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Reactions of N_2O and CO on neutral $Rh_{10}O_n$ clusters: a density functional study†

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Density functional theory calculations were performed to identify product, reactant and intermediate dissociative/associative structures for the oxygen abstraction and addition reactions: $Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} + CO_2$, n = 1-5 and $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} + N_2$, n = 0-4 reactions. In the case of the oxygen abstraction reactions, the energetics of the reaction path were very similar in energy regardless of the number of oxygen atoms on the $Rh_{10}O_n$ cluster, whereas for the addition of oxygen to the $Rh_{10}O_n$ cluster, the reaction was found to become significantly less exothermic with each successive addition of oxygen.

1 Introduction

Transition metal clusters have been of interest for several decades as both models of surfaces¹ and entities in their own right, where their remarkable size-² and structure-dependent³⁻⁵ properties make them highly attractive as potential heterogeneous catalysts.⁶⁻⁸ Late transition metal clusters, especially those of rhodium have maintained particular interest over that time with a multitude of studies employing a variety of spectroscopic and computational methods to probe their structure and behaviour.

Many of these studies have been motivated by the use of rhodium in automotive three-way catalytic converters and the reactions of N_2O , $^{10-12}$ NO, $^{13-22}$ $CO^{23,24}$ and CO_2^{25} have been extensively investigated on both pure and doped $^{26-28}$ rhodium clusters. Given that spectroscopic methods such as infrared multiple photon dissociation spectroscopy (IRMPD) 29 use mass spectroscopy to identify the species of interest, they generally provide information only about the products of reactions and reactivity studies therefore often include computational investigation, 30,31 typically using density functional theory (DFT), but sometimes including higher level methods such as CCSD(T).

Two particular reactions that are of interest for automotive catalytic converters are the reduction of NO_x to N_2 and the oxidation of CO to CO_2 . Both of these reactions have specifically

been investigated on rhodium^{11,26} and platinum^{33,34} clusters. Aviles *et al.* undertook calculations to elucidate the N₂O reduction mechanism on small Rh_n clusters (n = 2–4), and determined that the N₂–O bond was readily broken by the Rh_n cluster, but the N₂ bond was not.¹² Yamamoto *et al.* used both IRMPD and DFT methods to determine reaction paths for the reduction of N₂O by Pt₆ and Pt₇ clusters.,³⁵ while Hernández and coworkers used DFT to elucidate the reaction of N₂O on Pt₈.³⁶

Yet despite these successes, computational investigation remains challenging, especially on large and/or late transition metal clusters: Even employing frozen core/effective core potential approaches, clusters still possess a large number of electrons. A wide range of spin multiplicities need to be considered. And many clusters have multiple, very different geometries that all occur within a small energetic range, 5,37,38 that is thermally accessible and/or within the error of computational methods. For some clusters, the manifold of low-energy geometries may be so dense and the ease of interconversion so facile, that the clear identification of a ground state is challenging. Rh₁₀ has been identified as one such fluxional molecule.³⁹ Such a dense manifold of structures, energies and spin multiplicities is additionally challenging when employing reaction path following methods such as Quasi-Synchronous transit and intrinsic reaction coordinate methods, as reaction paths may follow a comparatively broad path, where spin and cluster geometry may differ throughout a reaction.

In 2012, Yamada *et al.* undertook a mass-spectroscopic study on neutral rhodium clusters, Rh_n (n = 10-28). ⁴⁰ By injecting both N_2O and CO into their cluster source, they were able to demonstrate a full catalytic cycle and observed that the distribution of oxide products Rh_nO_m was determined by an equilibrium between the oxygen being transferred to the Rh_nO_m

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[†] Electronic supplementary information (ESI) available: Spreadsheet of reaction path energies, zipfile of reaction path structures. Structures are additionally available at https://zenodo.org/record/8428605. See DOI: https://doi.org/10.1039/d3cp04929a

m = 0-4 cluster from N₂O and oxygen being abstracted from the Rh_nO_m m = 1-5, and that the rate constants for the oxygen abstraction reaction were 2-3 orders of magnitude higher on clusters where $m \geq 4$ than those clusters with three or less oxygen atoms. They concluded that oxidized rhodium clusters were more effective as catalysts compared to bare clusters. Yamada's study used experimentally derived rate constants and did not include any calculated structures or reaction paths. Indeed the size of the rhodium clusters employed by Yamada et al., Rh_n (n = 10-28) makes DFT challenging.

In this work, we revisit the catalytic cycle presented by Yamada et al. for the smallest rhodium cluster size, Rh₁₀. As reaction path following on such flat potential energy surfaces (PES) is difficult, we employ a fully stochastic search procedure followed by DFT calculations to identify the reactants, products and the bond forming/breaking intermediate for both the oxygen transfer and oxygen abstraction reactions on the Rh₁₀ cluster.

2 Computational method

The geometry of Rh₁₀ was determined using the Kick³ stochastic structure generator supplied with 10 Rh atoms. 41,42 Following optimization, the lowest energy Rh₁₀ geometry was then employed as a fragment in further $Kick^3$ runs with n oxygen atoms in order to determine the lowest energy Rh₁₀O_n structures, for n = 1-5. A key point on each $Rh_{10}O_n \rightarrow Rh_{10}O_{n\pm 1}$ reaction path, is the species where the $N_2 \cdots O$ bond is being broken, or where the O···CO bond is being formed. To identify these intermediate species directly, N₂O and OCO fragments with elongated NO and CO bonds respectively, were used as input to Kick³ to identify the $Rh_{10}O_n$ CO and $Rh_{10}O_n$ N_2 intermediate species. 1000 stochastic geometries were generated for each stoichiometry.

The stochastically generated structures were optimized in a two-stage procedure: Firstly using the semi-empirical GFN1-xTB method, 43 maintaining the geometry of all fragments employed in the stochastic structure generation, before a full optimization using BP86/DZP and then re-optimization using the TZ2P basis set. Up to the 4p electrons of rhodium were treated using a frozen core. Structures were initially optimized in the octet multiplicity before re-optimization with each multiplicity from doublet to 20-tet. 18- and 20-tet structures were found to lie very high in energy and were consequently not considered in analysis. All calculations were undertaken using AMS2022.1 using scalar relativivity and applying an electronic temperature of 373.15 K.44

To confirm that the BP86^{45,46} functional is sufficient, we also tested the TPSS⁴⁷ functional with the D3(BJ)^{48,49} dispersion correction. The choice of functional made no significant difference to the relative ordering of different multiplicity Rh₁₀O_n structures, as shown by difference-of-difference values in the ESI.† GGA functionals are expected to only give qualitative information about barrier heights, due to their self-interaction error⁵⁰ and this is evident, with BP86 intermediate relative energies for oxygen abstraction reactions being consistently \approx 0.4 eV higher in energy than the TPSS-calculated values and BP86 product energies being consistently ≈ 0.5 eV lower than the TPSS-calculated values. For the oxygen addition reaction paths, the TPSS relative energies for intermediate and product species are lowered w.r.t. the BP86 paths by ≈ 0.6 and 0.3 eV respectively.

After optimization of Rh₁₀O_n, Rh₁₀O_n CO and Rh₁₀O_n N₂ structures, $Rh_{10}O_n \rightarrow Rh_{10}O_{n\pm 1}$ reaction paths were determined by geometric RMSD. For each reaction, the lowest energy $Rh_{10}O_n$ reactant was chosen, and the nearest $Rh_{10}O_{+1}$ geometric match was determined $(Rh_{10}O_{n+1} \text{ for } Rh_{10}O_n + N_2O \rightarrow$ $Rh_{10}O_{n+1}+ N_2$ and $Rh_{10}O_{n-1}$ for $Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} +$ CO₂). The intermediate species was determined in the same manner, by determining the RMSD with the reactant and then confirming RMSD with the product. After the addition of the first two oxygen atoms to Rh₁₀, multiple oxygen abstraction reaction paths are possible, corresponding to abstraction of different oxygen atoms, the path presented in this work is the one containing the most favourable (lowest energy) intermediate geometry.

It is not necessarily the case that the lowest energy $Rh_{10}O_n$ will lead to the lowest energy $Rh_{10}O_{n\pm 1}$ species, and because of the circular nature of the catalytic reaction, each $Rh_{10}O_n$ species appears as both a product and a reactant. Therefore, a second reaction path, beginning with the lowest energy reactant was also determined following the same geometric RMSD procedure. We denote the two reaction paths as lowest energy reactant (LER) and lowest energy product (LEP).

3 Results and discussion

 Rh_{10}

Several computational^{38,51} and experimental studies have investigated the structure of the Rh₁₀ cluster in neutral, cationic and anionic charge states, identifying a dense manifold of low-energy structures. Knickelbein et al. measured the static electric dipole polarizabilities of Rh_n n = 5-28 clusters and suggested that Rh₁₀ is a fluxional molecule.³⁹ The lowest energy Rh_{10} octet species identified in this work is a structure with C_{2v} symmetry and a binding energy of -1.7686006 a.u. and is shown in Fig. 1. Re-optimization of each identified Rh₁₀ isomer at other multiplicities (doublet to 20-tet) revealed the lowest energy isomer overall, is the same structure, but with 16-tet

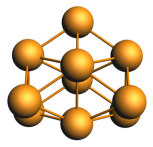


Fig. 1 Lowest energy geometry of Rh₁₀.

multiplicity and a binding energy of -1.7909319 a.u. This structure corresponds to the sc3 structure identified for Rh₁₀⁺ by Harding et al.51

$Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} + CO_2$ reactions

 $Rh_{10}O + CO \rightarrow Rh_{10} + CO_2$. Only the predetermined minimum Rh₁₀ structure is considered in this work, therefore there is only one $Rh_{10}O + CO \rightarrow Rh_{10} + CO_2$ reaction path, defined by the lowest energy Rh₁₀O species, which has a μ^2 -bound oxygen. The reaction path from this species is shown in Fig. 2. The $Rh_{10}O \cdots C$ distance in the intermediate state is 2.72 Å. Overall, the reaction path is very flat with the separated products (Rh₁₀ + CO₂) lying only 0.15 eV above the reactants and the intermediate Rh₁₀O···CO structure being effectively isoenergetic with the Rh₁₀O + CO reactants. For the reactant and intermediate structures, all multiplicities (doublet to 16-tet) were found within a range of 0.35 eV. Different multiplicities of the Rh₁₀ product span an energetic range of 0.71 eV.

 $Rh_{10}O_2 + CO \rightarrow Rh_{10}O + CO_2$. Both the LER (Fig. 3) and LEP (Fig. 4) reaction paths arrive at the same μ^2 Rh₁₀O product – *i.e.* the lowest energy Rh₁₀O₂ can follow a reaction path that results in the lowest energy Rh₁₀O. The lowest energy Rh₁₀O₂ structure has two μ²-bound oxygen atoms in symmetry-equivalent positions. The intermediate CO-bound species, removing either of these oxygen atoms is -1.93 eV lower in energy than the separated reactants and has a Rh₁₀O···C distance of 3.19 Å.

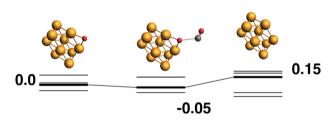


Fig. 2 $Rh_{10}O + CO \rightarrow Rh_{10} + CO_2$ reaction path for the lowest energy Rh₁₀O species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

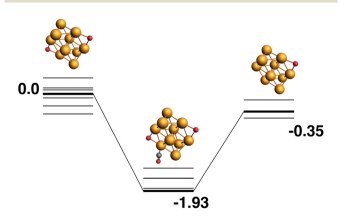


Fig. 3 $Rh_{10}O_2 + CO \rightarrow Rh_{10}O + CO_2$ reaction path for the lowest energy Rh₁₀O₂ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin

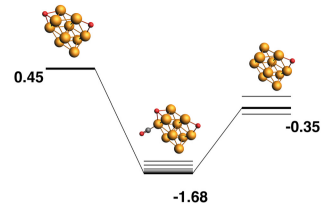


Fig. 4 $Rh_{10}O_2 + CO \rightarrow Rh_{10}O + CO_2$ reaction path for the lowest energy product (Rh₁₀O) species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy Rh₁₀O₂ reactant, as shown in Fig. 3. The octet surface is bolded, all other multiplicities are shown with thin lines

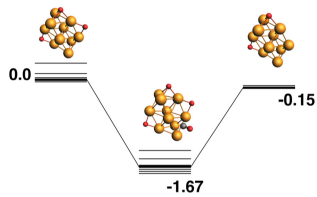


Fig. 5 Rh₁₀O₃ + CO \rightarrow Rh₁₀O₂ + CO₂ reaction path for the lowest energy Rh₁₀O₃ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

Beginning with the lowest energy μ^2 Rh₁₀O product and searching for a reaction path towards any Rh₁₀O₂ reactant, yields a $\mu^{3}\mu^{2}$ reactant 0.45 eV higher in energy than the lowest energy Rh₁₀O₂ and a corresponding Rh₁₀OO·CO intermediate. The intermediate structure removes the µ3-bound oxygen atom and is at -1.68 eV, 0.25 eV higher in energy than the intermediate in the LER pathway.

 $Rh_{10}O_3 + CO \rightarrow Rh_{10}O_2 + CO_2$. The reaction path from the lowest energy Rh₁₀O₃ structure is shown in Fig. 5. The lowest energy Rh₁₀O₃ structure has the same two μ²-bound oxygen atoms as in Rh₁₀O₂ plus a μ^3 -bound oxygen atom that shares a Rh atom with one of the μ^2 -bound oxygen atoms. In the Rh₁₀O₂O·CO intermediate structure, the incoming CO approaches the side of the cluster with both the μ^3 and μ^2 oxygen atoms, but abstracts the μ^2 -bound oxygen. The intermediate is at -1.67 eV and the Rh₁₀O···C distance is 3.39 Å. The Rh₁₀O₂ product is at -0.15 eV. Removal of the μ^3 -bound oxygen instead of the μ^2 -bound oxygen yields the lowest energy Rh₁₀O₂ structure at -0.59 eV, and this pathway is shown in Fig. 6. The intermediate in

Paper

-1.37

Fig. 6 $Rh_{10}O_3 + CO \rightarrow Rh_{10}O_2 + CO_2$ reaction path for the lowest energy product (Rh₁₀O₂) species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy Rh₁₀O₃ reactant, as shown in Fig. 5. The octet surface is bolded, all other multiplicities are shown with thin lines

this pathway is at -1.37 eV, 0.30 eV higher in energy than the LER path intermediate and the Rh₁₀O···C distance is 2.91 Å.

 $Rh_{10}O_4 + CO \rightarrow Rh_{10}O_3 + CO_2$. The lowest energy $Rh_{10}O_4$ structure has three μ^3 -bound oxygen atoms and a μ^2 -bound oxygen. The reaction path from this species is shown in Fig. 7 and removes one of the μ^3 oxygen atoms. The products (Rh₁₀O₃ + CO₂) are nearly isoenergetic with the reactants. As the oxygen atoms in the Rh₁₀O₄ cluster are all bound (in part) to the same four Rh atoms, the approaching CO is likely to interact with more than one oxygen atom. In the lowest energy intermediate structure identified, the CO could possibly abstract one of two a μ^3 -bound oxygen atoms, with Rh₁₀O···C distances of 3.08 and 3.22 Å respectively.

The lowest energy $Rh_{10}O_3$, as above, possesses two μ^2 -bound oxygen atoms and one μ^3 -bound oxygen, so it is not a possible product (without rearrangement) from the lowest energy Rh₁₀O₄ structure. The path ending with the lowest energy Rh₁₀O₃ structure beings with a high energy Rh₁₀O₄ structure (+1.25 eV), which has two μ³-bound oxygen atoms and two μ^2 -bound oxygen atoms (Fig. 8). The close Rh₁₀O···C distances in the intermediate structure, are 3.16 Å for the μ^3 -bound oxygen that is abstracted by the CO and 3.39 Å for the μ^2 bound oxygen that remains.

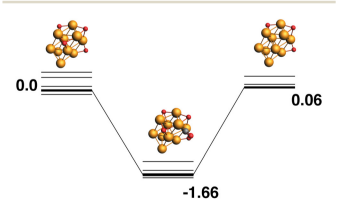


Fig. 7 $Rh_{10}O_4 + CO \rightarrow Rh_{10}O_3 + CO_2$ reaction path for the lowest energy Rh₁₀O₄ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

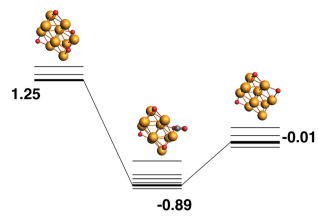


Fig. 8 Rh₁₀O₄ + CO \rightarrow Rh₁₀O₃ + CO₂ reaction path for the lowest energy product (Rh₁₀O₃) species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy Rh₁₀O₄ reactant, as shown in Fig. 7. The octet surface is bolded, all other multiplicities are shown with thin lines

 $Rh_{10}O_5 + CO \rightarrow Rh_{10}O_4 + CO_2$. The lowest energy $Rh_{10}O_5$ structure has four u³-bound oxygen atoms and a u²-bound oxygen. The reaction path from this species is shown in Fig. 9 and removes one of the \(\mu^3 \) oxygen atoms. Similarly to the $Rh_{10}O_4$ to $Rh_{10}O_3$ LER reaction path, the products $(Rh_{10}O_4 +$ CO₂) are nearly isoenergetic with the reactants, the relative energy of the Rh₁₀O₅·CO intermediate species is also similar to that of the $Rh_{10}O_4$ ·CO intermediate (Fig. 7), at -1.61 eV. The approaching CO abstracts a μ^3 -bound oxygen atom from 3.04 Å.

The LEP reaction path is shown in Fig. 10. The Rh₁₀O₄ product has a relative energy of -0.96 eV and the lowest energy reaction path removes a μ^{1} -bound oxygen atom that is bound to a previously bare Rh atom.

$$Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} + N_2$$
 reactions

 $Rh_{10} + N_2O \rightarrow Rh_{10}O + N_2$. Only the predetermined minimum Rh₁₀ structure is considered in this work, therefore there is only one $Rh_{10} + N_2 \rightarrow Rh_{10}O + N_2$ reaction path, defined by

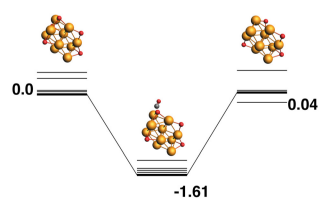


Fig. 9 $Rh_{10}O_5 + CO \rightarrow Rh_{10}O_4 + CO_2$ reaction path for the lowest energy Rh₁₀O₅ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines

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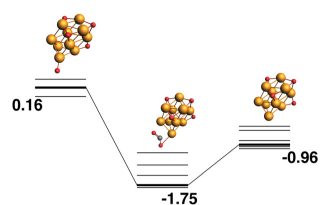


Fig. 10 $Rh_{10}O_5 + CO \rightarrow Rh_{10}O_4 + CO_2$ reaction path for the lowest energy product $(Rh_{10}O_4)$ species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy $Rh_{10}O_5$ reactant, as shown in Fig. 9. The octet surface is bolded, all other multiplicities are shown with thin lines.

the lowest energy $Rh_{10}O$ species, which has a μ^2 -bound oxygen. The reaction path from this species is shown in Fig. 11. The energetic profile of the reaction path is quite different from the reduction reactions, with the $Rh_{10}O.N_2$ intermediate structure being -4.54 eV below zero energy and the $Rh_{10}O+N_2$ products being found at -3.57 eV. The $Rh_{10}O.N_2$ separation is 3.07 Å.

 $Rh_{10}O + N_2O \rightarrow Rh_{10}O_2 + N_2$. The lowest energy $Rh_{10}O$ structure, as previously seen, has a μ^2 -bound oxygen. The reaction path from this species is shown in Fig. 12 and deposits a μ^3 oxygen atom such that no Rh atoms bind both oxygen atoms. The intermediate structure has a relative energy of

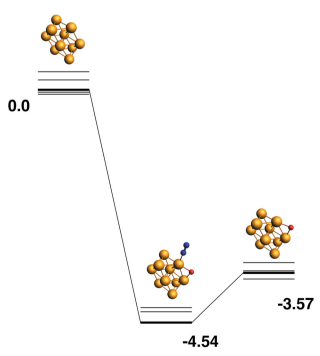


Fig. 11 Rh₁₀ + N₂O \rightarrow Rh₁₀O + N₂ reaction path for the lowest energy product (Rh₁₀O) species. Relative energies are shown for the octet surface in eV. Zero energy is the defined Rh₁₀ reactant, as shown in Fig. 1. The octet surface is bolded, all other multiplicities are shown with thin lines.

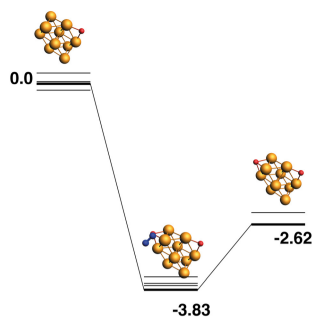


Fig. 12 $Rh_{10}O + N_2O \rightarrow Rh_{10}O_2 + N_2$ reaction path for the lowest energy $Rh_{10}O$ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

-3.83 eV and the Rh₁₀O·N₂ distance is 3.96 Å. The products are found at -2.62 eV.

The LEP reaction path is shown in Fig. 13. The $Rh_{10}O_2$ product has a relative energy of -3.07 eV and deposits the second of the two symmetry-equivalent μ^2 -bound oxygen atoms.

 $Rh_{10}O_2 + N_2O \rightarrow Rh_{10}O_3 + N_2$. Both the LER (14) and LEP (15) reaction paths form the lowest energy $Rh_{10}O_3$ product, which has two μ^2 and one μ^3 -bound oxygen atom. This product $(Rh_{10}O_2 + N_2)$ has an energy of -2.83 eV relative to the lowest energy $Rh_{10}O_2 + N_2O$. The intermediate structure connecting

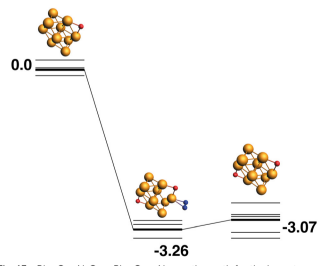


Fig. 13 $Rh_{10}O + N_2O \rightarrow Rh_{10}O_2 + N_2$ reaction path for the lowest energy product $(Rh_{10}O_2)$ species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy $Rh_{10}O$ reactant, as shown in Fig. 12. The octet surface is bolded, all other multiplicities are shown with thin lines.

these two lowest energy Rh₁₀O₂ and Rh₁₀O₃ structures, must therefore deposit the μ^3 -bound oxygen atom. This intermediate is found at -3.83 eV and the Rh₁₀O⋅N₂ distance is 3.96 Å. If instead, of depositing the μ^3 -bound oxygen atom, the intermediate is chosen to deposit one of the two μ^2 -oxygen atoms, the reaction path in Fig. 15 is observed. The Rh₁₀O₂ reactant and Rh₁₀O₃·N₂ intermediate structures are found at +0.44 and -3.83 eV respectively - *i.e.* the intermediate structure is 0.3 eV higher in energy than the LER intermediate and has a Rh₁₀O·N₂ distance of 2.99 Å.

 $Rh_{10}O_3 + N_2O \rightarrow Rh_{10}O_4 + N_2$. The LER reaction path for the $Rh_{10}O_3 + N_2O \rightarrow Rh_{10}O_4 + N_2$ reaction is shown in Fig. 16. The $Rh_{10}O_4$ product at -2.16 eV has two μ^2 and two μ^3 -bound oxygen atoms and the corresponding intermediate structure at -3.23 eV has a Rh₁₀O·N₂ distance of 3.20 Å, depositing the second μ^3 oxygen atom. The path leading to the lowest energy Rh₁₀O₄ product is shown in Fig. 17. The Rh₁₀O₄ product has three μ^3 and one μ^2 -bound oxygen atom, and the lowest energy intermediate structure deposits the third µ³-bound oxygen atom. The geometrically matching Rh₁₀O₃ reactant with two μ^3 and one μ^2 -bound oxygen, is very close in energy to the lowest energy Rh₁₀O₃ structure (two $\times \mu^3$ and $1 \times \mu^2$), with a relative energy of only 0.07 eV. The Rh₁₀O·N₂ distance in the intermediate structure is 3.56 Å.

 $Rh_{10}O_4 + N_2O \rightarrow Rh_{10}O_5 + N_2$. The reaction path from the lowest energy Rh₁₀O₄ structure is shown in Fig. 18. The Rh₁₀O₅ product lies at -2.41 eV has two μ^2 and three μ^3 -bound oxygen atoms and the corresponding intermediate structure, depositing the second μ^2 oxygen atom, lies very close in energy at -2.48 eV and has a Rh₁₀O⋅N₂ distance of 3.38 Å. The lowest energy Rh₁₀O₅ structure has one μ^2 and four μ^3 -bound oxygen

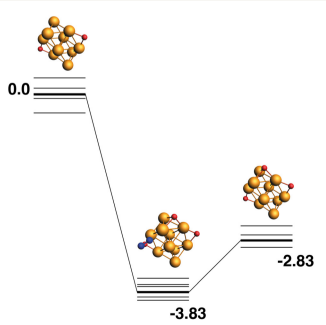


Fig. 14 $Rh_{10}O_2 + N_2O \rightarrow Rh_{10}O_3 + N_2$ reaction path for the lowest energy Rh₁₀O₂ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

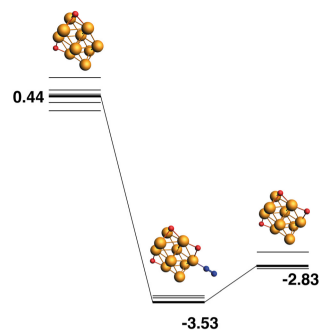


Fig. 15 $Rh_{10}O_2 + N_2O \rightarrow Rh_{10}O_3 + N_2$ reaction path for the lowest energy product (Rh₁₀O₃) species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy Rh₁₀O₂ reactant, as shown in Fig. 14. The octet surface is bolded, all other multiplicities are shown with thin lines

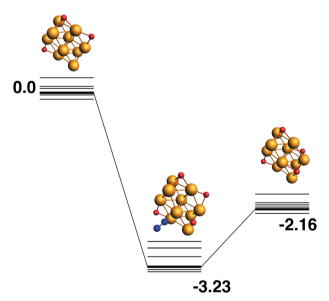


Fig. 16 $Rh_{10}O_3 + N_2O \rightarrow Rh_{10}O_4 + N_2$ reaction path for the lowest energy Rh₁₀O₃ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines

atoms and the path leading to this product is shown in Fig. 19. The lowest energy matching intermediate structure at -3.75 eV corresponds to depositing the fourth μ³-bound oxygen atom, but the corresponding Rh₁₀O₄ reactant is high in energy, at +1.00 eV. The Rh₁₀O·N₂ distance in the intermediate structure is 3.94 Å.

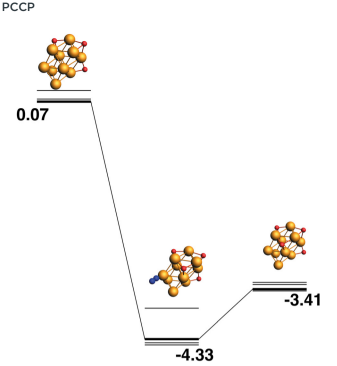


Fig. 17 $Rh_{10}O_3 + N_2O \rightarrow Rh_{10}O_4 + N_2$ reaction path for the lowest energy product $(Rh_{10}O_4)$ species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy $Rh_{10}O_3$ reactant, as shown in Fig. 16. The octet surface is bolded, all other multiplicities are shown with thin lines

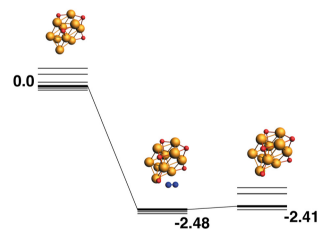


Fig. 18 Rh₁₀O₄ + N₂O \rightarrow Rh₁₀O₅ + N₂ reaction path for the lowest energy Rh₁₀O₄ species. Relative energies are shown for the octet surface in eV. The octet surface is bolded, all other multiplicities are shown with thin lines.

Comparison of the $\mathrm{Rh_{10}O_n}+\mathrm{CO} \to \mathrm{Rh_{10}O_{n-1}}+\mathrm{CO_2}$ reaction paths for n=1–5 reveals a largely similar energetic landscape regardless of n, as shown in Fig. 20 and 21 for the LER and LEP reaction paths respectively. For the pathways beginning with the lowest energy reactant, with the exception of the last abstraction of oxygen, resulting in a bare rhodium cluster (*i.e.* $\mathrm{Rh_{10}O}+\mathrm{CO} \to \mathrm{Rh_{10}O}+\mathrm{CO_2}$, Fig. 2), where the intermediate structure lies very close to the energy of both the products and

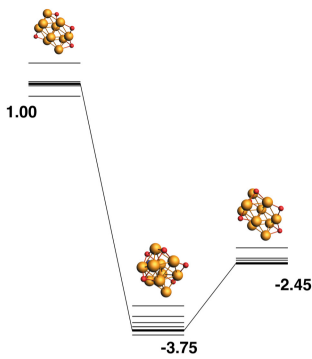


Fig. 19 $Rh_{10}O_4 + N_2O \rightarrow Rh_{10}O_5 + N_2$ reaction path for the lowest energy product $(Rh_{10}O_5)$ species. Relative energies are shown for the octet surface in eV. Zero energy is the lowest energy $Rh_{10}O_4$ reactant, as shown in Fig. 18. The octet surface is bolded, all other multiplicities are shown with thin lines.

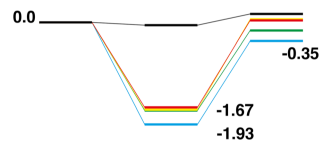


Fig. 20 Rh₁₀O_n + CO \rightarrow Rh₁₀O_{n-1} + CO₂, n = 1–5 LER reaction paths overlaid. The Rh₁₀O +CO₂ path is black, the Rh₁₀O_n paths are coloured blue, green, yellow and red, for n = 2–5 respectively.

reactants, the four reaction paths have almost the same energetics, with the $\mathrm{Rh_{10}O_{n}}$ -CO intermediate structure being found in a narrow range between -1.67 and -1.93 eV relative to the reactants. The $\mathrm{Rh_{10}O_{n-1}}$ products are found in a similarly narrow energy range, from -0.35 eV for the $\mathrm{Rh_{10}O_2}$ reaction to +0.06 eV for the $\mathrm{Rh_{10}O_4}$ reaction. Considering the reaction paths arriving at the lowest energy products, shows similar behaviour, with intermediate structures consistently 2 eV lower in energy than the corresponding reactants. The ΔE (products-reactants) shows some deviation with values of 0.80, 0.59, 1.26 and 1.17 eV for n=2-5 respectively, indicating that the $\mathrm{Rh_{10}O_{n-1}}$ products are more stable for the $\mathrm{Rh_{10}O_4}$ and $\mathrm{Rh_{10}O_5}$ reactants. This is consistent with the higher rates observed for the $\mathrm{Rh}_m\mathrm{O}_n+\mathrm{CO}\to\mathrm{Rh}_m\mathrm{O}_{n-1}+\mathrm{CO}_2$ reactions for n=4,5 by Yamada et $al.^{40}$

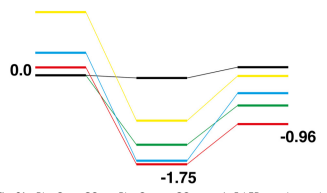


Fig. 21 $Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} + CO_2$, n = 1-5 LEP reaction paths overlaid. Zero energy is the lowest energy Rh₁₀O reactant, as shown in Fig. 2. The $Rh_{10}O + CO_2$ path is black, the $Rh_{10}O_n$ paths are coloured blue, green, yellow and red, for n = 2-5 respectively.

Comparison of the $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} + N_2$ reaction paths for n = 0-4, shows a very different pattern to the abstraction of oxygen reactions. Considering first the LER paths, shown overlaid in Fig. 22. As the $Rh_{10}O_n$ reactant is increasingly oxidized (i.e. as n increases), the relative energy of both the intermediate and product increases. With increasing oxygen content, the energy of the intermediate species rises by approximately 0.7 eV. The relative energy of the $Rh_{10}O_{n+1}$ product also rises, but less linearly, with the Rh₁₀O₄ and Rh₁₀O₅ products switching order. A change in behaviour is clearly seen for the $Rh_{10}O_4 + N_2O \rightarrow Rh_{10}O_5 + N_2$ reaction, where the intermediate and product species are almost isoenergetic, compared to the n = 0-3 reactions, where the products are less stable than the intermediate by approximately 1 eV. The LEP reaction paths, shown overlaid in Fig. 23 show broadly similar energy profiles. The most notable feature is the difference in the relative energies of the intermediate and product species, which

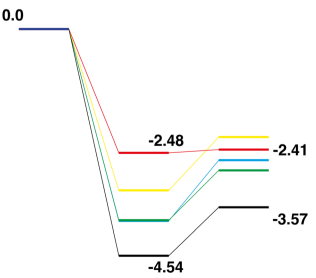


Fig. 22 $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} + N_2$, n = 0-4 LER reaction paths overlaid. The Rh₁₀ + N₂O path is black, the Rh₁₀O_n paths are coloured blue, green, yellow and red, for n = 1-4 respectively.

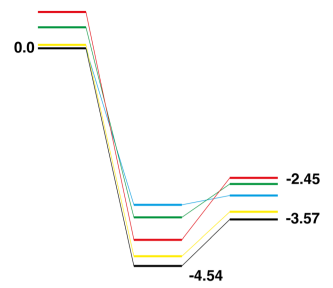


Fig. 23 $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} + N_2$, n = 0-4 LEP reaction paths overlaid. Zero energy is the lowest energy Rh₁₀ reactant, as shown in Fig. 11. The $Rh_{10} + N_2O$ path is black, the $Rh_{10}O_n$ paths are coloured blue, green, yellow and red, for n = 1-4 respectively.

increases from 0.195 eV for the Rh₁₀O + N₂ reaction, to 0.71, 0.93 and 1.29 eV for the n = 2-4 paths respectively.

4 Conclusion

We have determined the key points on the reaction paths of $Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} + CO_2$ and $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1}$ + N₂ reactions at multiplicities ranging from doublet up to 16-tet. The vast majority of structures were found to lie within a relatively small energetic range, indicating that most multiplicities would be accessible in reaction conditions.

For the $Rh_{10}O_n + CO \rightarrow Rh_{10}O_{n-1} + CO_2$ reactions, the relative energy of the O···CO intermediate. Structure was found to be remarkably consistent as the oxidation of the Rh₁₀ cluster increased, lying between -1.93 and -1.67 eV below the energy of the reactants. Similarly, the relative energy of the $Rh_{10}O_{n-1}$ product was found to lie in a narrow range of only 0.35 eV.

The oxygen addition reactions, $Rh_{10}O_n + N_2O \rightarrow Rh_{10}O_{n+1} +$ N₂, showed significantly different behaviour, with the energy of the $O \cdot \cdot \cdot N_2$ intermediate rising by approximately 0.7 eV with each addition oxygen on the $Rh_{10}O_n$ cluster. The relative energy of the Rh₁₀O_{n+1} product also rose significantly, by ≈ 0.3 eV, though a linear relationship was not seen for these species.

These reaction paths combined suggest that the rate of oxygen addition to the cluster surface is governed by both thermodynamic and kinetic effects, whereas, the rate of oxygen abstraction reaction is largely governed by access to oxygen on the cluster surface. Paths defined by the lowest energy reactant and lowest energy product show similar energetic profiles, suggesting that the experimentally observed reaction rates are unlikely to be significantly affected by the presence of a mixture of $Rh_{10}O_n$ isomers.

Author contributions

ATD, VM and OA: investigation MAA: conceptualization of this study, methodology, writing - review and editing.

Conflicts of interest

There are no conflicts to declare.

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