Chapter 6: Summary and Future Prospects

6.1 Summary

In recent times, the increasing energy requirements resulting from technological advancements and population growth, coupled with the diminishing availability of traditional fuels, have prompted a search for alternative options, particularly in the field of transportation. Hydrogen fuel stands out due to its abundance from various sources, eco-friendliness with minimal emissions and high energy density. However, using hydrogen faces hurdles: cost-effective production and compact storage. This thesis addresses challenges by examining advanced two-dimensional (2D) nanomaterials for hydrogen production through water-splitting and devising a viable compact hydrogen storage medium using state-of-the-art density functional theory (DFT) simulations.

Our study delves into hydrogen evolution reaction (HER) activity of pristine α -SiX (X=N, P, As, Sb, Bi), Ni anchored α -SiX (X=N, P, As, Sb, Bi) single-atom catalysts (SACs), pristine holey graphyne (HGY) and transition metals (Zr, Y, V, Sc, Mn, Co, Fe, Cr) anchored HGY SACs. This investigation is carried out through comprehensive dispersion corrected DFT simulations. Our results for the Gibbs free energy of H adsorption (ΔG_H) suggest that pristine form of α -SiX (X = N, P, As, Sb, Bi) and HGY monolayers are ineffective as catalysts for the HER activity. To enhance the HER activity, we have implemented a strategy of anchoring transition metals onto α -SiX (where X = N, P, As, Sb, Bi) and HGY monolayers in order to create SACs. Furthermore, we elucidate the alterations in the structural and electronic properties of α -SiX (X=N, P, As, Sb, Bi) and HGY monolayers due to anchoring with transition metals. This is achieved by considering orbital interactions, changes in band structures and partial density of states (PDOS) analysis. The significant bonding between Ni metal and α -SiX

(X = N, P, As, Sb, Bi) monolayers, as well as between transition metals and HGY monolayer, facilitated by p-d hybridization, enhances reactivity and conductivity. This is further attributed to a decrease in the band gap of the semiconducting α -SiX and HGY monolayers. Our DFT finding predicts that Ni@ α -SiX and Cr@HGY SACs emerge as highly efficient catalysts for HER, exhibiting remarkably low ΔG_H values of -0.04 eV and -0.05 eV, respectively. To validate their stability, we conducted extensive ab initio molecular dynamics (AIMD) simulations at 300K, affirming the integrity of Ni@ α -SiX and Cr@HGY SACs at room temperature. Given the exceptional electrochemical performance for HER activity and stability of Ni@ α -SiX and Cr@HGY SACs, they hold promise as viable catalysts for hydrogen production through the HER activity.

The dispersion corrected DFT simulations on pristine 2D orthorhombic diboron dinitride (o-B₂N₂) for HER activity show its limited suitability as a catalyst, as inferred from the Δ G_H calculation. However, the introduction of vacancy defects induces significant changes in the monolayer's structure and electronic properties, leading to the widening of the band gap from 0.66 eV to 0.99 eV in the case of the BN vacancy. Remarkably, the BN defected configuration exhibits Δ G_H close to zero compared to B and N defected as well as pristine configurations. This BN defect augmented HER activity by 77.34% in comparison to the pristine configuration at pH = 0. With the inclusion of C doping, the system's behavior transformed from a semiconductor to a metal. The Δ G_H of the C doped 2D o-B₂N₂ approached zero in both scenarios, signifying its suitability for HER. The C doping resulted in 86.71% boost in HER activity at the B site and 83.59% enhancement at the N site relative to the pristine configuration at pH = 0. This improvement was attributed to altered charge distribution following defect creation and C doping. Consequently, our findings propose that the metallic C doped 2D o-B₂N₂ could effectively act as an "electrocatalysts" for the HER activity within the pH range of 0 to 7. On the other hand, the BN defected 2D o-B₂N₂, characterized by 0.99 eV band gap, displays potential as a "photocatalyst" for HER activity at pH = 0. In addition, through AIMD simulations conducted at 300K, the structural integrity of 2D o-B₂N₂ with BN defects and C doping at room temperature is confirmed.

We explored the possibility of the pristine, defected and metals (including Na and K alkali metals; Be, Mg and Ca alkaline-earth metals and Sc, Ti, Y and Zr transition metals) decorated 2D o-B₂N₂ as a hydrogen storage medium through dispersion corrected DFT calculations. The findings indicate that the pristine, defected and metals Na, K, Be, Mg, Ca, Sc, Y and Zr decorated 2D o-B₂N₂ did not align favourably with the U.S. Department of Energy (DoE) requirements for an effective hydrogen storage medium. Nonetheless, promising results emerged from the Ti decorated 2D o-B₂N₂, suggesting its suitability as a hydrogen storage medium. Notably, the Ti atoms exhibited a strong and favourable bonding with the 2D o-B₂N₂, leading to a decrement in the band gap and an increase in conductivity. The adsorption energy for first hydrogen on the Ti decorated 2D o-B₂N₂ is -0.47 eV, satisfying the DoE's criteria. Furthermore, the charge donation from the Ti atom to the 2D $o-B_2N_2$ was observed through PDOS and Löwdin charge analysis. The two Ti decorated configuration demonstrated the capacity to bind a total of thirty-four hydrogen molecules, with average adsorption energy of -0.31 eV/H₂ and 396 K of desorption temperature. The calculated gravimetric storage capacity is 11.21%, notably surpassing the DoE's specified range. At a temperature of 396 K, our AIMD results convincingly demonstrate the excellent thermal stability of the Ti decorated 2D o-B₂N₂. As a summary, this research highlights the promising prospects of the Ti decorated 2D $o-B_2N_2$ as an efficient medium for high-capacity hydrogen storage.

6.2 Future Prospects

In this section, we will briefly discuss few future prospects. In connection with this thesis, the studied materials have not been examined under conditions such as strain (both expansive and compressive), transition metals (which are not addressed in the current thesis)

decoration or substitutional doping for applications in hydrogen production through water splitting and hydrogen storage medium. Moreover, a separate study could be conducted to predict the formation of various 2D allotropes of the studied materials by altering their chemical composition. This inspiration can be drawn from a thorough literature survey and state-of-the-art DFT simulations can be employed for this purpose. Exploring the landscape of 2D allotropes can uncover novel structures with distinct properties, potentially unlocking new avenues for applications beyond what is currently understood. In addition to the aforementioned aspects, the stable 2D materials that were examined within the scope of this thesis possess the potential to be further investigated for a diverse array of energy-related applications, such as solar cells and thermoelectric devices. Furthermore, their applicability extends to various other fields, including but not limited to bio-sensors and toxic gas sensors. The unique surface-to-volume ratio and high surface reactivity of 2D materials can enable highly sensitive and selective sensing platforms, revolutionizing the detection of biological molecules and hazardous gases. Beyond sensing applications, these materials could find utility in batteries and catalytic applications. Their high surface area and tailored electronic properties can contribute to improved energy storage and conversion, as well as catalytic processes like the oxygen evolution and oxygen reduction reactions in fuel cells. Hence, we hope that our theoretical data will serve as inspiration for experimentalists who aim to create cost-effective, high-performance HER catalysts for water-splitting and to design suitable materials for hydrogen storage applications. The integration of our insights with experimental efforts has the potential to drive innovation in the fields of renewable energy.