}_9]^{2-}\) and \([\text{W}_3\text{O}_9]^{2-}\) clusters are the only examples in which aromaticity comes from \(\sigma\)-bonding interactions, albeit with \(\sigma\) character.[21]

In the \([\text{Ta}_3\text{O}_3]\)^- cluster, we have found two new types of \(\delta\)-bonding interactions that lead to \(\pi\) and \(\delta\) aromaticity. The \(\delta\) aromaticity in this cluster is a new mode of chemical bonding that can only occur in multinuclear transition-metal systems. The current finding suggests that \(\delta\) aromaticity may exist in many cyclic transition-metal systems containing metal atoms in low oxidation states. The next challenge is to find \(\phi\) aromaticity, which may occur in multinuclear and cyclic f-metal systems.

Keywords: ab initio calculations · aromaticity · cluster compounds · metal–metal interactions · photoelectron spectroscopy

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[18] If the HOMO (4e') and the HOMO-3 (3a1') were composed of the same s–d hybrid functions they would cancel each other, thus resulting in negligible metal–metal σ bonding. However, the hybridization in the 4e' and 3a1' orbitals is somewhat different. Therefore, we cannot rule out some σ-bonding contribution in the Ta3 framework; that is, there should be some σ-aromatic character in [Ta3O3]4-.

[19] In the case of multiple aromaticity, the (4n+2) counting rule should be applied separately for each type of aromaticity encountered in a particular planar system, that is, separately for σ-, π-, δ-, and ϕ-type MOs.[14]

[20] Strictly speaking, the σ, π, δ, and ϕ notations for MOs are only appropriate for linear systems, where they are irreducible representations of the C∞ and D∞h point groups. However, it is customary in chemistry to use π notation in planar molecules for MOs that are formed by the p atomic orbitals and are perpendicular to the molecular plane, even though they do not belong to the π-irreducible representation. For example, the orbitals responsible for the aromaticity in the prototypical aromatic molecule C6H6 are called π orbitals. Following this tradition, one can introduce δ- or ϕ-type MOs in planar molecules formed from appropriate atomic orbitals.


