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Communication

CO oxidation on the heterodinuclear tantalum-nickel monoxide carbonyl complex anions

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ABSTRACT

the pulsed-laser vaporization source and characterized by mass-selected photoelectron velocity-map spectroscopy. During the consecutive CO adsorption, the μ^2 -O-bent structure initially is the most favorable for TaNiO(CO)₅⁻, and subsequently both μ^2 -O-bent and μ^2 -O-linear structures are degenerate for TaNiO(CO)₆⁻, then the μ^2 -O-linear structure is most preferential for TaNiO(CO)₇⁻, and finally the η^2 -CO₂-tagged structure is the most energetically competitive one for TaNiO(CO)₈⁻, *i.e.*, the CO oxidation occurs at n = 8. In contrast to the literature reported CO oxidation on heteronuclear metal oxide complexes generally proceeding via Langmuir-Hinshelwood-like mechanism, complementary theoretical calculations suggest that both Langmuir–Hinshelwood-like and Eley–Rideal-like mechanisms prevail for the CO oxidation reaction on $TaNiO(CO)_8^-$ complex. Our findings provide new insight into the composition-selective mechanism of CO oxidation on heteronuclear metal complexes, of which the

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The low-temperature oxidation of CO has received considerable attention from the environmental and material scientists. due to its important role in controlling of vehicle exhaust emissions and purification of gas streams derived from petrochemical industry [1–3]. Particularly interesting are the nanoscale catalysts, the composition and structure of which can be tailored to fulfill the desired chemical behaviors [4-6]. With the aid of state-of-art in-situ spectrum technologies, two dynamically distinct types of bimolecular surface reactions have been proposed in the heterogeneous catalysis of CO oxidation, which are now denoted as Langmuir-Hinshelwood (LH)- and Eley-Rideal (ER)-type mechanism, respectively [7]. In the LH mechanism, both the CO molecule and oxygen species are coadsorpted on the catalyst surface, and subsequently, the CO₂ is formed by either the rearrangement between chemisorbed CO and oxygen, or direct attachment of the

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CO ligand to the near-neighbor oxygen centers (intrasystem attack) [8]. On the contrary, in the ER mechanism, the CO molecule from the gas phase reacts directly with the chemisorbed oxygen species on the catalyst surface to give rise to CO_2 (intersystem attack) [8].

Microscopically, the real-life catalytic event usually takes place on a specific active site consisting of only a small number of atoms, of which the particular electronic, geometric, and bonding properties are the root of selectivity of CO oxidation mechanism. The gas-phased cluster reaction researches performed under isolated, size-controllable, and reproducible conditions provide an alternative route to capture clear molecular- and electronic-level mechanisms of the catalytic CO oxidation. The CO oxidation of the binary transition metal oxide (TMO) clusters serves as a welldefined model pertinent to mechanistic understandings of the surface catalysis. Recent researches of particular interest are the heteronuclear TMO clusters and highlight the potential to modulate chemical processes by selective cluster doping [9,10]. The different metallic fractions of heteronuclear TMO clusterscan mimic either the individual active sites or their supports of real-life catalysts [11]. The well similar behaviors paralleled with the condensed-phase CO oxidation by supported catalysts have been

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The series of heterodinuclear metal oxide carbonyls in the form of $TaNiO(CO)_n^-$ (n = 5-8) are generated in composition be tailored to fulfill the desired chemical behaviors.

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found in the gas-phase catalytic CO oxidation mediated by nanocatalysts doped by noble-metal single-atom and dimer [12–15]. The noble-metal-like behavior in many catalytic processes has also been found for a few noble-metal-free heteronuclear TMO clusters [14–16]. The high adsorption energy of CO on the atomic or dimeric dopants and dynamic nature of dopants in terms of the electron storage and release are found to be the driving force for the CO oxidation [17]. Due to the strong carbonyl binding to the doped TM, especially for the noble TM, the atomic TM dopant usually functions as a preferred trapping site and electron acceptor for CO adsorption and then acts as a deliverer of CO ligand for CO oxidation by the oxygen species on the heteronuclear TMO cluster. In this sense, available gas-phase experiments in the literature indicate that the CO oxidation reaction on heteronuclear TMO clusters preferentially proceeds *via* LH-like mechanism.

The recently reported literatures have successfully identified ligand-mediated reactivity in the CO oxidation of V-doped [16] and Nb-doped [18] nickel oxide complexes, on which the CO oxidation can occur by LH-like mechanism. Inspired by these facts, we extend our research to their tantalum congeners $TaNiO(CO)_n^-$ (n = 5-8). The details of experiment and computation can be found in the Supporting information. Fig. 1 illustrates the photoelectron spectra of the $TaNiO(CO)_n^-$ (n = 5-8) series obtained at 266 nm. A tabulated summary of the experimentally measured data is given in Table S2 (Supporting information), where they are compared with the theoretically predicted value at the BP86-D3BJ/def2-TZVP level. The spectral signatures are distinguishable from each other for the



Fig. 1. Photoelectron spectra of $TaNiO(CO)_n^-$ (n = 5-8) recorded at 266 nm (4.661 eV).

TaNiO(CO)_n⁻ (n = 5-8) series, despite the similarly broad bands. One broad band without discernible feature is observed in the spectrum of TaNiO(CO)5⁻, suggesting a lager structural change upon photodetachment for this anion. The main band in the spectrum of TaNiO(CO)₆⁻ is obviously split into two peaks located at 3.42 and 3.91 eV, with a slightly higher intensity in the second peak. The spectrum of $TaNiO(CO)_7^-$ is split into two peaks located at 3.74 and 4.29 eV, with the first peak becoming dominant. Moreover, the ground-state VDE¹ of TaNiO(CO)₇ $^-$ is blue-shifted by over 0.3 eV, as compared to those of TaNiO(CO)₅⁻ and TaNiO(CO)₆⁻. Such appreciable differences in the spectral feature imply that TaNiO(CO)₇⁻ should have different geometry from TaNiO(CO)_n⁻ (n = 5-6), as validated by the undermentioned theoretical calculations. However, only one broad band with the maximum located at 3.86 eV is revealed in the spectrum of $TaNiO(CO)_8^-$. Overall, the similarities and differences revealed in the experimental spectra, together with band shift, imply the emergence of geometric evolution during the consecutive CO adsorption on tantalumnickel monoxide complexes.

Quantum chemical calculations were carried out to elucidate the geometric and electronic properties of the TaNiO(CO)_n⁻ (n = 4-8) series. The ground state candidates and alternative structures optimized at the BP86-D3BJ/def2-TZVP level are all closed-shell species and depicted in Fig. S1 (Supporting information). As present in Fig. 2, three kinds of representative geometries with distinct structural characteristics are predicted to participate in the competition of the possible ground states during the continuous CO adsorption progress. The first kind structure in the form of $(CO)_xTa-(\mu^2-O)-Ni(CO)_{n-x}$ is labeled with *n*A and comprises of a



Fig. 2. Ground-state structures and selected low-lying isomers of the $TaNiO(CO)_n^-$ (n = 4-8) anions calculated at the BP86-D3BJ/def2-TZVP level. Relative energies are given in square brackets (eV).

triangular TaONi⁻ anion core and terminal CO ligands bonded to metal atoms. The oxygen atom in the TaONi⁻ core occupies the bridging position between Ta and Ni atoms. We define this type of geometry as a μ^2 -O-bent structure. The transformation of TaONi⁻ anion core from triangular to linear will create the second typed structure in the form of $(CO)_x$ Ta $-(\mu^2-O)$ -Ni $(CO)_{n-x}$. This kind of structure is defined as μ^2 -O-linear structure, which involves a linear TaONi⁻ anion core terminally bonded by multiple CO ligands. The third type geometry labeled with *n*C has a bent CO₂ ligand bonded to the Ta center in a side-on η^2 -C, O fashion, besides *n*-1 CO ligands either terminally or bridged bonded to the metal atoms. We define this type of structure as a η^2 -CO₂-tagged structure.

It can be seen from Fig. 2 that these three types of structures energetically compete with each other during the consecutive CO adsorption on tantalum-nickel monoxide complexes. For TaNiO(CO)₄⁻, the most stable isomer is the μ^2 -O-bent structure with triangle TaONi core (4A). Both the μ^2 -O-linear isomer (4B) involving linear Ta-O-Ni center and the η^2 -CO₂-tagged isomer (4C) containing a CO₂ ligand side-on bonded to Ta atomare thermodynamically unfavorable, which are 1.53 and 1.42 eV higher in energy than 4A, respectively. The large energy spacings hint at the overpowering stability of the μ^2 -O-bent isomer. Additional attachment of CO ligands to the metal center successively generates higher-coordinated TaNiO(CO)_n⁻ (n = 5-8) complexes with similar geometric characteristics. However, multiple adsorptions of CO ligands can vary both the relative stability and the energy separations among these three types of isomers. The energy spacing between μ^2 -O-bent structure and μ^2 -O-linear structure dramatically decreases when one CO terminally attaches to $TaNiO(CO)_4^-$. For $TaNiO(CO)_5^-$, the ground-state isomer is predicted to be a μ^2 -O-bent structure (5A), followed by μ^2 -Olinear isomer (5B) and η^2 -CO₂-tagged isomer (5C) being 0.14 and 1.03 eV higher in energy, respectively. Additional CO adsorption leads to a further decrease in energy separation and results in two near degenerate isomers 6A and 6B, with isomer 6B being slightly higher in energy. Therefore, the global minimums for $TaNiO(CO)_n^{-1}$ (n=4-6) are all μ^2 -O-bent structures, as similar with the congeneric NbNiO(CO)_n⁻ (n = 4-6) series [18]. However, an inversion of the relative stability of the μ^2 -O-bent structure and μ^2 -O-linear structure is achieved when one CO ligand attaches to TaNiO(CO)₆⁻. For n = 7, the thermodynamically most stable structure is the μ^2 -O-linear structure (7B), which is followed sequentially by μ^2 -O-linear isomer (7A) and η^2 -CO₂-tagged isomer (7C). The energy difference between 7B and 7A/7C is 0.20/0.46 eV. At this point, the situation for $TaNiO(CO)_7^-$ anion is different. For previously reported congener NbNiO(CO)7⁻ anion, it has already been transformed into the η^2 -CO₂-tagged structure [18]. Such an alteration in the structure occurs again when one more CO ligand is adsorbed onto TaNiO(CO)7⁻. The relative energetics of the isomers suggest that the ground state of $TaNiO(CO)_8^-$ is the most competitive isomer 8C, *i.e.*, η^2 -CO₂-tagged structure, which is 0.62 and 0.51 eV lower in energy than the μ^2 -O-bent isomer (8A) and μ^2 -O-linear structure (8B), respectively. In conclusion, during the consecutive CO adsorption, the μ^2 -O-bent structure theoretically is the most favorable for TaNiO(CO)_n⁻ (n = 4-6), then the μ^2 -O-linear isomer is more preferential for $TaNiO(CO)_7^-$, and finally the η^2 -CO₂-tagged geometry is predicted to be the most energetically competitive one for $TaNiO(CO)_8^{-}$.

Table S2 provides a comparison of the theoretical calculated VDEs and ADEs at the BP86-D3BJ/def2-TZVP level and the experimental data. Considerable reorganization energies (ROEs) are predicted for all of the complexes, correlating well with the experimentally observed broad band in the PES. The ROE is defined as the energy difference between the ground-state ADE and VDE, the amount of which roughly characterizes the anion-to-neutral

structural relaxation upon electron detachment. As an aside, it is interesting to note that the predicted VDEs for the *n*A isomers with a triangular TaONi⁻ anion core and *n*C isomers involving a bent CO₂ ligand are approximately located at around 3.4 and 3.7 eV, respectively. While for the μ^2 -O-linear structure, the VDE roughly increases with the multipleadsorptions of CO ligands.

For the PES of $TaNiO(CO)_n^-$ (*n* = 5–7) anions, the intensity of η^2 -CO₂-tagged isomer 5C, 6C and 7C are anticipated to be thermodynamically negligible and therefore can be excluded. due to their overestimated ADEs and VDEs and high relative energy of 1.03, 0.64 and 0.46 eV. Moreover, as can be seen from Fig. S2 (Supporting information), their simulated spectra disagree with the experimental PES. The major competition occurs between the μ^2 -O-bent structure and μ^2 -O-linear structure. The ground-state structure of TaNiO(CO)₅⁻ is μ^2 -O-bent isomer 5A, of which the predicted ADE and VDE are 3.01 and 3.45 eV, respectively, in accordance with experimental values. The ADE and VDE of isomer 5B are estimated to be 3.17 and 3.27 eV, respectively. The comparatively small ROE suggests that the μ^2 -O-linear structure is relatively rigid. The situation is slightly more complicated for TaNiO(CO)₆⁻. The μ^2 -O-bent isomer 6A and μ^2 -O-linear isomer 6B are predicted to be energetically near-degenerate, with 6A locating slightly lower in energy. The ground-state ADE and VDE of isomer 6A are 3.10 and 3.41 eV, respectively, in good agreement with the experiment. The theoretical first excited state VDE² (4.11 eV) is in line with the second peak observed in the PES. The VDE of isomer 6B is predicted to be 3.64 eV, which is a little bit blue-shifted with respect to the experimental data. For TaNiO(CO)₇, only the ground-state ADE and VDEs of isomer 7B are excellently consistent with the experimentally measured values. Moreover, the predicted VDE (4.26 eV) of the first excited state for isomer 7B excellently agrees with the second peak in the experimental PES. However, the ADE and VDE of isomer 7A are underestimated with respect to the experiment, while the ones for isomer 7C are overestimated. For TaNiO(CO)₈⁻, the experimental PES can be assigned to the structure 8C, for which the predicted ADE and VDE are in good agreement with the experimental data. Alternative isomers, such as μ^2 -O-bent isomer 8A and μ^2 -O-linear isomer 8B, are thermodynamically unfavorable and therefore excludable. Furthermore, the VDE of isomer 8A is severely underestimated relative to the experimental value, while the ADE of isomer 8B is overestimated.

Additionally, the spectra have been simulated based on the theoretically modeled excitation energies of the three types of isomers using time-dependent density functional theory (TDDFT). The simulated spectra are depicted in Fig. S2 and compared with the experimental PES. For the competitive isomers 5A and 5B, only the simulated PES of isomer 5A matches the experimental one. The fit PES of 5B has two separate peaks, which disagrees with the experimental PES. Thus, the observed PES is derived from the photodetachment of isomer 5A anion. For $TaNiO(CO)_6^-$, both of competitive isomers 6A and 6B have two partially overlapping peaks. As compared with isomer 6B, the first peak of isomer 6A fit more with the observed peak X, while the secondary peak is blueshifted with respect to the experimental peak A. Moreover, the ratio of the simulated two peaks mismatches the experimental PES. For the case of 6B, both two peaks are blue-shifted with respect to the experimental peaks. However, despite that, the ratio of the fitted two peaks agrees with the experiment. In this sense, both of the near-degenerate isomers potentially coexist and contribute to the experimental spectrum. Similarly, two separate peaks are modeled for both isomers 7A and 7B. However, only the spectral pattern of isomer 7B is in accordance with experimental PES of TaNiO(CO) $_7^-$, while the second of two peaks is dominant for isomer 7A, which deviates from the observed spectrum. Thus, both the predicted ADE and VDE and simulated PES suggest μ^2 -O-linear structure 7B as the ground-state structure of TaNiO(CO)₇⁻. For TaNiO(CO)₈⁻, only the simulated spectrum of ground-state 8C reasonably coincides with experimental observation. While for the isomers 8A and 8B, the simulated spectrum has either two separate peaks or only one deviated peak, which is found to be incompatible with the experiment. Therefore, the observed PES belongs to the n^2 -CO₂-tagged isomer 8C. Comprehensive comparison between the experiment and theory coincidentally demonstrate that, for the consecutive CO adsorption process of the TaNiO(CO)_n⁻ (n = 5-8) series, the μ^2 -O-bent structure is initially preferential, then the μ^2 -O-linear structure is thermodynamically favorable, and finally, the η^2 -CO₂-tagged structure becomes dominant. The details of comparison can be found in the electronic supplementary information. It is demonstrated that structural transformation occurs during the successive CO adsorption, and the CO oxidation happens at n = 8.

To rationalize the experimental results and provide insights into the dynamic reactivity of $TaNiO(CO)_n^-$ complexes toward CO oxidation, the structural isomerizations among three types of isomers were theoretically analyzed at the BP86-D3BJ/def2-TZVP level. For $TaNiO(CO)_n^-$ (n = 5-7), the potential energy profiles of the structural isomerizationsare shown in Fig. 3. The left part of Fig. 3 (shown in red) are potential energy profiles of the isomerizations

Fig. 3. DFT calculated potential energy profiles for the isomerization reactions $(nB \rightarrow nA \rightarrow nC)$ of TaNiO(CO)_n⁻ (n = 5-7). The zero-point vibration corrected energies (eV) with respect to the separated reactants (n-1)A and CO are given. The gray dot-dashed lines indicate the energetic baselines ((n-1)A + CO).

between the μ^2 -O-bent isomer and the μ^2 -O-linear isomer. The isomerization mainly involves a geometrical transformation of the TaONi core, namely, the variation of TaONi bond angle or Ta-Ni bond length. The low energy barriers for the isomerization step correlate well with the small Mayer bond orders of Ta-Ni bond summarized in Table S3 (Supporting information). There is almost no barrier for the isomerization reaction from isomer 5B to the more stable isomer 5A. Whereas, there is no barrier for the inverse reaction at *n* = 7. *i.e.*, the isomerization from isomer 7A to the more stableisomer 7B. These explain the absence of isomer 5B and isomer 7A in the PES of TaNiO(CO)₅⁻ and TaNiO(CO)₇⁻, respectively. For n = 6, however, the isomerization reaction is almost thermoneutral between the degenerate isomers 6A and 6B, which are separated by the transition state TS_6^1 with a barrier of 0.13 eV. Both of them may present in the anionic beam of $TaNiO(CO)_6^-$. It is demonstrated that the presence of μ^2 -O-bent isomer and/or μ^2 -O-linear isomer is a consequence of quantitatively different reaction barriers and relative thermodynamic stability between these isomers.

The right part of Fig. 3 (shown in blue) illustrates the isomerizations from μ^2 -O-bent structure to η^2 -CO₂-tagged structure. The proposed pathways proceed via the LH-like mechanism, in which one CO molecule is firstly adsorbed onto one metal atom of the μ^2 -O-bent precursor, and then the resultant μ^2 -O-bent structure is transformed into a η^2 -CO₂-tagged structure through an intrasystem CO attack on the bridging oxygen atom. The isomerization pathway involves one transition state TS_n^2 leading to the intermediate I_n^1 , in which a CO₂ subunit is asymmetrically bound to the bridging position of the Ta-Ni axis, and second transition state TS_n^3 leading to the final η^2 -CO₂-tagged structure, in which the CO₂ moiety is only bound to the Taatom. Generally, the CO oxidation reaction proceeding via a LH mechanism has to satisfy two requirements: the sufficiently large carbonyl binding energy and the lower energetics of the transition states involved in the subsequent CO₂ moiety formation than the initial reactants. As can be seen from Fig. 3, the attack of a chemisorbed CO ligand on the bridging O atom to form the intermediate I_n^1 is the bottleneck for the CO oxidation by the heteronuclear $TaNiO(CO)_n^-$ complexes. This key step of the isomerization process involves the formation of one C-O bond along with the weakening of Ta-O bond. The next step, *i.e.*, the migration of the CO₂ moiety from the unsymmetrical bridging position of the Ta-Ni axis to the Ta site, involves the breaking of the Ni-O bond. That is to say, the activation of the TaONi core upon CO chemisorption is vitally important for the oxygen migration so as to facilitate the CO oxidation. However, despite the large exothermicity of CO chemisorption on the TaNiO(CO)_n⁻ (n = 5-7) complexes, the formation of η^2 -CO₂-tagged structures is impeded by the relatively high barrier of the transition states, some of which are even higher with respect to the separate reactants. This explains why the η^2 -CO₂-tagged isomer is not observed experimentally for TaNiO(CO)_n⁻ (n = 5-7).

The pathways of CO oxidation on the TaNiO(CO)₇⁻ complex are also evaluated to understand the evolution from the CO association to the CO oxidation at n = 8. Fig. 4a shows the potential energy profiles of the CO oxidation reaction on the isomer 7B. Two distinct reaction channels are identified for the formation of the CO₂ unit. The very first reaction step is the formation of the weakly bound complex, in which the fresh CO molecule can be either chemisorbed onto the Ta atom to form isomer 8B or physisorbed onto 7B to generate intermediate I_8^1 . As opposed to the strongly exothermic association of CO onto TaNiO(CO)_n⁻ (n = 4-6) complexes, the CO adsorption on isomer 7B is almost thermoneutral, suggesting that the adsorption of CO on TaONi core presumably reaches saturation. These CO association complexes are separated





Fig. 4. DFT calculated potential energy profiles for the CO oxidation reactions starting from 7B + CO (a) and 7A + CO (b). The zero-point vibration corrected energies (eV) with respect to the separated reactants are given. The dot-dashed line indicates the energetics of reactants 7A and CO, while the dashed line stands for the energetics of the isomer 7B and CO. For the sake of distinction, the C and O atoms of the freshly adsorbed CO ligand are depicted in different colors from the rest of CO ligands.

by transition states TS_8^1 and TS_8^2 , respectively, from the same intermediate I_8^2 in which the freshly adsorbed CO is obliquely attached to the bridging oxygen atom to form a bent CO₂ unit. The freshly chemisorbed CO ligand to the Ta atom bends forwards to the bridging oxygen atom, leading to the formation of C-O bond in the transition state TS_8^1 . The transition state TS_8^2 has a structure related to the initial complex with a physisorbed CO ligand much closer to counterpart. Apparently, the pathway for CO chemisorption leading to the formation of intermediate. I_8^2 involves an internal CO attack and belongs to the LH-like mechanism, while the other pathway for CO physisorption involves an intersystem CO attack and can be attributed to the ER-like mechanism. Note that the CO₂ subunit has already been formed at this critical juncture of intermediate I_8^2 . As compared with those of TaNiO(CO)_n⁻ (n = 5-7), the reaction energy barriers for the attack of CO ligand on the bridging O atom are dramatically reduced, suggesting that the TaONi core becomes significantly reactive towards CO oxidation upon additional CO chemisorption.

Then the reaction proceeds further to form the more stable CO_2 -tagged structure through the CO_2 migration. By the rotation of CO out of the TaONi core plane and shortening of Ta–Ni distance, the structure. I_8^2 can be further isomerized to important intermediate. I_8^3 , which requires surmounting a low barrier of 0.19 eV (TS₈³). Thereafter, the reaction proceeds following two different channels, which both involve the migration of CO_2 and the conversion of two CO ligands from the terminal to the bridging coordination. The coordination pattern evolutions of two CO ligands can happen step by step in one reaction channel, or simultaneously in the other reaction channel. The energetically

more favorable pathway involves firstly an almost no barrier step (TS_8^4) to form CO singly bridging stereochemistry, followed by a deviation of the CO₂ unit away from the Ni atom and a transition from terminal to bridging coordination for another CO ligand, which is connected with the barrier of 0.14 eV (TS⁶₈). The second reaction pathway involves the breaking of the Ni-O bond and the simultaneous formation of the doubly CO bridging stereochemistry. The corresponding transition state (TS⁵₈) is 0.24 eV higher in energy than the initial reactants 7B and CO. At first glance, the oxidation reaction might be prevented due to the transition states with barriers slightly higher than the separate reactants 7B and CO. However, the heights of the involved activation barriers are much smaller with respect to the cases of TaNiO(CO)_n⁻ (n = 5-7), which can be readily surmounted by thermal collision in the supersonic molecular beam [19]. In addition, the whole CO oxidation reaction is thermodynamically exothermic by 0.45 eV. Therefore, the CO oxidation on TaNiO(CO)₇⁻ via both ER- and LH-like mechanisms are facile.

For comparison, the CO oxidation reactions starting from 7A and CO are also calculated, provided that 7A exists, and the corresponding potential energy profiles are present in Fig. 4b. Firstly, the CO molecule can be physisorbed or chemisorbed to generate complex I_8^5 or 8A, respectively. For complex 8A, the CO oxidation reaction proceeds further following the similar pathways as TaNiO(CO)_n⁻ (n = 5-7), *i.e.*, firstly the attack of a chemisorbed CO ligand on the bridging O atom to form CO₂ subunit and secondly the change of coordination patterns of CO₂ moiety from a bridge-type coordination to a side-on coordination on only Ta atom. While the further approaching of freshly physisorbed CO in the complex I_8^5 to the bridging oxygen atom will give rise to the same

intermediate I_8^3 , which can react following the same pathways as the abovementioned (Fig. 4a). It is demonstrated that the important intermediate I_8^3 can be generated either from the 7A and CO *via* an ER-like mechanism or from 7B and CO *via* both an ERlike mechanism and a LH-like mechanism. Obviously, the reactions occur through an ER-like mechanism is more favorable, whereas the reactions *via* a LH-like mechanism may still not easily proceed, as the CO molecule will experience quite a higher reaction barrier (TS⁸₈) when the adsorbed CO approaches to the bridging oxygen atom.

As shown in Table S3 (Supporting information), the Mayer bond orders for the Ta-O bonds (1.17-1.77) in the oxygen-bridged isomers are larger than those of Ni-O bonds (0.45-0.78). The stronger the TM-O bond strength is, the higher the reaction barrier will be. The differences in the bond orders are closely correlated with the distinct energy barriers of the two transition states TS_n^2 and TS_n^3 (n = 5-7) involved in the structural rearrangement from CO association to CO oxidation, suggesting that the activation of the Ta-O bond involved in the first step of the proposed pathway is critical. The natural population analysis indicates that both Ta and Ni atoms behave as electron acceptors upon the consecutive CO adsorption. The partial charges of Ni and Ta atoms reach their minimal value at n = 7 and n = 8, respectively, while the partial charges on the bridging O atom are almost unchanged. As a result, the charge difference between the Ni atom and the bridging O atom has a sudden change at n = 7, while the abrupt decrease in the charge difference between the bridging O atom and the Ta atom occurs at n = 8. The decrease of charge separation in TM-O subunits, leading to smaller ionic character of the TM-O bonds, may decrease atomic oxygen dissociation energy [19], and thus facilitates the atomic oxygen migration and the corresponding CO oxidation occurring at n = 8. The variation of natural populations correlates well with the change in Mayer bond orders. The consecutive CO adsorption has minimal perturbation on Ni-O bond in terms of bond length and bond order until the TaONi core chemisorbs up to 7 CO ligands. Despite significant reduction in bond order of Ni—O bond already occurring at n = 7, the oxidation reaction is still difficult, because CO multi-adsorption do not dramatically weaken the Ta-O bond until CO adsorption reaches the critical number (n = 8). It is demonstrated that, in comparison with the congeneric NbNiO(CO)_n⁻ complexes at n = 7[18], one more CO ligand adsorption is needed to promote the CO oxidation on the TaNiO(CO) $_n^-$ series. The current work further sheds light on the concept of CO self-promotion in the CO oxidation.

The CO adsorption energy retained by the TMO clusters is deem to be the driving force for CO oxidation [20]. The gold oxide clusters with different charge states, bearing great diversity in CO binding energy, express a preference for an ER-like or a LH-like mechanism for CO oxidation on them [20]. The LH-like mechanism is generally considered to be preferential for the CO oxidation in the gas phase [21], because the gained energy from the initial CO chemisorption in the LH-like mechanism usually is larger than that from the initial CO physisorption in ER-like mechanism. For the initial process of consecutive CO adsorption, the CO physisorption energies on $TaNiO(CO)_n^-$ complexes are so small that the ER-like mechanism becomes uncompetitive relative to the LH-like mechanism. However, as shown in Fig. S3 (Supporting information), the step binding energies of CO to the O-bridged isomers monotonically decrease with the consecutive CO adsorption, and finally converge to the values close to the physisorption energies. Consequently, the chemisorption loses its competitive edge at n = 8, and both ER- and LH-like mechanisms prevail for the CO oxidation reaction on $TaNiO(CO)_8^-$ complex.

Herein we identified the compositional dependence of reactivity and selectivity of the reaction mechanism for the heteronuclear complexes, which is rooted in the inherent nature of transition metal components. This dramatic difference in reactivity can be attributed to the stronger TM-O bonds of the 5d metals with respect to those of the lighter 3d and 4d congeners. The dissociation energy of diatomic TaO (8.30 eV) [22] has been confirmed to be larger than the values of 6.48 and 7.53 eV for lighter VO [23] and NbO [24]. The high-level *ab initio* calculations with the relativistic and correlation effects taken into account have demonstrated that the fascinating chemistry of 5d metals with regard to various chemical processes such as CO oxidation and C-H bond activation is to a great extent caused by relativity [25,26]. This same reason was invoked to elaborate on the discrepant maximum carbonyl-coordination number of the vanadium family (V⁺, Nb⁺, Ta⁺) [27]. The relativistic mass velocity contraction of the 6s orbital of Ta results in the contracted Ta-O bond (Table S3 in Supporting information) with respect to the corresponding Nb-O bond [18]. As a result of the 6s orbital stabilization and 5d orbital destabilization for Ta, the ⁴F (3d³6s²) ground state of Ta is in contrast to the ⁶D (3d⁴6s¹) ground state of Nb. Consequently, the electron affinity of Ta atom (0.323 eV) [28] is significantly lower than those of lighter homologsV atom (0.526 eV) [28] and Nb atom (0.894 eV) [28]. Furthermore, the 6s orbital stabilization and 5d orbital destabilization for Ta are complemented by a significantly better overlap between the π orbitals of the CO ligand with the $5d_{\pi}$ orbitals of Ta and therefore greater π back-donation of electron density from the 5d_{π} orbitals of Ta, when compared to the $3d_{\pi}$ and $4d_{\pi}$ counterparts of V and Nb. Accordingly, the capability of net electron storage of Ta atom in $TaNiO(CO)_n^-$ complexes is less competitive with respective to that of Nb atom in congeneric NbNiO(CO) $_{n}^{-}$ [18]. As a consequence of the stronger Ta-O bond and the less competitive electron storage ability of Ta, the TaNiO(CO) $_n^-$ complexes are less reactive towards CO oxidation than corresponding NbNiO(CO)_n⁻ congeners and additional CO ligand adsorption is needed to drive the CO oxidation on TaNiO(CO)_n⁻ complexes. Note that the calculated Mayer bond orders of Ta-O bond in μ^2 -O-bent structures of $TaNiO(CO)_n^-$ (n = 4–7) fall in the range of 1.32–1.52, while the those for the μ^2 -O-linearisomers are even larger and range from 1.60 to 1.77, suggesting the double-bond character of the Ta-O bond. As compared to the Ta⁺ cation forming a seven-coordinate complex [27], the maximum carbonyl-coordination number of Ta atom in the anionic TaNiO(CO) $_n^-$ complexes should decrease to six. The Ni atom in TaNiO(CO)7⁻ has reached its chemisorption saturation limit and its minimal value of partial charge. Consequently, beyond TaNiO(CO)7⁻, additional CO ligand can either be physisorbed in order to retain the Ta-O double bond, or be accommodated via chemisorption so that the Ta-O double bond becomes a single bond to reach the chemisorption saturation limit. In other word, both the chemisorption and physisorption on $TaNiO(CO)_7^-$ are so comparable with each other that both ER- and LH-like mechanisms prevail for the CO oxidation reaction on $TaNiO(CO)_8^-$ complex.

In summary, the structure evolution of the TaNiO(CO)_n⁻ (n = 5-8) series is characterized by the photoelectron velocity-map spectroscopy combined with DFT calculations. Three different types of structures participate in the competition of ground state, with the μ^2 -O-bent structure firstly being most favor, then the μ^2 -O-linear structure being most preferential, and finally, the η^2 -CO₂-tagged structure being dominant. In contrast to the CO oxidation on congeneric V-doped [15] and Nb-doped [22] nickel oxide complexes, which proceed *via* an LH-like mechanism as a result of the multi-adsorption of CO ligands, complementary theoretical calculations reveal that both ER- and LH-like mechanisms generally become favorable and lead to the self-promoted

CO oxidation on $TaNiO(CO)_8^-$. Our findings shed new light on the role of the composition of heteronuclear metal complexes in the regulation of the reactivity and selectivity of the CO oxidation mechanism, which potentially benefits the rational design and development of the high-efficiency catalysts.

Declaration of competing interest

The authors declare no competing financial interest.

Acknowledgments

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at https://doi.org/10.1016/j. cclet.2020.05.029.

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