Theoretical Insights into the Hydrogen Evolution Reaction on the Ni$_3$N Electrocatalyst

Russell W. Cross, Sachin R. Rondiya and Nelson Y. Dzade *

Abstract: Ni-based electrocatalysts are attractive alternatives to noble metal electrocatalysts for the hydrogen evolution reaction (HER). Herein, we present a dispersion-corrected density functional theory (DFT-D3) insight into HER activity on the (111), (110), (001), and (100) surfaces of metallic nickel nitride (Ni$_3$N). A combination of water and hydrogen adsorption was used to model the electrode interactions within the water splitting cell. Surface energies were used to characterise the stabilities of the Ni$_3$N surfaces, along with adsorption energies to determine preferable sites for adsorbate interactions. The surface stability order was found to be (111) < (100) < (001) < (110), with calculated surface energies of 2.10, 2.27, 2.37, and 2.38 Jm$^{-2}$, respectively. Water adsorption was found to be exothermic at all surfaces, and most favourable on the (111) surface, with $E_{ads}$ = −0.79 eV, followed closely by the (100), (110), and (001) surfaces at −0.66, −0.65, and −0.56 eV, respectively. The water splitting reaction was investigated at each surface to determine the rate determining Volmer step and the activation energies ($E_a$) for alkaline HER, which has thus far not been studied in detail for Ni$_3$N. The $E_a$ values for water splitting on the Ni$_3$N surfaces were predicted in the order (001) < (111) < (110) < (100), which were 0.17, 0.73, 1.11, and 1.60 eV, respectively, overall showing the (001) surface to be most active for the Volmer step of water dissociation. Active hydrogen adsorption sites are also presented for acidic HER, evaluated through the $AG_{H}$ descriptor. The (110) surface was shown to have an extremely active Ni–N bridging site with $AG_{H}$ = $-$0.05 eV.

Keywords: hydrogen evolution reaction (HER); nickel nitride (Ni$_3$N); water splitting; density functional theory (DFT)

1. Introduction

Hydrogen is fast becoming the best conceivable fuel to replace our damaging fossil fuel technologies. The transition to a hydrogen economy, however, requires efficient and economical ways of large-scale production of hydrogen to provide resources to all. Currently, steam reforming leads the way for industry; however, this is not a carbon-free process and results in the production of CO and CO$_2$ [1]. One promising method of obtaining hydrogen is through electrochemical water splitting [2,3]. This redox reaction breaks down to two half equations: the hydrogen evolution reaction (HER) and oxygen evolution reaction (OER). Pt electrodes are, at present, the best performing for HER; however, this expensive noble metal is not abundant enough to provide water splitting cells on a worldwide scale [4,5]. This introduces the urgent need for cheaper and more abundant alternatives. Much research has gone into using earth abundant transition metals to significantly reduce the cost of water splitting cells. Some promising materials have surfaced which have come close to the performance of Pt metals, such as transition metal sulphides (MoS$_2$ [6,7], WS$_2$ [8,9]), phosphides (FeP [10], CoP [11], Ni$_2$P [12]), (oxy)hydroxides (Ni(OH)$_2$ [13,14], CoO(OH)$_2$ [15]), and transition metal carbides (Mo$_2$C [16]). Transition metal nitrides (Mo$_2$N [17], TiN [18], Ni$_3$N [19]) have also been suggested recently, and much traction has grown with research of these materials.
Pure nickel catalysts such as Ni(111) have been shown to be active, although they do not present activities close to the benchmark Pt/C catalyst, but have prompted research into Ni-based subsidiaries [12]. Ni-based materials are strong candidates to replace expensive Pt for the hydrogen evolution reaction (HER) due to their low cost and strong corrosion resistance in alkaline media [20–24]. For instance, Strasser’s group recently derived dealloyed metal–oxide hybrid IrNi@IrOx core–shell NPs, which provide substantial advances towards more efficient and less expensive electrocatalytic water splitting [25]. Dai et al. also synthesized a NiO/Ni-CNT heterostructured catalyst, which showed good resistance to alkaline media, performing consistently for over 24 h [26]. Several Ni-hydroxies [27], oxides [28], phosphides [29], nitrides [30], sulphides [31,32], and selenides [33] have shown the capability of providing efficient output for full water splitting compared to the benchmark Pt- and IrO2-based electrocatalysts.

Nickel nitride (Ni3N) is one such example of a Ni-based material which has shown good HER characteristics, having consistent performance over 12 h of runtime and > 5000 cycles [34]. Although it has been shown to be promising in fields such as lithium-ion batteries [35,36], it was first tested for HER activity by Shalom et al., who found it exhibited an extremely low overpotential of 50 mV when supported on Ni foam [19]. It was suggested this activity could be due to a presence of Ni(OH)2 upon the surface. Ni3N has since been investigated in acidic media, with an H2SO4 electrolyte by Ding et al. [34] who used XPS to show satellite Ni–O bonds present on the catalyst surface. However, this was removed after etching of the surface, and XRD confirmed Ni3N was the major product phase. Ding et al. also studied Ni3N with DFT calculations and ascribed the inherent activity to the Gibbs free energy of hydrogen adsorption (ΔG*H*) values for hydrogen adsorption, whereby a value closer to zero is correlated to a higher activity catalyst. When a nanosheet with exposed (001) facets was tested, an optimal value of 0.065 eV was found for ΔG*H*.

Whilst ΔG*H* is a good descriptor for acidic environments, the initial Volmer step of adsorption is different when comparing with alkaline environments—it is instead water, which is dissociatively adsorbed onto the surface as OH* and H*, as opposed to a simple H adsorption [37,38]. Thus far, no studies have been dedicated to unravelling the mechanism of water adsorption and dissociation kinetics (the Volmer step) of HER on Ni3N surfaces. These insights are, however, essential to understand the HER activity of Ni3N and rationally engineer it to achieve improved HER performance. The present study aims to fill these gaps by investigating both molecular and dissociative water adsorption to the commonly observed low-index (111), (110), (001), and (100) surfaces of Ni3N [39,40]. Water adsorption is found to be exothermic atop all surfaces, binding via O–Ni interactions at all surfaces. From the water dissociation of the Volmer step, activation energies have been predicted, finding the (001) surface to be most active, whilst the (100) is the least. Hydrogen adsorption has also been investigated at all surfaces, with the most active sites for the Heyrovsky and Tafel steps presented.

2. Results and Discussion

2.1. Bulk and Surface Characterisation of Ni3N

Ni3N crystallised in the hexagonal Fe3N-type structure (Figure 1a) of the space group P6322. From full geometry optimisation, the unit cell parameters were predicted to be a = b = 4.571 Å and c = 4.262 Å, which were in good agreement with previous experimental and DFT studies [34,41]. The material contained Ni–N bonds of length 1.87 Å, which also corresponded well to experimental values of 1.89 Å [41]. From the partial density of states (PDOS) diagram (Figure 1b), it can be seen that Ni3N is a metallic conductor, with Ni-d orbitals dominating states at the Fermi level [34].
The low-index (111), (110), (001), and (100) Ni₃N surfaces (Figure 2) created from the fully relaxed bulk structure were optimised to ascertain their relative stabilities. The Ni₃N (111) and (110) are commonly observed through experimental syntheses and are therefore shown to be active for the hydrogen evolution reaction alongside the (001) surface [39,40,42]. The Ni₃N(100) surface is also presented, as its surface energy (\(\gamma\)) was calculated to be competitive with that of the (111) and (110) surfaces. The surface energies were found in the order of (111) < (100) < (001) < (110), with values of 2.10, 2.27, 2.37, and 2.38 J m\(^{-2}\), respectively.

The Ni/N surface ratio of the (111) surface was 5:1, leading to an electron-deficient surface. The (110) surface, although Ni-rich, contained low-coordination, adatom-like nitrogen sites. This site was, therefore, predicted to be heavily preferable for adsorption.
of water and hydrogen. Two terminations existed for the (001) surface, whereby the Ni-rich termination was chosen over the N-terminated, as previous studies have shown its increased activity in acidic conditions [34]. As such, only Ni sites were accessible across the slab. The Ni$_3$N(100) surface had a flat structure, and contained a higher concentration of N-sites than the (111), (110), and (001) surfaces, with a surface Ni/N atomic ratio of 2:1. Each Ni was in an equivalent position and bound to a single surface N.

2.2. Molecular Water Adsorption to Ni$_3$N(111), (110), (001), and (100) Surfaces

The most stable H$_2$O adsorption configuration for each surface is shown in Figure 3a–d. Water was found to preferentially adsorb to the Ni top site at all surfaces via the O atom. The ZPE-corrected adsorption energies, structural parameters including bond distances, bond angles, and O−H stretching vibrational modes for molecular water adsorption, are listed in Table 1. Water adsorption at the Ni$_3$N(111) surface was calculated to be exothermic, releasing $E_{ads}^o = -0.77$ eV, and contained a Ni−O bond length of 2.145 Å between adsorbed water and the surface site. Both O−H bonds remained close to values of molecular water at 0.983 and 0.973 Å. One O−H was translated towards a surface Ni to form a weak interaction, hence stabilising the system. This also caused a slight increase in the H−O−H angle from 104.5° to 105.6°. The stretching O−H modes of adsorbed water were also calculated to be 3818 and 3687 cm$^{-1}$, corresponding to the symmetric and asymmetric modes, respectively. A decrease in the O−H stretching modes compared to the gas phase H$_2$O molecule (3987 and 3865 cm$^{-1}$) also indicated that O−H bonds were somewhat activated. Bader charge analyses of the interacting surface site (q(Ni)) and of the adsorbed water (q(H$_2$O), Table 1) showed that the H$_2$O molecule transferred a charge of 0.031 e$^-$ to the surface upon adsorption. Consistent with the small charge transfers, our differential charge density isosurface analysis (Figure 3e–h) revealed only small changes in the surface electron density, with a small accumulation at one hydrogen. This is suggestive of weak hydrogen bonding interactions with the surface. Since the first and rate-determining step of HER is the removal of hydrogen from water (Volmer step), a strong interaction between hydrogen and the surface, and slight activation of water, may prove to increase activity.

![Figure 3. Lowest energy molecular water adsorption geometry on Ni$_3$N (111), (110), (001), and (100) surfaces (a–d), with corresponding ZPE-corrected adsorption energies (E$_{ads}^o$). Each image is displayed with isosurface below (e–h), where accumulation is in cyan, and depletion in yellow. The depletion/accumulation levels were ± 0.0224 eV/Å$^3$, as implemented in VESTA.](image-url)
Table 1. ZPE-corrected adsorption energies ($E_{ads}^0$), distance between water and surface Ni (Ni–O), O–H bond length in water, bond angles of water ($\phi$), and vibrational frequency of the symmetric and asymmetric stretch O–H stretches of water (where the molecular values were 3988 and 3866 cm$^{-1}$). Bader charges of the Ni active site before (clean) and after ($H_2O_{ads}$) adsorption are included, as well as the Bader charge of adsorbed water ($q(H_2O)$), where $q(H_2O)$ of molecular water was 0.

<table>
<thead>
<tr>
<th>Surface</th>
<th>$E_{ads}^0$ (eV)</th>
<th>d(Ni–O) (Å)</th>
<th>d(O–H) (Å)</th>
<th>$\phi$(H–O–H) (°)</th>
<th>$\nu$ (cm$^{-1}$)</th>
<th>$q$(Ni) (e$^-$)</th>
<th>$q$(H$_2$O) (e$^-$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(111)</td>
<td>−0.77</td>
<td>2.145</td>
<td>0.973, 0.983</td>
<td>105.6</td>
<td>3818, 3687</td>
<td>0.450</td>
<td>0.548</td>
</tr>
<tr>
<td>(110)</td>
<td>−0.62</td>
<td>2.047</td>
<td>0.981, 0.987</td>
<td>103.8</td>
<td>3744, 3640</td>
<td>0.214</td>
<td>0.354</td>
</tr>
<tr>
<td>(100)</td>
<td>−0.60</td>
<td>2.122</td>
<td>0.980, 0.982</td>
<td>104.1</td>
<td>3811, 3699</td>
<td>0.241</td>
<td>0.396</td>
</tr>
<tr>
<td>(001)</td>
<td>−0.56</td>
<td>2.085</td>
<td>0.983, 0.986</td>
<td>104.5</td>
<td>3660, 3532</td>
<td>0.184</td>
<td>0.340</td>
</tr>
</tbody>
</table>

Molecular adsorption of water on the Ni$_3$N(110) surface at the Ni site released an adsorption energy of −0.62 eV. The interacting Ni–O distance was calculated at 2.047 Å, along with the O–H distances at 0.987 and 0.981 Å and $\phi$(H–O–H) angle of 103.8°. The O–H stretching vibrational frequencies decreased to 3744 and 3640 cm$^{-1}$, indicating a weakening of O–H bonds. Water adsorbed to an N top site on the Ni$_3$N(001) Ni terminated surface, with an exothermic adsorption energy of $E_{ads}^0 = −0.56$ eV. The Ni–O bond distance was 2.085 Å, similar to geometries at the previously simulated surfaces. There was also minimal change at the O–H distances, which were predicted to be 0.983 and 0.986 Å, which was the same case for the H–O–H angle at 104.5°. It is worth noting that focus was on the Ni terminated surface due to the inability of the N terminated surface to adsorb water. A physisorbed state is preferred, yielding exothermic adsorption energies in the range 0.09–0.29 eV.

The adsorption of water to the Ni$_3$N(100) surface released an adsorption energy of $E_{ads}^0 = −0.60$ eV, with the interacting Ni–O distance predicted at 2.122 Å. The O–H bond lengths were not disturbed or changed from molecular water, remaining at 0.980 and 0.982 Å, and the $\phi$(H–O–H) angle remained similar (104.1°) to that of the gas phase. The symmetric and asymmetric O–H stretching modes, however, were slightly reduced to 3811 and 3699 cm$^{-1}$, respectively. The lesser extent of activation of the O–H bond at the (100) surface was likely to affect its water splitting ability.

2.3. Dissociated Water Adsorption on Ni$_3$N

From each molecular water adsorption system, we identified the preferred dissociative adsorption geometries (OH$_{ads} +$ H$_{ads}$) of water by moving one of the hydrogen atoms to an adjacent site. The most stable dissociative water adsorption geometries are shown in Figure 4 (FS, Figure 4i–l). At the Ni$_3$N(111) surface, the OH species remained adsorbed at the top Ni site, whereas the dissociated H atom preferentially bound to an Ni$_3$ site, releasing an adsorption of $−0.65$ eV (Figure 4a,e,i). The interacting Ni–O distance of the adsorbed OH species was reduced significantly from the molecularly adsorbed water (2.145 Å) to 1.839 Å, with the O–H distance calculated at 0.974 Å. The average Ni–H bond amongst the Ni$_3$ site was 1.700 Å. Overall, the dissociation process was slightly endothermic, with a reaction energy ($E_{rxn}$) of 0.14 eV. NEB calculations were performed to ascertain the transition state (Figure 4e–h, TS) along the minimum energy pathway to the dissociated state. Activation energy ($E_a$) barriers for the dissociative step were also found (Figure 5) and were adjusted by the ZPE as with the molecular water systems. The TS in each case was confirmed to have a single imaginary frequency.

$E_a$ for water splitting was found to be 0.73 eV at the (111) surface. Hydrogen diffused away from water, whereby the transition state (TS) had hydrogen midway between O and Ni, before reaching the Ni$_3$, hollow site (Figure 4e). Hydrogen was found to split to an adjacent N top site at the (110) surface (Figure 4i) and was likely favoured due to the low coordination of surface-N. O–H remained adsorbed at the top Ni site (Ni–O = 1.805 Å and O–H = 0.981 Å). The dissociated proton diffused to a N top site with a converged N–H bond distance of 1.024 Å. $E_{rxn}$ was calculated to be 0.07 eV, thereby showing molecular
adsorption to be slightly favoured and an endothermic path overall for the Volmer step. A higher activation barrier of 1.11 eV had to be overcome for this reaction to occur. The TS structure had hydrogen midway between oxygen and nitrogen, partially stabilised by an adjacent surface Ni atom.

Figure 4. Overall water splitting pathways: from (a–d) the initial molecular water adsorption to (e–h) the corresponding transition state, and (i–l), the final H + OH co-adsorption on each of the Ni$_3$N(111), (110), (001), and (100) surfaces.

Water dissociation on the (001) surface also led to an Ni$_3$ site for hydrogen diffusion, as with the (111) surface mechanism. This was due to the Ni-rich nature of the surface. An extremely stable binding site was found, with $E_{\text{rxn}}$ found to be highly exothermic in nature at $-0.64$ eV. $E_{\text{rxn}}$ was further reduced not just by the stability of the H site, but OH also moved to form a bridging site, hence becoming more stable compared to the final (111) and (110) structures. The activation barrier was also extremely small, with $E_a = 0.17$ eV. The low activation energy can be attributed to the inherent stabilization that occurred as hydrogen diffused to the binding site, with surface Ni acting to lower the energy of the transition state by interacting during diffusion. The Volmer step atop the Ni$_3$N(100) surface had the hydrogen atom adsorbed at a N top site, with an N–H bond length of 1.028 Å. The OH species adsorbed at a bridge Ni site as with the (001) mechanism (N–O bonds were 1.917 and 1.922 Å). Compared to the (111) and (110) surface, water dissociation on the (100) surface was also found to be exothermic ($E_{\text{rxn}} = -0.47$ eV) due to the strong interaction of OH at the bridge Ni site. Although the reaction was exothermic and thermodynamically favourable, it was kinetically limited due to the high activation barrier of 1.60 eV that had to be overcome. This suggests an extremely sluggish Volmer step ($H_2O + e^- \rightarrow OH^- + ...
H_{\text{ads}}) on the (100) surface, and thus poor HER activity, having the highest activation energy of the four surfaces.

Figure 4. Overall water splitting pathways: from (a–d) the initial molecular water adsorption to (e–h) the corresponding transition state, and (i–l), the final H^+ OH co-adsorption on each of the Ni_3N(111), (110), (001), and (100) surfaces.

2.4. Hydrogen Adsorption to Ni_3N Surfaces

Although the Volmer step is the first for HER in alkaline conditions, study of the consequential Tafel and Heyrovsky steps (combination of H^* to evolve H_2) are, however, important to gain insights into the overall HER activity. The Gibbs free energy of hydrogen adsorption ($\Delta G_{H^*}$) serves as a good descriptor for the HER activity, especially in acidic conditions [4]. Thermoneutral $\Delta G_{H^*}$ values close to 0 eV indicate that hydrogen will stick neither too strongly nor weakly, hence allowing for recombination to H_2 via the Tafel or Heyrovsky steps to be facile. The $\Delta G_{H^*}$ values at the (111), (110), (100), and (001) surfaces were obtained via the following equation: $\Delta G_{H^*} = \Delta E_{H^*} + \Delta ZPE - T\Delta S$, where $\Delta E_{H^*}$ is the 0 K, DFT calculated hydrogen adsorption energy, and $\Delta ZPE$ and $\Delta S$ are the zero-point energy and entropic corrections, respectively. Here, $\Delta ZPE - T\Delta S$ was estimated to be 0.24 eV, on the basis of previous studies, with T = 300 K [43].

The two most favourable structures for hydrogen adsorption at each surface were found and are shown in Figure 6. The only stable site found for the (001) surface was the Ni_3 site, due to the lack of available N atoms at the termination. Hydrogen adsorption at the Ni_3N(111) surface showed strong exothermic, with $\Delta G_{H^*} = -0.60$ eV at a top N site (Figure 6a) and $-0.83$ eV at the Ni_3 bridging site (Figure 6b). These two sites were highly unfavourable for HER as they may lead to a poisoning of the active sites when strongly adsorbed. The Ni_3 site is a poisoning site for HER in various transition metal-based materials, such as Ni_3P [12,43–46]. These values indicate poor activity for the Tafel step of HER due to the strong adsorption to the catalyst surface.
Heyrovsky steps can occur. The study was found to be the (001) surface; however, it was not suggested as a highly active anode in the water splitting cell due to the sub-optimal ΔG values of −0.83, −0.62, and −0.66 eV at the hydrogen resting spots. The lowest E values, calculated at −0.09 eV, as observed previously [34].

The Ni3N(111) surface, a low ΔGH* value was calculated at an Ni bridge site (−0.26 eV). This was vastly better than for the similar state at the Ni3N(111) surface. This was due to the reduced coordination number of the Ni site seen on the (111) surface leading to a stronger adsorption. An N top site was also tested; however, it converged to an N–Ni bridge site, with a highly optimal ΔGH* value of −0.05 eV. This was the most active site for acidic HER found across the four surfaces tested in this study. Due to the pure Ni nature of the Ni terminated (001) surface, hydrogen adsorption was limited to the favoured Ni3 site. This produced a strong binding site and low activity ΔGH* predictor of −0.66 eV, showing that although the Volmer step for the (001) surface was low, the activity was limited due to a possible poisoning of the site. Adsorption was also tested at the N terminated (001) surface, finding the extremely optimal Ni bridging site with ΔGH* = −0.09 eV, as observed previously [34].

The Ni3N(100) surface also yielded competitive ΔGH* values, calculated at −0.16 and −0.38 eV for the N top and Ni bridge sites, respectively. However, they did not compete with the ±0.10 eV value of platinum [47]. It is noteworthy that our studies here also accounted for the London dispersion forces, which have been neglected in previous studies, leading to a more accurate picture of the binding interaction between the surface and hydrogen [34,48].

From the calculation of the most favourable HER routes for alkaline HER, the final hydrogen positions for the Volmer step at the (111), (110), (001), and (100) surfaces were the Ni bridge (Figure 4i), N topsite (Figure 4j), Ni bridge (Figure 4k), and N topsite (Figure 4l), respectively. The hydrogen adsorption was found to be a severe limitation towards high HER activity on the (111), (110), and (001) surfaces due to the highly negative ΔGH* values of −0.83, −0.62, and −0.66 eV at the hydrogen resting spots. The lowest E2 value in this study was found to be the (001) surface; however, it was not suggested as a highly active anode in the water splitting cell due to the sub-optimal ΔGH* value, resulting in hydrogen sticking and poisoning the catalyst surface, hence reducing the rate at which the Tafel or Heyrovsky steps can occur.

These results suggest that efforts need to be made to engineer the surface properties of the Ni3N electrocatalyst in order to achieve improved HER activity. Recent experimental efforts have shown that substitutional doping of Ni with other transition metals could vastly improve HER activity by providing alternative hydrogen adsorption sites to the (001) surface [49–51]. The formation of heterostructures by coupling Ni3N with other materials, such as transition metal oxides [52,53] or nitrides [54,55], have also shown great promise. A NiMoN/Ni3N heterostructure synthesized by Wang et al. [55] was demonstrated to exhibit

Figure 6. Two most optimal free energy of hydrogen adsorption (ΔGH*) structures on each of the Ni3N surfaces. Where (a,b) are for the (111) surface, (c,d) for the (110), (e,f) for the (100), and (g) for the (001) surface.
an extremely low overpotential of 28 mV at 10 mA cm\(^{-2}\), owing to the synergistic effect between the two materials that favoured charge transfer processes for water dissociation.

3. Conclusions

In conclusion, we provide a comprehensive DFT mechanistic insight into the Volmer step of HER on the Ni\(_3\)N electrocatalyst through investigation of molecular and dissociative water adsorption on the (111), (110), (100), and (001) surfaces. Water was found to bind via the O atom to surface Ni sites in all cases. Adsorption was found to be exothermic for all surfaces; however, it was characterised by minimal interaction between the two, shown by our charge differential plots and small Bader charge values for H\(_2\)O: 0.031, 0.042, 0.045, and 0.027 e\(^-\) for the (111), (110), (100), and (001) surfaces, respectively. Activation energies for the rate-limiting Volmer step were found, and the Ni terminated (001) surface was shown to be the lowest at 0.17 eV, whilst it was found to be largest on the (100) surface, at \(E_a = 1.60\) eV. Hydrogen was found to split to a threefold Ni hollow site at the (001) site, whilst an adjacent N site for (100) was found. Furthermore, hydrogen adsorption was modelled at various Ni and N sites to yield valuable insight to the Tafel and Heyrovsky steps of the hydrogen evolution reaction. The resulting systems showed thermoneutral \(\Delta G^*\) values which were optimal for given HER applications, such as at the Ni–N bridge site atop the (110) surface, with \(\Delta G^* = -0.05\) eV. Large negative values, such as \(-0.83\) and \(-0.66\) eV for the Ni bridging sites atop the (111) and (001) surfaces, resulted in a sticking of hydrogen and poisoning of the catalyst. Hence, we found it to be a source of inactivity as hydrogen will proceed to stick at these sites. Heterostructure formation and transition metal doping have been shown to improve the HER activity of Ni\(_3\)N, and an ongoing DFT work aims to provide further detailed insight to the reasons behind these trends, as well as suggesting new combinations for facile hydrogen production.

4. Computational Details

Plane-wave-based DFT calculations were initiated through the Vienna ab initio Simulation Package (VASP) [56], using the Perdew Burke Ernzerhof (PBE) exchange correlation functional correction. A high energy cut-off of 600 eV and 10\(^{-6}\) eV difference between ionic steps of the self-consistency process was used to represent convergence. Grimme’s DFT-D3 scheme [57] was used to account Van der Waal’s dispersion forces across each system. When optimising bulk Ni\(_3\)N, we used a 5 \(\times\) 5 \(\times\) 5 Monkhorst-Pack K-point mesh to sample the Brillouin zone. For geometry optimisation of the surface slabs and the adsorbate surface systems, we utilized a 5 \(\times\) 5 \(\times\) 1 grid [58].

The METADISE code [59] was employed to obtain low miller index surfaces (111), (110), and (100) from the relaxed bulk material. For the (111), (100), and (001) surfaces, a (2 \(\times\) 2) slab was used to ensure no adjacent interactions between supercells in the x-y-plane, while a (2 \(\times\) 1) supercell was used for the (110) surface. Each surface contains a 15 Å vacuum region to ensure no interaction with periodic images in the z-direction, as well as a thickness of 10 Å to accurately mimic the catalytic material. The (001) surface was obtained from the bulk structure directly by adding a vacuum gap. For each surface termination, we obtained the surface energy (\(\gamma\)), which indicated the most stable arrangement for each surface. The following equation was used to calculate \(\gamma\):

\[
\gamma = \frac{E_{\text{surface}} - nE_{\text{bulk}}}{2A}
\]

where \(E_{\text{surface}}\) is the energy of the naked surface, \(n\) is the number of repeating unit cells in the z-direction, \(E_{\text{bulk}}\) is the energy of the bulk system, and \(A\) is the surface area of the relaxed system. \(\gamma\) (Jm\(^{-2}\)) indicates the energy required to cleave an infinite crystal to expose the termination; hence, a lower \(\gamma\)-value leads to a more favourable termination. The
following equation was used to determine the adsorption energies to the Ni$_3$N surfaces, whereby the ZPE was included:

$$E_{ads} = (E_{surface+adsorbate} + ZPE) - E_{surface} - (E_{adsorbate} + ZPE)$$

(2)

where $E_{surface+adsorbate}$ is the energy of the adsorbed water or hydrogen to the Ni$_3$N slab, $E_{surface}$ is the energy of the naked surface, and $E_{adsorbate}$ is the energy of the respective adsorbate in the gas phase. Therefore, a negative $E_{ads}$ (eV) value represents an exothermic, favourable adsorption, and vice versa.

Nudged elastic band (NEB) calculations were used to find transition states between molecular and dissociated water upon Ni$_3$N. For each of the (111), (110), (001), and (100) surfaces, the most favourable molecular and dissociated systems, indicated by the most exothermic respective $E_{ads}$ values, were taken as the start and end point with five images in between. The RMM-DIIS algorithm was then utilised, with a spring constant of $-5$ eV/Å between each image to find the transition state. Each was confirmed with vibrational frequency calculations as having a single imaginary frequency.

**Author Contributions:** Data curation, R.W.C.; formal analysis, R.W.C., S.R.R. and N.Y.D.; investigation R.W.C. and N.Y.D.; writing—original draft preparation, R.W.C.; writing—review and editing, S.R.R. and N.Y.D.; supervision, N.Y.D.; project administration, N.Y.D.; funding acquisition, N.Y.D. All authors have read and agreed to the published version of the manuscript.

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**Data Availability Statement:** Information on the data that underpins the results presented here, including how to access them, can be found in the Cardiff University data catalogue at http://doi.org/10.17035/d.2021.0136161101, accessed on 8 June 2021.

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