A Blyholder mechanism in the chemisorption of N\textsubscript{2}O on Ni(111) – studied with Auger-photoelectron coincidence spectroscopy

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A B L I N E  A R T I C L E  I N F O

Keywords:
Auger-photoelectron coincidence spectroscopy
Blyholder model
Heterogeneous catalysis

A B S T R A C T

In heterogeneous catalysis the surface-adsorbate bond strength is critical for the function of the system. Here we study a series consisting of bilayer and monolayer N\textsubscript{2}O on Ni(111) and employ Auger-photoelectron coincidence spectroscopy (APECS) to study the interaction between the molecule and the substrate directly. We observe intensity in the nitrogen Auger spectra that arise from the interaction between molecule and surface (not observed in free molecules) whereas the oxygen spectra are thickness-independent. Since the two nitrogen atoms of N\textsubscript{2}O are chemically inequivalent we can assign the intensity present in the bilayer and monolayer cases to orbitals centered on the terminal nitrogen which is closest to the Ni(111) surface. Using \textit{ab initio} molecular dynamics and solid-state density functional theory calculations we infer a Blyholder model of the surface bond as consisting of donation from the terminal nitrogen lone-pair valence orbital with back-donation from the metal into the unoccupied orbitals on that nitrogen. This coincidence technique can readily be used to study substrate–adsorbate interactions directly with chemical and orbital specificity — this opens up prospects to study fundamental steps of molecular adsorption and heterogeneous catalysis with unprecedented detail.

1. Introduction

The interaction between metal surfaces and molecules constitutes a challenging and important area of study not the least for applications in catalysis. The oxides of nitrogen (NO\textsubscript{x}) play a particularly important role as they are products from combustion and thus contribute to the anthropogenic concentration in the atmosphere of these gases. The main source of NO\textsubscript{x} is from surface emissions of nitrous oxide, N\textsubscript{2}O, which also is the third most significant greenhouse gas [1]. N\textsubscript{2}O as an ozone-depleting agent is predicted to remain the main driver of ozone depletion during the 21st century [2].

N\textsubscript{2}O is a triatomic molecule with the configuration N⋯N⋯O that is linear in the isolated case with the terminal (N\textsubscript{T}) and central (N\textsubscript{C}) nitrogen atoms being chemically inequivalent. The N\textsubscript{T}⋯N\textsubscript{C} bond is 1.126 Å and the N\textsubscript{C}⋯O bond is 1.191 Å. [3] N\textsubscript{2}O is an obvious candidate for heterogeneous catalysis since it is thermodynamically unstable with respect to N\textsubscript{2} and O\textsubscript{2} (standard enthalpy of formation \(\Delta H^*\) at 82 kJ/mol) but with a high activation energy barrier [4].

Catalytic conversion of N\textsubscript{2}O into non-pollutant components is an active challenge for the surface science community [5,6] and the development of catalytic converters in combustion engines (reviewed by [7]) is a success story for the field. If new fuels (e.g. NH\textsubscript{3} as an alternative to H\textsubscript{2} [8]) or energy conversion systems like fuel cells [9] are to be employed the problem of removing pollutants remain.

Nitrous oxide has been extensively studied in its own regard using electron spectroscopy. Since the N\textsubscript{T} and N\textsubscript{C} have different chemical surrounding, both core level and unoccupied orbitals can be studied, with electrons [10] and tunable X-rays (e.g. from synchrotrons) to obtain information about site specific excitations into different orbitals in the electronic structure [11–13].

An interesting application for this is that N\textsubscript{2}O decomposition on Ni surfaces can be used to measure Ni particle sizes in catalyst via pulse titration of the gas over the catalyst [14].

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https://doi.org/10.1016/j.apsusc.2024.160340
Received 7 March 2024; Received in revised form 17 May 2024; Accepted 19 May 2024
Available online 21 May 2024
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The adsorption of N₂O on various metal surfaces has been extensively studied as reviewed by Zeigarnik [15]. From early studies on Ni [16] to molecular orientation after physisorption on Ni(111) surface [3], and further: monolayers on Ag [17] and ab initio modeling of the self-assembly on these surfaces [18]. Similar to NO₂ and N₂, the decomposition of N₂O has also been studied on various surfaces including Ni(110) [19]. Kodama et al. have also shown that N₂O does not decompose on fcc(111) faces, but on the vicinal (755) surface of Ni(111) it occurs while step-sites are available and when those are saturated by O, one of the decomposition products, further adsorption leave the N₂O intact [20].

The chemisorption bond of molecules on metal surfaces may in some instances be conceptualized by the Blyholder model, where donation of HOMO (highest occupied molecular orbital) electron density from the molecule to the surface is compensated by a back-donation of electron density from the surface into the LUMO (lowest unoccupied molecular orbital) to keep the system charge neutral. A classical example is CO chemisorption on Pt(111) where the 2Ө and 5Ө are involved in the back-donation and donation [21].

The electronic ground state of N₂O is

\[ 1σ^2 2σ^2 3σ^2 4σ^2 5σ^2 6σ^2 1π^2 7σ^2 2Σ^+ \]

with 1σ, 2σ and 3σ corresponding to localized O 1s, N_C 1s and N_F 1s respectively.

The Auger spectrum of free N₂O was analyzed already in 1976 by Connor et al. using restricted Hartree–Fock and limited configuration interaction wave functions coupled to the one-center model for Auger intensities [22]. These authors could grossly assign features to localization of orbitals and to the spread of singlet and triplet multiplet states.

The non-equivalent N atoms also make for ideal studies with coincidence spectroscopy. With studies including the dissociation of N₂O using electron–ion coincidences [23], site-specific fragmentation [24], multi ion–electron coincidences [25], and site-selective Auger electron spectra recorded using photoelectron-Auger electron coincidence spectroscopy (APECS) [26]. All of the mentioned coincidence studies were performed in the gas phase and to the best of our knowledge no APECS studies have to date been published on adsorbed N₂O.

Herein we present the first APECS study on adsorbed N₂O on Ni(111). This particular surface was chosen since the weak chemisorption leaves the molecular adsorbate intact [15]. Together with ab initio calculations we characterize the surface-adsorbate bond mechanism in detail.

2. Materials and methods

2.1. Samples and preparation

The Ni(111) crystal (99.995% MaTecK Gmbh) was cleaned with repeated cycles of argon sputtering (1.5 kV, 10 mA, 5.10⁻⁶ mbar) for 5–10 min each followed by annealing to 1070 K for 3 min. The surface was then checked for contamination using X-ray photoelectron spectroscopy (XPS) revealing a clean surface with O 1s and C 1s contributions within the noise of a survey spectrum. After cooling the crystal using liquid helium, N₂O (5N, Messer) was dosed using a leak valve into the experimental chamber. The thermocouple readout during deposition was 31 K (with the temperature on the sample probably higher as discussed in the results below) with N₂O at a pressure of 1.10⁻⁷ mbar. For the thick film a total of 100L N₂O was dosed and for the intermediate and monolayer films 5L was dosed prior to heating to desorb all but the desired thickness, details in the text below.

![Fig. 1. (a) shows the evolution of the N 1s core level while heating the sample. At 650 s the thermocouple readout was 39 K and at 800 s 42 K with all molecules desorbed, further discussed in the text. The bottom two panels shows 3 typical spectra of N 1s (b) and O 1s (c) of multilayer, bilayer and monolayer coverage. With dashed lines highlighting the binding energy shifts of the different coverages and red arrows the binding energy of dissociated O and N.](image-url)
spectrometer is a considerable increase in the coincidence count rate as seen in Table 1.

During a APECS measurement, events when both spectrometers record an electron within a given time window, are recorded as a coincidence event. This includes both true and accidental coincidences. Accidental coincidence events are when electrons from different ionization events hit the detectors in the same time window. The true events are extracted from the total coincidence counts by subtracting the accidental coincidences, as described in detail in [27,29]. In all coincidence spectra below, only the true coincidence events are plotted.

The statistics for the coincidence measurements are presented in Table 1. The binding energies were calibrated using the Fermi level of the Ni(111) crystal as a reference and the kinetic energies were calibrated using the Fermi level plus the work function on Ni(111), 5.23 eV [30]. For details on the APECS technique at the CoESCA station see [27,29,31,32].

2.3. Theoretical methods

Density functional theory (DFT) calculations were carried out using projector augmented-wave (PAW) formalism [33] as implemented in the Vienna ab initio simulation package (VASP) [34,35]. The revised Perdew–Burke–Ernzerhof (rPBE) [36] and Heyd–Scuseria–Ernzerhof (HSE06) [37] exchange–correlation functionals with the Grimme D3 correction [38] to weak disperse interactions were utilized to optimize the geometry and calculate electronic structure. Mostly the rPBE functional was used unless otherwise specified. The plane wave kinetic energy cutoff of 450 eV was used. The first Brillouin zone was sampled with the Monkhorst–Pack scheme [39] with 12 × 12 × 1 k-points. The convergence tolerances of the force and electronic minimizations were 10−4 eV/Å and 10−6 eV, respectively. A vacuum region of > 10 Å was set to avoid artificial interactions between images. Atomic structures were visualized using The Visualization for Electronic and Structural Analysis (VESTA) software [40] and the ”vaspkit” code was employed for postprocessing the results [41].

In addition to solid state calculations we have carried out ab initio calculations on N₂O and N₂O-Ni using the MOLCAS program suite [42], with the purpose of having a theoretical characterization of the backdonation. We focused first on the electronic structure of the ground state with the purpose of having a theoretical characterization of the backdonation orbitals. The convergence of the wave-function was based on a state-average procedure considering a configuration interaction between the lowest 10 states.

For the calculations of the Auger transition intensities we firstly note that these are governed by two-electron Hamiltonian elements that couple to the continuum function of the outgoing Auger electron. As the precise determination of the Auger electron continuum function in the anisotropic molecular potential is a quite daunting task only a few methods have been put forward at this level of theory and applied for the smallest free molecules. Therefore a more practical approach adopted here is to consider the local nature of Auger transitions and use so-called one-center models [44,45], where two-electron coupling to the continuum is replaced by atomic continuum integrals weighted by the local on-atom atomic orbital expansion coefficients [46] of the involved (two) molecular orbitals (MOs) for the atom in question. Providing a quick inspection of the spectra, one can utilize an even simpler model based on orbital population, and only qualitatively consider the variation of the coupling to the continuum. In this view, combinations of orbital populations can guide the Auger spectrum even for surface adsorbates. It is notable that there are both singlet and triplet spin couplings for the two-hole Auger final states, but that a propensity rule [44,45] favors the singlets (owing, respectively, to constructive and destructive interference between Coulomb and exchange continuum integrals). Finally, we note that the two hole states are subject to ”Coulomb explosion” leading to dissociative final states and broad features, possibly except for one or two of the lowest lying final two-hole states [47]. It is to be noted that second quantization derived intensity expressions for Auger transitions between core ionized and double hole final states in a non-orthogonal basis contains many terms [48], but that the common Wentzel one-term ansatz [49] is still almost always assumed in actual calculations. That raises the question of which orbital basis is to be preferred. We have here in the below the presentation of the spectra using the one-center intensity model, assuming both ground and core hole optimized orbitals, bearing in mind that rather significant differences may arise.

Table 1

<table>
<thead>
<tr>
<th>System</th>
<th>True counts</th>
<th>True/Acc</th>
<th>Countrate (c/s)</th>
</tr>
</thead>
<tbody>
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<td>24595</td>
<td>0.71</td>
<td>4.3</td>
</tr>
<tr>
<td>Bilayer</td>
<td>84149</td>
<td>1.96</td>
<td>3.5</td>
</tr>
<tr>
<td>Monolayer</td>
<td>155890</td>
<td>1.04</td>
<td>4.4</td>
</tr>
<tr>
<td>Oxygen Multilayer</td>
<td>84582</td>
<td>0.56</td>
<td>15.7</td>
</tr>
<tr>
<td>Bilayer</td>
<td>64735</td>
<td>0.85</td>
<td>12.2</td>
</tr>
<tr>
<td>Monolayer</td>
<td>57182</td>
<td>1.13</td>
<td>1.7</td>
</tr>
</tbody>
</table>

3. Results and discussion

3.1. XPS and coverage

The N 1s core level photoelectron spectrum of N₂O consists of two contributions, one at higher binding energy from the N₂, atom and one at lower binding energy from the N₂ [11]. As can be seen in the multilayer spectra, blue spectrum in Fig. 1b, the contributions are well separated and the binding energy positions and peak widths fit well with earlier studies [16]. The multilayer O 1s primarily consists of one component as can be seen in Fig. 1c with a small contribution at lower binding energies that arises from dissociated O at the surface [3,20,29]. The presence of dissociated O also suggests a presence of dissociated N which could explain the shoulder seen at 401 eV binding energy for all coverages in the N 1s spectra in Fig. 1 [3,50]. The two dissociated species are marked with red arrows in Fig. 1. At minute coverages N₂O is known to dissociate at steps at vicinal Ni(755), a process that is terminated when step-sites are saturated with O [20]. The dissociation observed here occurs at defects on our Ni(111) crystal and the core level spectra suggest that it is in small quantities. As noted, the binding energy position of the dissociated N is well separated from those of N₂O [50]. The KLL Auger spectra from N₂ and N₂O will overlap in our kinetic energy region of interest; the main feature of N₂ Auger spectrum may be found around 360 eV kinetic energy [51] and being a minor contribution to the total spectrum and indistinguishable. This and the contribution to the O Auger spectrum are discussed in detail below.

In order to investigate the interaction between the substrate and the N₂O, thinner films were produced by heating the sample to control desorption. After exposing the cooled Ni surface to 5L of N₂O, it was slowly heated using resistive heating while simultaneously recording
Fig. 2. Panels a–f show the N coincidence maps and integrated spectra from the regions marked with colored lines in the coincidences maps. Panel a also shows a comparison to the coincidence spectra from gas-phase N$_2$O, the gas-phase data is taken from Ref. [26], note the top energy axis for the gas-phase reference. Panels g–l shows the corresponding O coincidence data.

the N 1s and O 1s photoelectron spectra using the two ARTOF spectrometers. Monitoring the N 1s spectra while heating, as seen in Fig. 1a, shows a clear point of the multilayer desorption with a sharp decrease in the peak intensities followed by a small shift ($\Delta = 0.52$ eV) and the appearance of a doublet peak for each N atom, the green spectrum in the middle panel of Fig. 1. With further heating one peak of the doublet disappears with the other remaining unchanged, the orange spectrum in the middle panel of Fig. 1, until all molecules are desorbed and no N signal can be seen. The O 1s spectrum shows a similar trend as seen in the bottom panel of Fig. 1.

A similar behavior is discussed by Väterlein et al. [3] where their thermal desorption spectra revealed multilayer desorption at 70 K, monolayer desorption at 92 K and a third species which they attributed to a dilute monolayer that desorbed at 100 K containing 10% of a monolayer. They also claim no significant binding energy shift for the monolayer and dilute monolayer. This is similar to our results with three distinct steps in the desorption process but we observe a binding energy shift between the two lower coverage cases and that the low coverage case consists of 50% of the intermediate case. However, we observe this with significantly lower temperature read-out indicating that the actual sample temperature in our experiment is higher than the read-out, probably due to the location of the thermocouple. We assign the intermediate state with the doublet peak structure (green in Fig. 1) as bilayer coverage whereas the single peak structure (orange in Fig. 1) is assigned to monolayer coverage, with a molecular coverage of $\theta = 5/9$ determined from calculations as described below. This would follow a Frank–Van der Merwe layer-by-layer growth structure.

3.2. Auger-Photoelectron Coincidence Spectroscopy

3.2.1. Multilayer films

In Fig. 2d the N 1s/N KVV coincidence map of a multilayer N$_2$O film on Ni(111) can be seen. The map is dominated by two distinct intensity regions distributed in binding energy that are marked with blue (406–411 eV) and green (402–406 eV) lines. Looking at the binding energy positions for these regions the intensity comes from the two different N atoms in the molecule and they have seemingly different Auger spectra. By integrating the region around the binding energies for the two different N atoms the Auger spectra for each can be extracted.
These spectra can be seen in Fig. 2a where the blue and green spectra are the integrated region marked with the blue and green lines in the map below. The blue corresponds to N$_T$ and the green to N$_C$. The N$_T$ Auger spectrum contains more distinct features compared to the N$_C$. The Auger spectra of both N atoms are similar to that of gas-phase N$_2$O measured in coincidence by Bolognesi et al. [26] which is also plotted in Fig. 2a for comparison, note the separate kinetic energy axis for the gas phase spectra. The main features of the gas phase spectra are reproduced in the thick film albeit with a shift induced from the solid structure and by the substrate. The two energy axes are aligned so that the main Auger feature at 373.5 eV for both solid and gas end up at the same position in the figure. With these aligned it can be seen that other Auger feature also show some small shifts but are generally well reproduced.

Using a mean free path ($\lambda$) of 11 Å (calculated with the TPP-2M method [52]) for both photo- and Auger electrons (close to the same kinetic energy for this experiment) the effective mean free path for the APECS experiment can be calculated to 5.5 Å or about 2 monolayers [53]. Since the film is thicker than this, the APECS measurement is of pure solid N$_2$O and does not probe the interface with the Ni surface. The O 1s/O KVV coincidence map of the multilayer film can be seen in Fig. 2. It is dominated by 4 features in the 536–538 eV binding energy range, noticeably these exhibit a slight tilt. This correlation of the kinetic energies of the Auger- and photoelectron pairs can only be revealed in a coincidence measurement. While the well known energy sharing between an Auger and a photoelectron (e.g. [54]) leads to intensity stripes at constant kinetic energy sum, here we observe an opposite trend, where the stripes are rather at constant kinetic energy difference. The tilt is most apparent in the multilayer case and completely absent in the monolayer case (Fig. 2). This could be explained by a change in the dielectric function with different film thicknesses resulting in different binding energies for spectra from different depths [55].

3.3.1. Auger spectra of free N$_2$O

The Auger coincidence spectra, Fig. 2, show some salient features spreading over an interval of about 15 eV in kinetic energy. It is clear that for both bi- and monolayer N$_2$O there is a shift of intensity towards higher kinetic energies for N$_T$ compared to the N$_C$ spectra. The former shows an additional weaker structure towards the high energy end for N$_T$ not at all represented for N$_C$. It is notable that the N Auger spectrum of free N$_2$O [22], which is a composite of N$_T$ and N$_C$ transitions, show three main bands in this energy interval, but shifted towards lower kinetic energies by a few eV than the spectra in Fig. 2. The results of the RASSCF/PT2 calculations in terms of leading configuration and energy of the outer Auger states are given in Table 2. Here the energy of the outermost state (2σ$^*$ − 1 2𝛱$^*$) is used as a reference point. By inspecting the core separated spectra in Fig. 2d-f it is clear that the outermost band (at high Auger electron kinetic energies) of the composite N spectrum covering 0–6 eV is dominated by Auger transitions connecting to the N$_T$. The separated N$_T$ spectrum shows two bands in this region. The computational assignment is here clear — the outermost of these two bands refers to terminally localized 2σ and 7ε orbitals, while the inner one of the two also receives contributions from the more central N localized 1x orbital, n.b. in combination with 2ε MO so generating the 1x−1 2𝛱−1 1 𝜓 and 3 𝜓 eigenstates. We note, as is the general case for molecular valence Auger spectra, that there is a great deal of breakdown of the molecular orbital picture in analyzing the spectrum [47,56]. Thus only for the outer MOs (1x, 7ε, 2σ) a one-to-one assignment can be made between two MOs and (spin-coupled) single configuration eigenstates. Here a division between inner–inner-, inner–outer and outer–outer valence regions is appropriately based on the inclusion of the atomic 2s or 2p into the MOs. It is often the case that inner–inner valence states suffer a complete breakdown, the inner–outer a partial breakdown, while only the outer–outer valence states may receive an orbital assignment including only two MOs [47]. This means that both orbital localization and the breakdown effects come into play when analyzing the N$_2$O Auger spectrum. Thus going to lower Auger energies and the second, N$_C$, dominated broad band the breakdown effects become progressively more important.

3.3.2. Bi- & monolayers

The bi- and monolayer coverages were produced by stopping the heating of a thick film (5L exposure) when the desired spectral features were observed and subsequently the sample was cooled down to base temperature again for the duration of the measurements. APECS measurements were then performed on these two coverages. The resulting APECS maps and integrated region spectra can be seen in Fig. 2 for N 1s/N KVV and O 1s/O KVV for the few- and monolayer coverages.

Comparing the N coincidence maps in Fig. 2 for the different coverages the shift seen in the XPS spectrum from the desorption is also seen in the Auger although much larger, \( \approx 2.5 \) eV. Particularly the N$_T$ Auger spectra are comparatively different from the multilayer and gas phase spectra. The monolayer spectrum contains sharp features whereas the bilayer case contains both the features of the monolayer interaction with the substrate and the interaction with the molecular layers above it. As discussed previously, the signal in the bilayer case is dominated by the top-most layers owing to the shallow information depth in APECS.

The O 1s/O KVV coincidence map, similar to the N 1s/N KVV case, contains two different binding energy separated Auger spectra where the one at high binding energy has a distinct structure similar to that seen in the N spectra. The Auger spectra from the 534–537 eV binding energy resemble those of the electron–ion coincidence gas phase Auger spectrum [24]. The low binding energy part contains a single broad peak in the Auger kinetic energy and binding energy region where we expect dissociated O to be present [3,20,29].

The most striking feature in the monolayer case is the intensity found above 380 eV kinetic energy for N$_T$. It is also present in the bilayer spectra but with lower intensity as expected since the signal from the monolayer is damped by the presence of the molecular layer on top of it and it is not observed in the thick film case (Fig. 2j).
Fig. 3. Diagram of the active space used for describing the ground state of the N₂O+Ni idealized structure. The MOs are disposed according to the symmetry components of the Cᵥ point group, and they are arranged by orbital energy from left to right. The vertical dashed line divides the occupied orbitals (left) from the unoccupied orbitals (right). Numbers in parenthesis denote natural occupation numbers.

Table 2
Relative energies, in eV, of the triplet and singlet Auger states of N₂O. The reference value corresponds to the ground state 3ς⁻ of N₂O²⁺.

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<tr>
<th>State</th>
<th>Config</th>
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<tbody>
<tr>
<td>³ς⁻</td>
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</tr>
<tr>
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<td>2ς⁻</td>
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</tr>
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</tr>
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<td>½ς⁺</td>
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<td>6.94</td>
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Table 3
N₂O MO coefficients for the valence σ orbitals.

<table>
<thead>
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<th>sym. adpat. basis</th>
<th>7ς</th>
<th>6ς</th>
<th>5ς</th>
<th>4ς</th>
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<td>N₆ 1ς</td>
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<td>+0.5690</td>
<td>-0.1261</td>
<td>-0.1877</td>
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</table>

The orbital characterization is also becoming less relevant due to the above mentioned MO breakdown effects. Furthermore, analyzing the role of the surface, see below, it is clearly the outer 2p region which matters most.

3.3.2. Auger spectrum of N₂O-Ni

Going from free to surface adsorbed N₂O the most salient spectral feature in the Auger spectra is thus the addition of a structure at 385 eV kinetic energy appearing in the N₆ spectrum but not for N₆C. Evidence that this feature relates to backdonation is reflected in the electronic structure of the ground state. Therefore, in order to have a meaningful description of the backdonation, we have considered an idealized model of a system composed of one molecule of N₂O plus a single Ni atom, all disposed in a linear structure.

5 eV region. We note from Tables 3 and 4 that both 7ς and 2ς orbitals have strong N₆C 2p contributions, and the strong composite N₆ spin band at 370–375 eV should be assigned to these two states. Correspondingly the N₆C 2p contributions to these orbitals are substantially smaller, explaining the much lower intensity in this region. Going further down in the spectra, into the 2ς/2ς region, there is a rather mixed representation of N₆ and N₆C in terms of atomic orbital composition, and we also observe rather smeared out "look-alike" features in the two spectra.
The intensity of these transitions as a response to the changes on the MO a marginal effect, the presence of the nickel atom change the relative lower intensities as discussed above. Although it can be considered as peaks in the experimental spectrum (taking $N_1$ corresponds to a double ionization of the nickel atom. With our active ionization of the Nickel atom. We thus understand that this is a first corresponding to the extra structure at higher kinetic energies corresponds to states involv-

The local nature of the Auger transitions suggests that the doubly ionized atomic nickel states have very low intensities, for they would involve non-local electron decays. On the other hand, those states involving singly ionization of the nickel atom together with singly ionization of the $N_2O$ molecule would have larger intensities, although smaller than the main peaks. From Mulliken charge analysis, the states were divided into the ones involving doubly charged Ni (Ni$^{+2}$) and the ones corresponding to singly ionized Ni plus singly ionized $N_2O$ (Fig. 5f). The blue solid lines stand for the states that would provide extra structure at 380–385 eV in panel (b), showing density of doubly charge states with energies below $^1\Sigma^−$ is here divided into double ionized Ni (red) and singly ionized Ni plus singly ionized $N_2O$ (blue). Note that, in contrast to panel c.

Within this linear model, we can understand the effect of the nickel atom as (i) the influence on the Auger transitions corresponding to the states of isolated $N_2O$ and (ii) the introduction of new transitions (mainly due to the backbonding). For the first part, a partial Auger spectra was calculated based on the one-center approximation (Fig. 4). Here we look at the same final states of single $N_2O$, and we report the spectra calculated for $N_2O$ and for $N_2O+Ni$. The assignments of the final states respect the labels of single $N_2O$ molecule, also shown in Table 2. It is worth pointing out that the lowest-lying $3\Sigma^-$ state has no intensities in the one-center approximation. The reason is that the Coulomb and exchange integrals cancel each other out in this case. For all three core-holes we observe three main bands in this energy interval, the first corresponding to the $1\Delta$ plus $1\Sigma^+$ states, and the second and third related to the $1\Pi$ and $1\Sigma^-$ states. The triplet $\Pi$ states have lower intensities as discussed above. Although it can be considered as a marginal effect, the presence of the nickel atom change the relative intensity of these transitions as a response to the changes on the MO coefficients, when compared to the isolated $N_2O$ (Tables 5 and 6).

Our interpretation of the overall spectra is that (i) the most intense peaks in the experimental spectrum (taking $N_2$, for instance, the peaks from 365–375 eV) correspond to the transitions on $N_2$, principally the ones linked to the singlet states as discussed above, and (ii) the extra structure at higher kinetic energies corresponds to states involv-
ing ionization of the Nickel atom. We thus understand that this is a consequence of the latter effect of the nickel atom (the introduction of new transitions). This interpretation is supported by our RASPT2 calculations for the doubly charged $N_2O+Ni$. In fact, the lowest-lying doubly charged state lies below the $^3\Sigma^−$ state by more than 13 eV, and corresponds to a double ionization of the nickel atom. With our active space, more than 300 roots were found in this interval, below the $^3\Sigma^−$, which was set as the reference energy. Besides, the density of states shows a broad peak between $−4$ eV and $−10$ eV, which is in a very good agreement with the experimental value of the extra structure (Fig. 5).

Table 4
$N_2O$ MO coefficients for the valence $\pi$ orbitals. The $\pi$ MOs are $s$/$y$-doubly degenerate ($h_\pi/h_\Sigma$).

<table>
<thead>
<tr>
<th>sym. adpat. basis</th>
<th>$2\pi$</th>
<th>$1\pi$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_2$ 2$p_x$/2$p_y$</td>
<td>−0.5659</td>
<td>+0.3449</td>
</tr>
<tr>
<td>$N_2$ 2$p_x$/2$p_y$</td>
<td>−0.2162</td>
<td>+0.5742</td>
</tr>
<tr>
<td>$O$ 2$p_x$/2$p_y$</td>
<td>+0.7743</td>
<td>+0.4849</td>
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</table>

Table 5
$N_2O$+Ni MO coefficients for the valence $\pi$ orbitals.

<table>
<thead>
<tr>
<th>sym. adpat. basis</th>
<th>$N_2O$ 7$x$</th>
<th>$N_2O$ 6$x$</th>
<th>$Ni$ 3$d_{x^2}$</th>
<th>$Ni$ 3$d_{y^2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_2$ 1$x$</td>
<td>−0.048</td>
<td>−0.020</td>
<td>0.000</td>
<td>0.030</td>
</tr>
<tr>
<td>$N_2$ 2$x$</td>
<td>−0.799</td>
<td>−0.006</td>
<td>0.000</td>
<td>0.271</td>
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<tr>
<td>$N_2$ 2$p_x$</td>
<td>+0.461</td>
<td>+0.069</td>
<td>0.000</td>
<td>−0.045</td>
</tr>
<tr>
<td>$O$ 1$x$</td>
<td>+0.000</td>
<td>+0.015</td>
<td>0.000</td>
<td>−0.001</td>
</tr>
<tr>
<td>$O$ 2$x$</td>
<td>−0.036</td>
<td>+0.039</td>
<td>0.000</td>
<td>−0.003</td>
</tr>
<tr>
<td>$O$ 2$p_x$</td>
<td>+0.028</td>
<td>+0.657</td>
<td>0.000</td>
<td>−0.020</td>
</tr>
<tr>
<td>$Ni$ 3$d_{x^2}$</td>
<td>+0.000</td>
<td>+0.000</td>
<td>0.997</td>
<td>+0.000</td>
</tr>
<tr>
<td>$Ni$ 3$d_{y^2}$</td>
<td>−0.138</td>
<td>−0.015</td>
<td>+0.000</td>
<td>−0.929</td>
</tr>
<tr>
<td>$Ni$ 4$s$</td>
<td>−0.189</td>
<td>−0.037</td>
<td>+0.000</td>
<td>+0.086</td>
</tr>
</tbody>
</table>

Within this linear model, we can understand the effect of the nickel atom as (i) the influence on the Auger transitions corresponding to the states of isolated $N_2O$ and (ii) the introduction of new transitions (mainly due to the backbonding). For the first part, a partial Auger spectra was calculated based on the one-center approximation (Fig. 4). Here we look at the same final states of single $N_2O$, and we report the spectra calculated for $N_2O$ and for $N_2O+Ni$. The assignments of the final states respect the labels of single $N_2O$ molecule, also shown in Table 2. It is worth pointing out that the lowest-lying $3\Sigma^-$ state has no intensities in the one-center approximation. The reason is that the Coulomb and exchange integrals cancel each other out in this case. For all three core-holes we observe three main bands in this energy interval, the first corresponding to the $1\Delta$ plus $1\Sigma^+$ states, and the second and third related to the $1\Pi$ and $1\Sigma^-$ states. The triplet $\Pi$ states have lower intensities as discussed above. Although it can be considered as a marginal effect, the presence of the nickel atom change the relative intensity of these transitions as a response to the changes on the MO coefficients, when compared to the isolated $N_2O$ (Tables 5 and 6).

Table 6
$N_2O$+Ni MO coefficients for the valence $\pi$ orbitals. The $\pi$ MOs are $s$/$y$-doubly degenerate ($h_\pi/h_\Sigma$).

<table>
<thead>
<tr>
<th>sym. adpat. basis</th>
<th>$N_2O$ 2$x$</th>
<th>$N_2O$ 1$x$</th>
<th>$Ni$ 3$d_{x^2}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$N_2$ 2$p_x$</td>
<td>+0.628</td>
<td>+0.386</td>
<td>+0.011</td>
</tr>
<tr>
<td>$N_2$ 2$p_y$</td>
<td>+0.034</td>
<td>+0.609</td>
<td>+0.002</td>
</tr>
<tr>
<td>$O$ 2$p_x$</td>
<td>−0.706</td>
<td>+0.412</td>
<td>+0.005</td>
</tr>
<tr>
<td>$Ni$ 3$d_{x^2}$</td>
<td>−0.044</td>
<td>−0.030</td>
<td>−0.995</td>
</tr>
</tbody>
</table>
noticed that the band denoted by the blue line is roughly centered between $-5$ eV and $-8$ eV, which means 7 to 10 eV below the most intense $1\overline{4}$ peak and agrees well with the experimental $N_2$ spectrum. In Tables 5 and 6 we show the coefficients of the Ni 3d/atomic orbitals in the outer valence MOs. The Ni 3d$_{5/2}$ orbital in fact interacts with the 2$s$ orbital of the $N_2$. That led us to conclude that the extra peak is built up principally of transitions involving Ni 3d$_{5/2}$ plus $N_2$ 2$s^{-1}$. Furthermore, we can also use the same argument to explain why this structure cannot be clearly observed in the $N_2$ spectrum.

3.3.3. Adsorption geometry

To examine the monolayer electronic structure we have made ab initio molecular dynamics simulations (AIMD) of the monolayer and density of state calculations of isolated and adsorbed $N_2O$. The equilibrium geometry at 0 K revealed a $3 \times 3$ overlayer structure ($5 N_2O$ molecules per unit cell) with $N_2$ closest to the surface on-top of the Ni atoms as seen in Fig. 6. The molecules demonstrate nearly the same preferable orientation as in the case of a single molecule on-top of Ni (discussed below) with some small bending. Thus reproduce the single molecule on Ni(111) by other authors [57]. In the adsorbed monolayer, the HOMO (which is contributed only by the $N_2O$ atoms bonded with the surface) is located between $-5.8$ and $-3.8$ eV and LUMO (contributed by both atoms) is between 1 and 3 eV.

To reveal the geometry of $N_2O$ monolayer on the Ni(111) surface at finite temperature, we employed ab-initio molecular dynamics simulations (Fig. 7) at 60 and 300 K within NVT thermostat and a time step of 1 fs. It was found that at 60 K, part of the molecules are desorbed from the surface, while some of them change their positions from top to hcp/fcc and demonstrate some bending compared to the molecules adsorbed in the on-top position. At 300 K, we found that the number of desorbed molecules increased while the diversity in the geometries of adsorbed molecules became greater. For example, some molecules are adsorbed in a “lying down” geometry by binding both O and N atoms with Ni(111).

3.3.4. Density of states analysis

To investigate the density of states (DOS) in different bonding configurations we compare the isolated molecule to those adsorbed on-top and in a lying-down configuration. Fig. 8 from top to bottom demonstrates DOS of isolated $N_2O$, $N_2O$ adsorbed in top position (lowest in energy without taking into account ZPE, zero-point energy, and entropy factors), and in lying down position with $N_2O$ adsorbed by O (top) and N (hcp) which is the second in energy, but only predicted to be 0.05 eV higher compared to the top standing $N_2O$. We thus have two bonding situations that give very different DOS and therefore Auger spectra. Here we clearly notice states corresponding to the two uppermost occupied orbitals - $1\overline{1}s$, $7\sigma$, $2e^\prime$ - while above the $2e^\prime$ there is a HOMO-LUMO gap to the unoccupied $3\sigma$ orbital. Going upwards in energy we see a gap to the higher lying orbitals, in a region represented by the breakdown effects according to Table 2. This plot thus represents a one-particle interpretation of the ionization spectrum of $N_2O$. The energy levels and localization, e.g. with the $N_2$ localization of the two uppermost states, are in accordance with the Hartree–Fock (HF) analysis present in Table 5. The up and down plots represent spin-up and spin-down DOS. They are indeed equal for free $N_2O$ as expected for a closed shell molecule. That result is also comparable well with the Hartree–Fock results for free $N_2O$, and is not surprising as Koopmans theorem (HF) and Janaks theorem (DFT) both equalize orbital energies with ionization potentials to a large degree of accuracy. For isolated molecules, it is thus quite clear that HOMO is localized on $N_2O$, while LUMO is localized on both N atoms. Adsorption leads to interaction with the Ni(111) surface and as a result prominent shift of states to lower energies caused by electron transfer from Ni to the molecule. At the top position, HOMO is at $-4.4$ eV and LUMO is broadened between 1.5 and 4.4 eV due to proximity-induced effects. One can also see small but non-zero spin polarization of $N_2O$ states.

The electronic structure of $N_2O$ adsorbed in the complex configuration reveals a significant broadening of orbitals and spin polarization resulting from stronger interaction with the Ni(111) surface. This also can be confirmed by charge transfer analyzed by the Bader approach. The isolated $N_2O$ reveals that the O atom gains 0.32 e from the $N_2$ part. Adsorption in the top position keeps the charge of O the same as in the isolated molecule while N gain 0.2 e transferred from $N_2O$. Complex adsorption leads to significant charge transfer from the surface to both N and O parts. For instance, O gains additionally 0.24 e as compared to isolated $N_2O$, while 0.53 e is transferred from the metal surface to N. This results in significant induced spin polarization of the molecule as demonstrated in DOS.
The DOS results indicate a shifted N₂O structure but with an additional broad structure appearing a few eV above the Fermi surface, replacing the LUMO level of free N₂O. This structure represents the surface to N₂O (3d- π*) backbonding situation with the broad feature related to a splitting of the 3d band of Nickel by the N₂O bonding. Here we see that the free N₂O DOS results are very much compromised for the HOMO and LUMO levels, showing neither the preserved N₂O structure nor the 3d- π* backbonding indicated by the standing up N₂O case in the middle panel of Fig. 8a.

Finally, in Fig. 8b we show the case of standing N₂O on the surface for core ionized (N₆ or N₇) N₂O density of states, where the core hole potential is represented by the equivalent core (Z+1) approximation. As the core ionized state (CS) is the initial state of the Auger transition it can be relevant to compare it with the ground state results — here one sees that N₂O orbital structure is only minorly varied but that the 3d- π* backbonding band complex is significantly stabilized in energy.

3.4. Blyholder or not

So considering the standing position it seems that the excess intensity in the high kinetic energy part of the Auger spectrum of the Ni₇ atom (in comparison with the Ni₆ atom) is due to charge transfer from metal to the π* of the Ni₇ atom. This is similar to the Blyholder back–donation model for CO / N₂ on transition metals, thus that in the ground state, there is back donation from metal 3d to the (otherwise empty) LUMO on the N atoms, something that strengthens the bonding of the adsorbate to the surface [21,58]. This back-donation has also been suggested by Kiss et al. for the N₂O on Pt(111) system [59] and from DFT analysis on N₂O on Fe-doped graphene [60].

The kinetic energy position of this extra intensity has been observed of N₂O in the gas phase both as a shake-up from core ionization [61] and resonant Auger [62]. In Fig. 9 we present backdonation into the LUMO from the nickel surface and the donation from the 7σ lone pair orbital of the N₇. This explains the chemisorbing nature of the N₂O/Ni(111) interaction and the extra intensity seen in APECS at high kinetic energy for the thin coverages that is not present in the multilayer film.

As indicated by the density of states analysis there is a broad extra structure around the Fermi level that can be interpreted as due to a N₂O (3d- π*) backbonding situation. It is also possible to gain insight into the backdonation in the ground electronic structure from the idealized model considered above comprising a system composed of one molecule of N₂O plus a single Ni atom, all disposed in a linear structure, see further information in the computational section. Direct analysis of the Mulliken charges of the ground state indicates a slight total charge transfer from the Ni atom to the N₂O molecule. In fact, the total atomic charge on Ni is −0.05, and the charges of the N₆, N₇, and O atoms are, respectively, −0.27, +0.78 and −0.47. By large this agrees well with the analysis of the DOS in Fig. 8. When considering the electronic charge occupying only the 3d and 4s atomic orbitals of Ni we obtained −9.59, against the expected −10 for the isolated Ni. On top of that, the occupation number of the orbitals also presents valuable information on the backdonation. First, the sum of the MO coefficients squared associated with Ni atomic orbitals of the orbital labeled as 7σ - occupation number 1.99 - is 0.07, which is a way of quantifying the hybridization of the 7σ orbital. Besides, the hybridization between the N₂O LUMO orbital (3ε) and the 3d4σ atomic orbitals of Ni produces a 0.13 population on the 3ε orbital (second and third lines of Fig. 3), thus supporting the notion of a 3d- π* backbonding.
4. Conclusions

APECS represents an experimental method where backdonation in surface–adsorbate bond may be directly measured. Additionally since we are able to distinguish the Auger spectra from chemically nonequivalent atoms, we can in cases when those are well separated determine which orbital at which atom receives the backdonated electron density.

We have compared multilayer, bilayer and monolayer coverage of N$_2$O on a Ni(111) surface where the multilayer film in many regards is similar to a kinetic energy shifted and broadened gas-phase spectra and we can see that the monolayer contains features not found in the gas-phase. For the N coincidences this new intensity feature arises from an interaction with the nickel substrate whereas in the O case the spectrum is not dependent on coverage. The observed behavior is explained with calculations that demonstrate that the N$_2$O is standing up with the N$_2$-closest to the surface with a surface bond built up from donation and backdonation of electron density involving the Ni surface and the N$_2$-atom.

The APECS methodology outlined here suggests that the method should be applied to problems in catalysis where small molecules interact with a substrate to make a detailed assessment of how they interact. In the present case for instance, we could identify the decomposition products and differentiate them from intact species. The present work also illustrates the insight into the electronic structure and dynamics of the surface adsorption gained by combining APECS and theoretical analysis — solid state and quantum ab initio.

CRediT authorship contribution statement

Fredrik O.L. Johansson: Writing – review & editing, Writing – original draft, Visualization, Methodology, Investigation, Formal analysis, Conceptualization. Lucas M. Cornetta: Writing – review & editing, Writing – original draft, Visualization, Investigation, Formal analysis. Elin Berggren: Writing – review & editing, Investigation. Artem Kuklin: Writing – original draft, Visualization, Investigation, Formal analysis. Yi-Chen Weng: Writing – review & editing, Investigation. Swarnshikha Sinha: Writing – review & editing, Investigation. Danilo Kühn: Writing – review & editing, Methodology, Investigation. Alexander Föhlisch: Writing – review & editing, Methodology, Investigation. Hans Ågren: Writing – review & editing, Writing – original draft, Methodology, Investigation, Formal analysis. Andreas Lindblad: Writing – review & editing, Writing – original draft, Supervision, Investigation, Formal analysis, Conceptualization.

Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

Acknowledgments

F.O.L.J acknowledges support from the Swedish Research Council (Grant 2020-06409). L.M.C acknowledges the Brazilian funding agency FAPESP, under processes nr. 2020/04822-9 and 2021/06527-7. A.K. acknowledges Olle Engkvist Byggmästarskapet Foundation for the support under contract no. 212-0178. We thank the Helmholtz-Zentrum Berlin für Materialien and Energie for the allocation of synchrotron radiation beamtime. The authors also acknowledge the Swedish National Infrastructure for Computing (NAISS 2023-3-40 and NAISS 2024-5-73) at the National Supercomputer Centre of Linköping University (Sweden) partially funded by the Swedish Research Council through grant agreements no. 2022-06725 and no. 2018-05973.

References


