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Single Pt Atom Decorated Graphitic Carbon Nitride as an

Efficient Photo-catalyst for the Hydrogenation of

Nitrobenzene into Aniline

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Abstract

The hydrogenation of nitrobenzene into aniline is one of industrially important reactions,

but still remains great challenge due to the lack of highly active, chemo-selective and eco-

friendly catalyst. By using extensive density functional theory calculations, herein we predict

that single Pt atom decorated g-C₃N₄ (Pt@g-C₃N₄) exhibits excellent catalytic activity and

selectivity for the conversion of nitrobenzene into aniline under visible light. The overall

activation energy barrier for the hydrogenation of nitrobenzene on single atom Pt@g-C₃N₄

catalyst is even lower that that of the bare Pt (111) surface. The dissociation of N-O bonds on

single Pt atom is triggered by single hydrogen atom rather than double hydrogen atoms on

the Pt (111) surface. Moreover, the Pt@g-C₃N₄ catalyst exhibts outstanding chemoselectivity

towards the common reducible substituents, such as phenyl, -C=C, $-C\equiv C$ and -CHO groups

during the hydrogenation. In addition, the doped single Pt atom can significantly enhance

the photoconversion efficiency by broadening the light absorption of the pristine g-C₃N₄ to

visible light region. Our results highlight an interesting single-atom photocatalyst (Pt@g-

C₃N₄) for efficient hydrogenation of nitrobenzene to aniline under a sustainable and green

approach.

Keywords: chemoselective hydrogenation, single-atom catalyst, photocatalyst,

nitrobenzene, aniline

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Introduction

Hydrogenation are among the most important and challenging technological reactions in modern chemical industry¹. The high activity and chemoselectivity are often very difficult to control especially when more than one reducible groups are present during the hydrogenation process²⁻³. An important example is the catalytic hydrogenation of nitrobenzene into aniline, a high-value chemical raw material which can be widely used in dyes, pigments, rubber, agrochemical and pharmaceutical industry.⁴⁻⁶ Currently, the catalytic hydrogenation of nitro groups to produce anilines mainly relies on noble metal complexes catalysts, such as palladium, rhodium, ruthenium and iridium ⁷⁻⁸. Although the precious metals have the advantage of high catalytic activity, they still suffer from low selectivity and high expense. Therefore, the exploration of environmentally benign and cost effective catalysts for aniline synthesis is thus of great fundamental as well as practical interest ⁹⁻¹¹.

Great research efforts toward improving the activity and selectivity has been made by developing a variety of catalysts, such as noble metal catalysts Au^{10, 12-13}, AuPt¹⁴, CoPd¹⁵, Pd/CeO₂², non-noble metal catalysts Ni¹⁶, Ni-Co as well as Ni-Fe alloys¹⁷⁻¹⁸ and so on. However, the problem often arising is that these catalyts have to be modified by additives or combining with other metal oxides and molecules to achieve high selectivity¹⁹⁻²². In most cases, the well-chosen additives will cause enviormental problems and at the same time reduce their activity²³. Very recently, cobalt or iron based complexes catalysts have been developed as inexpensive catalysts with excellent chemoselectivities for the hydrogenation of nitroarens, but the reaction conditions are relatively demanding with more than 12h reaction time, high temperature and high hydrogen pressure 24-25. In order to find a more gentle and economical protocol, some researchers tend to use semiconductor photocatalysys such as the gold nanoparticles or clusters (Au/TiO₂ and Au/ZrO₂) ²⁶⁻²⁹. Some of them demonstrated excellent performance in degradation of organic pollutants and various organic transfermations³⁰. However, hydrogen atoms can easily diffuse to the subsurface region of metal nanoparticles or clusters during the hydrogenation reaction and the formed sub-hydrogen atoms could cause over-hydrogenation of the intermediates and thus lower the selectivity towards the desired products 31-32. Interestingly, some studies have revealed that the chemoselectivity towards the hydrogenation of -C=C and -C≡C groups is highly sensitive to the particle size, while the hydrogenation of nitro groups appears to be size insensitive over metal nanopaticles ³³. Inspired by these results, we assume that we can tune the chemoselectivity by reducing the size of metal particles ^{22, 34}. The limitation for downsizing metal particles is to use isolate single atom on the supports, namely 'single-atom catalyst (SAC)'³⁵. Most recently, single atom catalysis has been extensively explored since the seminal work of Pt₁/FeO_x reported by Zhang and co-workers in 2011 ³⁶. The efficient use of the metals and superior activity are rendering SAC very popular in water-splitting^{37–38}, metal-air battery³⁹, CO oxidation^{36, 40}, CO₂ reduction⁴¹ and N₂ fixation⁴². However, few works have been reported for the use of SACs in the selective hydrogenation reactions.

Experimentally, optically active graphitic carbon nitride (g-C₃N₄) has been widely used as an excellent substrate for the design of single-atom photocatalysts⁴³⁻⁴⁵. The best example is the single Pt atom supported on g-C₃N₄ which has been successfully fabricated in recent experiment, showing excellent photocatalytic performance⁴⁶. Although some non-noble metal catalysts have been developed recently, Pt is still regarded as the dominant catalyst in the hydrogenation of various nitroaromatics due to its excellent catalytic properties⁴⁷. In this work, we for the first time propose to use single-atom Pt atom supported on g-C₃N₄ as a green single atom catalyst to address the activity, selectivity and economical issues during the hydrogenation of nitrobenzene into aniline. Our results indicate that the nitrobenzene can selectively adsorb on the Pt atom through the phenyl group and be further hydrogenated into aniline with a rather low activation barrier of 0.73 eV, which is even lower than that on the widely studied Pt (111) surface⁴⁸. Meanwhile, unlike over Pt surfaces, the monodispersed Pt atom prevented the coupling of multiple reactants to form the azo-compounds as byproducts due to the strong steric hindrance. Additionally, the Pt@g-C₃N₄ catalyst exhibits excellent selectivity towards some common reducible substituents, such as phenyl, -C=C, - $C \equiv C$ and -CHO groups. Most importantly, the supported Pt atom can singnificantly extended the absorption edge of g-C₃N₄ to the visbile light region which enable the photodriven hydrogenation of nitrobenzene into aniline.

Computational Details

The Vienna ab initio simulation package (VASP) code was used to perform all the geometric optimizations and electronic structures calculations⁴⁹⁻⁵⁰. The core-valence interactions was accounted by using the functional of generalized gradient approximation (GGA) in the

Perdew-Burk-Emzerhof (PBE) form combined with the projector augmented wave (PAW) method^{49, 51}. The spin-polarization was employed through all the calculations and the energy cutoff for plan-wave expansions was set to 500 eV. The electronic relaxation for self-consistency was set to 10⁻⁵ eV and the residual forces were limited to less than 0.01 eV/Å. A Monkhorst-Pack k-point mesh of 3×3×1 grid was used during the calculations. To avoid the interaction between two periodic units, the vacuum space was set to more than 15 Å. DFT+D3 was used to treat the long range van der Waals interaction. The climbing nudged elastic band (CI-NEB) method was adopted to search for the saddle point and the optimal reaction pathway⁵²⁻⁵³. We have taken all the possible elementary reaction steps which may happen on the catalyst into consideration. The energies changes related to these reaction steps were calculated using the following equation:

$$\Delta E = E_{adsorbate+catalyst} - E_{catalyst} - E_{adsorbate}$$

where $E_{adsorbate+catalyst}$, $E_{catalyst}$ and $E_{adsorbate}$ stand for the total energy of the Pt@g-C₃N₄ with adsorbate, Pt@g-C₃N₄ catalyst and isolate adsorbate, respectively.

Results and discussions

Single Pt atom can be stably anchored into the host cavity of g-C₃N₄ by coordinating with two N atoms, as shown in Figure 1a. The binding energy was calculated to be -3.15 eV, which is consistent with other reports^{45, 54}. The adsorption mode of the reactants on the catalyst surface has significant impact on the catalytic selectivity during the heterogeneous catalysis⁵⁵⁻⁵⁶. Thus, we first investigated all the possible adsorption configurations of nitrobenzene on the Pt@g-C₃N₄ catalyst. The nitrobenzene can be adsorbed on the single Pt atom through two different types, one is via the phenyl-group and the other one is via nitrogroup. Four different adsorption geometries including Pt-phenyl, Pt-O-O, Pt-C-N and Pt-N-O were taken into consideration. Geometry optimization was carried out and binding energies were calcualted for all these configurations. For the Pt-C-N configuration, the nitrobenzene spontaneously changed to Pt-phenyl adsorption mode during the optimization. As shown in Figure 1b, c and d, the Pt-phenyl configuration owns the lowest binding energy of -1.96 eV compared to -1.7 eV for Pt-O-O and -1.42 eV for Pt-N-O. Accordingly, nitrobenzene prefers to bind on the single Pt atom through phenyl group. We further calculated the differential charge densities of the Pt-phenyl adsorption mode, as shown in

Figure S₁ in the Supporting Information. The adsorption behavior of nitrobenzene on Pt atom is similar to that on the Pt (111) surface⁵⁷⁻⁵⁸. There is a significant charge transfer between the Pt atom and nitrobenzene, indicating a strong interaction between the reactant and catalyst. There is a charge depletion around the Pt atom with 1.12 e^- lost, while charge was accumulated arount the nitro-group with 0.99 e^- gained based on the Bader charge analysis⁵⁹. These results clearly demonstrated that nitrobenzene can be stably adsorbed on the catalyst which is crucial for the subsequent hydrogenation reaction.

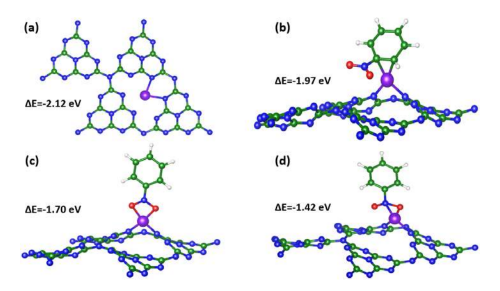


Figure 1 The top view of (a) the single Pt atom supported on g-C₃N₄ and side view of three typical adsorption configurations of (b) Pt-phenyl, (c) Pt-O-O and (d) Pt-N-O.

After figuring out the most stable adsorption configuration of nitrobenzene on the single Pt atom, we continue to investigate hydrogenation reaction of the nitrobenzene (Ph-NO₂) to aniline (Ph-NH₂). The overall hydrogenation can be expressed as a six-electron process: $PhNO_2 + 6H^+ + 6e^- \rightarrow PhNH_2 + 2H_2O$. Two hydrogenation mechanisms were well known, i.e. the direct and condensation reaction pathways (see Scheme S1 in the Supporting Information) as proposed by Haber in 1898⁶⁰. The difference between these two reaction pathways is whether azo-compounds can be formed during the hydrogenation reaction. Therefore, we first examined the possibility of the two reaction intermediates (*Ph-NO and *Ph-NHOH) interacting with each other to generate *Ph-N=NO-Ph* intermediate. The optimized structure showed that *Ph-NO intermediate could not bind with *Ph-NHOH intermediate due to the large distance between the adjacent active sites. Moreover, unlike the metal clusters or surfaces which could provide sufficient active sites for the co-adsorption of more

than one nitrobenzene molecules, single atom active site can only adsorb one nitrobenzene molecule at one time due to the large steric hindrance. Under this circumstance, it is very hard for the intermediates to couple into azo-compounds on single Pt atom as by-products and thus production efficiency of aniline could be greatly improved. Therefore, single atom $Pt@g-C_3N_4$ catalyst could offer signficant advantage over the metal cluster or surface. In the following, we specifically investigated the direct reaction routes as shown in Scheme 1.

Scheme 1 Proposed possible reaction pathways for the reduction of nitrobenzene into aniline through direct routes (the reaction pathway in blue is the optimal one).

Table 1 Calculated reaction energies (ΔE , in eV) and activation barriers (E_a , in eV) of the elementary reaction steps involved the hydrogenation of nitrobenzene to aniline on the Pt/C_3N_4 catalyst.

Reactions	ΔΕ	Ea
$*C_6H_5NO_2 + H^* \rightarrow *C_6H_5NOOH$	0.28	0.48
$^*C_6H_5NOOH+ H^* \rightarrow ^*C_6H_5N(OH)_2$	-0.48	0.46
$^*C_6H_5NOOH+ H^* \rightarrow ^*C_6H_5NO + H_2O$	-1.66	0.24
$^*C_6H_5N(OH)_2+H^* \rightarrow ^*C_6H_5NOH$	-2.42	0.19
+H₂O		
$*C_6H_5NO+H^* \rightarrow *C_6H_5NOH$	-1.25	0.09
$^*C_6H_5NOH+ H^* \rightarrow ^*C_6H_5N + H_2O$	-0.95	0.73
$^*C_6H_5NOH + H^* \rightarrow ^*C_6H_5NHOH$	0.15	0.99
$^*C_6H_5NHOH+ H^* \rightarrow ^*C_6H_5NH + H_2O$	-2.83	1.18
$^*C_6H_5N+H^* \rightarrow ^*C_6H_5NH$	-1.72	0.53
$^*C_6H_5NH+ H^* \rightarrow ^*C_6H_5NH_2$	0.24	0.66

We then move to investigate the catalytic activity of single atom Pt@g-C₃N₄ catalyst for the conversion of nitrobenzene into aniline. The reaction process mainly involves the dissociation of N-O bonds and the formation of N-H bonds while accompanying by the generation of two water molecules as the by-products. As we can see from the Scheme 1, four possible reaction pathways will be fully explored to indentify the most appropriate route. The calculated reaction energies and the activation energies for all the possible intermidates are summarized in Table 1. Notably, most reaction energies are negative and the most postive one is only 0.28 eV, suggesting favorable thermodynamic for the hydroenation of nitrobenzene into aniline on the Pt@g-C₃N₄ catalyst. Then we calculated the activation barrier for every reaction step using the NEB method. The energy profiles for the most optimal reaction pathway is presented in Figure 2. All the other three less favorable routes are included in Figure S2-S4 in the Supporting Information. For all the four reaction pathways, the first step for hydrogenation of nitro-group is that one proton-electron pair attack on one of the O atom to form *Ph-NOOH intermediate, which is basically an uphill reaction with a small postive reaction energy of 0.28 eV and needs to overcome a low activation barrier of 0.48 eV. The next proton-electron pair can either bind with another O atom to form *Ph-(OH)₂ intermediate or attack the newly formed OH group to generate a H₂O molecule. It is apparent that the reaction pathway with the broken N-O bond is more preferred with quite negative reaction energies (-1.66 eV vs -0.48 eV) and small activation barriers (0.24 eV vs 0.46 eV). Such low activation barriers mainly attribute to the adsorption of H atom that elongated the length of N-O bond from 1.21 Å to 1.42 Å and thus facilitated the dissociation of N-O bond to generate a H₂O molecule. Next, the third proton-electron pair comes up to bond with the O atom to form *Ph-NOH intermidate with a ultralow barrier of 0.09 eV. The fourth proton-electron pair will either combine with the N atom or break the second N-O bond. The caculated reaction energies for the formation of *Ph-NHOH and *Ph-N intermidates are 0.15 eV and -0.95 eV, respectively, suggesting the *Ph-NOH→ *Ph-NHOH reaction path may be more difficult to occur than the *Ph-NOH→ *Ph-N + H₂O path. The former reaction path needs to overcome a higher activation barrier of 0.99 eV (*Ph-NOH→ *Ph-NHOH) compared to 0.73eV for the later one (*Ph-NOH→ *Ph-N + H₂O). Hence the perferred reaction path in this step is to break the second N-O bond and releasing another H₂O molecule. This reaction (*Ph-NOH→ *Ph-N + H₂O) was the overall rate-determining step during the whole hydrogenation reaction. It is worth noting that previous works also revealed that the dissciation of N-O bonds are the key steps during the reduction of nitrogroup⁶¹⁻⁶³. With another two proton-electron pairs get transferred, the nitrobenzene was eventually converted into aniline.

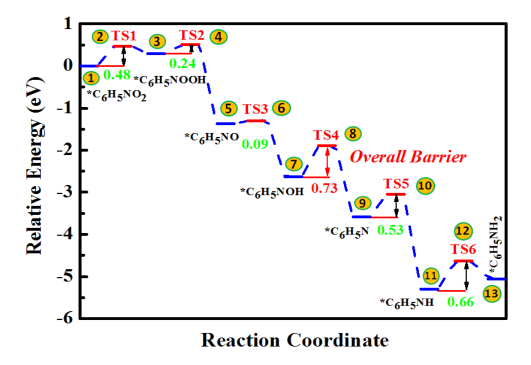


Figure 2 Energy profiles of the optimal reaction pathway for the hydrogenation of nitrobenzene into aniline on single atom Pt@g-C₃N₄ catalyst.

Figure 3 presented all the optimzied reaction intermidates (odd-numbered configurations) and the transition states (even-numbered configurations) along the most favorable hydrogenation reaction pathway (see Fig. 2). Here we focused more on examining the transition state geometries of the dissociations of two N-O bonds (TS2 and TS4). It was found that the N-O bonds were first elongated more than 0.2 Å when the first and the third H* were transferred to O atom. That means the N-O bonds can be easily activated on the Pt@g-C₃N₄ catalyst by the adsorbed single proton. On the basis of reaction and activation energies shown in Table 1, the Ph-NOOH \rightarrow Ph-NOH \rightarrow Ph-NOH \rightarrow Ph-N pathway is more favorable than the reaction (Ph-NOOH \rightarrow Ph-N(OH)₂ \rightarrow Ph-NOH \rightarrow Ph-NHOH). Therefore, unlike conventional double H-indued reaction pathway on the Pt (111) surface⁶³, single hydrogen induced activation of N-O bond plays a more important role on the Pt@g-C₃N₄ catalyst during the deoxygenation reactions. Hence the mechanism for overall hydrogenation of nitrobenzene on the Pt@g-C₃N₄ catalyst can be identified as: Ph-NO₂ \rightarrow Ph-NOOH \rightarrow Ph-NO \rightarrow Ph-NOH \rightarrow Ph-NOH \rightarrow Ph-NH \rightarrow Ph-NH₂. The highest activation

energy occurs at the dissociation of the second N-O bond ($\mathfrak{T} \rightarrow \mathfrak{D}$ as shown in Fig. 2) with a activation energy barrier of 0.73 eV, which is even smaller than that on the Pt(111) surface. Thus the catalytic activity of single atom Pt@g-C₃N₄ is superior to that of the Pt (111) surface. More importantly, the use of single Pt atom significantly reduced the load of precious metal to the greatest extent.

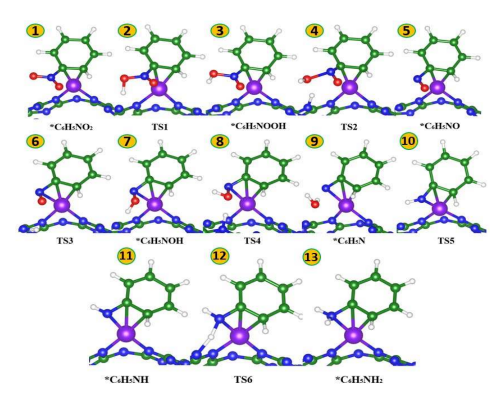


Figure 3 The optimized geometries for the reaction intermediates and transition states for the most favorable pathway shown in Fig. 2 during the hydrogenation process.

Besides the high catalytic activity, the chemoselectivity is also another primary factor to evaluate the hydrogenation performance for single atom Pt@g-C₃N₄ catlayst. Generally some important nitroarenes during the organic synthesis are usually functionalized with a variety of other reducible groups⁶⁴. These substituents will compete with the nitro-group and produce waste during the hydrogenation of nitrobenzene into aniline. For this reason, we further investigated the chemoselectivity of Pt@g-C₃N₄ catalyst towards some common reducible groups including phenyl, -C=C, $-C\equiv C$ and -CHO substituents. The reaction pathways for hydrogenating these functional groups as depicted in Figure 4. It can be clearly seen that the hydrogenatations of phenyl, -C=C, $-C\equiv C$ and -CHO substituents on single atom Pt@g-C₃N₄ catlayt possess remarkably high barriers of 1.05, 1.49, 1.29 and 1.53 eV,

respectively. It is worth noting that the overall activation energy for the hydrogenation of nitrobenzene was only 0.73 eV. The large diffrence of the activation energy between the above reactgions indicated the hydrogenation of nitronbenze is highly feasible at the room temperature, while the hydroenation of other phenyl, -C=C, $-C\equiv C$ and -CHO substitutents is rather difficult. Therefore, the overall hydrogenation reaction will give preference for the single $Pt@g-C_3N_4$ catalyst to attack the nitro-group to yield aniline products even in the presence of the above four reducible groups. This indicated an ideal high chemoselectivity for the hdyrogenation of nitrobenzen into aniline, which is in sharpt contrast other widely studied metal cluster or surface catalysts where some environmentally-unfriendly additives are normally used to improve their chemoselectivity, but often at the price of sarificing the catalytic activity to a great extent.

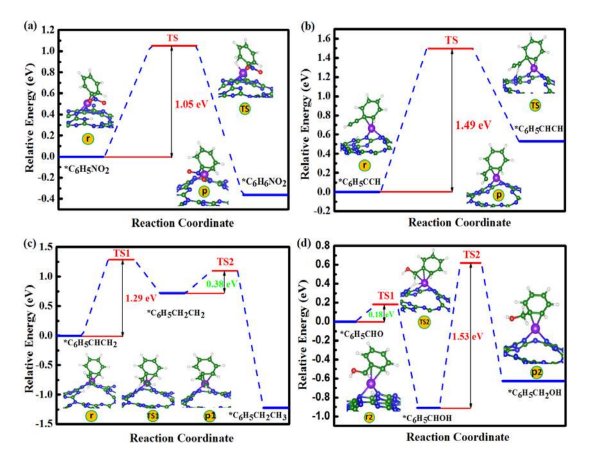


Figure 4 Energy profiles for the hydrogenation of phenyl, -C = C, $-C \equiv C$ and -CHO substituents on single atom $Pt@g-C_3N_4$ catalyst. Insets are the structures of reactants, products and transition states during the hydrogenation process. The green, blue, purple and white balls reprsent C, N, Pt and H atoms, respectively.

Experimentally, g- C_3N_4 -based photocatalysts have been widely studied for energy conversion and pollutants degradation⁶⁵⁻⁶⁶. The pristine g- C_3N_4 has a relative large band gap of 2.7 eV, rendering minimal absorption of visible light⁶⁷. As shown in Fig. 5, the light adsorption for the pristine g- C_3N_4 mainly located in the ultraviolet region. However, single Pt atom decorated g- C_3N_4 could significantly improve the optical absorption activity (see Fig. 5). The light absorption of the Pt@g- C_3N_4 composite can be extented to visible and infrared light region due to a narrowed band gap (0.72 eV)⁴⁵. Therefore, the Pt@g- C_3N_4 catalyst is able to hydrogenate nitrobenze into aniline under visible/infrared light irradiation compared the pure g- C_3N_4 catalyst.

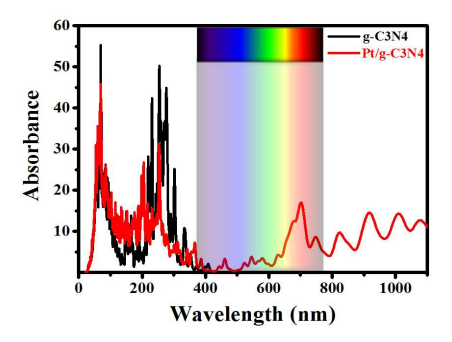


Figure 5 The optical absorption spectra for the pristine g-C₃N₄ and the Pt/g-C₃N₄.

Conclusions

In summary, we propose a novel single atom photocatalyst, $Pt@g-C_3N_4$, with outstanding photo-catalytic activity and chemoselectivity for the hydrogenation of nitrobenzen into aniline. The optimal hydrogenation pathway is identified as: $Ph-NO_2 \rightarrow Ph-NOOH \rightarrow Ph-NO$ $\rightarrow Ph-NOH \rightarrow Ph-NH \rightarrow Ph-NH_2$. The most energy demanding step among the hydrogenation process is the dissociation of the second N-O bond with a activation barrier of 0.73 eV only. The mechanism of single H-induced activation of N-O bond plays a cruical role in the hydrogenation reaction. Moreover, activation barriers for hydrogenation of

phenyl, -C=C, $-C\equiv C$ and -CHO substituents are rather high compared with the nitro-group, indicating that single atom $Pt@g-C_3N_4$ catalyst possesses excellent chemoselectivities towards various reducible groups. Additionally, single Pt atom supported on $g-C_3N_4$ can greatly broadened the range of light absorption, allowing the hydrogenation of nitrobenzene into aniline to be driven under visible/infrared light. The high catalytic activity, outstanding selectivity towards various reducible groups, excellent photocatalytic performance and efficient use of the noble metal render $Pt@g-C_3N_4$ a promising environmentally benign single atom photocatalyst for the hydrogenation of nitrobenzene into aniline.

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Single Pt Atom Decorated Graphitic Carbon Nitride as an Efficient Photo-catalyst for the Hydrogenation of Nitrobenzene into Aniline

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1. Two hydrogenation mechanisms, i.e. the direct and condensation reaction pathways proposed by Haber.

Scheme S1 The Proposed mechanism of hydrogenation of nitrobenzene into aniline.

2. The difference charge density of nitrobenzene absorbed on Pt@g-C₃N₄.

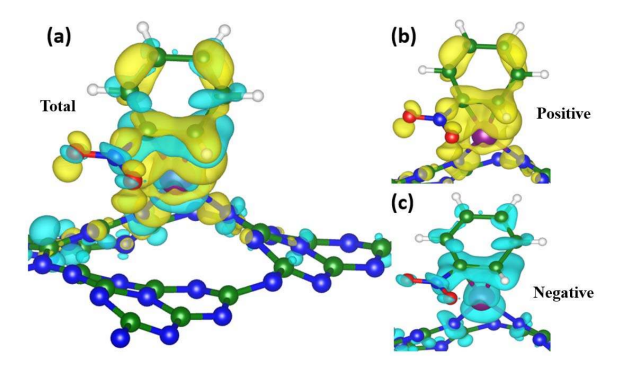


Figure S1. Difference charge density of $Pt@g-C_3N_4$ with the adsorption of nitrobenzene, where the isosurface value is set to be 0.005 e/Å and the positive and negative changes are shown in yellow and cyan, respectively. Green, blue, red, purple and white balls represent the C, N, O, Pt and H atoms, respectively.

Three difference reaction pathway for the hydrogenation of nitrobenzene into aniline on single atom Pt@g-C3N4 catalyst.

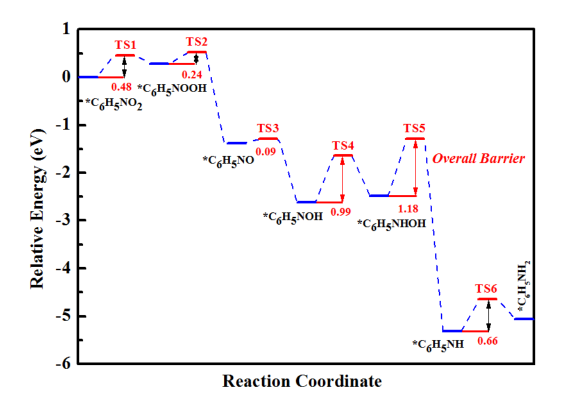


Figure S2 Energy profiles of the reaction pathway through *Ph-NHOH intermediate for the hydrogenation of nitrobenzene into aniline on single atom Pt@g-C₃N₄ catalyst.

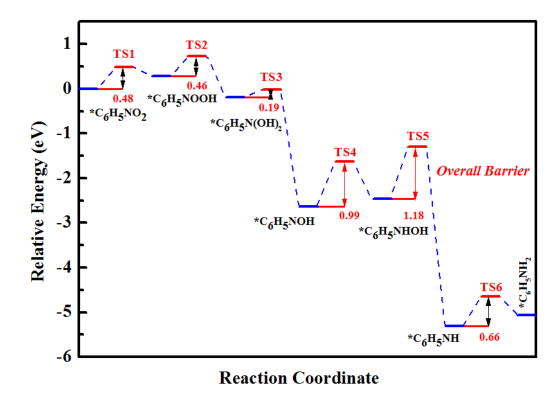


Figure S₃ Energy profiles of the reaction pathway through *Ph-N(OH)₂ and *Ph-NHOH intermediates for the hydrogenation of nitrobenzene into aniline on single atom $Pt@g-C_3N_4$ catalyst.

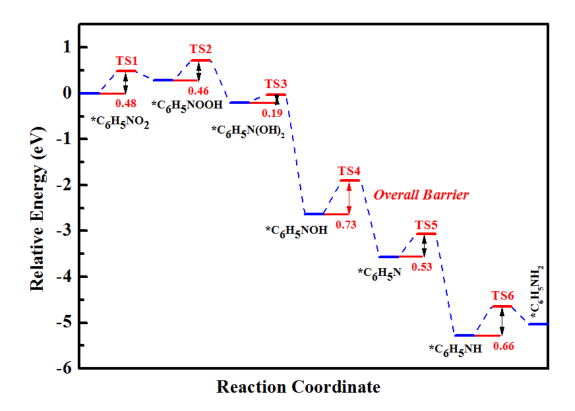


Figure S4 Energy profiles of the reaction pathway through *Ph-N(OH)₂ and *Ph-N intermediates for the hydrogenation of nitrobenzene into aniline on single atom $Pt@g-C_3N_4$ catalyst.