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# PAPER

Cite this: *Phys. Chem. Chem. Phys.*, 2024, 26, 209

# f-Block reactions of metal cations with carbon dioxide studied by inductively coupled plasma tandem mass spectrometry

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f-Block chemistry offers an opportunity to test current knowledge of chemical reactivity. The energy dependence of lanthanide cation (Ln<sup>+</sup> = Ce<sup>+</sup>, Pr<sup>+</sup>, Nd<sup>+</sup>-Eu<sup>+</sup>) and actinide cation (An<sup>+</sup> = Th<sup>+</sup>, U<sup>+</sup>-Am<sup>+</sup>) oxidation reactions by  $CO<sub>2</sub>$ , was observed by inductively coupled plasma tandem mass spectrometry. This reaction is commonly spin-unallowed because the neutral reactant (CO<sub>2</sub>,  ${}^1\Sigma^+_g$ ) and product (CO<sub>,</sub>  ${}^{1}\Sigma^{+}$ ) require the metal and metal oxide cations to have the same spin state. Correlation of the promotion energy  $(E_p)$  to the first state with two free d-electrons with the reaction efficiency indicates that spin conservation is not a primary factor in the reaction rate. The  $E<sub>p</sub>$  likely influences the reaction rate by partially setting the crossing between the ground and reactive states. Comparison of  $\text{Ln}^+$  and  $\text{An}^+$  congener reactivity indicates that the 5f-orbitals play a small role in the  $An<sup>+</sup>$  reactions. **PAPER**<br> **(a)** Check for updates<br> **EVALUATE COCK reactions of metal cations with carbon dioxide studies through the studies of the** 

Received 30th August 2023, Accepted 29th November 2023

DOI: 10.1039/d3cp04180h

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# Introduction

The f-block, or elements with f-electrons in the valence shells, represents an interesting frontier in chemistry. While understanding f-block chemistry is vital for the nuclear industry to ensure safe and efficient operation, the f-block represents an opportunity to confirm or revise our current understanding of chemical reactions because most of our understanding of chemical theory is derived from work with s, p, and d-blocks. The radial distribution of the 4f-orbitals suggests that their participation in bonding will be minimal, but the 5f-orbitals have the correct distribution to participate. Nevertheless, the parameters of when the 5f-orbitals participate in bonding are unclear.

While many current studies focus on condensed phases, gas phase studies offer an attractive alternative. Gas phase studies eliminate many perturbations in condensed phases like solvent effects so that the fundamental interaction of the analyte species can be probed. Indeed, mass spectrometry has proven adept at observing the reactions of actinides (An) and lanthanides (Ln). Fourier transform ion cyclotron resonance mass spectrometry (FTICR-MS) has been utilized to study  $An^+ (An^+ =$ Th<sup>+</sup>-Cm<sup>+</sup>) with many small molecules.<sup>1-6</sup> Similar reactions of the  $Ln<sup>+</sup>$  with small molecules have been studied using selected ion flow tube mass spectrometry  $(SIFT-MS)$ <sup>7-9</sup> Unfortunately, FTICR-MS and SIFT-MS are thermal methods where only barrierless, exothermic or thermoneutral reactions can be observed, somewhat limiting the impact of the studies and the reactions that can be probed. Furthermore, observation of a reaction at a single energy point can mislead conclusions. For example, the low efficiency reaction observed between  $Th^+ + CH_4$ in FTICR-MS experiments was originally attributed to spinrestrictions;<sup>4</sup> however, it was shown later in energy dependence studies by guided ion beam tandem mass spectrometry (GIBMS) to be caused by a small barrier $10$  where the low efficiency reaction was observed from a small population of reactants within the energy distribution with sufficient energy to overcome the reaction barrier.

 $GIBMS<sup>10-27</sup>$  oxidation studies shed light on the reaction by observing it over a wide range of kinetic energies. The energy dependence of the reaction often simplifies the interpretation of results, as noted in the GIBMS study of  $Th^+ + CH_4$ ,<sup>10</sup> and allows the observation of endothermic reactions or reactions with a barrier. However, many of the An elements are significantly radioactive so that they are difficult to obtain and work with safely. Consequently, studies of the energy dependence of transuranic species have not been reported in the gas phase. Yet there remain examples of low efficiency reactions, for example  $Pu^+ + CO_2$  from FTICR-MS studies,<sup>1</sup> that may require knowing the energy dependence of the reaction to correctly interpret.

Additionally, the reactions of  $Ln^+$  and  $An^+$  with  $CO_2$  is an interesting system to study, in part because  $SIFT-MS<sup>7</sup>$  and FTICR-MS<sup>1,2</sup> studies provide examples of exothermic reactions where no or limited product is observed. Furthermore, atmospheric  $CO<sub>2</sub>$  also remains a large reservoir of carbon that can



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potentially be tapped with the correct catalyst. However,  $CO<sub>2</sub>$  is difficult to crack because activating the  $C=O$  bond according to reaction (1) is formally spin-forbidden:

$$
CO2(1\Sigmag+) \rightarrow CO(1\Sigma+) + O(3P)
$$
 (1)

The spin allowed pathway lies 1.97 eV higher in energy dissociating to the CO + O  $(^1D)$ .<sup>28</sup> Nevertheless, developing an economical catalyst that can break the  $C=O$  bond could revolutionize manufacturing and lead to a promising method to remove excess carbon from the atmosphere. While transuranics are unlikely to be that catalyst, they offer an extreme system to test chemical theory upon.

For  $CO<sub>2</sub>$ , reaction (2) is often spin-forbidden because unpaired electrons in the metal cation  $(M<sup>+</sup>)$  and oxygen usually pair when forming MO<sup>+</sup> to lower the spin in the metal oxide cation product:

$$
M^{\dagger}({}^{s}L) + CO_2({}^{1}\Sigma^+) \rightarrow MO^{\dagger}({}^{s-2}\Lambda) + CO({}^{1}\Sigma^+) \tag{2}
$$

where s is the spin-multiplicity. This spin-restriction is often cited as a reason that reactions proceed inefficiently.<sup>1,4</sup> However, it is unclear why spin-conservation should apply to large metals. Indeed, recent work by Cox et  $al$ ,<sup>22</sup> which analyzed the observed efficiency of  $\text{Ln}^+$  and  $\text{An}^+$  cations reactions with  $\text{O}_2$ ,  $\text{CO}_2$  and  $\text{H}_2\text{O}$ at thermal energies, $1-3,5,7-9$  concluded that the primary influence on the reaction rate was the promotion energy  $(E<sub>p</sub>)$  from the ground state to the first state with two free nd-electrons, where  $n = 5$ , 6. This correlation appears to be caused because the free nd-electrons are necessary to form an efficient  $\text{LnO}^+$  or  $\text{AnO}^+$ bond; therefore, the rate is at least partially influenced by the crossing between the potential energy surfaces evolving from the ground state and the more reactive higher energy state at  $E_{\rm p}(n{\rm d}^2)$ .

Here we present the energy dependence of An<sup>+</sup> (An<sup>+</sup> = Th<sup>+</sup>,  $U^+$ -Am<sup>+</sup>) reactions with CO<sub>2</sub> studied using an Agilent 8900 inductively coupled plasma tandem mass spectrometry (ICP-MS/MS) over an extended energy range. The goal of this study is to understand the role of the  $An<sup>+</sup>$  valence electrons, and specifically the 5f-orbitals, in the reaction through observing the reactivity trend across the An series. For comparison as a baseline when f-electrons are not expected to contribute, $29,30$ the energy dependence of  $Ln^+$   $(Ln^+ = Ce^+, Pr^+, Nd^+, Sm^+, Eu^+)$ reactions are also presented.

# Experimental methods

Caution: The An used in this study are all radioisotopes with varying activities and half-lives. All work was done within the radiological protection controls of specialized laboratories at Pacific Northwest National Laboratory.

Experiments were conducted using an Agilent 8900 ''triple quadrupole'' mass spectrometer (ICP-MS/MS; Agilent Technologies, Santa Clara, CA, USA) located within a radiological facility at Pacific Northwest National Laboratory.31 This instrument utilized an ICP ion source equipped with a quartz double-pass spray chamber and 100  $\mu$ L min<sup>-1</sup> perflouroalkoxy alkane (PFA) nebulizer. The instrument utilizes a quadrupole mass filter (1 amu resolution) to mass

Table 1 Average electronic energy (in eV) of  $Ln<sup>+</sup>$  and  $An<sup>+</sup>$  from the ICP source assuming a Boltzmann distribution<sup>a</sup>

$M+$ (isotope) Mass (amu) 300 K 700 K 5000 K 8000 K 9000 K 10 000 K							
$Ce^+(140)$	139.90	0.00	0.04	0.58	0.69	0.71	0.74
$Pr^+(141)$	140.91	0.01	0.03	0.42	0.54	0.57	0.61
$Nd^+(146)$	145.91	0.01	0.03	0.57	0.79	0.85	0.90
$Sm^{+}(149)$	148.92	0.02	0.06	0.43	0.59	0.64	0.69
Eu' (153)	152.92	0.00	0.01	0.28	0.68	0.81	0.93
$Th^+(232)$	232.04	0.00	0.02	0.66	0.85	0.90	0.93
$U^{+}$ (238)	238.05	0.01	0.03	0.51	0.67	0.71	0.74
$Np^{+}$ (237)	237.05	0.00	0.01	0.23	0.28	0.30	0.31
$Pu^+(242)$	242.06	0.00	0.01	0.70	0.87	0.91	0.95
Am <sup>+</sup> (243)	243.06	0.00	0.00	0.19	0.52	0.63	0.75

 $a<sup>a</sup>$  Masses and electronic states taken from ref. 28 ([https://physics.nist.](https://physics.nist.gov/PhysRefData/Handbook/periodictable.htm) [gov/PhysRefData/Handbook/periodictable.htm](https://physics.nist.gov/PhysRefData/Handbook/periodictable.htm)).

select the reactant ion beam that is passed into a collision cell contained within an octopole ion guide where the reactant ion is reacted with the neutral gas,  $CO<sub>2</sub>$ . Products and unreacted ions drift to the end of the octopole where they are focused through a second quadrupole mass filter, mass selected, and subsequently detected at a standard electron multiplier detector.

Stock multi-element standard solutions containing 1 ng  $g^{-1}$ of Ce–Nd, Sm, Eu, Th, and U in  $2\%$  HNO<sub>3</sub> were prepared. To minimize high activity radioisotopes within the instrument, a multi-element standard solution of 1 pg  $g^{-1}$  of Np-Am in 2%  $HNO<sub>3</sub>$  was also prepared. Table 1 lists the isotopes used for each M<sup>+</sup>. Carbon dioxide (CO<sub>2</sub>; 99.99%; Oxarc) was used as the reaction gas. The flow rates ranged from  $0.06-0.13$  mL  $min^{-1}$ , which corresponds to estimated pressures of 1.4 and 2.8 mTorr. Tuning parameters were optimized to provide maximum sensitivity for the high mass range using a 1 ng  $g^{-1}$  Th and U solution. The octopole bias was adjusted in intervals from  $+7$  V to  $-45$  V while keeping other cell parameters constant: octopole rf peak-to-peak voltage of 180 V, axial acceleration of 2.0 V, and a kinetic energy discrimination (KED, the voltage difference between the octopole bias in the CRC and the second quadrupole) of  $-10.0$  V. Data were acquired in triplicate using 1 s acquisition times for the 1 ng  $g^{-1}$ solutions and 4 s for the 1 pg  $g^{-1}$  solutions. Paper<br>
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### Kinetic energy dependent cross section

Reaction cross sections  $(\sigma)$  are calculated from the raw signal intensities using a Beer's Law analog:<sup>32</sup>

$$
I = I_0 e^{-\rho \sigma l} \tag{3}
$$

where  $I$  is the reactant intensity exiting the collision cell,  $I_0$  is the reactant intensity entering the collision cell,  $\rho$  is the number density of the neutral reactant in the collision cell, and  $l$  is the effective length of the collision cell.  $I_0$  is estimated from the sum of all ion signals observed after the second quadrupole from the mass selected in the first quadrupole.  $l$  is estimated as the physical length of the collision cell, 10 cm, in the Agilent 8900. This is not strictly correct because a pressure gradient will exist at the entrance and exit apertures that will lengthen the effective cross section. The difference is anticipated to be  $\leq$ 20% and is incorporated into the absolute uncertainty of the cross section. Individual product cross sections  $(e.g., MO<sup>+</sup>)$  are calculated as a

percentage of the overall reaction cross section; however, only  $\mathrm{MO}^+$  or  $\mathrm{MO_2}^+$  products were observed. Because the Agilent 8900 operates under multi-collision conditions, to compare the observed cross section to the reaction collision limit, the cross sections observed at 1.4 and 2.8 mTorr were extrapolated to zero pressure for rigorous single collision conditions. Absolute uncertainties in the cross section are estimated to be  $\pm 50\%$ with relative uncertainties of  $\pm 10\%$ .

The energy in the laboratory frame is estimated from the octopole bias by the relationship: 33

$$
E_{\text{Lab}} = V_{\text{p}} + \frac{m}{m_{\text{Ar}}^2} \frac{5}{2} k_{\text{B}} T_{\text{P}} - V_{\text{oct}}
$$
 (4)

where  $V_p$  is the plasma potential ( $\sim$  2 V), *m* is the reactant ion,  $M^+$ , mass,  $m_{Ar}$  is the mass of argon,  $k_b$  is Boltzmann's constant,  $T<sub>p</sub>$  is the ion temperature entering the octopole, and  $V<sub>oct</sub>$  is the octopole bias. The temperature of the ion exiting the plasma is expected to be the plasma temperature, 8000–10 000 K. Collisional cooling in a differential pumping region between the source and first quadrupole is expected to drop the average temperature entering the octopole, so  $T<sub>p</sub>$  is conservatively estimated to be 5000–10 000 K. Table 1 lists the average electronic energy for a Boltzmann distribution in these temperature ranges. Above 5000 K, average electronic energy can be significant, and this difference likely affects the reactivity observed where excited states may be more or less reactive than the ground state. Because of the great number of overlapping electronic states, it is assumed that a pathway exists from any given excited state to the lowest energy reaction surface leading to products. Consequently, the average electronic energy is treated as energy available for reaction. The energy in the center-of-mass  $(E_{CM})$  frame represents the kinetic energy available for a chemical reaction. The relationship between  $E_{\text{LAB}}$  and  $E_{\text{CM}}$  is described by eqn (5):<sup>32</sup> PCCP<br>
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$$
E_{\text{CM}} = E_{\text{LAB}} \times M/(M+m) \tag{5}
$$

where *M* is the mass of the neutral reactant partner,  $CO_2 = 44.01$ amu and  $m$  is the mass of the metal ion.

### Calculating reaction efficiency

Reaction efficiency is calculated relative to the collision limit  $(k<sub>c</sub>)$ . The collision limit is estimated using the Langevin–Gioumousis–Stevenson (LGS) model<sup>34</sup> with a polarizability of 2.59  $\times$  $10^{-24}$  cm<sup>3</sup> for CO<sub>2</sub>.<sup>35</sup> Two approaches were taken to calculating the reaction efficiency. The first approach was to calculate the efficiency at the lowest observed energy  $(E_{CM})$  observed, 0.15– 0.3 eV. This is similar to single energy methods, SIFT-MS or FTICR-MS, calculations, although the corresponding temperatures here are 1600–2300 K. The second approach was to average the calculated efficiencies for all results below 1 eV. Deviations in the cross section reaction energy dependence have been observed for the Th<sup>+</sup> reaction with  $O_2$  in GIBMS experiments.<sup>13</sup> Nevertheless, no deviations above 10% of the mean efficiency were observed calculating the efficiency in this manner with the exception to  $Sm^+$  and  $Pu^+$ , where the cross sections suggest a weak barrier in excess of the reaction enthalpy. The minimum efficiency reported was  $k/k_{\text{col}} = 0.01$ .

# Experimental results

## $Ln^+ + CO<sub>2</sub>$

The absolute cross section as a function of kinetic energy for reaction (2)  $(M^+ = Ce^+, Pr^+, Nd^+, Sm^+, Eu^+)$  are presented in Fig. 1. Only two products,  $MO<sup>+</sup>$  and  $MO<sub>2</sub><sup>+</sup>$ , were observed, although  $MC^+$  and  $MCO^+$  were explicitly looked for and not observed. Of these products,  $MO_2^+$  had a clear dependence on  $CO<sub>2</sub>$  pressure suggesting that it is formed through the secondary reaction, reaction (6):

$$
MO^{+} + CO_{2} \rightarrow MO_{2}^{+} + CO
$$
 (6)

A minor pressure dependence was also observed for reaction (2), likely associated with  $MO<sup>+</sup>$  depletion from reaction (6). The cross sections shown in Fig. 1 has been extrapolated to zero pressure to remove all dependencies from the  $CO<sub>2</sub>$  pressure.

The cross sections of CeO<sup>+</sup>, PrO<sup>+</sup>, and NdO<sup>+</sup> formed in reaction (2) initially decrease with increasing energy consistent with a barrierless exothermic reaction until energies near 2.4, 1.9, and 1.2 eV, respectively. Beyond the initial decline, the cross sections increase peaking near the  $OC=O$  bond dissociation energy,  $D_0(\text{OC}-\text{O}) = 5.453 \text{ eV}^{36}$  where sufficient energy is available for  $MO<sup>+</sup>$  to dissociate. It should be noted that the initial low energy feature obscures where this increase begins making it likely that the more efficient the reaction, the more likely the threshold of the higher energy feature is obscured. Conversely, the cross sections of  $SmO^+$  and  $EuO^+$  increase with increasing energy. For SmO<sup>+</sup>, where  $D_0(Sm^+$ -O) = 5.596  $\pm$  0.004 eV,<sup>37</sup> the observed cross section indicates a barrier to this reaction, which is exothermic by 0.143 eV. This behavior has been observed previously in GIBMS experiments.<sup>15</sup> In the present experiment, the apparent threshold is shifted to lower energies than that



Fig. 1 Reaction cross section of  $Ln^{+}$  [ $Ln^{+}$  = Ce<sup>+</sup> (red circles), Pr<sup>+</sup> (dark red octagons), Nd<sup>+</sup> (blue triangles), Sm<sup>+</sup> (purple squares), Eu<sup>+</sup> (teal diamonds)] +  $CO<sub>2</sub> \rightarrow \text{LnO}^{+} + \text{CO}$  as a function of kinetic energy in CM frame (lower x-axis). The black line represents the Langevin–Gioumousis–Stevenson (LGS) model cross section.  $E_{Lab}$  of  $^{146}Nd^+$  is represented on the upper x-axis.

observed in the GIBMS experiment, indicating the presence of excited states. Given the typical operating temperature of the ICP plasma is 8000–10 000 K, the observation of excited states from this source is expected. A second increase in cross section magnitude is also observed for all  $Ln<sup>+</sup>$  peaking around 10 eV. This may be an effect from ion focusing caused by the interaction between differing radio frequencies between the two quadrupoles and octopole regions within the instrument No effort is made to interpret or characterize this feature.

The reaction efficiencies  $(k/k_{\text{col}})$  of the Ln<sup>+</sup> reactions relative to the Langevin–Gioumousis–Stevenson (LGS) collision limit $34$  $(k_{\text{col}})$  are listed in Table 2. Two approaches were taken to calculate the reaction efficiency. The first approach was to average the  $k/k_{\text{col}}$  for all energies below 1 eV. The second approach was to calculate the  $k/k<sub>col</sub>$  for the lowest observed energy, 0.15–0.2 eV, which corresponds to measuring the reaction efficiency at  $\approx$  1600–2300 K. For the Ln $^+$ , the difference between each approach is minimal. For comparison, the reaction efficiencies of GIBMS,  $15,17,19$  selected ion flow tube mass spectrometry (SIFT-MS),<sup>7</sup> and FTICR-MS<sup>38</sup> efficiencies are also listed in Table 2. In general, the ICP-MS/MS values here are within the combined uncertainties of the previous measurements. The largest observed difference was for  $Pr<sup>+</sup>$  where the GIBMS and SIFT-MS value are half of the ICP-MS/MS mean (but still within the combined measurement uncertainty). This difference can likely be attributed to the difference in the metal cation electronic distribution. Smaller differences for the other  $Ln<sup>+</sup>$  can also be attributed to electronic distributions. Public on the cutset or public on the cutset of the cutset are also that in the cutset of the common and components

### $An^+ + CO<sub>2</sub>$

Absolute cross sections as a function of kinetic energy for reaction (2)  $(M^+ = Th^+, U^+$ –Am<sup>+</sup>) are presented in Fig. 2. Like the  $\text{Ln}^+$ , only  $\text{MO}^+$  and  $\text{MO}^+_2$  products were observed. After extrapolating to zero pressure, only  $\mathrm{MO}^+$  was observed. Th $\mathrm{O}^+$ ,  $UO<sup>+</sup>$ , and NpO<sup>+</sup> cross sections decrease with increasing energy consistent with a barrierless exothermic reaction. After the initial decline in the cross sections, like the  $Ln^{+}$ , an increase in the cross section is observed  $\approx$  2.0, 1.9, and 1.3 eV,



Fig. 2 The reaction cross sections of An<sup>+</sup> [An<sup>+</sup> = Th<sup>+</sup> (red circles), U<sup>+</sup> (blue triangles),  $Np^{+}$  (green inverted triangles),  $Pu^{+}$  (purple squares),  $Am^{+}$  (teal diamonds)] +  $CO_2 \rightarrow$  AnO<sup>+</sup> + CO as a function of kinetic energy in the CM frame (lower x-axis). The black line represents the Langevin–Gioumousis– Stevenson (LGS) model cross section.  $E_{lab}$  of <sup>238</sup>U<sup>+</sup> is represented on the upper x-axis.

respectively. However, for the An<sup>+</sup>, this feature is much less pronounced than observed in the  $Ln^+$ . This may be because the  $An<sup>+</sup>$  reactions are more efficient obscuring the magnitude of the second energy feature. Curiously,  $PuO<sup>+</sup>$  and Am $O<sup>+</sup>$  have increasing cross sections with increasing energy. Given  $D_0(Pu^+$ -O) = 6.75  $\pm$  0.20 eV,  $D_0(Am^+$ –O) = 5.80  $\pm$  0.29 eV<sup>39</sup> and  $D_0(OC$ –O) = 5.45 eV, $36$  these cross sections indicate a clear barrier in excess of the reaction enthalpy.

A comparison of  $k/k_{\text{col}}$  between the two methods indicates that the average efficiency below 1 eV differs from the efficiency determined from the lowest energy point. This difference largely disappears when comparing the second energy point to the average. Previous measurements of reaction efficiency (Table 2), are primarily limited to FTICR-MS measurements by Gibson and coworkers.<sup>1,2</sup> Additional measurements for Th<sup>+</sup> and





<sup>a</sup> k<sub>col</sub> is derived from the Langevin–Gioumosis-Stevenson model cross section for CO<sub>2</sub>. <sup>b</sup> See ref. 22 and references therein. <sup>c</sup> Average k/k<sub>col</sub> for all energies <1 eV.  $d$  k/k<sub>col</sub> for the lowest energy measured (0.15–0.20 eV). <sup>e</sup> Ref. 7. f Work in progress, see ref. 26. <sup>g</sup> Ref. 38. <sup>h</sup> Ref. 19. <sup>i</sup> Ref. 17. <sup>j</sup> No reaction observed.  $k$  A barrier in excess of the thermodynamic threshold was observed. See ref. 15. <sup>1</sup> Ref. 1.  $^m$  Ref. 11.  $^n$  Work in progress, see ref. 25.  $^{\circ}$  Ref. 2.

U<sup>+</sup> are available from FTICR-MS,<sup>38</sup> GIBMS (Th<sup>+</sup>),<sup>22</sup> and an early ion beam experiment  $(U^{\dagger})$ .<sup>11</sup> The Th<sup>+</sup> and U<sup>+</sup> efficiencies reported here compare well with the ion beam $11,22$  and FTICR-MS results from Cornehl et  $al.^{38}$  The FTICR-MS measurements of Gibson and coworkers<sup>1,2</sup> are lower than those observed here with the exception of  $Pu^+$  and  $Am^+$  where very inefficient reactions under nominally thermal conditions were observed in the FTICR-MS work. Note that the FTICR-MS measurements of Gibson and coworkers<sup>1,5</sup> are systematically lower than those observed in GIBMS experiments as observed for several reactions with Th<sup>+</sup>,<sup>13,18,22,40</sup> U<sup>+</sup>,<sup>23</sup> and Gd<sup>+</sup>.<sup>14,16</sup> The differences in  $k/k_{\text{col}}$  between measurements has been previously attributed to a different starting ion electronic energy distribution.<sup>10,13</sup>

### Comparison to GIBMS cross sections

GIBMS cross sections for reaction (2) have been reported for  $Pr<sup>+</sup>,<sup>19</sup> Nd<sup>+</sup>,<sup>17</sup> Sm<sup>+</sup>,<sup>15</sup> and Th<sup>+22</sup> by Armentrout and coworkers.$ Early ion beam cross sections have also been published for  $U^+$ by Armentrout and Beauchamp. $11$  As noted above, the efficiency for the  $Ln<sup>+</sup>$  species reported here, generally exceeds the efficiency reported in the GIBMS experiments. This could be caused by the use of higher energies in the calculation of reaction efficiency than are typically used in the GIBMS experiments (e.g.  $k/k_{\text{col}}$  for Pr<sup>+</sup> is calculated at 0.05 eV).<sup>19</sup> However, a direct comparison of cross sections indicate some differences in absolute magnitude. In the case of Th $^+$ , the absolute magnitude at the cross section peak centered at  $\approx$  5.5 eV is +50% for the ICP-MS/MS cross section compared to the GIBMS cross section.<sup>22</sup> This is similar to the difference in the peak magnitude between ICP-MS/MS and GIBMS for  $Nd^+$ .<sup>17</sup> The differences in the Sm<sup>+</sup> cross section are observed in the apparent threshold that is  $\approx$  1 eV lower in energy for the ICP-MS/MS cross section than the GIBMS cross section; $15$  nevertheless, the peak magnitude is very similar. Peak magnitude is also similar between ICP-MS/MS and GIBMS for Pr<sup>+</sup>.<sup>19</sup> The overall cross section shape for the ICP-MS/MS U<sup>+</sup> reaction is different than the early ion beam experiment.<sup>11</sup> The early energy magnitude of the ion beam experiment is slightly higher in magnitude, though. PCCP<br>
Or are available from FITGANS,<sup>32</sup> Danses (ht)<sub><sup>33</sup></sub> and an early <sup>3</sup> are are available of the cell super-<br>
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The differences here compared to the GIBMS experiments can likely be attributed (at least in part) to the starting electronic distribution. This is most evident in the shifted threshold or reaction  $(2)$  for Sm<sup>+</sup> that must indicate the presence of excited states. The Agilent instrument does not include a high pressure region ( $\sim$  500 mTorr) to collisionally cool reactants ions that the GIBMS<sup>41</sup> instrument does. Indeed, the SIFT-MS incorporates a similar high pressure drift tube.<sup>7</sup> Also, the present work was extrapolated to the single collision conditions where GIBMS naturally operates. Because the cross sections between the methods are within the combined uncertainties of those methods, this is probably an acceptable approach, but the cross section may retain some multi-collision character. In an ICP-MS/MS cross section of  $Co^+ + O_2$  at higher pressures (3.5) mTorr), these pressure effects have appeared as increased cross section magnitude by a factor of 2 compared to GIBMS experiments.<sup>31</sup> Another potential difference is that the octopole only spans the physical length of the collision cell in the Agilent instrument. Consequently, the effective cell length (from pumping gradients outside of the cell) will extend into regions pre and post octopole so that reactions or scattering outside of the collision cell may not be fully represented in the reaction cross section. Nevertheless, we anticipate that the relative trends from the work presented here are accurate.

## **Discussion**

Previous work $39,42$  has indicated that the bond dissociation energies (BDEs) of AnO<sup>+</sup> are correlated to the promotion energy of the first excited electronic state with two free 6d-electrons  $[E_p(6d^2)]$ . Similarly, Ln<sup>+</sup> BDEs have been correlated to the promotion energy to the first excited state with two free 5d-electrons  $[E_{\rm p} (5d^2)]^{42}$  Armentrout<sup>27</sup> has recently shown that a better correlation exists between the BDEs and the excitation to  $Ln^{2+} E_p(5d)$  +  $O^-$ , but this argument only shifts how the electrons are shared within the bond and not the orbitals involved (i.e. two 5delectrons are still needed to form the  $LnO<sup>+</sup>$  triple bond). Presumably this correlation exists because the four 2p-electrons in O can combine with the two free d-electrons to form a  $\sigma$ -bond and two  $\pi$ -bonds to form a robust triple bond. More recent work has also indicated a correlation of the rate of reaction (2) to  $E_p(n d^2)$ for  $n = 5$  and  $6<sup>22</sup>$  In this analysis, reaction efficiencies were taken from the SIFT-MS work of Bohme and coworkers<sup>7</sup> for the  $Ln^+$  and the FTICR-MS work of Gibson and coworkers<sup>1-3,5</sup> for the An<sup>+</sup>.

Fig. 3 presents a similar analysis from the current work for both the Ln<sup>+</sup> and An<sup>+</sup>, indicating a strong linear correlation between the  $k/k_{\text{col}}$  and  $E_p(nd^2)$ . Noticeably, the cross sections of  $SmO<sup>+</sup>$  and EuO<sup>+</sup> in Fig. 1 show clear barriers. The result for EuO<sup>+</sup> is not surprising because this reaction is endothermic by  $1.42 \text{ eV}^{42}$ . In the case of  $SmO^+$ , reaction (2) is exothermic by 0.143 eV, as



Fig. 3 Correlation of the reaction efficiency relative to the LGS collision rate (k/ $k_{\text{col}}$ ) to the promotion energy  $E_{\text{p}}$ (nd<sup>2</sup>) for  $n = 5$  (blue) and  $n = 6$  (red). The solid blue and red line represent the least-squares linear regression fits of the Ln<sup>+</sup> and An<sup>+</sup> experimental data ( $R^2$  = 0.97 and  $R^2$  = 0.99). Sm<sup>+</sup> has been excluded from the  $Ln<sup>+</sup>$  linear regression fit. The dashed lines represent  $E_p(n d^2)$  of the indicated cation with dark blue and dark red lines corresponding to Ln<sup>+</sup> and An<sup>+</sup> species, respectively. See also Table 1.

noted above. This barrier was observed previously in GIBMS experiments with carefully thermalized  $Sm^+$  reactant ions.<sup>15</sup> Ab initio calculations indicate that this barrier could be attributed to the crossing seam between the potential energy surface (PES) of reaction (2) evolving from the ground state reactant asymptote and the PES evolving from an excited state reactant asymptote lying  $E_{\rm p}$ (5d<sup>2</sup>) higher in energy, where the latter PES correlated with the formation of the  $SmO<sup>+</sup>$  ground state product. Here, similar barriers are also observed for  $PuO<sup>+</sup>$  and  $AmO<sup>+</sup>$  in Fig. 2, where reaction (2) is expected to be exothermic by  $1.3 \pm 0.2$  eV and  $0.35 \pm 0.29$  eV for Pu<sup>+</sup> (D<sub>0</sub>(Pu<sup>+</sup>-O) = 6.75  $\pm$  0.20 eV) and Am<sup>+</sup> (D<sub>0</sub>(Am<sup>+</sup>-O) = 5.80  $\pm$ 0.29 eV),<sup>39</sup> respectively. As for Sm<sup>+</sup>, the barriers observed for Pu<sup>+</sup> and  $Am<sup>+</sup>$  can may also be attributed to the energy at the crossing seam between the PESs evolving from the ground and reactive states. Interestingly, FTICR-MS reactions have indicated that for the much less thermodynamically favored reaction of Pu<sup>+</sup> with NO  $(D_0(N–O) = 6.51 \text{ eV})$ ,<sup>36</sup>  $k/k_{col} = 0.17$  was observed, even though  $\Delta H_r =$  $-0.24 \pm 0.20$  eV is only mildly exothermic. Because the reaction of  $M^+$  + NO ( ${}^{2}$ H) has less spin-restrictions than reaction (2), this observation may argue that spin conservation could still be important for heavy metals. Puper<br>
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Reaction (2) requires that  $M^+$  and  $MO^+$  have the same spin state for the reaction to conserve spin. Table 3 lists the ground states of  $M^+$  and the expected state of  $MO^+$  from theoretical calculations and indicates that for the majority of  $M^+$  in this study, the formation of ground state  $MO<sup>+</sup>$  from reaction (2) is indeed spin-forbidden. The exception is U<sup>+</sup> (<sup>4</sup>I,  $5f^37s^2)^{28}$  that likely forms UO<sup>+</sup> ( $^4\Gamma$ ,  $\pi\delta\phi$ )<sup>6</sup> (where the orbital configuration listed is for the unpaired, non-bonding f-orbitals). Because the ground level of Th<sup>+</sup> is actually a mixed  ${}^{4}F_{3/2}{}^{-2}D_{3/2}$  (6d<sup>2</sup>7s/ 6d7s<sup>2</sup>)<sup>43</sup> configuration, reaction (2) forming ThO<sup>+</sup> (<sup>2</sup> $\Sigma$ <sup>+</sup>,  $\sigma$ )<sup>44</sup> may also conceivably be considered spin-allowed. Given that reaction (2) is not spin-restricted for some of the  $An<sup>+</sup>$  but is for most, the strong correlation of reaction rate to  $E_{\rm p}({\rm 6d^2})$  in Fig. 3 suggests that  $E_p(6d^2)$  has a stronger effect on the reaction rates than spin conservation.

Unlike the  $An^+$ , reaction  $(2)$  is likely formally spin-unallowed for the  $Ln<sup>+</sup>$  in this study, Table 3. There is a noticeable change in reaction efficiency between the  $\mathrm{Ln}^+$  and  $\mathrm{An}^+$ , but many  $k/k_\mathrm{col}$ are within the combined experimental uncertainties for each  $An^+$  and  $Ln^+$  congener of similar excitation energy, so it is also unlikely that spin-conservation plays a significant role at least at the lower energies. Like the An $^+$ , the strong correlation of

 $E_{\rm p}$ (5d<sup>2</sup>) to the rates of reaction (2) in Fig. 3 strongly suggests that  $E_p(5d^2)$  more strongly influences the reaction rate than spin-conservation. Nevertheless, spin conservation may still play a role at higher energies.

A high energy feature is observed for CeO $^+$ , PrO $^+$ , and NdO $^+$ in Fig. 1 with apparent thresholds of  $\approx$  1–2.5 eV. Similar high energy features appear in the  $An<sup>+</sup>$  reactions, but these are less pronounced than those in the  $Ln<sup>+</sup>$  reactions. This feature was also observed in GIBMS studies of reaction (2) for  $Pr^+$  and  $Nd^+$ , and it was speculated that this higher energy pathway was the formation of an excited state  $LnO<sup>+</sup>$  by a spin-allowed reaction. This was further substantiated by quantum chemical calculations that indicated at least one excited state with the correct spin and consistent excitation energy to the observed threshold.<sup>17,19</sup> These features appear inversely correlated with  $E_{\rm p}(n{\rm d}^2)$ ; however, this is likely an artifact from the magnitude of the reaction cross section of the lower feature that obscures the start of the threshold region of the higher energy feature cross section. Indeed, when subtracting the low energy feature of the  $LnO<sup>+</sup>$  and  $AnO<sup>+</sup>$  cross sections in Fig. 1 and 2 by scaling the LGS cross section, the apparent thresholds are indistinguishable within the limits of uncertainty.

As noted above, it has been previously proposed that the role of  $E_p(nd^2)$  in modulating the reaction rate is by partially determining the energy relative to the reactants of the crossing seam between the PESs originating from the ground state M<sup>+</sup> and the excited state with two d-electrons.<sup>22</sup> Here, we consider a simple potential energy surface that consists of an initial intermediate formed, an association of  $CO<sub>2</sub>$  with ground state  $\text{M}^+$  , forming the  $\text{M}^+ \text{-}\mathrm{CO}_2$  adduct. This has been demonstrated for both Sm<sup>+</sup> and Gd<sup>+</sup>, where  $D_0(\text{Sm}^+\text{-}\text{CO}_2) = 0.42 \pm 0.03 \text{ eV}^{15}$ and  $D_0(Gd^+$ -CO<sub>2</sub>) = 0.38  $\pm$  0.05 eV<sup>16</sup> have been measured in collision induced dissociation (CID) reactions. A second intermediate presumably exists where the metal inserts into activated carbon dioxide bond forming O-M<sup>+</sup>-CO. This is likely equivalent to an association complex between ground state MO+ and CO. Indeed, CID reactions of such species yield  $D_0(\text{OSm}^+\text{-CO})$  = 0.97  $\pm$  0.09 eV<sup>15</sup> and  $D_0(\text{OGd}^+\text{-CO})$  = 0.57  $\pm$ 0.05  $e, V^{16}$  bond energies consistent with a weakly bound iondipole interaction. Thus, Table 3 indicates that  $M^+$ -CO<sub>2</sub> and O-M<sup>+</sup>-CO will usually have different spin states. Because formation of the  $MO<sup>+</sup>$  ground state requires two *n*d-electrons from  $M^*$ , the surface that includes the second intermediate



<sup>a</sup> Ref. 28 (<https://physics.nist.gov/PhysRefData/Handbook/periodictable.htm>).  $\frac{b}{c}$  Ref. 26.  $\frac{c}{c}$  Ref. 17.  $\frac{d}{c}$  Presumed based on calculation trends. <sup>f</sup> Ref. 15. <sup>g</sup> Ref. 43 (<https://www.lac.universite-paris-saclay.fr/Data/Database/>). <sup>h</sup> See also ref. 13 and ESI. <sup>i</sup> Unless otherwise noted, ref. 6.  $<sup>j</sup>$  Ref. 44.</sup>

must evolve from the excited state asymptote  $E_{\rm p}(n{\rm d}^2)$  higher in energy than the ground state  $\text{M}^{\text{+}}$ . The MO $^{\text{+}}$  bond energy for An $^{\text{+}}$ and  $\text{Ln}^+$  also depend upon  $E_\text{p}(\textit{nd}^2)$  because of the requirement of the two d-electrons to form the ground state bond so that the energy of the second intermediate is also partially set by this energy (the remaining influence is the strength of the  $OM<sup>+</sup>-CO$ bond). The ground and excited state surfaces interact to form a crossing seam that must be traversed to form ground state products from the ground state reactants. Therefore, when  $E_{\rm p}(nd^2)$  is sufficiently high, as is the case for Sm<sup>+</sup>, Eu<sup>+</sup>, Pu<sup>+</sup>, and Am<sup>+</sup>, the barrier exceeds the reactants energy. When  $E_{\rm p}(nd^2)$  is sufficiently low, as is the case for Ce<sup>+</sup>, Pr<sup>+</sup>, Nd<sup>+</sup>, Th $^+$ , U $^+$ , and Np $^+$  the barrier is localized between the two intermediates. Notably, the observed reaction rate is the equilibrium between the  $\text{M}^{\text{+}}\text{-}\mathrm{CO}_2$  falling apart forming the reactants and the forward reaction forming the products. Thus, the height of the crossing seam relative to the  $M^+$ –CO<sub>2</sub> adduct defines how competitive the forward reaction is compared to the backward reaction, and, therefore, the rate limiting step. Fig. 4 illustrates this concept using reaction (2) surfaces for  $U^+$ and Pu $^+$ . Fig. 4 is semiquantitative estimating intermediate well depths from  $Sm<sup>+</sup>$  CID reactions<sup>15</sup> and product energies from  $UO<sup>+</sup>$  and PuO<sup>+</sup> BDEs.<sup>23,39</sup> PCCP<br>
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The use of the  $Sm<sup>+</sup>$  data is likely reasonable assuming that the intermediates are only electrostatic interactions between the ground state reactant and the ground state product and the corresponding ligand  $(CO<sub>2</sub>$  and CO). However, there may be fundamental differences between  $Sm^{+}$  and the  $An^{+}$  that make this analysis, Fig. 4, semiquantitative. Most notably, f-orbital



**Fig. 4** Semiquantitative potential energy surface of  $M^+ + CO_2 \rightarrow MO^+ +$ CO indicating that  $E<sub>p</sub>$  and the intermediate well depth determine the crossing seam between the surface originating from the ground state reactant asymptote (black) and the surface leading to the ground state products. For species like Pu<sup>+</sup> (blue) that have a sufficiently high promotion energy,  $E_p = 2.14 \text{ eV}^{39}$  the crossing between surfaces exceeds the reactant energy and a barrier is observed. For species like  $U^+$  (blue) that have a lower promotion energy,  $E_p = 0.57 \text{eV}^{\frac{39}{2}}$  is lower than the reactant energy forming a localized barrier between the intermediates that affects the observed reaction efficiency. The intermediates are considered electrostatic interactions between ground state  $M^+$  and  $CO_2$  ( $M^+$ – $CO_2$ ) and ground state  $MO^+$  and  $CO$  ( $O-M^+$ –CO). Values are estimated from  $Sm^+$ CID reactions.<sup>15</sup> Product asymptotes are taken from the BDEs from Marcalo and Gibson<sup>39</sup> or Zhang et al.<sup>23</sup>

participation cannot be ruled out with the present data. Nevertheless, there are key similarities between  $Sm^+$  and  $Pu^+$  that suggests that this explanation is plausible. Both  $Sm^{+}$  and  $Pu^{+}$ have similar promotion energies,  $E_p(Sm^+) = 2.35$  eV and  $E_{\rm p}({\rm Pu}^+)$  = 2.14 eV,<sup>28</sup> and both M<sup>+</sup> have barriers to an otherwise exothermic reaction as shown in Fig. 1 and 2. Computational modelling of the potential energy surface may provide additional information, but that is beyond the scope of this text.

The comparison of the  $Ln<sup>+</sup>$  and  $An<sup>+</sup>$  reaction (2) cross sections are also instructive. In general, the absolute magnitude of the  $AnO<sup>+</sup>$  cross sections are consistently higher than their Ln<sup>+</sup> counterparts. Because the primary difference between the  $Ln<sup>+</sup>$  and the  $An<sup>+</sup>$  is the possibility of the f-orbitals participating in the reaction, this difference may indicate 5f-orbital participation for the  $An^+$ . As noted above, the rate limiting step, the crossing seam energy, appears to be influenced by  $E_p$ and the insertion intermediate, O-M<sup>+</sup>-CO. The interaction of the carbonyl with  $M^+$  may indicate whether the 5f-orbitals participate in the O-An<sup>+</sup>-CO through  $\pi^*$  back-bonding; however, CID reactions of ThCO<sup>+</sup>  $[D_0(\text{Th}^+$ -CO $)$  = 0.94  $\pm$  0.06 eV] indicate that even when back-bonding does occur, $^{21}$  it may not be sufficiently strong to distinguish between cases where it does not. Consequently, the carbonyl interaction is unlikely to greatly affect the energy of the crossing seam. The energy of the O-M<sup>+</sup>-CO intermediate's energy is also set by the ground state product asymptote. Previous work has suggested an intrinsic BDE model<sup>39,42</sup> for both the  $LnO<sup>+</sup>$  and  $AnO<sup>+</sup>$ . This model assumes that the orbitals used throughout both series to bind O are constant (*i.e.* the d-orbitals). Thus any  $\text{LnO}^+$  or AnO<sup>+</sup> BDE could be calculated with knowledge of the intrinsic BDE  $[D_0(M^+\!\!-\!\!O)^\ast]$  and the  $M^+$   $E_\mathrm{p}(n\mathrm{d}^2)$  according to eqn (7):<sup>42</sup>

$$
D_0(M^+ - O) = D_0(M^+ - O)^* - E_p(nd^2)
$$
 (7)

Such a model suggests that the d-orbitals are primarily involved in forming the  $MO^+$  bond. A careful comparison of the  $LnO^+$ and  $AnO<sup>+</sup>$  BDEs trends suggests that the  $AnO<sup>+</sup>$  BDEs may include some 5f-orbital participation. This is observed by the slope of the  $LnO<sup>+</sup>$  trend vs. the slope of the  $AnO<sup>+</sup>$  trend in Fig. 5, where the latter deviates from the predicted slope  $(i.e.$  unity) in eqn (7). This suggests that 5f-orbital participation may be present in reaction  $(2)$  for the An<sup>+</sup>. Notably, this is not an unambiguous indication of 5f-orbital participation. The  $AnO<sup>+</sup>$  BDEs in Fig. 5 are primarily from high temperature experiments extrapolated to 0 K, so the difference may only be experimental error. Likewise, other effects such as actinide contraction may explain some differences in observed trends. Nevertheless, the magnitudes of the AnO<sup>+</sup> cross sections compared to their  $Ln<sup>+</sup>$  congeners suggest some minor influence of the 5f-orbitals on the reaction rate.

Finally, Fig. 1 and 2 establish the presence of excited states in the current work when comparing the previous GIBMS work.15,17,19 This is most notably evident in the threshold region of SmO<sup>+</sup>, where the apparent threshold of the ICP-MS/ MS work is approximately 1.5 eV lower in energy than the GIBMS work.<sup>15</sup> Excited states are expected in the current work because the ICP source temperature is 8000–10 000 K, and there is limited ability in the ICP-MS/MS to quench excited states



**Fig. 5** LnO<sup>+</sup> (blue) and AnO<sup>+</sup> (red) BDEs as a function of  $E_p(nd^2)$  for  $n = 5$ and 6. Vertical dashed lines are  $E_{\sf p}$ (nd<sup>2</sup>) for the indicated M<sup>+</sup>. The solid lines represent the least-squares linear regression fit of each series. The LnO<sup>+</sup> fit is 8.41–1.01 $E_p$ (5d<sup>2</sup>),  $R^2$  = 0.98. The AnO<sup>+</sup> fit is 8.53–0.79 $E_p$ (6d<sup>2</sup>),  $R^2$  = 0.98. Ln<sup>+</sup>  $E_p$  and LnO<sup>+</sup> BDEs are taken from Gibson<sup>42</sup> or Armentrout and coworkers. $^{14,17,19,27,37}$  An $^+$   $E_{\sf p}$  and AnO $^+$  BDEs are taken from Marcalo and Gibson<sup>39</sup> or Armentrout and coworkers.<sup>13,23</sup> PaO<sup>+</sup> is not shown but included in the fit.

prior to injection into the reaction cell. By contrast, the GIBMS work utilizes a 1 m tube pressurized to  $\approx 0.5$  Torr of buffer gas where ions undergo  $\approx 10^5$  collisions.<sup>41</sup> A conservative estimate of the ion electronic distribution in the GIBMS experiments is 700  $\pm$  $400$  K.<sup>45-49</sup> Likewise, the SIFT-MS experiments utilize a drift tube pressurized to 0.35 Torr with the reactive gas introduced downstream.7 The Gibson and coworkers FTICR-MS work thermalized ions produced by laser desorption/ionization by cooling periods in  $10^{-5}$  Torr Ar.<sup>1-3,5</sup> Notably, the FTICR-MS data is consistently low compared to GIBMS data. The difference is unclear. Nevertheless, GIBMS and SIFT-MS  $k/k_{\text{col}}$  for CO<sub>2</sub> are in good agreement, Table 2. This is likely due to similar thermalization processes of the ions prior to reaction that presumably yield similar electronic energy distributions. Assuming a 700 K distribution, the average electronic energy  $(E_{el})$  is 0.00–0.06 eV, Table 1. By contrast, at 5000 K,  $E_{el}$  is 0.19-0.66 eV.

Excited states can, in principle, be more or less reactive than the ground state depending on how the PES evolves from that state. When many surfaces exist with crossing seams, then presumably a pathway exists for the reaction to proceed to product despite the contour of the starting PES. In this limit,  $E_{el}$  would act as additional energy for reaction. Because more energy is available for reaction, a more efficient reaction is observed. The Ln and An have many low-lying states, $^{28}$  so presumably the density of states in the reactant channel allows the extra  $E_{el}$  to act as additional energy available for reaction. Consequently, the higher electronic distribution in the ICP-MS/MS experiments likely leads to a higher observed reaction efficiency than the more thermalized GIBMS and SIFT-MS experiments, Table 2.

## Conclusion

The energy dependence of the reactions of  $Ln^+ (Ln^+ = Ce^+, Pr^+,$  $Nd^+$ ,  $Sm^+$   $Eu^+$ ) and  $An^+$   $(An^+ = Th^+, U^+$ – $Am^+)$  with  $CO_2$  were studied by ICP-MS/MS where the primary product observed was  $MO^+$ .  $k/k_{\text{col}}$ , compared to the LGS collision limit, for the barrierless exothermic reactions observed are correlated to  $E_{\rm p} (5 {\rm d}^2)$  for the  ${\rm Ln}^+$  and  $E_{\rm p} (6 {\rm d}^2)$  for the  ${\rm An}^+$  suggesting that the d-electrons are primarily involved in setting the rate of reaction (2). Barriers in excess of the reaction enthalpy were also observed for the reactions of Sm<sup>+</sup>, Pu<sup>+</sup>, and Am<sup>+</sup>. These barriers can likely be attributed to a crossing seam between PESs originating from the ground state reactants and the excited state surface that leads to the ground state products. The energy of the crossing seam is likely determined by the energy of the reaction intermediates and  $E_p$ . For reaction (2), the energy of the more influential O-M<sup>+</sup>-CO intermediate is heavily influenced by the product asymptote because it is a weakly bound complex between  $MO<sup>+</sup>$  and CO. Because the enthalpy of the reaction is also dependent on  $E_p(nd^2)$ , the promotion energy has a large impact so that even very exothermic reactions like Pu<sup>+</sup> + CO<sub>2</sub> ( $\Delta H_2$  =  $-1.30$   $\pm$ 0.20 eV) may have a barrier if  $E_p$  (where  $E_p(Pu^+) = 2.14 \text{ eV}^{22}$ ) is sufficiently high. This suggests that the *n*d-orbitals are primarily involved in reaction (2). Nevertheless, differences in the absolute reaction cross sections between the An<sup>+</sup> and Ln<sup>+</sup>, where the former is generally greater than the latter, suggest that the 5f-orbitals play at least a minor role. This role is likely in forming the  $AnO<sup>+</sup>$  bond. Paper  $G_n^c = G_n^c = P_1^c$  Motion  $Q_n^c = G_n^c$  Motion 2023. Downloaded by Research Commons Articles. Published are commons and the the number primary produce of the resulting produce of the resulting produce of the resulting

Notably, reaction (2), where  $CO<sub>2</sub>$  and CO are both singlets, is often spin-forbidden for  $M^+$  all metals considered here except Th<sup>+</sup> and U<sup>+</sup>. The correlation of the reaction rate to  $E_{\rm p}(nd^2)$ regardless of whether the reaction is spin-allowed indicate that for the  $Ln<sup>+</sup>$  and  $An<sup>+</sup>$ , spin-restrictions do not appear to be a strong influence on the reaction rate.

# Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

This research was supported by the Open Call Initiative under the Laboratory Directed Research and Development (LDRD) Program at Pacific Northwest National Laboratory (PNNL). PNNL is a multiprogram national laboratory operated for the U.S. Department of Energy (DOE) by Battelle Memorial Institute under Contract No. DE-AC05-76RL01830. The authors also thank Prof. Peter Armentrout, University of Utah for helpful advice while formulating the arguments in this manuscript. The authors also thank Prof. Armentrout for providing measurements from guided ion beam tandem mass spectrometry experiments of the reactions  $Ce^{+}/U^{+}$  +  $CO<sub>2</sub>$ . These works are currently in progress.

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