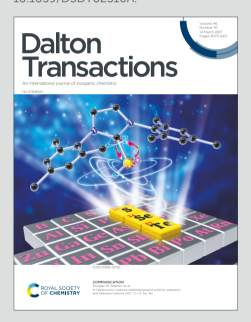
Dalton Transactions

An international journal of inorganic chemistry

Accepted Manuscript

This article can be cited before page numbers have been issued, to do this please use: S. Bellomi, I. Barlocco, S. Tumiati, P. Fumagalli, N. Dimitratos, A. Roldan and A. Villa, *Dalton Trans.*, 2023, DOI: 10.1039/D3DT02310A.



This is an Accepted Manuscript, which has been through the Royal Society of Chemistry peer review process and has been accepted for publication.

Accepted Manuscripts are published online shortly after acceptance, before technical editing, formatting and proof reading. Using this free service, authors can make their results available to the community, in citable form, before we publish the edited article. We will replace this Accepted Manuscript with the edited and formatted Advance Article as soon as it is available.

You can find more information about Accepted Manuscripts in the <u>Information for Authors</u>.

Please note that technical editing may introduce minor changes to the text and/or graphics, which may alter content. The journal's standard <u>Terms & Conditions</u> and the <u>Ethical guidelines</u> still apply. In no event shall the Royal Society of Chemistry be held responsible for any errors or omissions in this Accepted Manuscript or any consequences arising from the use of any information it contains.



View Article Online

View Journal

View Article Online DOI: 10.1039/D3DT02310A

ARTICLE

Effect of oxygen functionalities on the hydrous hydrazine decomposition over carbonaceous materials

Received 00th January 20xx, Accepted 00th January 20xx

DOI: 10.1039/x0xx000000x

Silvio Bellomi^a, Ilaria Barlocco^a, Simone Tumiati^b, Patrizia Fumagalli^b, Nikolaos Dimitratos^{c,d}, Alberto Roldan^{e*}, Alberto Villa ^{a*}

Metal-free heterogeneous catalysis is promising in the context of H_2 generation. Therefore, establishing structure-activity relationships is a crucial issue to improve the development of more efficient catalysts. Herein, to evaluate the reactivity of the oxygen functionalities in carbonaceous materials, functionalized commercial pyrolytically stripped carbon nanofibers (CNFs) were used as catalysts in the liquid-phase hydrous hydrazine decomposition process and compared its activity to the one of pristine CNF material. Different oxygenated groups were inserted by treating CNFs with hydrogen peroxide for 1h (O1- H_2O_2) and H_2O_3 and H_3 for 1h (O1- H_3) and 6h (O6- H_3). An increase in activity was observed as a function of the oxidizing agent strength (H_3) and H_2 0 and the functionalization time (H_3) and the functionalization time (H_3) and the functionalization time (H_3) and H_3 0 of H_3 0 of H_3 0 of H_3 0 of H_3 1 of H_3 2 of H_3 3 of H_3 3 of H_3 4 of H_3 5 of H_3 6 of H_3 6 of H_3 6 of H_3 7 of H_3 8 of H_3 9 of

Keywords: Hydrogen, Hydrous hydrazine, Heterogeneous catalysts, Metal-free, Carbon materials

Introduction

The search for sustainable chemical processes is crucial for achieving a resilient society. Alongside this, the global energetic demand is steadily increasing, raising the necessity for higher energy generation capacities, better management and efficiency of generated energy, and lower dependence on non-renewable fossil fuels.¹

Molecular hydrogen is recognized as one of the most promising clean energy sources due to its high intrinsic energy density and innocuous by-products upon utilization. Hydrogen can be produced through several methods, including photocatalytic and electro-catalytic systems, steam reforming and biomass-, coal- or natural gas-based processes.¹ Even though the benefits

of hydrogen are apparent, its direct usage is limited by the lack of economically sustainable and safe storage and transportation technologies.² Typically, storage procedures require pressurized tanks, both for gaseous or liquified hydrogen, whose applicability is restricted by low efficiencies and high costs.^{3,4}

Liquid Hydrogen Carriers (LHCs) are chemical compounds easily storable and transportable with a high intrinsic hydrogen content, which can be released chemically, desirably at mild conditions. Many LHCs exist, but the need to reduce CO_2 emissions has led to a focus on nitrogen-hydrogen compounds. N_2H_4 possesses a hydrogen content of 12.5 wt.%.⁵ Nonetheless, given its intrinsic hypergolic nature, i.e., serious risk of explosions in the presence of certain metals,⁶ it is handled in the hydrated form, hydrous hydrazine ($N_2H_4 \cdot H_2O$, 8 wt.% hydrogen content).^{7,8} $N_2H_4 \cdot H_2O$ is reported to decompose catalytically through two different pathways:⁹

$$N_2H_4 \rightarrow N_2(g) + 2H_2(g)$$
 $\Delta H^0 = -95.4 \, kJmol^{-1}$ (1)

$$3N_2H_4 \rightarrow N_2(g) + 4NH_3(g)$$
 $\Delta H^0 = -157.0 \ kJmol^{-1}$ (2)

The first reaction (Equation 1) represents the complete reforming pathway, where molecular hydrogen and nitrogen are produced. The second path (Equation 2) produces nitrogen and ammonia and is the thermodynamically favoured process. Several studies focused on using noble^{10,11} and non-noble^{12,13} metal-based catalysts to promote hydrazine decomposition.

^a Dipartimento di Chimica, Università degli Studi di Milano, via Golgi 19, I-20133, Milano. Italv.

^b Dipartimento di Scienze della Terra Ardito Desio, Università degli Studi di Milano, via Mangiagalli 34, Milano I-20133, Italy

^c Dipartimento di Chimica Industriale "Toso Montanari", Alma Mater Studiorum Università di Bologna, Viale Risorgimento 4, Bologna 40126, Italy

d Center for Chemical Catalysis-C3, Alma Mater Studiorum Università di Bologna, Viale Risorqimento 4, Bologna 40136, Italy

^e Cardiff Catalysis Institute, School of Chemistry, Cardiff University, Main Building, Park Place, CF10 3AT, Cardiff, United Kingdom.

^{*} alberto.villa@unimi.it

^{*} RoldanMartinezA@cardiff.ac.uk

[†] Footnotes relating to the title and/or authors should appear here. Electronic Supplementary Information (ESI) available: [details of any supplementary information available should be included here]. See DOI: 10.1039/x0xx00000x

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence Access Article. Published on 26 September 2023. Downloaded on 10/4/2023 2:29:08 PM

ARTICLE Dalton Transactions

These investigations highlighted a dependence of the reaction selectivity upon the catalyst's nature and the reaction parameters, e.g., pressure, temperature, and stirring. 11,14-17 Despite the catalytic activity presented, the carbon fingerprint and scalability of the metal catalysts are questionable due to their mining and manufacturing methods and scarcity. Therefore, it is advised to design new metal-free catalysts for a sustainable future.18

Carbon-based materials are attracting significant attention as catalysts due to their high tunability. Facile changes in their lattice structure allow significant variations of, for instance, surface area, porosity, functional groups and topological defects.¹⁹ Of particular interest are those metal-free carbocatalysts that successfully generated hydrogen in the liquid phase from different substrates, including alkenes, alkanes and formic acid. 20,21

Several works focused on the hydrazine-mediated reduction of different target substrates.22-24 Remarkable results were obtained for the reduction of nitroarenes performed over carbo-catalysts.²⁵ Fujita and co-workers²⁶ complemented these results unveiling the effect of N and O functionalities. The introduction of dopants induced changes in the surface properties of the carbons, enhancing the catalytic properties of the carbo-catalysts, i.e., conversions and selectivity of the target substrates.

Previous investigations in the activity of carbonaceous defects materials' intrinsic for hydrous hvdrazine decomposition unveiled full that pyrolytically stripped carbon nanofibers (CNFs) exhibited the best performances among the tested carbo-catalysts thanks to a high degree of intrinsic defects, resulting in complete conversion of hydrous hydrazine up to 6 h of reaction with 89 % H₂ selectivity.²⁷ Despite previous works and to the best of our knowledge, the literature lacks investigations rationalizing the effects of oxygen functionalities in metal-free carbonaceous materials for the decomposition of hydrous hydrazine.

The work presented here focuses on the liquid-phase oxygen functionalization of CNFs through different procedures and unveils their role in hydrous hydrazine decomposition. The nature of oxygen groups was modulated by employing different oxidizing agents (H₂O₂ and HNO₃) at variable functionalization times (1 and 6 h), resulting in O1-H₂O₂, O1-HNO₃, and O6-HNO₃. The functionalized carbo-catalysts were tested as catalysts under mild conditions, and their activity was compared with the pristine CNFs. To understand the effect of the functionalization procedures on the carbonaceous materials, fresh and spent catalysts were thoroughly characterized through ICP, Raman and XPS in combination with computational techniques.

Experimental

Materials and chemicals

Pyrolytically Stripped carbon nanofibers (CNFs) were purchased from the Applied Science Company. Hydrous hydrazine (N₂H₄·H₂O, 98 %), sodium hydroxide (NaOH, ≥98 %), nitric acid (HNO₃, 65 %) and hydrogen peroxide $P(H_2O_2,03)O^3\%$ where acquired from Sigma-Aldrich.

Catalyst synthesis

In a typical functionalization procedure, 1 g of CNFs were dispersed in 200 ml of the desired oxidizing agent (HNO₃ 65 % or H₂O₂ 30 %) in a 500 ml round bottom flask. Two different oxidizing agents were employed, namely HNO₃ and H₂O₂. For the HNO₃ treatment, a 353 K temperature and 800 rpm stirring rate were employed, and the mixture was left for the desired time (1 or 6 h), resulting in O1- and O6- HNO3, respectively. Differently, for the H₂O₂ treatment, the functionalization was performed at RT and with a stirring rate of 800 rpm for 1 h, leading to O1-H₂O₂. A longer functionalization time with H₂O₂ (6 h) did not produce significantly different kinetic results than the 1 h functionalized carbons; thus, we excluded O6-H₂O₂ from the proposed investigation.

Catalytic tests

Liquid phase N₂H₄·H₂O decomposition was performed at a constant reaction temperature of 323 K using a 50 mL threenecked round bottom flask, with one of the flask's necks connected to a burette for the volumetric analysis of the gaseous products. Typically, the desired amount of catalyst (hydrous hydrazine/catalyst wt. ratio 16/1) was added in the reactor in the presence of 16.0 mL of a 0.5 M NaOH aqueous solution, afterwards equilibrated at a stirring rate of 1400 rpm. Then, 600 µL of a 3.3 M aqueous solution of hydrous hydrazine was injected. Hydrous hydrazine conversion was analysed through a Jasco V-730 spectrophotometer, using a 1 cm quartz cell, based on the quantitative reaction of N₂H₄·H₂O with 4dimethylaminobenzaldehyde (4-DMAB) in dilute hydrochloric acid (HCl), resulting in a p-quinone structure that strongly absorbs radiation at 456 nm (Figure 1).28 The volume of gas produced was measured using the water displacement method. In a typical experiment, the reactor was connected with a tube fed under a flipped burette, filled with water, and placed in a beaker containing H₂O. During the evolution, the gases were allowed to pass in a 0.05 M HCl aqueous solution to ensure the removal of NH₃, if any. Therefore, we quantified the evolved gaseous products (H₂ and N₂) from the volume of water displaced within the burette.27

Catalysts characterization

Functionalized carbon materials were characterized by Raman spectroscopy and X-ray Photoelectron Spectroscopy (XPS).

Figure 1 Reaction of 4-DMAB and hydrous hydrazine to obtain the UV-Vis detectable pquinone.

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 26 September 2023. Downloaded on 10/4/2023 2:29:08 PM

ARTICLE Dalton Transactions

Raman spectroscopy was executed at the Department of Earth Sciences - University of Milan, using a Horiba LabRam HR Evolution micro-Raman spectrometer equipped with a green solid-state laser (532 nm) focused through a 100× objective, giving a spatial resolution of approximately 1 µm. The micro-Raman system was set with 600 lines per mm grating and a hole of size 300 µm; the spectrum was collected with a final laser power of about 0.05 mW. The sample surface was measured through a hand-held power meter. Spectra were calibrated using the 520.7 cm⁻¹ line of a silicon wafer. The sample was scanned at an exposure time of 300 s, and two accumulations were performed, giving a spectrum. XPS measurements were performed with a Thermo Scientific K-alpha+ spectrometer. The samples were analysed using a monochromatic Al X-Ray source operating at 72 W, with the signal averaged over an oval-shaped area of 600×400 μm. Data were recorded at 150 eV for survey scans and 40 eV for high-resolution (HR) scans with a 1 eV and 0.1 eV step size, respectively. CASAXPS (v2.3.17 PR1.1) was used to analyse the XPS raw data. The C1s and O1s XPS core levels were fitted in agreement with models previously reported in the literature.²⁹ The presence of possible residual metal was analysed by inductively coupled plasma optical emission spectroscopy (ICP-OES) using a PerkinElmer Optima 8000 emission spectrometer.

Computational details

Plane-wave density functional theory (DFT) calculations were performed through the Vienna Ab-initio Simulation Package (VASP). 30,31 A generalized gradient in the revised Perdew-Burke-Ernzerhof approximation functional (rPBE)³² was employed to evaluate the correlation-exchange electronic contributions, with a kinetic energy cutoff of 500 eV chosen for the expansion of the Kohn–Sham valence states plane-waves.³³ rPBE is found to be efficient in modelling surface adsorption processes.³² In addition, the VASP implementation of the rPBE functional improves the previous revPBE implementation³⁴, fulfilling the Lieb-Oxford criterion globally by construction without any additional fitted parameter. All the calculations included the long-range dispersion correction approach by the Grimme DFT-D3 methods.^{35,36} The implicit polarizable continuum solvation model, as implemented in VASPsol, 37,38 was included in all the calculations. The optimization thresholds were 10⁻⁵ eV and 0.02 eV/Å for electronic and ionic forces relaxation. For the Brillouin zone (BZ) sampling, a Γ-centred k-point mesh generated through the Monkhorst–Pack method of dimensions $5 \times 5 \times 1$ was employed.³⁹ A first-order Methfessel-Paxton method was used with an energetic width value of 0.1 eV to ease the BZ sampling and improve the wavefunction convergence. All carbon-based materials were modelled starting from a single layer slab of a 6 x 6 pristine graphene supercell, where a single vacancy (SV) was introduced and functionalized with different O-containing groups, namely carbonylic (C=O), esteric (C-O-C), hydroxylic (C-OH) and carboxylic (COOH) moieties (Figure S1), in agreement with the XPS fitting model employed²⁹. The supercell is in a hexagonal crystalline system with unit cell vectors a and b lying in surface plane (γ =120°) and perpendicular to c axis.

Both, a and b, were optimized at 2.46 Å, in well agreement with experimental values obtained by HRPTEM039603DTCARBON materials.40 A polarizable continuum dielectric bath of ~16 Å perpendicular to the C-surface was introduced to avoid spurious periodic interactions. The adsorption energy (Eads) was defined as the difference between the combined system and the isolated species. Gibbs energies of adsorption (Gads) were evaluated by adding zero-point energy corrections and entropy terms (details can be found in SI).41 The first were calculated in the harmonic approximation, while the entropy of hydrous hydrazine was obtained from the international tables⁴². The entropies of solid-state species were considered negligible. 43 All thermal corrections were made at the reaction temperature

Results and discussion

Structure

The effect of oxygen functionalities on the hydrous hydrazine decomposition was investigated by modulating the nature of the oxidizing agent and the functionalization time on pristine carbon nanofibers (CNFs). As reported in the literature, a strong oxidizing agent such as nitric acid (HNO₃) can preferentially carboxylic/carbonylic introduce (C=O/COOH) proportionally to the functionalization time. Whereas a milder oxidizing strength compound, e.g., hydrogen peroxide (H2O2), increases the hydroxylic/esteric (C-OH/-C-O-C) group content.44 All the catalysts were thoroughly characterized through ICP-OES, Raman and XPS to disclose the catalysts' structure-activity relationship. Results from the ICP-OES analysis ensured the absence of metal impurities catalysing the hydrous hydrazine decomposition. $^{10,45}\,\mbox{Raman}$ spectroscopy evaluated the effect of the functionalization procedures on the catalyst morphology.²⁷ Commonly, Carbon materials present two characteristic bands at around 1600 cm⁻¹ (G band) and 1350 cm⁻¹ (D band), and their intensity ratio (I_D/I_G) is commonly used as a descriptor of the defect density. In fact, the G band intensity correlates to structurally ordered graphite domains. The D band is symmetry forbidden and only allowed in structural defects or in-plane substitutional heteroatoms. 46 The comparable I_D/I_G ratios between the functionalized carbons and CNFs allowed us to infer that the functionalization procedures did not induce modification in the carbon framework, i.e., substitutional heteroatoms, Figure S2 and Table S1, excluding the influence of additional intrinsic defects for the hydrous hydrazine decomposition. XPS provided information on the oxygen functionalities from analyses of the C1s and O1s core levels. The treatment with 6 h in HNO₃ introduced the highest quantity of oxygen groups (O/C = 0.157, O6-HNO₃), followed by the treatment for 1 h in HNO_3 (O/C = 0.141, O1- HNO_3) and the 1h in H_2O_2 (O/C = 0.141, O1- H_2O_2). All the procedures increased the number of oxygen functionalities with respect to the pristine material (O/C = 0.122, CNFs). The results are summarised in Table S1.

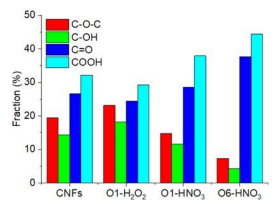


Figure 3 HR-XPS analysis of the O1s region, speciation of the oxygen-containing functional groups for the different catalysts.

We carried out a quantitative assessment of surface oxygen through high-resolution (HR) XPS analyses, focused on the O1s and C1s core levels to obtain more insights into the type of oxygen and surface content of each group. 47-49 Based on previously reported models, five peaks were considered for the O1s region as a function of their binding energies (BE).²⁹ Peaks at 531.1-531.8 eV can be assigned to carbon-oxygen double bond (C=O), 532.4-532.7 eV to carbon-oxygen ether-like single bonds (C-O-C), and 533.3-533.6 eV to carbon-oxygen bond in hydroxylic moieties (C-OH). Carboxylic moieties (COOH) are in the BE region of 532.0-533.0 eV, presenting an intermediate character between single and double bonds. A broad peak at about 535.2 eV is assigned to residual adsorbed water on the catalyst's surface. The results of the O1s core level fitting are summarised in Figure 2. The C1s core levels, Figure S4 and Table S3, agree with the O1s data.

Functionalizing CNFs with H_2O_2 for 1 h (O1- H_2O_2) increased the ether-like (C-O-C) and hydroxylic (C-OH) groups to 23.1 and 18.2% compared to 19.5 and 14.3% in the initials CNFs, Figure 2 and Table S2. Differently, the HNO₃ functionalization of CNFs, at both 1 h (O1-HNO₃) and 6 h (O6-HNO₃) led to the preferential formation of carbonylic (C=O, 28.6 and 37.6% vs. 26.6% in the original CNFs) and carboxylic groups (COOH, 37.8 and 44.4% vs. 32.2 % in CNFs). These results align with the higher oxidation strength of HNO₃ than H_2O_2 , Table S2, and with previous reports. 44,50

Activity

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence

Open Access Article. Published on 26 September 2023. Downloaded on 10/4/2023 2:29:08 PM

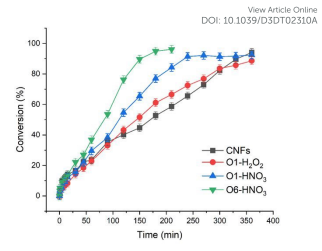
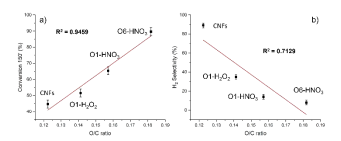


Figure 2 Conversion trend in the hydrazine decomposition reaction for the different catalysts: CNFs (black), O1-HNO $_3$ (blue), O6-HNO $_3$ (green) and O1-H $_2$ O $_2$ (red). All the tests were performed at 323 K and 1400 rpm, using 600 μ L of a hydrazine solution 3.3 M in 16 mL of NaOH 0.5 M and a N $_2$ H $_4$ ·H $_2$ O: catalyst weight ratio of 16:1.

The hydrous hydrazine decomposition on each catalyst was studied in a batch reactor at mild conditions (323 K and 1 atm). The conversions were quantified through the UV-Vis analysis of hydrous hydrazine quantitatively reacted with a colorimetric reagent (Figure 1). At the same time, the selectivity towards $\rm H_2$ production was calculated via the water displacement method

Figure 4 The trends of a) conversion at 150' and b) H_2 selectivity concerning the oxygen content (O/C ratio) derived from XPS analyses. Insets are the catalyst labels and R^2

when gas evolution reached a plateau, i.e., the end of the decomposition reaction. The kinetic results are represented in Figure 3, showing that at 150 minutes of reaction, O6-HNO3 resulted in the highest catalytic conversion (89 %), followed by $O1-HNO_3$ (65 %), $O1-H_2O_2$ (52 %), and CNFs (45 %). The O6-HNO₃ sample completed the reaction within 210 minutes of reaction, much faster than other catalysts (360 minutes). The enhanced activity in the presence of oxygen functionalities agrees with previous works on functionalized carbons for the hydrazinemediated H transfer to nitroarenes.^{25,26} Differently, we observed a decrease in the H₂ production concerning the oxidizing agent strength and functionalization time (O6-HNO₃, 8 % < O1-HNO₃, 14 % < O1-H₂O₂, 35 % < CNFs, 89 %). Therefore, we postulate that the reported enhanced hydrogen transfer in the presence of oxygen functionalities^{25,26} is due to the hydrous hydrazine decomposition intermediates, capable of transferring



H more readily to the reduction targets.

Structure-Activity Relationship

Dalton Transactions ARTICLE

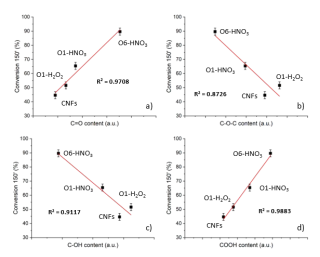


Figure 5 Trends of conversion with respect to the total surface content of a) C=O, b) C-O-C, c) C-OH and d) COOH, as quantified from the XPS analysis. Insets are the catalyst labels and R² factor of the linear fit.

The oxygen content is linearly correlated with the catalytic activity (89 %, O6-HNO₃ > 65 %, O1-HNO₃ > 52 %, O1-H₂O₂ > 52 %, CNFs). Thus, confirming the active role of the oxygen groups in the hydrous hydrazine decomposition, Figure 4a. However, the greater the number of oxygen functionalities, the lower the selectivity towards H₂ (8 %, O6-HNO₃ < 14 %, O1-HNO₃ < 35 %, O1-H₂O₂ < 89 %, CNFs), although no linear correlation has been observed, Figure 4b. Table S1 summarises structural features (O/C ratio from XPS and I_D/I_G from Raman) of the different catalysts considered and catalytic conversions at 150 minutes and H₂ selectivity at the end of the reaction Insets are the catalyst labels and R² factor of the linear fit.. We plotted the N₂H₄ conversion as a function of the total surface content of each functional group in Figure 5 to identify the active site for the reaction. A positive correlation was found between the conversion and the concentrations of C=O (R^2 = 0.9708) and COOH (R2 = 0.9883). On the other hand, a content increase of C-O-C ($R^2 = 0.8726$) and C-OH ($R^2 = 0.9117$) decreases the catalytic performances. These results lead to the conclusion that carbonoxygen double bond groups, i.e., C=O and COOH, are the active sites for the hydrous hydrazine decomposition reaction.

Atomistic Modeling

Systematic DFT simulations were employed to unravel the role of oxygen functionalities at the atomistic scale and rationalize the catalytical performances and kinetic trends in relation to the surface species. Represented in Figure S1 are the four different graphitic surfaces modelled as derived from XPS, i.e., C=O, C-O-C, C-OH and COOH. All the structures were optimized from relaxed graphene layers with a single vacancy to which the desired functional group was attached.⁵¹

Previous works on hydrazine highlighted the cis configuration of N_2H_4 as the most favorable for adsorption processes. 12,27,52 Therefore, cis- N_2H_4 was brought close to the functional group on the surface, so that the N atoms would be able to interact properly with the dangling bonds The most favourable adsorptions (Figure 6 and S5) led to trends as a function of the

oxygenated species: C-OH (-0.82 eV) < C=O (-1.39 eV) < CQOH c 1.34 eV) < C-O-C (-1.62 eV). Figure 6 shows that the N2H interaction with C=O, C-OH and COOH promoted the N-N dissociation, leading two NH₂ groups saturating nearby C-dangling bonds. In addition, for all the surfaces, anti and gauche configurations were tested, favoring the interaction of the dangling bonds with the N-H moieties, but the process was found endothermic for all the structures.

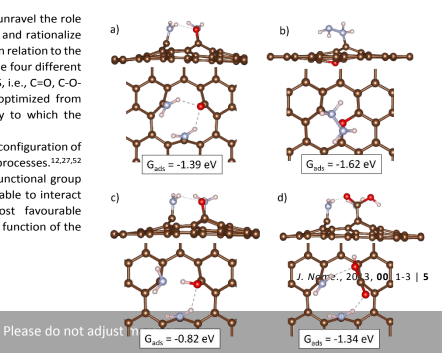
The initial exploration of the N_2H_4 interaction with the functionalized surface, in tandem with the previous experimental results, indicates Gibbs energies of \sim -1.3/1.4 eV to be neither too strong nor too weak to promote the N_2H_4 decomposition. 53 The strong N_2H_4 molecular adsorption on C-O-C sites may lead to surface poisoning, a hypothesis confirmed by the systematic decrease of C-O-C moieties observed in the O1s core level of the spent catalysts, Table S4. The orientation of the adsorbates and the functional C=O and COOH groups also points to a possible H-transfer and enolate formation as part of the catalytic cycle, which C-OH cannot undertake. The exothermic adsorption on the active sites (C=O and COOH) agrees with the experimental enhanced catalytic activity observed at increasing oxygen content. In addition, we previously demonstrated that the scission of the N-N bond in

Figure 6 . Top and side view of the most favourable adsorption configurations upon interaction with N_2H_4 : a) C=O, b) C-O-C, c) C-OH and d) COOH. The carbon atom is labelled brown, oxygen in red and hydrogen in white. Inset are the Gibbs adsorption energies (G_{ads}).

the first adsorption step of hydrazine favours NH_3 formation (Equation 2)²⁷ and previous works on hydrazine mediated H-transfer highlighted enhanced reduction of target substrates in presence of oxygenated carbonaceous materials.^{25,26} Therefore, we explain the selectivity results of the active sites as a combined N-N dissociation and preferential H-transfer, suppressing H_2 generation as a whole.

Conclusions

A combination of computational and experimental studies was employed to elucidate the role of oxygen functionalities in carbonaceous materials for the hydrous hydrazine decomposition reaction. Three different oxygen functionalized catalysts were synthesized (O1- H_2O_2 , O1- HNO_3) and O6- HNO_3)



This journal is © The Royal Society of Chemistry 20xx

Jalton Transactions Accepted Manuscril

ARTICLE Dalton Transactions

using different oxidizing agents (HNO₃, H₂O₂) and variable times (1 and 6 h). These samples were compared to the pristine nanofibers. An enhanced catalytic activity (89 %, O6-HNO₃ > 65 %, O1-HNO₃ > 52 %, O1-H₂O₂ > 52 %, CNFs) and decreased H₂ production (8 %, O6-HNO₃ < 14 %, O1-HNO₃ < 35 %, O1-H₂O₂ < 89 %, CNFs) were observed concerning the oxidizing agents' and functionalization time. Α characterization of the catalyst through Raman and XPS unveiled a positive linear correlation between the oxygen content and the catalytic activity ($R^2 = 0.9459$). In contrast, a negative trend was observed for hydrogen production, although there is no linear correlation ($R^2 = 0.7129$). A targeted analysis of O1s and C1s core levels and kinetic data assigned carbon-oxygen double bond groups (COOH and C=O) as active sites. The adsorption of hydrazine on graphitic surfaces resembling those derived from XPS was simulated using DFT, resulting in a preferential N-N breakage on the active sites (C=O and COOH). These groups may form enolate intermediates, favouring the NH₃ and explaining the experimental trends in H₂ selectivity.

Author Contributions

Conceptualization, A.V; Validation, A.V, A.R.; Formal analysis, investigation and writing-original draft preparation, S.B. Writing-review and editing A.R., I.B., P.F, S.T. All authors have read and agreed to the published version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research is part of the project "One Health Action Hub: University Task Force for the resilience of territorial ecosystems", supported by Università degli Studi di Milano -PSR 2021 - GSA. We acknowledge computing time on the facilities of Supercomputing Wales and the Advanced Research Computing Cardiff (ARCCA) at Cardiff University.

Notes and references

- 1 T. N. Veziroğlu and S. Şahin, Energy Convers. Manag., 2008, 49, 1820-1831.
- 2 P. Nikolaidis and A. Poullikkas, Renew. Sustain. Energy Rev., 2017, 67, 597-611.
- 3 H. T. Hwang and A. Varma, Curr. Opin. Chem. Eng., 2014, 5, 42 - 48.
- D. Pukazhselvan, V. Kumar and S. K. Singh, Nano Energy, 4 2012, 1, 566-589.
- 5 J. K. Norskov and C. H. Christensen, Science (80-.)., 2006, **312**, 1322 LP - 1323.
- 6 Z. Zhang, S. Zhang, Q. Yao, X. Chen and Z.-H. Lu, Inorg. Chem., 2017, 56, 11938-11945.

- S. J. Cho, J. Lee, Y. S. Lee and D. P. Kim, Catal. Letters, 2006. **109**. 181-186.
- A. K. Singh, M. Yadav, K. Aranishi and Q. Xu, Int. J. Hydrogen Energy, 2012, 37, 18915-18919.
- P.-X. Zhang, Y.-G. Wang, Y.-Q. Huang, T. Zhang, G.-S. Wu and J. Li, Catal. today, 2011, **165**, 80–88.
- 10 S. Bellomi, I. Barlocco, X. Chen, J. J. Delgado, R. Arrigo, N. Dimitratos, A. Roldan and A. Villa, Phys. Chem. Chem. Phys., 2023, 25, 1081-1095.
- D. Motta, I. Barlocco, S. Bellomi, A. Villa and N. Dimitratos, Nanomaterials, 2021, 11, 1340.
- 12 S. S. Tafreshi, A. Roldan and N. H. de Leeuw, Phys. Chem. Chem. Phys., 2015, 17, 21533-21546.
- W. Kang, H. Guo and A. Varma, Appl. Catal. B Environ., 13 2019, 249, 54-62.
- 14 L. Zhou, X. Luo, L. Xu, C. Wan and M. Ye, Catalysts, , DOI:10.3390/catal10080930.
- L. He, B. Liang, L. Li, X. Yang, Y. Huang, A. Wang, X. Wang 15 and T. Zhang, Acs Catal., 2015, 5, 1623-1628.
- 16 K. S. Sanjay, X. B. Zhang and Q. Xu, J. Am. Chem. Soc., 2009, **131**, 9894-9895.
- 17 X. Lu, S. Francis, D. Motta, N. Dimitratos and A. Roldan, Phys. Chem. Chem. Phys., 2020, 22, 3883-3896.
- N. Gupta, O. Khavryuchenko, A. Villa and D. Su, 18 ChemSusChem, 2017, 10, 3030-3034.
- 19 M. S. Shafeeyan, W. M. A. W. Daud, A. Houshmand and A. Arami-Niya, Appl. Surf. Sci., 2011, 257, 3936-3942.
- 20 I. Barlocco, S. Capelli, X. Lu, S. Tumiati, N. Dimitratos, A. Roldan and A. Villa, Nanoscale, 2020, 12, 22768-22777.
- 21 J. Zhang, D. S. Su, R. Blume, R. Schlögl, R. Wang, X. Yang and A. Gajović, Angew. Chemie - Int. Ed., 2010, 49, 8640-8644.
- T. Hirashima and O. Manabe, Chem. Lett., 1975, 4, 259-22
- 23 G. R. Srinivasa, K. Abiraj and D. Gowda, .
- D. Cantillo, M. M. Moghaddam and C. O. Kappe, J. Org. Chem., 2013, 78, 4530-4542.
- 25 J. W. Larsen, M. Freund, K. Y. Kim, M. Sidovar and J. L. Stuart, Carbon N. Y., 2000, 38, 655-661.
- S. Fujita, H. Watanabe, A. Katagiri, H. Yoshida and M. Arai, J. Mol. Catal. A Chem., 2014, 393, 257-262.
- 27 I. Barlocco, S. Bellomi, S. Tumiati, P. Fumagalli, N. Dimitratos, A. Roldan and A. Villa, Phys. Chem. Chem. Phys., 2022, 24, 3017-3029.
- C. Gojon and B. Dureault, J. Nucl. Sci. Technol., 1996, 33, 28 731-735.
- 29 R. Arrigo, M. Hävecker, S. Wrabetz, R. Blume, M. Lerch, J. McGregor, E. P. J. Parrott, J. A. Zeitler, L. F. Gladden, A. Knop-Gericke, R. Schlögl and D. S. Su, J. Am. Chem. Soc., 2010, 132, 9616-9630.
- 30 G. Kresse and J. Furthmüller, Phys. Rev. B - Condens. Matter Mater. Phys., 1996, 54, 11169-11186.
- 31 G. Kresse, J. Non. Cryst. Solids, 1995, 192-193, 222-229.
- 32 B. Hammer, L. B. Hansen and J. K. Nørskov, Phys. Rev. B, 1999, 59, 7413-7421.
- 33 N. D. Mermin, Phys. Rev., 1965, 137, 1-3.
- 34 Y. Zhang and W. Yang, Phys. Rev. Lett., 1998, 80, 890.

View Article Online DOI: 10.1039/D3DT02310A

Dalton Transactions ARTICLE

35 R. Sure, J. Antony and S. Grimme, *J. Phys. Chem. B*, 2014, **118**. 3431–3440.

- 118, 3431–3440.
 S. Ehrlich, J. Moellmann, W. Reckien, T. Bredow and S. Grimme, *ChemPhysChem*, 2011, 12, 3414–3420.
- 37 R. Sundararaman and K. Schwarz, *J. Chem. Phys.*, , DOI:10.1063/1.4976971.
- 38 K. Mathew, R. Sundararaman, K. Letchworth-Weaver, T. A. Arias and R. G. Hennig, *J. Chem. Phys.*, , DOI:10.1063/1.4865107.
- J. D. Pack and H. J. Monkhorst, *Phys. Rev. B*, 1977, 16, 1748–1749.
- 40 Y. W. Tan, H. L. Stormer, P. Kim, K. S. Novoselov, M. L. Cohen, S. G. Louie, X. Wang, L. Zhang, S. Lee, H. Dai, Y. Kobayashi, K. Fukui, M. Fujita, G. Dresselhaus, M. S. Dresselhaus, M. A. Pimenta, B. R. A. Neves, A. Jorio, Y. Zhang, M. Mailman, P. M. Ajayan, S. K. Nayak, C. H. Park, Y. W. Son, S. P. Lu, S. Piscanec, A. C. Ferrari, G. Dobrik, P. Lambin, A. Oberlin, T. Solid, K. Suenaga, S. Iijima, P. Hermet, V. Meunier, L. Henrard, D. Gunlycke, C. T. White, S. Chen, B. I. Yakobson and S. Gradecak, *Science* (80-.)., 2009, 323, 1705–1708.
- I. Barlocco, L. A. Cipriano, G. Di Liberto and G. Pacchioni, J. Catal., 2023, 417, 351–359.
- 42 M. W. Chase and N. I. S. O. (US), *NIST-JANAF* thermochemical tables, American Chemical Society Washington, DC, 1998, vol. 9.
- L. A. Cipriano, G. Di Liberto and G. Pacchioni, ACS Catal.,
 2022, 12, 11682–11691.
- S. Sengodan, R. Lan, J. Humphreys, D. Du, W. Xu, H. Wang and S. Tao, *Renew. Sustain. Energy Rev.*, 2018, **82**, 761–780.
- 45 S. K. Singh, A. K. Singh, K. Aranishi and Q. Xu, J. Am. Chem. Soc., 2011, 133, 19638–19641.
- F. Tuinstra and J. L. Koenig, J. Chem. Phys., 1970, 53, 1126–1130.
- 47 H. Murphy, P. Papakonstantinou and T. I. T. Okpalugo, *J. Vac. Sci. Technol. B Microelectron. Nanom. Struct. Process. Meas. Phenom.*, 2006, **24**, 715–720.
- V. Datsyuk, M. Kalyva, K. Papagelis, J. Parthenios, D. Tasis,
 A. Siokou, I. Kallitsis and C. Galiotis, *Carbon N. Y.*, 2008, 46,
 833–840
- 49 S.-Y. Yang, C.-C. M. Ma, C.-C. Teng, Y.-W. Huang, S.-H. Liao, Y.-L. Huang, H.-W. Tien, T.-M. Lee and K.-C. Chiou, *Carbon N. Y.*, 2010, 48, 592–603.
- K. A. Wepasnick, B. A. Smith, K. E. Schrote, H. K. Wilson, S.
 R. Diegelmann and D. H. Fairbrother, *Carbon N. Y.*, 2011,
 49, 24–36.
- 51 C. H. Lau, R. Cervini, S. R. Clarke, M. G. Markovic, J. G. Matisons, S. C. Hawkins, C. P. Huynh and G. P. Simon, *J. Nanoparticle Res.*, 2008, **10**, 77–88.
- 52 X. Lu, S. Francis, D. Motta, N. Dimitratos and A. Roldan, *Phys. Chem. Chem. Phys.*, 2020, **22**, 3883–3896.
- 53 Q. Zhai, Z. Xia and L. Dai, *Catal. Today*, 2023, **418**, 114129.