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alumina-supported palladium catalyst: an infrared spectroscopic study†

The adsorption of nitrobenzene over an

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As part of an on-going programme of development of an aniline synthesis catalyst suitable for operation at elevated temperatures, the geometry of the adsorption complex for nitrobenzene on a 5 wt% Pd/ Al₂O₃ catalyst is investigated by infrared (IR) spectroscopy. Via an appreciation of the reduced site symmetry resulting from adsorption, application of the metal surface selection rule, and observation of in-plane modes only, the adsorption complex (Pd-nitrobenzene) at 28 °C is assigned as occurring vertically or tilted with respect to the metal surface, adopting $C_s\sigma_v(yz)$ symmetry. Moreover, adsorption occurs via a single Pd-O bond. Single molecule DFT calculations and simulated IR spectra assist vibrational assignments but indicate a parallel adsorption geometry to be energetically favourable. The contradiction between calculated and observed structures is attributed to the DFT calculations corresponding to an isolated molecule adsorption complex, while IR spectra relate to multi molecule adsorption that is encountered during sustained catalytic turnover. Residual hydrogen from the catalyst reduction stage leads to aniline formation on the Pd surface at low nitrobenzene coverages but, on increasing nitrobenzene exposure, the aniline is forced on to the alumina support. A reaction scheme is proposed whereby the nitrobenzene adsorption geometry is inherently linked to the high aniline selectivity observed for Pd/Al₂O₃ catalysts.

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1. Introduction

The heterogeneously catalysed hydrogenation of nitrobenzene is a commonly employed route to produce aniline, which in

turn is a key component in the formation of methylene diphenyl diisocyanate (MDI) utilised throughout the polyurethane industry.1 Supported Pd catalysts are regularly described as achieving high aniline selectivity from nitrobenzene hydrogenation²⁻⁹ owing to the high activity and propensity of Pd to reduce functional groups in proximity to aromatic structures. The current investigation aims to investigate nitrobenzene adsorption over a reference Pd/Al₂O₃ catalyst via infrared (IR) spectroscopy to better understand the origins of this favourable aniline selectivity.

Nitrobenzene and aniline both possess a phenyl ring and a functional group moiety, permitting the possibility of various interactions between the molecules and a catalyst surface. It follows that each of these potential interactions may result in different preferential orientations of nitrobenzene and aniline on metal surfaces - a parameter which may be directly related to catalytic outputs. A DFT study by Chen et al. highlighted the possible relationship between observed chemical pathways and adsorbate geometry, where 100% product selectivity during hydrogenation of p-chloronitrobenzene using platinum nanoparticles encapsulated into zeolite Y as a catalyst was reported. 10 This perfect selectivity was attributed to an end-on orientation of p-chloronitrobenzene enforced by steric constraints of the zeolite pore size.

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[†] Electronic supplementary information (ESI) available: Fig. S1: Diagram of nitrobenzene with respect to a specified xyz-coordinate (blue = nitrogen; red = oxygen), with the z axis defining the principal component axis of the molecule; Fig. S2: Form of the vibrational modes for a mono-substituted benzene in C_{2v} symmetry; Table S1: Approximate wavenumber ranges for the Mi vibrations; Table S2: Vibrational transition energies, infrared intensities and assignments for nitrobenzene in the solid state as calculated by periodic-DFT; Fig. S3: Single beam DRIFTS measurement of activated GU-1; Table S3: Experimental and calculated geometry of nitrobenzene; Fig. S4: Wide scan DRIFTS spectra depicting increasing nitrobenzene exposure; Fig. S5: Visualisation of the $\nu_{\rm S}({
m NO_2})$ and $\delta_{\rm oop}({
m CH})$ nitrobenzene modes during a (a) and (b) parallel orientation and (c) and (d) a tilted adsorption orientation over a metal surface with respect to the molecules xz-plane; Fig. S6: Diagram visualising nitrobenzene adsorption to Pd(111) in a tilted orientation with respect to the yz-plane depicting (a). O positioned in a 2fold hollow site with H in registry with on-top and (b). the H positioned in a 2-fold hollow site with O in registry with the on-top site. See DOI: https://doi.org/ 10.1039/d3cp03028h

The geometry of adsorbates may be inferred from IR spectra with reference to the metal surface selection rule (MSSR):¹¹ molecular vibrations that yield a dipole parallel to a metal surface are unobservable, whilst vibrations yielding a nonparallel dipole to a metal surface are observable. 12,13

The IR spectrum of nitrobenzene (C_{2y} symmetry) has been extensively described, 14-17 with two high intensity peaks centred around 1530 and 1350 cm⁻¹ assigned to the in-plane asymmetric (B2 symmetry) and symmetric (A1 symmetry) NO2 stretching modes of nitrobenzene ($\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$), respectively, being the most prominent features. In addition, several out-of-plane B₁ symmetry modes are observed, including a moderate intensity band at ca. 705 cm⁻¹ that corresponds to the out-of-plane NO_2 deformation mode $(\delta_{oop}(NO_2))$ and smaller features centred at ca. 935 and 793 cm⁻¹, both arising from out-of-plane C-H deformation modes $(\delta_{oop}(CH))$. ^{14,15}

Surface science studies are insightful. Richardson and coworkers used reflection-absorption infrared spectroscopy (RAIRS) to investigate the adsorption of nitrobenzene on Cu(110) over a range of temperatures. 18 For temperatures <223 K multilayer adsorption was observed, with no preferential orientation of the nitrobenzene molecule relative to the metal surface. Adsorption at 300 K led to dissociation of the nitrobenzene. A subsequent vibrational electron energy loss spectroscopy study established the retained moieties to be phenyl species and atomic oxygen. 18 Koel and co-workers used RAIRS to investigate the adsorption of nitrobenzene on Au(111) at low temperatures. 19 For monolayer coverages, only two bands were observed: an out-of-plane CH deformation at 793 cm⁻¹ and a NO₂ deformation at 710 cm⁻¹. Assuming C_{2v} symmetry, these modes exhibit B_1 symmetry, indicating the nitrobenzene molecule to be aligned parallel to the surface. On increasing coverage, modes with A_1 and B_2 symmetry were observed, indicative of a change of molecular orientation. As neither in-plane nor out-of-plane modes are screened completely, the molecule was assumed to adopt a tilted geometry. Temperature-programmed desorption measurements showed the monolayer to exhibit a desorption maximum at 290 K.¹⁹ Bridging the gap between surface science and heterogeneous catalysis, Corma and co-workers have examined the adsorption of nitroaromatics on Au single crystal surfaces and supported Au particles.²⁰ For the case of nitrostyrene adsorption, the Au/support interface was reported to be the active site of Au/TiO₂, which exhibits high chemo-selectivity via preferential activation of the nitro group.

Density functional theory (DFT) is widely used to investigate adsorbate orientation, and has been utilised to investigate the adsorption geometry of nitrobenzene on varying systems. A recent study by Hajiahmadi et al. considered the likelihood of six different nitrobenzene adsorption configurations, three vertical and three parallel, over Pd(111).²¹ They found a parallel nitrobenzene adsorption geometry with N atoms located on top of Pd atoms to be most favourable for adsorption to Pd(111). Additionally, Boronat et al. conducted a DFT investigation into nitroarene adsorption primarily over non-noble metals, with the inclusion of Pd(111) for comparison, and report parallel

nitrobenzene adsorption over Pd(111) via the aromatic ring with the NO₂ group twisted relative to the aromatic moiety.²² Interestingly, Boronat and co-workers reported strictly perpendicular nitrobenzene adsorption over the non-noble metal Cu(111) and used this specified adsorbate geometry to propose a possible non-noble metal catalyst candidate for high selectivity aniline synthesis. Contrarily for Pd studies, a DFT investigation of nitrobenzene adsorption on bimetallic Pd₃/Pt(111) found vertical nitrobenzene adsorption over Pd crystallites to be preferable to parallel adsorption.²³ Comparison of these opposing outcomes indicates that nitrobenzene adsorption orientation is substrate dependent. This perspective meshes with a recent spectroscopic and computational study by Millán and co-workers on nitrobenzene activation on non-noble metals-based mono- and bimetallic catalysts.24

This article concentrates on the adsorption of nitrobenzene over a 5 wt% Pd/Al₂O₃ catalyst (GU-1) that is a model for low metal loading Pd/Al2O3 aniline synthesis catalysts intended for operation at elevated temperatures.8 The performance of this catalyst in vapour phase nitrobenzene hydrogenation has previously been described,8 as has the morphology of the Pd crystallites.²⁵ The approach adopted is as follows. A complete vibrational analysis of nitrobenzene is undertaken with reference to measured solid and liquid phase spectra (inelastic neutron scattering, infrared spectroscopy, and Raman scattering). The infrared spectrum of nitrobenzene adsorbed on GU-1 is compared to simulated spectra corresponding to DFT calculations for the adsorption of a single unit of nitrobenzene on a Pd(111) surface, providing information on the geometry of the adsorption complex. The unidentate nature of the interaction between the nitro group and the Pd surface is consistent with established reaction schemes. A small quantity of aniline is formed at the Pd surface at low nitrobenzene exposures, but it is forced on to the alumina support on increasing nitrobenzene exposure. A reaction scheme is proposed whereby the nitrobenzene adsorption geometry is inherently linked to the high aniline selectivity observed for Pd/Al₂O₃ catalysts.

2. Experimental

A 5 wt% Pd/γ-Al₂O₃ catalyst was obtained from Alfa Aesar (ref.: 11713). The catalyst has been comprehensively characterised²⁵ and is referred to here, and in previous communications from this group, 8,25 as GU-1.

2.1 Computational methods

The Gaussian09²⁶ and GaussView software packages were used to derive the vibrational spectrum of gas phase nitrobenzene. Density functional theory (DFT) was utilised with the B3LYP functional and the 6-311G++(3df.2p) basis set. A DFT scaling factor of 0.967 was used on calculated wavenumbers.²⁷ Solid state calculations used the plane wave pseudopotential method as implemented in CASTEP.28 The generalized gradient approximation (GGA) was used with the PBE functional.²⁹ The Tkatchenko and Scheffler-dispersion correction was used to

account for the long-range interactions. 30 For nitrobenzene, the initial structure used the experimental crystal structure that was determined at 103 K.31 The plane wave cut-off was 1020 eV and the Brillouin zone sampling used a $12 \times 4 \times 4$ (48 k-points) Monkhorst-Pack grid. The adsorption of nitrobenzene on Pd(111) was investigated using a three-layer slab with (111) truncation. A 15 Å vacuum gap was used. For both the parallel and vertical orientations, the plane wave cut-off was 1020 eV and the Brillouin zone sampling used a 11 \times 11 \times 1 (36 k-points) Monkhorst-Pack grid. After geometry optimisation, the vibrational transition energies were calculated with density functional perturbation theory.³² To reduce the time required for the calculation, only the internal modes of the nitrobenzene were calculated. The computational outputs include the amplitude of motion of each atom in the vibrational mode, this enables mode visualisation and enables the inelastic neutron scattering (INS) spectrum to be calculated using the program AbINS.33

In Section 3.2.3 Gaussian calculations are used to simulate the IR spectra for nitrobenzene adopting parallel and vertical adsorption geometries relative to the Pd surface. Here, with consideration of point group theory and the MSSR, simulated spectra for parallel $(C_s\sigma_v(xz))$ and mono-dentate vertical $(C_s\sigma_v(yz))$ nitrobenzene adsorption over a metal surface were produced by adjusting the intensity of non-accessible modes to 0 a.u.

2.2 Vibrational spectroscopy

Infrared spectra (256 scans, 4 cm⁻¹ resolution, eight-times zerofilling) of nitrobenzene were measured with a Bruker Vertex 70 Fourier transform infrared (FT-IR) spectrometer. The liquid at room temperature was measured by attenuated total internal reflection (ATR) using a Bruker Diamond ATR accessory. Variable temperature (210-296 K) spectra were recorded with a low temperature SpecAc Golden Gate ATR accessory. Raman spectra were recorded with a Bruker Fourier transform Raman spectrometer (64 scans, 4 cm⁻¹ resolution, eight-times zerofilling, 500 mW laser power at 1064 nm). The sample was held in a quartz cuvette and the liquid measured at room temperature and the solid after immersion of the cuvette in liquid nitrogen. INS spectra of the solid were recorded at 10 K from the sample held in a flat-plate indium wire sealed aluminium can using the broad band, high resolution spectrometer TOSCA34 at ISIS.35

2.3 DRIFTS measurements: nitrobenzene adsorption and temperature-programmed IR spectroscopy

In situ DRIFTS measurements were performed with a Bruker Vertex 70 FT-IR spectrometer fitted with a MCT detector. GU-1 was supplied as a powder and was used directly. Ca. 50 mg of GU-1 was placed into a Harrick Praying Mantis DRIFTS reaction chamber. A Harrick ATC heater permitted control of cell temperature. Activation of Pd crystallites occurred in situ using a flow of helium (35 mL min⁻¹, BOC gases, 99.9%) and hydrogen (15 mL min⁻¹, BOC gases, 99.8%) while heating to 110 °C. This temperature was held for 30 minutes. The temperature was then increased to 200 °C for one hour, with the hydrogen flow stopped after 30 minutes and the sample allowed to return to ambient temperature in flowing He. A He purge of 18 hours was utilised to minimise the levels of retained hydrogen within GU-1 from the reduction process, which would otherwise result in nitrobenzene hydrogenation to aniline. Post-purge, a background spectrum was collected at 28 °C.

Nitrobenzene was supplied via a nitrobenzene bubbler system which supplied $61.2 \, \mu mol_{(NB)} \, min^{-1} \, g_{(cat)}^{-1}$ to the reaction chamber in the vapour phase using He as a carrier gas. A 5-minute pulse duration was selected, so that 1 pulse corresponded to a nitrobenzene exposure of 0.31 mmol_(NB) $g_{(cat)}^{-1}$. After each pulse, the cell was purged with He for 10 minutes to evacuate unbound nitrobenzene before spectral acquisition.

For desorption experiments, the catalyst was heated in situ under a flow of He and maintained at the designated temperature for 30 minutes before cooling to 28 °C for spectral acquisition. This process was repeated for 50, 100, 120, 140, 160, 180 and 200 °C. All spectra were recorded at 28 °C for 520 scans at 4 cm⁻¹ resolution. Spectra are presented as difference spectra, where the spectrum of the activated catalyst has been subtracted from that of a nitrobenzene dosed spectrum. No additional spectrum treatment was performed. A minor aniline impurity was present in the feed stream to the IR cell, which could not be eliminated. Though vexatious, its minimal presence in the IR spectra can be accounted for and will be suitably considered in the following sections.

Results

Molecular symmetry and DFT: assigning key nitrobenzene modes

In the gas and liquid phases nitrobenzene exhibits $C_{2\nu}$ symmetry,³⁶ see Fig. S1 in the ESI.† In the solid state, there are four molecules in the primitive cell, each on a site with C_1 symmetry, however, the C_{2v} symmetry is largely retained.³¹ Reference to the C_{2v} character table confirms modes possessing A_1 , B_1 and B_2 symmetry are infrared active.³⁷

Fig. 1 shows the infrared and Raman spectra in the liquid phase, the infrared, Raman, and INS spectra in the solid state, together with the calculated infrared and INS spectra for the solid state. Table 1 presents experimental wavenumber values and DFT-derived frequencies for nitrobenzene in the liquid phase. The vibrational spectra of mono-substituted benzenes have been extensively studied over the decades and the various schemes are lucidly discussed by Gardner and Wright.³⁸ Based on DFT calculations, they proposed some revision to the assignments and designated the modes as M1-M30. The form of the modes and their usual ranges are reproduced from ref. 38 in the ESI† (Fig. S2 and Table S1). We have adopted their assignment scheme and the vibrational assignments and associated symmetry designations are also listed in Table 1.

In the solid state, the presence of four molecules in the primitive cell results in factor group splitting. However, periodic-DFT calculations of the complete unit cell show that

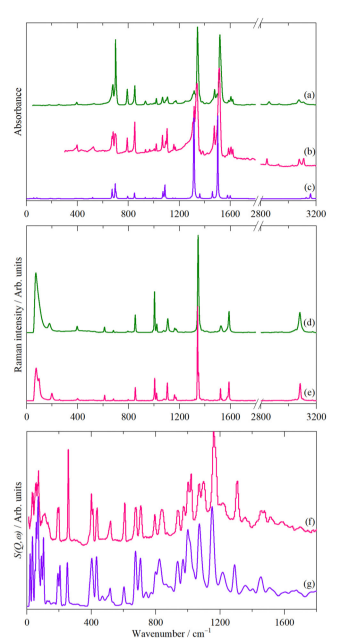


Fig. 1 Experimental and calculated vibrational spectra of nitrobenzene. Top panel: infrared spectra of (a) liquid nitrobenzene at room temperature, (b) solid nitrobenzene at 210 K, (c) calculated spectrum for solid nitrobenzene. Middle panel: Raman spectra of (d) liquid nitrobenzene at room temperature, (e) solid nitrobenzene. Bottom panel: INS spectrum (f) solid nitrobenzene at 10 K and (g) calculated spectrum for solid nitrobenzene.

for most of the internal modes this is 10 cm $^{-1}$ or less, although there are exceptions (see Table S2, ESI†). These are the Ph–NO $_2$ torsion and some of the out-of-plane C–H bending modes. As can be seen from Fig. 1, the calculated infrared and INS spectra are in good agreement with the experimental data (cf. Fig. 1b, c and f, g).

As mentioned in the Introduction, owing to their high intensity, 14,21 the B_2 symmetry $\nu_{\rm AS}({\rm NO_2})$ and A_1 symmetry $\nu_{\rm S}({\rm NO_2})$ modes at 1531 and 1348 cm⁻¹ respectively, represent key nitrobenzene modes for orientational diagnostics. The IR spectrum of activated GU-1 is presented in Fig. S3 (ESI†) and

Table 1 Comparison of experimental and DFT-derived wavenumbers (cm⁻¹) of nitrobenzene. DFT calculations performed utilising the B3LYP method and 6-311G++(3df.2p) basis set. Vibrational assignments are based on DFT calculations

Mode	Sym.										
1	A_2	52 ^c	57	τ(Ph-NO ₂)							
2	$\mathbf{B_1}$	179	164	$\delta_{\rm oop}({\rm Ph\text{-}NO}_2)$	M20	oop					
3	B_2	256	249	$\delta_{ m ip}({ m Ph-NO}_2)$	M30	ip					
4	A_1	396	383	Ring deformation	M11	ip					
5	A_2	423	404	Ring deformation	M14	oop					
6	$\mathbf{B_1}$	$454^{\ c}$	432	Ring deformation	M19	oop					
7	B_2	529	508	NO ₂ rock		ip					
8	\mathbf{B}_2	608	607	$\delta_{ m ip}({ m CCC})$	M29	ip					
9	B_1	677	663	$\delta_{\rm oop}({ m CH})$	M18	oop					
10	A_1	680	671	$\delta_{\rm ip}(\rm ring) + \delta_{\rm ip}(\rm NO_2) + \nu(\rm CN)$	$\mathcal{M}10$	ip					
11	$\mathbf{B_1}$	700	704	$NO_2 \text{ wag} + \delta_{\text{oop}}(CH)$	$\mathcal{M}17^d$	oop					
12^e	$\mathbf{B_1}$	793	791	$NO_2 \text{ wag} + \delta_{\text{oop}}(CH)$	$M17^d$	oop					
13	\mathbf{A}_2	837	831	$\delta_{\rm oop}({ m CH})$	M13	oop					
14	A_1	851	839	NO ₂ scissors		ip					
15^e	B_1	935	936	$\delta_{\rm oop}({ m CH})$	M16	oop					
16	A_2	975	972	$\delta_{\rm oop}({ m CH})$	M12	oop					
17	A_1	977	977	$\delta_{ m ip}({ m ring})$	M9	ip					
18	B_1	991	988	$\delta_{\rm oop}({ m CH})$	M15	oop					
19	A_1	1003	1006	$\delta_{ m ip}\!(m ring)$	M8	ip					
20	\mathbf{B}_2	1069	1063	$\delta_{\mathrm{ip}}(\mathrm{CH})$	M28	ip					
21	A_1	1094	1078	$\nu(\text{Ph-NO}_2)$	$\mathcal{M}6$	ip					
22	\mathbf{B}_2		1145	$\delta_{\mathrm{ip}}(\mathrm{CH})$	M27	ip					
23	A_1	1162	1157	$\delta_{\mathrm{ip}}(\mathrm{CH})$	M7	ip					
24	\mathbf{B}_2	1248	1294	$\delta_{ m ip}({ m CH})$	M26	ip					
25	B_2	1308	1305	$\nu(CC)$	M25	ip					
26^e	A_1	1344	1348	$\nu_{ m S}({ m NO}_2)$		ip					
27	B_2	1415	1439	$\nu(\text{CC})$	M24	ip					
28	A_1	1478	1462	$\nu(\text{CC})$	M5	ip					
29^e	B_2	1528	1531	$\nu_{AS}(NO_2)$		ip					
30	A_1	1588	1572	$\nu(CC)$	$\mathcal{M}4$	ip					
31	B_2	1606	1596	$\nu(CC)$	M23	ip					
32	A_1	3029	3066	ν (CH)	M3	ip					
33	B_2	3077	3079	ν (CH)	M22	ip					
34	A_1		3086	ν (CH)	M2	ip					
35	B_2	3107	3116	ν (CH)	M21	ip					
36	A_1	3107	3117	ν (CH)	M1	ip					

^a Experimental wavenumbers for nitrobenzene are from Fig. 1a, or otherwise indicated. ^b Scaling factor for DFT frequencies was 0.967. ⁴⁰ Ref. 41. ^d This mode appears twice because it is mixed with the NO₂ wag. ^e Key diagnostic modes for orientational assignments in the present investigation.

exhibits a spectral cut-off due to strong Al–O phonon modes of the alumina support³⁹ at about 1100 cm⁻¹. With respect to Table 1, in the presence of the catalyst, several B_1 symmetry diagnostic nitrobenzene modes are inaccessible; for example, the $\delta_{\rm oop}({\rm NO_2})$ feature present at 704 cm⁻¹. However, rather fortuitously, GU-1's γ -alumina displays a narrow transmission window in the 970–745 cm⁻¹ range (Fig. S3, ESI†), which enables access to two $\delta_{\rm oop}({\rm CH})$ B_1 symmetry modes at 791 and 936 cm⁻¹, respectively. Therefore, on this basis, vibrational transitions 12, 15, 26 and 29 (Table 1) were selected as key modes for orientational diagnostics.

3.2 DRIFTS: adsorbate geometry

3.2.1 Nitrobenzene adsorption on KBr at 28 °C. Fig. 2b presents the spectrum of nitrobenzene adsorbed on KBr

at 28 °C. The spectrum is indicative of physisorbed nitrobenzene with no specific orientation, and yields all four key nitrobenzene modes (Table 1): the B_2 symmetry $\nu_{AS}(NO_2)$ mode at 1529 cm⁻¹, the A_1 symmetry $\nu_{\rm S}({\rm NO_2})$ mode at 1347 cm⁻¹ and the two B_1 symmetry $\delta_{\text{oop}}(\text{CH})$ modes at 935 and 793 cm⁻¹. ^{14,15} This measurement confirmed the capability of the DRIFTS set-up utilised to observe all A_1 , B_1 and B_2 symmetry key diagnostic modes.

3.2.2 Saturation nitrobenzene coverage on GU-1 at 28 °C. Fig. 2a presents the IR spectrum for a saturation coverage of nitrobenzene over GU-1 at 28 °C. As observed for KBr (Fig. 2b), the A_1 and B_2 symmetry modes at 1529 and 1347 cm⁻¹ are present but, intriguingly, the B_1 symmetry modes at 935 and 793 cm⁻¹ were not observed.

The observation of A_1 and B_2 key modes and absence of B_1 modes in spectra associated with nitrobenzene adsorption to the catalyst provides information on the form of the adsorption complex. However, prior to assignment of molecular geometry utilising the MSSR, it is essential to consider the impact of binding to a metal surface on the molecular symmetry of nitrobenzene; thus, reference to the C_{2v} correlation table is necessary. C_{2v} symmetry correlates to four lower symmetry point groups:³⁷ C_1 , C_2 , $C_8\sigma_v(xz)$ and $C_8\sigma_v(yz)$, where $C_8\sigma_v(xz)$ and $C_s\sigma_v(yz)$ represent a molecule possessing C_s symmetry with a single mirror plane in the molecule's xz and yz-planes, respectively.

Table 2 considers the accessibility of each key nitrobenzene diagnostic mode during nitrobenzene adsorption to a metal surface in a vertical or parallel orientation for nitrobenzene exhibiting C_{2v} , C_2 , $C_s\sigma_v(xz)$ and $C_s\sigma_v(yz)$ symmetry. With reference to this table, at a saturation coverage of nitrobenzene, Fig. 2a indicates that nitrobenzene exhibits $C_s \sigma_v(yz)$ symmetry

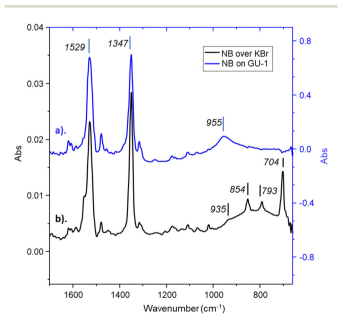


Fig. 2 DRIFTS spectra of nitrobenzene adsorption over (a) GU-1 (blue) and (b) KBr (black) at 28 °C. Wavenumber range: 1700-660 cm⁻¹. Note the differences in y-axes scales for both spectra.

Table 2 Assignment, wavenumber values and symmetric representations for key nitrobenzene diagnostic modes

		Symmetric Representations											
	Expt.a		C_{2v}			C_2			$C_{\rm s}\sigma_{ m v}$	(xz)		$C_{\rm s}\sigma_{ m v}$	(yz)
Assign.	(cm^{-1})		\perp^{b}	$\ ^c$		\perp^{I}) c		$\perp^{\it b}$	\parallel^c		$\perp^{\it b}$	\parallel^c
$\overline{\text{NO}_2 \text{ wag} + \delta_{\text{oop}}(\text{CH})}$	794	B_1	N	Y	В	N	Y	A'	Y	Y	A"	N	d
$\delta_{\rm oop}({ m CH})$	935	B_1	N	Y	В	N	Y	${\bf A}'$	Y	Y	$A^{\prime\prime}$	N	d
$\nu_{\rm S}({ m NO}_2)$	1351	A_1	Y	N	A	Y	N	\mathbf{A}'	Y	N	${\bf A}'$	Y	d
$\nu_{\rm AS}({ m NO}_2)$	1527	\mathbf{B}_2	N	N	В	N	N	$A^{\prime\prime}$	Y	N	\mathbf{A}'	Y	d

^a Experimental wavenumbers for nitrobenzene are from Fig. 1a. ^b Indicates if mode is allowed during vertical nitrobenzene adsorption over a metal surface, as per the MSSR. ¹¹ N = no; Y = yes. ^c Indicates if mode is allowed during parallel nitrobenzene adsorption over a metal surface, as per the MSSR. ¹¹ N = no; Y = yes. ^d Corresponding orientation is not possible for nitrobenzene exhibiting stated symmetric representation.

and is positioned in a vertical orientation over the metal surface; a deduction that contradicts previously introduced DFT investigations. 20,21

For nitrobenzene to exhibit $C_s\sigma_v(yz)$ symmetry adsorption must occur via only one Pd-O bond (Fig. 3(c and d)); as binding to the Pd surface via both oxygens of the nitro group would retain the C_{2v} symmetry of nitrobenzene (Fig. 3(a and b)). For vertical nitrobenzene bidentate adsorption (C_{2v} symmetry) the $\nu_{\rm S}({\rm NO_2})$ and $\nu_{\rm AS}({\rm NO_2})$ modes exhibit A_1 and B_2 symmetry, respectively. For the $A_1 \nu_{\rm S}({\rm NO}_2)$ (Fig. 3b) the resulting dipole

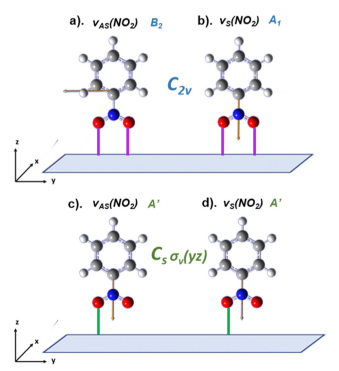


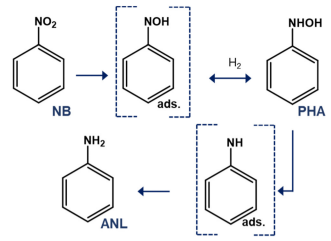
Fig. 3 Visualisation of $\nu_{\rm S}({\rm NO_2})$ and $\nu_{\rm AS}({\rm NO_2})$ nitrobenzene modes during vertical adsorption over a metal surface via (a) and (b) bidentate and (c) and (d) monodentate binding. Orange arrows indicate the dipole derivative unit vector associated with each mode. Grey parallelograms symbolise a nonspecific Pd crystallite. White, grey, blue and red balls represent hydrogen, carbon, nitrogen and oxygen, respectively.

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(Fig. 3, orange arrow) is positioned perpendicularly to the metal surface, thus with consideration of the MSSR, this mode would be observed in IR spectra, 11,13 as is the case during adsorption to GU-1. However, the dipole associated with the B_2 symmetry $\nu_{\rm AS}({\rm NO_2})$ mode (Fig. 3a) is aligned parallel to the metal surface during bidentate vertical adsorption, and resultingly, would not be observable in IR spectra. Thus, a bidentate vertical adsorption of nitrobenzene to the metal surface, and thus retention of C_{2v} symmetry during binding, is not possible as both the A_1 $\nu_{\rm S}({
m NO_2})$ and B_2 $\nu_{\rm AS}({
m NO_2})$ symmetry modes of nitrobenzene were clearly observed during adsorption to GU-1 (Fig. 2a). Conversely, for monodentate vertical nitrobenzene adsorption $(C_S\sigma_v(yz))$ both the $\nu_S(NO_2)$ and $\nu_{AS}(NO_2)$ modes correlate to A' symmetry, and so are both totally symmetric in-plane modes (Fig. 3(c and d)). The resulting dipole for each A' mode during a vertical nitrobenzene adsorption would be positioned perpendicular to the metal surface and would therefore be MSSR allowed. Thus, nitrobenzene adsorbs to the Pd crystallites of GU-1 in a vertical orientation via monodentate binding, and not bidentate binding of the metal group. Moving forward, nitrobenzene modes will be referred to as exhibiting either A' or A'' symmetry as encountered for $C_s\sigma_v(yz)$ symmetry (Table 2).

The known chemistry of metal catalysed nitrobenzene hydrogenation is supportive of the proposed geometric structure. A scenario in which both O atoms of nitrobenzene simultaneously interacted with the metal would result in a rapid and simultaneous one-step hydrogenation of the nitro group with no observable intermediates. However, as reported by Gelder et al., hydrogenation of the nitro moiety occurs in a stepwise fashion, and intermediates such as phenylhydroxylamine (PHA) are accessible during nitrobenzene hydrogenation with Pd (Scheme 1).42 Moreover, this observable intermediate, PHA, exhibits $C_s\sigma_v(yz)$ symmetry, further strengthening the hypothesis of nitrobenzene exhibiting the same symmetry during adsorption to GU-1. Thus, we assert that the assignment of $C_s\sigma_v(yz)$ symmetry and monodentate binding for nitrobenzene adsorption over the catalyst is consistent with the reported chemistry.

The authors acknowledge the totally vertical orientation depicted in Fig. 3(c and d) may be an over-simplification, as it is not possible to entirely differentiate if nitrobenzene exhibiting $C_s\sigma_v(yz)$ symmetry adsorbs vertically, or if the molecule exhibits some degree of tilting in the yz-plane. Fig. 4(a and b) presents visualisation of the in-plane A' symmetry $\nu_{\rm S}({\rm NO_2})$ and out-of-plane A" symmetry $\delta_{\rm oop}({\rm CH})$ modes of nitrobenzene whilst exhibiting $C_s\sigma_v(yz)$ symmetry in a tilted orientation, and highlights the respective perpendicular and parallel alignment of resulting dipoles with respect to the metal surface. The non-parallel orientation of the A' symmetry dipole will permit observation of the $\nu_{\rm S}({\rm NO_2})$ mode in IR spectra (Fig. 4a), while the parallel orientation of the out-of-plane A'' dipole will yield the $\delta_{\text{cop}}(CH)$ mode unobservable (Fig. 4b). Thus, nitrobenzene adsorption orientation may be visualised as occurring vertically, or with some degree of tilting in the molecules yz-plane. For completeness, Fig. S4 (ESI†) presents a visualisation of nitrobenzene aligned with respect to the molecule's xz-plane and eliminates the option of either vertical or tilted variants in that plane.



Scheme 1 A schematic representation for aniline (ANL) formation via the stepwise hydrogenation of nitrobenzene (NB) to phenylhydroxylamine (PHA) over a supported Pd catalyst as reported by Gelder and coworkers. Blue hashed boxes represent surface species. The figure has been adapted from Chemical Communications. 42

Small frequency shifts were observed when comparing wavenumber values for the key $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes during nitrobenzene adsorption over KBr (Fig. 2b) to adsorption over GU-1 (Fig. 2a) from 1529 to 1530 cm^{-1} and 1347 to 1352 cm⁻¹, respectively. A larger wavenumber shift upon adsorption was anticipated; not least because Section 3.2.5 reveals an appreciable enthalpy of adsorption over GU-1. One possible scenario is that dipole coupling effects⁴³ are contributing to the spectrum of the bound nitrobenzene via two opposing effects: a dipole coupling shift which acts to increase wavenumber values for high coverages, and a chemical shift that results in a lowering of wavenumber values. 43 RAIRS spectra reported by Koel and co-workers depicting increasing nitrobenzene coverage over Au(111) obtained via RAIRS do not exhibit any discernible frequency shift either.19

An additional broad feature centred at 955 cm⁻¹ is clearly discernible for the IR spectrum depicting nitrobenzene adsorption to GU-1 (Fig. 2a); this peak cannot be assigned to any nitrobenzene vibrational mode (Table 1). The origins of this band will be addressed further in Section 3.2.4.

3.2.3 Computational studies. In the previous section it was proposed that nitrobenzene is chemisorbed on the Pd nanoparticles in a vertical/tilted configuration via one Pd-O bond. In this section we investigate by periodic-DFT the extreme possibilities, i.e. where the plane of the phenyl ring is oriented perpendicular or parallel to the metal surface. As a model system, we have used a three-layer slab of Pd, cleaved along (111) with a 15 Å vacuum gap. For both orientations, there is a mirror plane perpendicular to the surface. The results are shown in Fig. 5. Parallel adsorption results in a significant distortion of the molecule with the nitro group and the hydrogens bent away from the plane of the molecule (see Table S3, ESI†), with the molecule exhibiting $C_s\sigma_v(xz)$ symmetry. This is reminiscent of the ligand distortions found in metal cyclopentadienyl (e.g. ferrocene⁴⁴) and metal arene complexes (e.g. bisbenzene chromium⁴⁵).

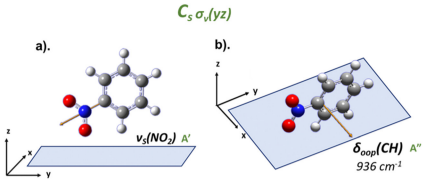


Fig. 4 Visualisation of (a). $V_S(NO_2)$ and (b). $\delta_{OOD}(CH)$ nitrobenzene modes during a tilted adsorption orientation over a metal surface with respect to the molecules yz-plane. Orange arrows indicate the dipole derivative unit vector associated with each mode. Grey parallelograms symbolise a non-specific Pd crystallite. White, grey, blue and red balls represent hydrogen, carbon, nitrogen and oxygen, respectively.

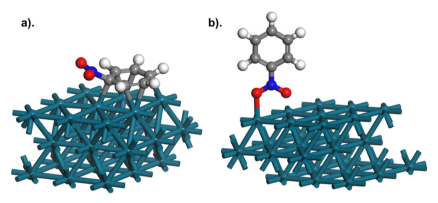


Fig. 5 Geometry optimised periodic-DFT structures of nitrobenzene adsorbed parallel (a) and perpendicular (b) on the Pd(111) facet. White, grey, blue and red balls represent hydrogen, carbon, nitrogen and oxygen, respectively

The optimised structure for nitrobenzene adsorbed on Pd(111) was recently reported by Millán and co-workers24 as occurring in a parallel orientation via binding of the phenyl group and via a single N-O···Pd, with the C-H bonds distorted from the plane of the molecule and the NO2 group twisted with respect to the phenyl ring. In contrast, our calculations show perpendicular adsorption results in only minimal changes in the molecular geometry. Consistent with the infrared studies (Section 3.2.2), monodentate adsorption via a N-O···Pd interaction is found.

The adsorption energy $(E_{ads}, kJ mol^{-1})$ can be calculated from:

$$E_{\text{ads}} = [\text{Pd}(111) + \text{nitrobenzene}] - [\text{Pd}(111)] - [\text{nitrobenzene}],$$
(1)

where the terms in square brackets refer to the energies of the adsorbate complex (i.e., as shown in Fig. 3), the clean surface and the free molecule, respectively. This gives $E_{ads} = -366$ and -99.3 kJ mol⁻¹ for the parallel and perpendicular adsorption geometries, respectively.

That the parallel geometry is the lowest energy state is a surprising result. Fig. 6 presents the experimental spectrum of nitrobenzene adsorption over GU-1 and the simulated spectra for parallel $(C_s\sigma_v(xz))$ and monodentate vertical $(C_s\sigma_v(yz))$ nitrobenzene adsorption over a metal surface. The simulated spectra

are obtained via the process described in Section 2.1. For a parallel adsorption, the simulated spectrum (Fig. 6a) depicts the absence of the in-plane $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes usually visualised at ca. 1350 and 1530 cm⁻¹ as per the MSSR. However, these modes are clearly observed during adsorption to GU-1 (Fig. 2a). In distinct contrast, the simulated spectrum for a monodentate nitrobenzene adsorption complex does present these modes and, crucially, it does not include any out-of-plane A" symmetry modes, as observed experimentally (Fig. 2a). Thus, comparison between the simulated $C_s\sigma_v(yz)$ symmetry vertical nitrobenzene adsorption spectrum (Fig. 6b) and experimental outcomes herewith is distinct, and clearly identifies nitrobenzene adsorption over GU-1 to result in an adsorption complex in which nitrobenzene exhibits $C_s\sigma_v(yz)$ symmetry, and not the proposed $C_s\sigma_v(xz)$ symmetry from simulated outcomes. Additionally, the pronounced distortion that nitrobenzene undergoes on parallel adsorption is not intuitive to a favourable geometry.

Relative to the ideal C_{2v} geometry of gas phase nitrobenzene, the parallel form of nitrobenzene is 113.1 kJ mol⁻¹ higher in energy, while the perpendicular form is only 7.9 kJ mol⁻¹ higher. The stabilisation of the parallel form presumably reflects that there are six Pd-C bonds, whereas there is a single Pd-O bond for the perpendicular form. Thus, we have

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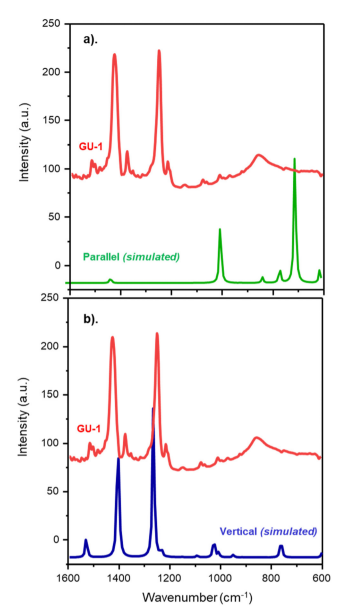


Fig. 6 DRIFTS spectrum of nitrobenzene adsorption to GU-1 (red) alongside the simulated spectrum for (a) parallel and (b) monodentate vertical nitrobenzene adsorption over a metal surface. Wavenumber range: $1600-600 \text{ cm}^{-1}$.

an interesting result and an apparent conundrum: the measured IR spectrum strongly correlates with nitrobenzene adopting a vertical/tilted geometry relative to the metal surface with binding occurring *via* a single O atom (Fig. 5b), whilst the energetics emanating from the calculations strongly indicate a preference for a distorted parallel geometry (Fig. 5a).

The authors note that the calculations correspond to a single adsorption event that does not account for any ensemble effects from intermolecular interactions between multiple molecular adsorptions; a scenario that is not wholly indicative of elevated nitrobenzene exposures as encountered in catalytic turnover. Specifically, GU-1 possesses a Pd surface density of $1.153 \times 10^{20} \ \mathrm{Pd}_{(\mathrm{s})} \ \mathrm{g}^{-1}^{25}$ and the incident nitrobenzene flux is

 3.68×10^{19} NB molecules min $^{-1}$ g $^{-1}$ (Section 2.3) that corresponds to a continuous incident flux of 0.32 NB molecules min $^{-1}$ Pd $_{(s)}^{-1}$. The lowest nitrobenzene exposure examined (0.62 mmol $_{(NB)}$ g $_{(cat)}^{-1}$) corresponds to a cumulative exposure of 1.87×10^{22} nitrobenzene molecules, which translates to 3234 nitrobenzene molecules Pd $_{(s)}^{-1}$. Thus, even the lowest nitrobenzene exposures considered here correspond to a multimonolayer coverage regime as encountered during catalytic turnover, which is differentiated from the single molecule coverage regime examined by the DFT calculations. Under these conditions, adsorbate–adsorbate interactions will prevail. Additionally, the Pd crystallites of GU-1, which represents a real working catalytic system, 8 may present a level of complexity not accessible via modelling on a perfect Pd(111) surface.

These reasonings represent suggestions for the encountered conundrum and highlight difficulties to be considered when applying a combination of *in situ* structural determination and isolated molecule DFT simulated structures within the field of heterogeneous catalysis. Summarising, nitrobenzene adsorption corresponding to aniline synthesis is thought to require the nitrobenzene to adopt an upright/tilted geometry with $C_s\sigma_v(yz)$ symmetry, as illustrated in Fig. 5b.

3.2.4 Surface coverage investigation: nitrobenzene adsorption at 28 °C. Fig. 7 presents spectra for nitrobenzene adsorption to GU-1 for the wavenumber range 1700–700 cm $^{-1}$. The full range for all spectra can be found in the supporting information (Fig. S5, ESI†). Characteristic nitrobenzene spectral features arising from the A' symmetry $\nu_{\rm AS}({\rm NO_2})$ and $\nu_{\rm S}({\rm NO_2})$ modes were observed post-exposure to 0.93 mmol_(NB) $g_{\rm (cat)}^{-1}$, centred at 1529 and 1347 cm $^{-1}$, respectively. IR bands arising

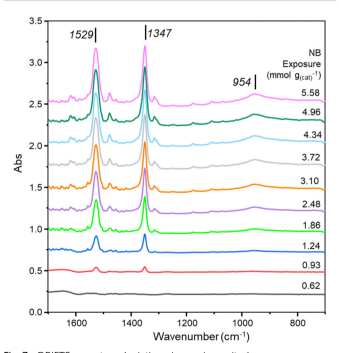


Fig. 7 DRIFTS spectra depicting increasing nitrobenzene exposure $(0.62-5.58 \text{ mmol}_{\text{(NB)}} \ g_{\text{(cat)}}^{-1})$ to the catalyst at 28 °C. The spectra have been offset by 0.25 au to facilitate viewing. Wavenumber range: 1700–700 cm $^{-1}$.

from A' symmetry modes associated with the aromatic moiety of nitrobenzene were first observable after introduction of 1.24 mmol_(NB) $g_{(cat)}^{-1}$. For example, ring stretching (ν (CC)) modes at 1607, 1585, 1476, 1415 and 1317 cm⁻¹; and a series of in-plane C-H bending modes ($\delta_{ip}(CH)$) at ca. 1300-1000 cm⁻¹. No IR features corresponding to out-of-plane A" symmetry nitrobenzene modes were observable for nitrobenzene exposures $\leq 5.58 \text{ mmol}_{(NB)} \text{ g}_{(cat)}^{-1}$.

As briefly considered in Section 3.2.2, a broad feature centred at 954 cm⁻¹ is observed at saturation coverages (Fig. 2a) that is not assigned to any nitrobenzene vibrational mode. Fig. 8 presents spectra for nitrobenzene exposures of 0.62 and 0.93 mmol_(NB) $g_{(cat)}^{-1}$ only and reveals a series of bands at 1647, 1271 and 954 cm⁻¹ after the initial 0.62 mmol_(NB) $g_{\text{(cat)}}^{-1}$ exposure. Aniline possesses a NH₂ bending mode $(\delta_{\text{oop}}(NH_2))$, a C-N stretching mode $(\nu(CN))$ and a series of C-H deformation modes (δ (CH)) at ca. 1628, 1278 and 996-883 cm⁻¹, 46,47 respectively. Thus, with reference to Fig. 8, we attribute the observation of stated wavenumber values at $0.62 \text{ mmol}_{(NB)} \text{ g}_{(cat)}^{-1}$ exposure to the formation of aniline on the catalyst, with the $\delta_{\text{oop}}(NH_2)$ mode exhibiting a notable shift in wavenumber, and assign the broad 954 cm⁻¹ feature observed at a saturation coverage of nitrobenzene (Fig. 2a and 7) to a δ (CH) mode of aniline. The intense and broad nature of this mode at nitrobenzene saturation coverage is anomalous and requires further investigation.

With reference to Fig. 7 and 8, the aniline features, excluding that of the broad band at ca. 955 cm⁻¹, are only observable at low nitrobenzene exposures. As they do not scale with nitrobenzene exposure ($\leq 0.62 \text{ mmol}_{(NB)} \text{ g}_{(cat)}^{-1}$); *i.e.*, no nitrobenzene modes are observed, the aniline is thought to occur via

nitrobenzene hydrogenation via reaction with a small reservoir of residual hydrogen left over from the catalyst reduction stage. Thus, we have two sources of aniline adsorption to GU-1: (i) hydrogenation of nitrobenzene via a limited reservoir of sub-surface and surface hydrogen from reduction, as evidenced in Fig. 8, and (ii). via a minor aniline impurity within the experimental set-up.

Fig. 9 presents the intensity profile for the main IR bands observed in Fig. 7. At an exposure of 0.62 $\text{mmol}_{(NB)} g_{(cat)}^{-1}$ no $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ features are observable but, thereafter, at higher exposures ($\geq 0.93 \text{ mmol}_{(NB)} \text{ g}_{(cat)}^{-1}$) nitrobenzene coverage increases and approaches a plateau for nitrobenzene loadings exceeding 4.96 mmol_(NB) $g_{(cat)}^{-1}$. The aniline intensity, as signified by the broad feature at 954 cm⁻¹, is seen to progressively increase as a function of nitrobenzene exposure. The separate profiles for nitrobenzene and aniline in Fig. 9 suggest separate adsorption sites on GU-1 for reagent and product. Summarising, for exposures of $\geq 0.93 \text{ mmol}_{(NB)} \text{ g}_{(cat)}^{-1}$, nitrobenzene adsorption on the Pd surface is seen to occur in a vertical/titled orientation in the yz-plane via one Pd-O bond independent of coverage, with the concomitant formation of aniline, which adsorbs in a different manner on GU-1.

3.2.5 Temperature-programmed nitrobenzene desorption from GU-1. Fig. 10 presents the temperature-programmed IR (TP-IR) spectra for a saturation dose of nitrobenzene over GU-1. No change in nitrobenzene adsorption geometry was observed with increasing desorption temperature; in-plane A' symmetry $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes remained observable with no emergence of any out-of-plane A" symmetry ($\delta_{\text{oop}}(CH)$) derived nitrobenzene IR features. Thus, the form of the nitrobenzene adsorption complex is unchanged throughout the desorption

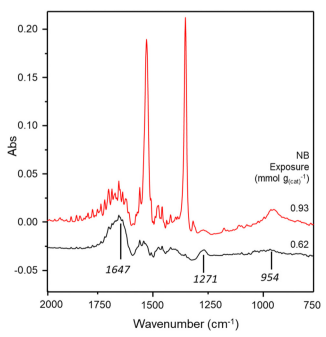


Fig. 8 DRIFTS spectra depicting nitrobenzene exposures of 0.62 (black) and 0.93 (red) $\text{mmol}_{(\text{NB})} \ {g_{(\text{cat})}}^{-1}$ to the catalyst at 28 °C. The spectra have been offset to facilitate viewing

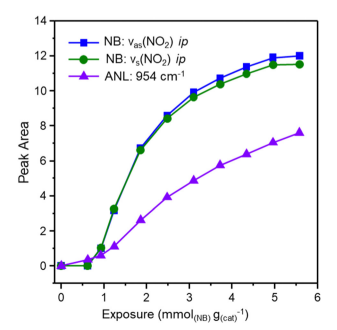


Fig. 9 Plot of peak area for nitrobenzene $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes and aniline δ (CH) mode (954 cm⁻¹) as a function of increasing nitrobenzene exposure to GU-1 at 28 °C.

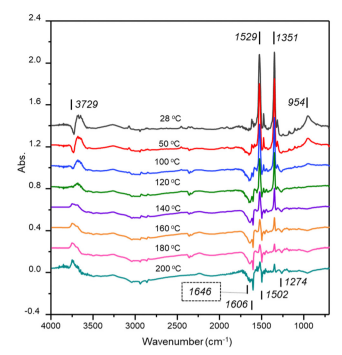


Fig. 10 DRIFTS spectra depicting nitrobenzene desorption from GU-1 with increasing temperature (28-200 °C). The spectra have been offset by 0.2 au to facilitate viewing

phase depicted in Fig. 10 and remains in a vertical/tilted orientation with $C_s\sigma_v(yz)$ symmetry. A change of orientation in the xz-plane (Fig. S4, ESI†) would be distinct in IR spectra, as described in the ESI†

Although orientation of the reagent did not change with increasing desorption, several spectral changes are evident in Fig. 10. Firstly, the saturation coverage spectrum (Fig. 10, 28 °C) exhibits a negative peak at 3729 cm⁻¹ and a positive peak at 3677 cm⁻¹. Surface hydroxyl groups of γ -Al₂O₃ include terminal hydroxyls (type I) at ca. 3764 cm⁻¹, bridging hydroxyls (type II) at ca. 3730 cm⁻¹, and tri-bridged hydroxyls (type III) at ca. 3675 cm^{-1} . 48,49 We therefore assign the negative IR feature at 3729 cm⁻¹ to bridging hydroxyl groups and the positive feature at 3677 cm⁻¹ to tri-bridging hydroxyl groups on the alumina support of GU-1, respectively.

All spectra presented have been background subtracted using a spectrum of the activated catalyst. Thus, the observed hydroxyl features suggest perturbation of the bridging and tribridged hydroxyl groups of the catalyst support. We propose the negative feature at 3729 cm⁻¹ observed at saturation coverage is indicative of a moiety forming a hydrogen bond to the bridging hydroxyl groups of the alumina support; this shifts the discrete ν (OH) to lower wavenumber, where it is submerged in the broad envelope of hydrogen-bonded hydroxyls. On background subtraction, this loss of the isolated $\nu(OH)$ leads to a negative feature. Similar observations have been reported over various aluminas during reagent adsorption. 49-51 As the desorption temperature is increased, the negative peak at 3729 cm⁻¹ decreases such that at 120 °C the negative hydroxyl feature is no longer observable, indicating desorption of the adsorbed

species from the bridging hydroxyl groups of the alumina support of GU-1 at this temperature. Crucially, at this temperature, the key in-plane A' symmetry modes ($\nu_{\rm S}({\rm NO_2})$ and $\nu_{AS}(NO_2)$) of nitrobenzene remain discernible, confirming that nitrobenzene adsorption occurred on the Pd crystallites of GU-1 and not the alumina support.

TP-IR measurements also revealed the emergence of derivative type features below 1700 cm⁻¹ with increasing desorption temperatures ≥100 °C. Four discernible negative intensity features were observed at 1646, 1606, 1502 and 1274 cm⁻¹ and are assigned to aniline: the broad feature at 1646 cm⁻¹ corresponds to a shifted out-of-plane NH₂ bending $(\delta_{oop}(NH_2))$ mode, sharp bands at 1606 and 1502 cm⁻¹ correspond to in-plane ring stretching ($\nu(CC)$) modes and the feature at 1274 cm⁻¹ corresponds to the in-plane C-N stretching (ν (CN)) mode.47 These negative aniline bands are derived from the small aniline impurity within the experimental set-up, which permitted a limited degree of aniline adsorption to GU-1 prior to background collection (see Fig. 8). This small population of aniline is believed to be a 'spectator' species that does not affect observables concerning the nitrobenzene adsorption process. Thus, as referred to in Section 3.2.4, IR spectroscopy revealed the presence of two categories of adsorbed aniline that require accounting for. In the first instance, a small amount of aniline held as an impurity in the gas lines is present in the IR background spectrum. Secondly, low nitrobenzene exposures to GU-1 react with a small degree of retained hydrogen to form aniline that is adsorbed on metal sites (Fig. 8), with aniline shifted to the alumina support with increasing nitrobenzene exposure and represented by a broad feature at 954 cm⁻¹.

Fig. 11 presents a plot of peak area for the A' symmetry nitrobenzene modes ($\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$) and the anilinederived mode observed at 954 cm⁻¹ as a function of increasing temperature and indicates different binding strengths of nitrobenzene and aniline on GU-1. The $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes of nitrobenzene remain observable in IR spectra after heating to the maximum desorption temperature (200 °C) utilised in this investigation, albeit in a significantly reduced capacity, while the aniline mode (954 cm⁻¹) is absent in the 120 °C spectrum; indicative of aniline desorption from GU-1 for the temperature range of 100-120 °C. Thus, under the stated reaction conditions, TP-IR spectra indicate nitrobenzene to have a greater binding strength to GU-1 than that of the product aniline, an outcome which is desirable for high selectivity catalysis.

The TP-IR temperature range for aniline desorption on GU-1 (100-120 °C) is illuminating. Previously in this section, we attributed adsorption of some species to the bridging hydroxyl groups of the alumina support up to a temperature of 120 °C. With the coincidence of these desorption temperatures (the absence of both the negative hydroxyl feature and the $954~{\rm cm}^{-1}$ IR feature at 120 °C), it is proposed that aniline binds to the alumina support of the catalyst via bridging hydroxyl groups with nitrobenzene adsorption occurring on the Pd crystallites in a vertical/tilted in the yz-plane orientation via monodentate binding. Furthermore, it is proposed that increasing

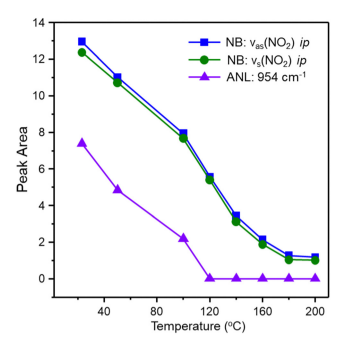


Fig. 11 Plot of peak area for nitrobenzene $\nu_{AS}(NO_2)$ and $\nu_{S}(NO_2)$ modes and aniline $\delta(\text{CH})$ at 954 cm^{-1} as observed on GU-1 as a function of increasing temperature

nitrobenzene exposure forces the aniline that is formed from nitrobenzene hydrogenation at 28 °C onto the support material, as indicated by the broad feature at 954 cm⁻¹ in Fig. 7.

4. Discussion

Previous reaction testing with GU-1 revealed near complete nitrobenzene conversion, good aniline selectivity (ca. 90% at 60 °C) and a prevalence for aniline derived vs. nitrobenzene

derived by-products during nitrobenzene hydrogenation.8 It is opportune to consider the observed catalytic behaviour with respect to the deduced adsorbate geometry and/or adsorbate location on GU-1, derived via DRIFTS measurements. Fig. 12 presents a schematic diagram depicting the $C_s\sigma_v(yz)$ symmetry monodentate vertical adsorption of nitrobenzene to the Pd crystallites of GU-1; in the presence of hydrogen this produces aniline, some of which adsorbs on the support material. For higher nitrobenzene exposures, no aniline adsorbs on the Pd.

Nitrobenzene is hydrogenated to yield predominantly aniline, with low levels of by-products reported at 60 °C (aniline derived by-products selectivity ≤10%; nitrobenzene derived by-products selectivity $\leq 0.2\%$)⁸. The authors propose that it is the nitrobenzene adsorption geometry over GU-1 that is key to explaining the observed low levels of by-products associated with the reagent (<0.2%). Specifically, it is the perpendicular/ tilted in the yz-plane nature of the adsorption of nitrobenzene to the metal surface that results in the distancing of the aromatic moiety of the molecule to the Pd crystallites, excluding a parallel arrangement that would otherwise favour ring hydrogenation. The minimal observation of nitrobenzene derived by-products cyclohexanol and cyclohexanone, which require aromatic ring hydrogenation,9 illustrate this chemical pathway to be minimised due to the proposed geometric reasons. This outcome, as observed with IR spectra, opposes those reported in Section 3.2.3. where simulated outcomes indicated a parallel adsorption of nitrobenzene over Pd(111) to be energetically favourable. However, it is crucial to re-iterate that these DFT calculations represent a single nitrobenzene molecule on a perfect Pd(111) surface - parameters of significance include (i) experimental coverages significantly exceed single molecule adsorption, and (ii) GU-1 is not a perfect Pd(111) surface. Additionally, IR measurements (Fig. 7, 9 and 10) indicate that high coverages of nitrobenzene force small quantities of aniline, formed via reaction with residual

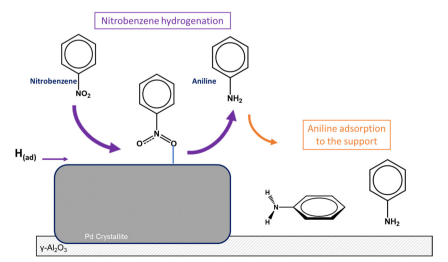


Fig. 12 Diagram visualising nitrobenzene adsorption to Pd in a vertical orientation via one Pd-O bond, with subsequent hydrogenation to aniline, and re-adsorption of some aniline to the support with no specific geometry identified. The solid grey box represents a non-specific Pd crystallite; the hashed box represents the Al₂O₃ support. Larger arrows for vertical NB hydrogenation to ANL depicts this as the major transformation.

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hydrogen, from the Pd surface onto the support material. This dynamic scenario is thought to be a contributory factor to high aniline selectivity for nitrobenzene hydrogenation over GU-18 and supported Pd catalysts in general.

5. Conclusions

Application of FT-IR spectroscopy, sequential nitrobenzene adsorption, TP-IR and consideration of the metal surface selection rule (MSSR) permitted determination of the adsorbate complex orientation for nitrobenzene adsorption over GU-1. The following conclusions can be drawn.

- Adsorption of nitrobenzene to GU-1 yielded DRIFTS spectra exhibiting IR features solely associated with monodentate vertical or tilted $C_s\sigma_v(yz)$ symmetry with in-plane A' symmetry modes observed, and the IR active out-of-plane A'' symmetry modes unobserved.
- Comparison of experimental outcomes and DFT simulations for a single adsorption event over Pd(111) present opposing conclusions. However, this conundrum is attributed to the DFT calculations representing single molecule adsorption to a perfect (111) surface, whereas cooperative effects at elevated coverages (exposures of \geq 0.93 mmol_(NB) $g_{(cat)}^{-1}$), as encountered during catalytic turnover, are attributed to the nitrobenzene adopting an upright geometry.
- The lowest nitrobenzene exposure (0.62 mmol_(NB) $g_{(cat)}^{-1}$) investigated via DRIFTS identified the presence of aniline on GU-1 that was formed via nitrobenzene reaction with residual hydrogen.
- TP-IR measurements from GU-1 indicated aniline to be interacting with bridging hydroxyl groups of the alumina support of the catalyst up to 120 °C.
- It is proposed that the vertical/tilted orientation of nitrobenzene in the yz-plane with respect to the metal surface limits the formation of nitrobenzene derived by-products due to the perpendicular positioning of the aromatic ring of the molecule with the Pd surface.
- The predominant re-adsorption of aniline to the γ -Al₂O₃ support of GU-1 as opposed to adsorption on the Pd is proposed to limit product over-hydrogenation, and thus contributes to the elevated aniline selectivity observed.

Data management statement

Data for this paper, including spectroscopic datasets, are available from the University of Glasgow Library via the following link, http://dx.doi.org/10.5525/gla.researchdata.1447.

Conflicts of interest

There are no conflicts of interest to declare.

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