Communication

Dehydro-Sulfurization Utilizing Doubly-Charged Negative Ions of Molybdenum Disulfide (MoS₂), Graphene-Flake (GR-28), Armchair Carbon Nanotube (CNT 6,6), and Fullerene (C-60) as Catalysts for Depollution, Natural Resource Mining, and the Chemical Functionalization of Novel Drone Technologies Inspired by the Biomimicry of Bee Species' Bio-Catalysis of Pollen Conversion to Organic Honey

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Abstract: The Sulfur Dioxide (SO₂) compound is a primary environmental pollutant worldwide, whereas elemental Sulfur (S) is a global commodity possessing a variety of industrial as well as commercial functions. The chemical relationship between poisonous SO₂ and commercially viable elemental S has motivated this investigation using Density Functional Theory calculation of the relative transition state barriers for the 2-step Dehydro-sulfurization oxidation-reduction reaction. Additionally, doubly-charged nanoscale platelet Molybdenum Disulfide (MoS₂), Armchair (6,6) Carbon Nanotube, 28-atom Graphene nanoflake (GR-28), and Fullerene C-60 are utilized as catalysts. The optimal heterogeneous and homogeneous catalysis pathways of the 2-step oxidation-reduction from SO₂ to elemental S are further inspired by the biomimicry of the honeybee species multi-step bio-catalysis of pollen conversion to organic honey. Potential applications include environmental depollution, the mining of elemental sulfur, and the functionalization of novel technologies such as the recently patented aerial and amphibious Lynchpin TM drones.

Keywords: fullerene; graphene; molybdenum disulfide; carbon nanotubes; catalysts; density functional theory; enzymes; honeybee biomimicry; drones; depollution; doubly-charged negative ions

1. Introduction

Environmental depollution and mineral synthesis continue to be areas of international interest given the dynamics of an everchanging analog-to-digital-to-cloud economy, developing and maintaining sustainable green industrial processes, and future pandemic avoidance [1]. Specifically, elemental sulfur synthesis and SO₂ reduction reactions remain integral to present and future commercial applications as well as adherence to clean environmental protocols [2,3]. Herein, we use Density Functional Theory to investigate the 2-step dehydro-sulfurization reaction. The oxidation-reduction for this 2-step reaction entails increased water molecules between steps one and two with the product of elemental Sulfur remaining as the final solute, and molecular water remaining as the solvent within the resulting solution. Therefore, the reaction can efficiently serve a two-fold

function as an environmental cleaning agent; while simultaneously creating the commercially profitable elemental Sulfur as shown by equation 1 and 2 [4,5]. Transition metals such as gold, palladium, and silver have been shown previously to be highly inert in the bulk scale. In this theoretical work we overcome the tendency for the chemical inert behavior that is typical of the transition metals by investigating semi-metallic and metallic systems at the nanoscale versus bulk scale as well as exciting the nanoscale catalysts by an electrical charge equivalent to two negative charges. The introduction of doublycharged negative ions results in single and double bond breakage in the transition state, and bond reformation in the final state or product in the desulfurization redox reaction [6-8]. Hence, we show the intrinsic value of optimizing the 2-step dehydro-sulfurization reaction by the addition of two negative charges to the catalysts Molybdenum Disulfide, Graphene-nanoflake (GR-28), (6,6) Armchair Carbon Nanotube, and Fullerene (C-60). The varying of homogeneous and heterogeneous nanoscale catalysts in each of the two steps of dehydro-sulfurization reaction is motivated by the biomimicry of the fundamental role that the four different enzymes invertase, amylase, glucose oxidase, and catalase play in the bio-catalysis of sucrose conversion to organic honey. We further propose that this study may have future applications in the realm of aero-amphibious drone natural resource mining technologies [27,28,31-33].

2. Results

The calculations predict the chemical tunability of 2-step transition state paths of the doubly-charged heterogenous and homogenous nanoscale catalysts for the potential application to toxic SO₂ purification systems [1,25]. As shown in Table 1 and Table 2, we report Step 1 transition state (TS-1) values of 4.44eV, 1.12eV, 0.49eV, and 0.44eV for CNT (6,6), MoS₂, GR-28, and C-60 respectively. The Step 2 transition state (TS-2) values are 4.36eV, 1.06eV, 4.59eV, 4.30eV for CNT (6,6), MoS₂, GR-28, and C-60, respectively. The introduction of doubly-charged negative ions to the dehydro-sulfurization reaction tunes the transition state reaction barriers of metallic and semi-metallic catalysts yielding calculation results that are consistent with previous international academic and commercial findings [7]. Figure 1 predicts that each step of the 2-Step ORR is optimizable via the sequential heterogenous or homogenous utilization of the doubly-charged negative planar catalysts GR-28 for TS-1 and MoS2 for TS-2 with the calculated respective relative optimal values of 0.49eV and 1.06eV. However, a similar TS-1 barrier minimization is realized when spherical C-60 and planar MoS2 are applied sequentially and step-wise with TS-1 and TS-2 predicted to be 0.44eV and 1.06eV [22]. Alternatively, catalysis via the homogenous usage of CNT (6,6) is predicted to slow down the reaction speed with values of 4.44eV and 4.36eV reported for TS-1 and TS-2. The sole use of the heterogeneous catalyst MoS₂ is to minimize the transition states to 1.12eV and 1.06eV for TS-1 and TS-2. These calculations support the application of the short-lived ion creation fundamental mechanism to multistep chemical reactions. The transition state barriers presented here suggest efficacy for reaction barrier optimization or "chemical tuning" for future depollution and elemental S acquisition applications [18,24,26].

The desulfurization reaction mechanism may serve as the fundamental process that mimics honeybee bio-catalysis. Moreover, the focus of this work is desulfurization that entails toxic SO₂ elimination with the simultaneous generation of elemental Sulfur commodity rather than organic honey production. It is interesting to note that the mechanism for the biochemical production of honey is documented as an approximated 4-step biocatalytic process involving the four primary enzymes invertase, amylase, glucose oxidase, and catalase as presented in figures 8-10[30,31]. The catalysts of study are analogously tailored to the 2-step desulfurization reaction in lieu of SO₂ elimination with simultaneous generation of elemental Sulfur thereby mimicking the varying of enzymes per each step of the bio-catalysis of organic honey.

This section may be divided by subheadings. It should provide a concise and precise description of the experimental results, their interpretation, as well as the experimental conclusions that can be drawn.





Figure 1. Conceptual rendering of the theoretical transition state 2-step reaction path for the dehydro-sulfurization reaction. The red curve indicates net chemical results from equation 1 and 2 where the activation energies (E_a) and (E_a') are the 2-transition state (TS) barriers calculated by the subsequent equation E_a or $\Delta TS = TS$ -initial state for barrier height values for each step represented by the green and dark red arrows respectively. The purple arrow represents the intermediate formation of 2 molar sulfur dioxide.



Figure 2. Calculated relative transition state barrier paths values for the 2-Step Oxidation-Reduction Reaction (ORR) of Sulfur dioxide by utilizing Armchair (6,6) Carbon Nanotube, Molybdenum Disulfide (MoS₂), 28-atomed graphene nanoflake (GR-28), and Fullerene (C-60) as catalysts represented by the inset purple, green, blue, and orange bars, respectively. The black dashed line identifies the optimal catalytic path for the transition states 1 and 2. The relative transition state barriers (Δ TS-1) and (Δ TS-2) for respective Steps 1 and 2 are defined by equation (3): Δ TS=Transition State – Initial State.

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Figure 3. Calculated relative initial, transition, and final state reaction barriers of Step-1 and Step-2 for the oxidation reduction of Sulfur dioxide to elemental Sulfur Utilizing Armchair Carbon Nanotube CNT (6,6) represented by the purple bars. Pink Inset: Geometrically optimized Final state 1 (Final-1) of SO₂ covalently bonding to CNT (6,6) surrounded by 2 water molecules. Sulfur, Carbon, Oxygen, and Hydrogen are represented by the yellow, gray, red, and white spheres, respectively.



Figure 4. Calculated relative initial, transition, and final state reaction barriers of Step-1 and Step-2 for the oxidation reduction of Sulfur dioxide to elemental Sulfur by utilizing the doubly-charged catalyst MoS₂ represented by the dark green bars. White Inset: Geometrically optimized Final State 2 (Final-2) of MoS₂ with noncovalent precipitated S₂ and H₂O. Sulfur, Carbon, Oxygen, and Hydrogen are represented by the yellow, gray, red, and white spheres, respectively.



Figure 5. Calculated relative initial, transition, and final state reaction barriers of Step-1 and Step-2 for the oxidation reduction of Sulfur dioxide to elemental Sulfur (S) by utilizing doubly-charged catalyst graphene flake (GR-28) represented by the blue bars. Light Blue Inset: Geometrically optimized Final State 2 (Final-2) of GR-28 with covalent bonded atomic S surrounded by 4 molar H₂O. Sulfur, Carbon, Oxygen, and Hydrogen are represented by the yellow, gray, red, and white spheres, respectively.



Figure 6. Calculated relative initial, transition, and final state reaction barriers of Step-1 and Step-2 for the oxidation reduction of Sulfur dioxide to elemental Sulfur (S) by utilizing doubly-charged catalyst fullerene C-60 represented by the brown bars. Light Purple Inset: Geometrically optimized Final State 2 (Final-2) of the C-60 catalyst with [2+1] cycloaddition elemental S to fullerene C-C bonds and 4 molar H₂O as the final product for Step-2. Sulfur, Carbon, Oxygen, and Hydrogen are represented by the yellow, gray, red, and white spheres, respectively.



Figure 7. Calculate initial, transition, and final states for geometrically optimized doubly charged AA stacked MoS₂ catalyzing the oxidation reduction of sulfur dioxide to elemental Sulfur. The Mo, S, O, and H atoms are represented by cyan, yellow, red, and white spheres, respectively. The curved arrows demonstrate the introduction of 2 electrons into the reactant states causing bond breakage in the predicted transition states of Step 1 and Step 2 which describes the doubly-charged catalytic process used for our calculations.



Figure 8. Bee biomimicry inspired rendering of both Steps of the initial, transition, and final states for geometrically optimized doubly- charged fullerene C-60 and AA stacked MoS₂ heterogeneously catalyzing the oxidation reduction of sulfur dioxide to elemental Sulfur. The Mo, S, O, and H atoms are represented by cyan, yellow, red, and white spheres, respectively. The curved arrows demonstrate the introduction of 2 electrons into the reactant states causing bond breakage in the predicted relative transition states of Step 1 and Step 2.



Figure 9. The biocatalytic step 1 conversion of the Sucrose molecule ($C_{12}O_{11}H_{22}$) contained in nectar and pollen to intermediate glucose ($C_6O_5H_{11}$) and fructose molecules ($C_6H_{12}O_6$). The orange arrow indicates the utilization of the enzyme invertase as the biocatalytic promoter of the reaction. The blue arrows indicate flow of two electrons that cause bond breakage and reformation. The gray, red, and white spheres represent Carbon, Oxygen, and Hydrogen, respectively.



Figure 10. The Step 2 and Step 3 mechanisms for glucose intermediate decomposition to glucose and the oxidation of glucose ($C_6H_{12}O_6$) to gluconic acid ($C_6H_{11}O_6$) and hydrogen peroxide. Step 2 uses the enzyme amylase whereas Step 3 has the enzyme glucose oxidase as the promotor of the reaction. The index "1" is used to indicate the position of the C=O double bond throughout the mechanism. The gray, red, and white spheres represent Carbon, Oxygen, and Hydrogen, respectively.

BIOCATALYTIC STEP 4



Figure 11. The Step 4 of the final step of the sucrose conversion to organic honey. The enzyme catalase decomposes 2 molar hydrogen peroxide to 2 molar water and diatomic oxygen. The red and white spheres represent Oxygen and Hydrogen, respectively.

2.3. Tables

Table 1. Direct comparison of the calculated relative transition state barrier differential heights from initial states (Δ TS =TS-initial state) for Step 1 and 2 of the oxidation reduction reaction to atomic Sulfur are represented by Δ TS-1 and Δ TS-2 with electro-volts (eV) as the representative units. Arm-chair (6,6) carbon nanotube (CNT (6.6)), Molybdenum disulfide (MoS₂), graphene flake containing 28 atoms (GR-28), and fullerene (C-60) are the doubly-charged catalysts.

Catalyst	ΔTS-1	ΔTS-2
(Doubly-Charged)	(eV)	(eV)
CNT (6,6)	4.44	4.36
MoS ₂	1.12	1.06
GR-28	0.49	4.59
C-60	0.44	4.30

Table 2. Calculated relative initial, transition, and final states of Step 1 and Step 2 of the desulfurization redox reaction with electro-volts (eV) as the representative units. The relative states have been calibrated to zero for the lowest calculated potentials that were found in Step 1 "or" Step 2 for Armchair (6,6) carbon nanotube, Molybdenum disulfide (MoS₂), graphene flake containing 28 atoms (GR-28), and fullerene (C-60) doubly-charged catalysts.

Catalyst	Initial 1	TS-1	Final 1	Initial 2	TS-2	Final 2
(Doubly-Charged)	(eV)	(eV)	(eV)	(eV)	(eV)	(eV)
CNT (6,6)	0	4.44	15.34	8.89	4.36	5.68
MoS ₂	10.09	11.21	10.16	0	1.06	0.95
GR-28	10.62	11.10	7.05	0	4.59	10.29
C-60	0	0.44	2.31	11.92	4.30	9.53

3. Discussion

The dehydro-sulfurization reaction is an oxidation reduction mechanism also referred to as redox reaction. The reactant includes the introduction of diatomic oxygen into sulfur dihydride. Our research approach further introduces 2 electrons into the chemical reaction in order to increase the catalytic reactivities of heterogeneous (moly-disulfide) MoS2 as well as the homogeneous molecules (6,6) armchair carbon nanotube, fullerene (C-60), and graphene (Gr-28). The resulting product is sulfur dioxide precipitated in 2 molar waters due to bond breakage and reformation as indicated by equation 1. Additionally, 2 more electrons are added to the catalysts in order to further break the bond arrangements of the reactant in step 2 containing sulfur dihydride and 2 molar sulfur dioxides. Hence, the oxygen is removed from the sulfur dioxide due to bond breakage to form 3 molar diatomic sulfurs and 4 molar waters in the final product of step 2. The addition of doubly-charged negative ions to the spherical, cylindrical, and planar nanoscale catalysts function as a "chemical tuning" agent for each step of the dehydro-sulfurization reaction [26, 27]. Moreover, the varying of the catalysts with each step of the redox reaction mimics the biocatalytic mechanism of sucrose conversion to organic honey through the flow of 2e- electrons that similarly cause bond breakage and bond reformation as shown in figures 8-10. The relative barriers generated by the 2e- electrons addition are compared with reference to the initial states according to equation 3. The 2-step chemical reaction can therefore be tailored with various levels of control at each step of the reaction by the sequential application of homogeneous and/or heterogeneous catalysts as indicated by figures 1-7.

Equations:

Catalyst (-2)

1) $2H_2S + 3O_2 = 2SO_2 + 2H_2O$ **STEP 1**

Catalyst (-2)

2) $4H_2S + 2SO_2> 3S_2 + 4H_2O$ STEP 2	2)	$4H_2S + 2SO_2>$	$3S_2 + 4H_2O$	STEP 2
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- 3) $6H_2S + 3O_2$ -----> $3S_2 + 6H_2O$ ("Net" Chemical Results for the Reaction)
- 4) $\Delta TS = Transition State Initial State (Relative T.S. Reaction Barriers)$

4. Materials and Methods

[2+1] Cyclo-addition is a typical phenomenon that occurs in organo-metallic catalytic systems, oxidation reduction reactions, and nitrene chemistry. In this work, [2+1] Cycloaddition is specifically used in the geometrical optimization of fullerene C-60 as shown in the inset of Figure 5[4,5,8,9]. However, 28-atomed graphene (GR-28), Armchair Nanotube (6,6), and MoS₂ catalysts are geometrically optimized by covalent and noncovalent bonding as indicated in the insets of Figures 2-5 and the entirety of Figure 6[4-6]. The doublycharged MoS₂, GR-28, CNT (6,6), and C-60 catalysis substrates function as reaction barrier tuning mechanisms for the desulfurization oxidation reduction reaction (ORR) as indicated in equations 1 and 2[10-13]. Oxidation-reduction speed is therefore predicted to be regulated by the size, type, and charge of any given sample of catalysts used to produce the elemental Sulfur and water products for both steps of the reaction. Hence, the quality, quantity, and overall efficiencies of the reactions can be tailored to the reactivity of each step of the ORR reaction as well as the type of nanoscale catalyst that is used [5,8]. Transition state theory is executed by performing Density Functional Theory approximations shown to be excellent for predicting chemical, physical, and electromagnetic properties of nanoscopic systems. The initial, transition, and final states are optimized, and are dependent on the application of two negative charges that activate the metallic and semi-metallic planar, cylindrical, and spherical catalysts that cause single and double bond breakage in the transition state consistent with the research findings [4-9, 20]. The GR-28 nano-flake configuration investigated is due to its similar geometrical structure and chemical function as the "Sulflower" discussed in the 2017 "MOLECULES OF THE YEAR" edition of the C&EN journal [6-8,21]. Geometrical optimization calculation efficiency for initial, transition, and final states is achieved, as indicated by the data in Table 2, with the usage of minimum basis sets and Self-Consistent Field (SCF) tolerances of 0.01Ha, and smearing values of 0.05Ha. We use Equation (3) in order to specifically compare the relative Transition State (Δ TS) values in Table 1 in order to highlight the most likely efficient heterogenous or homogenous catalyst configuration as indicated in Figure 1[17,18]. Whereas,

Figures 2-5 show that the calculated relative reaction barriers for the initial, transition, and final states are zeroed to the minimum potential for initial state calculated values occurring in Step 1 or Step 2 for each set of nanoscale catalysts. A 3-D triclinic lattice type with lattice lengths a:12.7548, b:12.7548, and c:17.2744 Angstroms with subsequent respective angles consisting of α :90, β :90, μ : 120 degrees is used for the AA stacked MoS₂ catalyst. The values of the MoS₂ are represented by a 1x1x1 Monkhorst-Pack grid as shown in the inset of Figure 3 as well as the entirety of Figure 6. Noncovalent bonding is used to predict TS-1 and TS-2 for MoS₂ versus the Cyclo-addition and covalent approaches utilized for CNT (6,6), GR-28, and C-60[17-23].

5. Conclusions

The transition states for spherical C-60, planar MoS₂ and graphene (Gr-28), and CNT (6,6) tubal shaped have been calculated. The results indicate efficacy for the chemical tuning of the dehydro-sulfurization reaction as inspired by natures biocatalytic solution to the biocatalytic conversion of pollen or nectar to organic honey. We further propose that our findings may assist in the understanding and development of future innovative technologies by utilizing tangential flight enabled amphibious Lynchpin [™] drones as patented and developed by T. Dashawn Howard et.al [28]. Some unique capabilities of the patented drone technology include aerial bird-like flock, bee swarm, and "school" of fish biomimicry that is tailorable to future multidisciplinary engineering solutions. For example, we postulate that the implementation of our doubly-charged negative-ion catalytic desulfurization predictions combined with the Lynchpin [™] drone design may offer promising SO₂ depollution remedies as well as the profitable commercial mining of elemental Sulfur [1,2,14-18,23,28-34].

Supplementary Materials: Not Applicable

Author Contributions: For research articles with several authors, a short paragraph specifying their individual contributions must be provided. The conceptualization, methodology, writing, validation utilizing Accelyrs® software were performed by K.S. Formal analysis, review, and editing were executed by K.S., D.S., and A.M. Supervision and funding acquisition were provided by A.M. All authors have read and agreed to the published version of the manuscript.

Funding: This work was partly supported by U.S. DOE, Division of Chemical Sciences, Office of Basic Energy Sciences, Office of Energy Research

Data Availability Statement: Not applicable.

Acknowledgments: The authors express gratitude to T. Dashawn Howard and his research team for their fruitful discussions on the potential future innovations of their Lynchpin[™] drone technologies.

Conflicts of Interest: The authors declare no conflict of interest." Authors must identify and declare any personal circumstances or interest that may be perceived as inappropriately influencing the representation or interpretation of reported research results. Any role of the funders in the design of the study; in the collection, analyses or interpretation of data; in the writing of the manuscript; or in the decision to publish the results must be declared in this section. If there is no role, please state "The funders had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; or in the decision to publish the results of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript; or in the decision to publish the results".

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