The Reactions of Carbon Monoxide Catalytic Oxidation on Ti and Zr-embedded Graphene, a DFT Study

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Density functional theory (DFT) calculations have been performed to explore the mechanistic aspects of the catalytic oxidation of CO over Ti and Zr-embedded graphene. The present results show that the title reactions start with the activation of an oxygen molecule as: O₂ → O_ads. The CO oxidation over the catalyst surface proceeds through the following elementary steps: (a) O_ads + CO → CO₂ + O_ads; (b) O_ads + CO → CO₂. Both the Ti and Zr-embedded graphene show good catalytic activity toward the CO oxidation via the ER mechanism with a three-step route. The present results may be helpful in understanding the mechanism of CO oxidation over metal-decorated graphene and further experimental design of low-cost catalyst in CO emission.

Keywords: DFT, NBO, titanium, zirconium, graphene.

1. INTRODUCTION

As one of the main poisonous gases from the combustion of fuel, vehicles and industrial process, carbon monoxide pose a serious threat to the health of living organisms and ecosystems. Nowadays, it has become one of the most challenging subjects to monitor and remove this toxic gas from industrial and domestic emissions quickly and efficiently. In the past many years, considerable attention has been paid to convert this species into nontoxic CO₂. Catalytic oxidation is one of the effective routes to remove this undesired hazardous molecule [1]. In previous years, some types of noble metals such as Pd, Pt, Ru, Rh, Ag, Au were used as effective catalysts for CO oxidation [2–12], but the high cost limits their widespread use in practical application. According to the recent studies, much interest has been focused on the experimental and theoretical studies of metal-free and non-noble metal-embedded graphene, which shows better performance for the CO oxidation but with a relatively lower cost [13–24], indicating such complexes have potential for practical application in oxidation of CO. Li et al. have investigated the catalytic oxidation of CO over Iron-embedded graphene by means of first-principles computations [15], they found the Fe-embedded graphene show good catalytic activity for the CO oxidation via the more favorable Eley-Rideal (ER) mechanism. Limtrakul et al. have theoretically investigated the oxidation of carbon monoxide using nitrous oxide as an oxidizing agent on Fe-graphene [16], their results show the high catalytic activities of Fe-embedded graphene toward CO oxidation, and the graphene sheet play an important role during the reaction progress. Esrafili et al. have examined the reaction of CO oxidation by O₂ over Al-, Si- and Ge-embedded graphene [18], their DFT calculations indicate that for the CO oxidation, the ER mechanism to be the most favorable channel, and all these materials can be used as excellent and low-cost catalysts due to relatively high catalytic ability for the CO oxidation.

As typical first and second-row early transition metals, titanium and zirconium have good catalytic activity toward many types of reactions. From our previous theoretical study [25–28], we found the naked titanium cation has strong catalytic performance of activating the C–S and C–O bonds in SCO molecule [25], zirconium atom shows good ability toward the activation of C–O, C–H, and C–C bonds in carbon dioxide, acetone and acetaldehyde [26–28]. Recently, Krasheninnikov et al. presented a DFT study of transition-metal atoms embedded in vacancy in a graphene [29], they found the bonding is very strong and the migration barrier of metal atoms on a single vacancy of metal-embedded complexes are relatively high. The metals-embedded graphene would be stable enough to be utilized in catalysis applications. To the best of our knowledge, there is no relative report on the mechanisms of the CO oxidation reaction over Ti and Zr-embedded graphene. Therefore, the research on CO oxidation over typical early metal-embedded graphene has important theoretical and experimental significances. The calculated results are expected to forecast further experimental findings and to provide some valuable reference for designing of graphene-based catalysts for monitoring and removing toxic gases.

2. METHODOLOGY

To understand the detailed reaction mechanism at the molecular level, quantum chemical calculations were performed in this study. All the geometries involved in the reaction were optimized by employing the B3LYP density functional theory method as implemented in the Gaussian09 program package [30, 31]. In all calculations, the effective core potentials (ECP) of Stuttgart basis set
was used for the zirconium, the 5s and 4d in zirconium were treated explicitly by a (8s7p6d) Gaussian basis set contracted to (6s5p3d) [32]. For Ti and non-metal atoms, a standardized 6-31G(d) basis set was used. Previous investigation proved that this method is an effective way to explore the graphene properties [33]. The nature of the stationary point was verified through a vibrational analysis (no imaginary frequencies). A hexagonal graphene super cell (4x4 graphene unit cell) containing 48 carbon atoms was chosen as the basic model for the calculations. A similar model used in previous theoretical study by Esrafil et al. has got satisfied results [18]. The metal-embedded graphene was then modeled by substituting a single metal atom with one carbon atom on the surface. The adsorption energy \( E_{\text{ads}} \) is calculated as:

\[
E_{\text{ads}} = E_{\text{adsorbate-substrate}} - (E_{\text{adsorbate}} + E_{\text{substrate}}),
\]

where a negative \( E_{\text{ads}} \) indicates an exothermic process. Electron density difference (EDD) was conducted using MultiWFN program package [34].

\section*{3. RESULTS AND DISCUSSION}

\subsection*{3.1. Ti and Zr-embedded graphene}

When the metal atom is placed above a vacancy in graphene, it moves outward the plane of graphene due to its relatively larger atomic radius than the carbon atom after geometry optimization. As it is clear from the structure of Ti and Zr-embedded graphene (IM0) in Fig. 1, the average bond length of metal atom with its neighboring carbon atoms are 1.928 and 2.039 Å respectively, which are about 0.518 and 0.629 Å larger than the average C–C bond length in the pristine graphene.

![LUMO](image1.png)  
**Fig. 1.** a – frontier molecular orbits of the Ti-embedded graphene; b – frontier molecular orbits of the Zr-embedded graphene

The calculated adsorption energy of the metal atom over the vacancy site of the graphene are \(-145.4\) and \(-179.8\) kcal/mol for Ti and Zr respectively, indicating a strong interaction between the dopant atoms and the surface. To gain deeper insight into the significant enhancement of metal adsorption at the graphene vacancy site, we plotted the frontier molecular orbits of IM0. The calculated electronic structures of the molecular orbits are shown in Fig. 1. Both the HOMO and LUMO locate on the metal atom dominantly, the electron clouds of the lowest unoccupied molecular orbits (LUMO) for the compound illustrate the strong d-orbital characteristics of metal atom allowing an incoming electron to occupy this state. All of these observations demonstrate that the metal atom might be the activate site for adsorbing an electrophile probe molecule. To further analyze the intermolecular interactions, the geometries of metal-embedded graphene were used to perform NBO analysis [35]. The result shows that about 1.236 and 1.203e (for Ti and Zr respectively) are transferred from metal atoms to the adjacent carbon atoms that lead to the strong bonding between metal atoms and their neighbors, due to the electronegativity difference between metal and carbon atoms. The strong electron interaction between metal atoms and surface moiety facilitate the transfer of electron in metal-embedded graphene composite. The NBO result indicates that the graphene sheet act as an electron withdrawing support. In addition, this can suggest that both the Ti and Zr atoms are activated to be more electrophilic by their support. For metal-embedded graphene composite molecule the calculated MESP map displays the electrophilic region near the metal center. The more positively charged metal atoms are expected to have a stronger interaction with the adsorption gas molecule. In the next section, the adsorption behaviors of \( O_2 \) and CO on the metal-embedded graphene and the catalytic oxidation processes for the CO will be discussed.

\subsection*{3.2. Adsorption of \( O_2 \) and CO molecules over Ti-embedded graphene}

Previous studies have been established that there are two possible mechanisms for CO oxidation, namely the Eley-Rideal (ER) mechanism and Langmuir-Hinshelwood (LH) mechanism [15]. In the ER mechanism, the \( O_2 \) molecule is firstly absorbed over the catalyst surface, then the CO approaches the activated \( O_2 \) to form \( CO_2 \). This reaction mode is quite different from the later, as in the LH mechanism, both \( O_2 \) and CO molecules co-adsorb on the surface and to generate \( CO_2 \) directly. But many previous studies indicate that it is almost impossible for the CO oxidation over metal-embedded graphene through LH mechanism. So, in the present study we emphasize the ER mechanism only.

First, we will discuss the reaction over Ti-embedded graphene. The adsorption structures on the Ti-embedded graphene are described in Fig. 1a. For \( O_2 \), the most energetically favorable configuration (IM1) is characterized by \( O_2 \) parallel to the graphene surface forming two chemical bonds with the Ti atom, the calculated adsorption energy is \(-56.19\) kcal/mol. According the NBO analysis, about 0.759e charge is
transferred from Ti-embedded graphene to O₂, which could occupy the antibonding π* orbital of O₂ and subsequently lead to the elongation of the O−O bond from 1.215 to 1.477 Å. The electron density difference (EDD) plots of the adsorbed configurations are depicted in Fig. 2, in which the full and dotted lines are related to the electron density accumulation and depletion areas respectively.

The EDD plot shows the charge transfer from the Ti-embedded graphene to the O₂, and the accumulation of electron density at the two Ti−O bonds. The EDD plot agrees well with the result of NBO analysis. With respect to the CO, its adsorption behavior over Ti-embedded graphene plays an important role in the following oxidation reaction. Calculation indicates the CO can also be adsorbed on the Ti-embedded graphene easily in the OC−Ti adsorbing direction with the adsorption energy of −20.63 kcal/mol, the calculated distance between Ti and C is 2.240 Å, and only 0.044e charge is transferred from CO to the surface. The above results reveal that both O₂ and CO have strong interactions with Ti-embedded graphene, but it is clear that O₂ has even more favorable binding energy. So we believe that if a mixture of O₂/CO is injected as the reaction gas, the Ti site will be covered by the adsorbed O₂ molecule dominantly.

3.3. Oxidation of CO by O₂ over the Ti-embedded graphene

The reaction mechanism is proposed in stepwise reactions, which are the activation of O₂ and followed by the oxidation of CO as presented in Eq. 2 – Eq. 4:

The adsorption and activation of O₂:

\[ \text{O}_2 \rightarrow \text{O}_{2\text{act}}; \quad (2) \]

The adsorption and oxidation of CO:

\[ \text{O}_{2\text{act}} + \text{CO} \rightarrow \text{CO}_2 + \text{O}_{\text{ads}}; \quad (3) \]
\[ \text{O}_{\text{ads}} + \text{CO} \rightarrow \text{CO}_2. \quad (4) \]

The activation reaction starts with the formation of the O₂-bound encounter complex IM1, which is −56.19 kcal/mol below the entrance channel IM0 + O₂. Starting from IM1, it can rearrange to form IM2, which undergoes a rupture of O−O bond via a transition state TS₁₂ that is 27.65 kcal/mol above IM1. As shown in Fig. 1a, the distance between two oxygen atoms is elongated by 0.587 Å. This fact indicates that the weak electrostatic interaction between Ti and O₂ has strengthened when it is converted into TS₁₂. The distances between Ti atom and two oxygen atoms are shortened from 1.850 to 1.720 Å and 1.827 to 1.778 Å respectively, which suggests that the two Ti−O bonds is forming. Simultaneously, the activated O−O bond is almost broken, and the bond length is elongated to 2.064 Å. The imaginary frequency of TS₁₂ is −504.1i cm⁻¹, and the normal mode corresponds to the rupture of O−O bond with the result of Ti atom inserting into O−O bond. The inserted complex IM2 is −53.79 kcal/mol lower in energy than the reactants (see Fig. 3 a). As shown in Fig. 1a, the O−O bond length in IM2 is elongated to 2.670 Å, which means this bond has ruptured thoroughly. The large amount of energy gained in the formation of the IM1 can drive the cleavage process complete easily. Clearly, the Ti-embedded graphene can activate adsorbed oxygen effectively. From the NBO analysis, during the reaction process from IM1 to IM2, about 0.576e negative charge on graphene is transferred to Ti atom and oxygen molecule. In IM2, the electron transferring from the Ti-embedded graphene increases the negative charge on the two adsorbed oxygen atoms from −0.759 to −1.175e, at the same time, the positive charge on the Ti atom is decreased from 1.257 to 1.097e. Obviously, from IM0 to IM2, the oxygen atom pulls the charge from the surface owing to its strong electronegativity, and the Ti-graphene substrate acts as the electron donor during the process of oxygen activation, while the titanium atom acts as the bridge of transferring electron due to only a negligible charge (0.139e) is transferred on it from IM0 to IM2.

As shown in Fig. 3 b, the first oxidation reaction channel starts with the formation of an encounter complex IM3, along this reaction pathway, the carbon atom can insert into the activated O−O bond. From Fig. 1 a, one can see the distances between the carbon atom and the two oxygen atoms in IM3 are 1.366 and 1.352 Å respectively, this indicates that the two C−O bonds are forming. Simultaneously, from Fig. 1a, the distances between Ti and two O atoms in IM3 are lengthened to 1.914 and 1.966 Å respectively. Along this reaction coordinate, one of the Ti−O bonds and a C−O bond rupture simultaneously to form IM4, with an energy barrier of 28.65 kcal/mol. From Fig. 3 b, one can see that via the transition state TS₃₄, the CO₂ molecule is generated and one activated atomic oxygen remains on the Ti site of the surface to form IM5. The binding energy between CO₂ and IM5 is calculated to be 6.69 kcal/mol. Obviously, from Fig. 3 b, this CO oxidation channel IM2 + CO → IM5 + CO₂ is spontaneous in energy, and exothermic by 5.78 kcal/mol. It should be pointed out that although several trials were undertaken to search for possible transition states that connect reactants and IM3.

![Fig. 2. Electron density difference (EDD) between the O₂ adsorption state and Ti-embedded graphene (the electron density depletion and accumulation sites are displayed in dashed and solid lines respectively)](image-url)
no such structures were obtained. Obviously, the formation of IM3 is a barrier-free exothermic reaction. Form the O-Ti-graphene (IM5) easily. Due to its strong electronegativity, the oxygen atom in IM5 can easily pull the charge from Ti. The NBO analysis shows in IM5, there is 1.244e on the Ti atom, –0.648e on the O atom, suggesting that the Ti-O site might be the active site for adsorbing and oxidation an probe molecule, such as CO.

Now we will discuss the second CO oxidation channel, which may lead to product (CO₂ + IM0) formation. As described in Fig. 3 c, this route starts with the adsorption of CO on both O and Ti sites of the O-Ti-graphene (IM5), formation of a three-membered ring structure, IM6. NBO analysis of this complex shows there exist strong interaction between the 4d orbital of Ti and the π* orbital of C–O bond, and the Ti can receive some electron from CO. Energetically, IM6 lies –11.14 kcal/mol lower than the separate reactants of CO + IM5, and the formation of IM6 is a barrier-free exothermic reaction. Along the reaction channel, the next step corresponds to the CO₂ formation followed by the non-reactive-dissociation. Throughout a transition state TS₆₇, the intermediate complex IM7 is yielded at last with an activation barrier of 11.37 kcal/mol. Geometrically, the C–Oads bond in TS₆₇ is shortened to 1.793 Å, synchronously, the distance between Ti and Oads is elongated to 1.758 Å. The imaginary frequency of TS₆₇ is 336.8i cm⁻¹, and the normal mode corresponds to the formation of C–O bond. One exit channel of IM7 is direct desorption of CO₂ to yield the product (CO₂ + IM0), but this is a greatly endothermic process with the desorption energy of 13.02 kcal/mol. Obviously, the surface of catalyst can be recovered after second CO oxidation. During the process of second CO oxidation, the graphene support plays an important role in the chemical properties for the Ti-embedded graphene, the negative charge of –0.548e can be lowered to –1.278e by electron transfer from the adsorption of CO. Similar to the first CO oxidation, the graphene act as an electron withdrawing group here.

3.4. The reaction over the Zr-embedded graphene

In addition, we also located possible CO oxidation channels on the Zr-embedded graphene. As shown in Fig. 3, the reaction mechanism on this surface is in general similar to that of the Ti-embedded one. The reaction starts with an oxygen molecular activation, followed by two possible CO oxidation pathways to form CO₂. In both cases, as all the intermediates, transition states and products involved in the reaction lie below the respective reactants, the reactions over Ti and Zr-embedded graphene are expected to occur spontaneously in energy. From Fig. 3 a, one can see for the O₂ activation over Ti-embedded graphene, both the initial intermediate (IM1), and transition state (TS₁₁₂) involved in the reaction, lie far below the analogues over the Zr-embedded one, we believe the Ti-embedded graphene can activate oxygen more effectively. But, for the next CO oxidation over surface with oxygen adsorption and O–Zr–graphene, the oxidation pathways are almost energetically preferable with respect to the corresponding routes over Ti. Obviously, Zr-embedded graphene appears to be more effective during the process of CO oxidation.
4. CONCLUSIONS

In the present study, the oxidation reactions of O\textsubscript{2} and CO over Ti and Zr-embedded graphene have been investigated detailedy. The reactions are expected to occur using the ER mechanism. For investigating the CO oxidation over Ti and Zr-embedded graphene in the presence of O\textsubscript{2}, a possible reaction is found and it includes the following steps.

\begin{align*}
O_2 & \rightarrow O_{\text{act}}; \\
O_{\text{act}} + CO & \rightarrow CO_2 + O_{\text{ads}}; \\
O_{\text{ads}} + CO & \rightarrow CO_2.
\end{align*}

The catalyst substrate acts both the electron donor and acceptor during different reaction processes. The present theoretical work is a beneficial complement to the oxidation reaction of CO over metal-embedded graphene, which plays an important role in promoting further experimental efforts to design novel catalysts for solving the environmentally harmful exhaust gases from industrial waste.

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