Graphene Quantum Dot-Modified Lipase for Synthesis of L-menthyl Acetate with Improved Activity, Stability and Thermostability

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Abstract

**Title:** Pseudomonas cepacia lipase modified with Graphene Quantum Dots (GQD) offers a higher activity, stability and thermostability compared to the lipase modified with graphene oxide.

**Background:** GQD presents extraordinary properties attracting extensive attention from scientists in chemistry, physics, materials, biology and other interdisciplinary sciences. However, no research involves the application in biocatalysts in non-aqueous media up to now. In the study, we for the first time reported a promising GQD-modified lipase for the synthesis of L-menthyl acetate.

**Methods and findings:** *Pseudomonas cepacia* lipase was modified with GQD and then used as a biocatalyst for the synthesis of L-menthyl acetate in 1-isobutyl-3-methylimidazolium hexafluorophosphate medium. As contrasts, the reaction was also carried out using bare lipase and graphene oxide-modified lipase as the catalyst. Besides, the modification method of enzyme, the amounts of GQD, the reaction temperature, molar ratio of the two substrates, and reuse of the enzyme were investigated. The result shows that the GQD-modified lipase as a biocatalyst was best among the three enzymes. Under optimal reaction conditions, the reaction reaches the equilibrium within 8 h with a high conversion of L-menthol (97.3%). Its initial enzyme activity and half-lifetime were more than 1.08 and 2.07-fold that of the bare lipase, and 1.04 and 1.66-fold that of the graphene oxide-modified lipase, respectively. The lipase was recycled 10 times without substantial diminution in activity. In addition, the GQD-modified lipase also offers a better thermostability compared with bare lipase. These improvements could be attributed to the unique small size and edge effect of GQD.

**Conclusion:** GQD-modified lipase showed a higher activity, stability and thermostability compared to GO-modified lipase. GQD was widely used as a promising material for enzyme modification due to its good biocompatibility and small size.

**Keywords:** Graphene Quantum Dot (GQD); Lipase; Biocatalyst; L-menthyl acetate; Synthesis

Introduction

Enzyme is a biological molecule that can regulate chemical reactions in biological process [1,2]. Lipase as a biocatalyst has been widely applied in various organic reactions, including hydrolysis, transesterification, resolution of racemic mixtures and stereoselective oligomerization [3-6]. The significant value of lipases mainly lies on their wide industrial applications in pharmaceutical synthesis, detergent formulation, cosmetics production and oil/fat degradation [7-10]. However, lipase-based catalysts are not always sufficient for industrial application due to their poor catalytic activity and stability, which largely limits their application [11]. To overcome the above limitations, many efforts have been made to improve enzyme catalytic performance to meet the need of practical applications, including the use of additives with stable effect, modification of structure by chemical methods or protein engineering methods and various immobilization strategies [12-15]. The enzyme immobilization or modifications have gained widespread attentions due to their advantages of enzyme reutilization, industrial-scale application and simplification of operation Process [16].

The performance of immobilized enzyme greatly depends on the support materials. Recently, nanostructured materials evolve as promising alternatives for enzyme modification. The advantages for nanomaterials standing out among other supporters are their ultrahigh surface area for biomolecule adsorption and access to functionalization of surface functional groups for interacting with biomolecules. Enzyme immobilized on nanostructured materials can enhance catalytic activity, stability. Besides, they have the advantages...
of repeated or continuous use, easy separation from the reaction mixture, prevention of protein contamination in the product. Accompanying the rapid development of nanotechnology, a great number of nanomaterials have been utilized for enzyme immobilization or modification, such as gold nanoparticles, magnetic nanoparticles and porous silica structures [17-20]. However, no reports refer on the modification of industrial enzyme by Graphene Quantum Dots (GQD) to improve the enzyme performance of nonaqueous catalysts.

GQD, one type of D graphene sheets with lateral size less than 100 nm, has attracted increasing attention owing to their robust chemical inertness, low cytotoxicity, excellent biocompatibility, high photostability and ease of preparation [21-23]. Great potential of GQD has been shown in various applications including biosensing, cellular imaging, photovoltaic devices and biomedicine [24-26]. Similar to Graphene Oxide (GO), GQD contains rich of oxygen-containing groups such as hydroxyl, carboxyl and epoxy groups. The characteristic makes it possible to modify enzymes without any surface modification or coupling reagents. Particularly, small size and controllable polarity make it more suitable for enzyme modification compared with GO. The physicochemical properties and structure suggest GQD has great potential for applications in nonaqueous reaction media.

In this study, we reported synthesis of nitrogen and sulfur co-doped GQD via pyrolyzing the mixture of citric acid and L-cysteine and its application in the modification of Pseudomonas cepacia lipase. The modified lipase was used as a biocatalyst for synthesis of l-menthyl acetate in the hydrophobic ionic liquid 1-isobutyl-3-methylimidazolium hexafluorophosphate. The modified lipase exhibits a better catalytic efficiency and reproducibility and operational stability compared to bare lipase. It can be widely used in various biocatalysis reactions for practical applications to improve the activity and stability of enzymes.

Methods

Materials and reagents

Pseudomonas cepacia lipase (PCL, 30 U mg\(^{-1}\), solid) and L-menthol were obtained from Sigma-Aldrich Chemical Company (Mainland, China). The ionic liquid, 1-isobutyl-3-methylimidazolium hexafluorophosphate ([I-C\(_4\)-min][PF\(_6\)]) was prepared according to the literature [4]. GO was synthesized from natural graphite powder by a modified Hummers method [27]. All other reagents employed were purchased from Shanghai Chemical Company (Shanghai, China) with analytical reagent grade. Ultrapure water (18.2 MQ cm) purified from a Milli-Q purification system was used throughout the experiment.

Apparatus

Transmission electron microscope (TEM) image was obtained by a JEOL 2100 FEG microscope at 200 KeV. The sample was prepared by dispensing a small amount of dry Val-GQDs in ultrapure water and followed by drop one drop of the suspension on 200 mesh copper TEM grids covered with thin amorphous carbon films. Scanning electron microscope (SEM) analysis was carried out using a S4800 field emission scanning electron microscope (Hitachi, Japan). X-ray photoelectron spectroscopy (XPS) measurement was carried out on a PHI 5700 ESCA spectrometer using Al HR radiation (hv=1486.6 eV). Infrared spectrum (IR) was recorded on a Nicolet FT-IR 6700 spectrometer. GC analysis was performed with a FULI 9790 instrument equipped with FID detector (Fuli Analytical Instrument Co. Ltd, China) and a chiral β-DEX 120 capillary column (30 m × 0.25 mm × 0.25 μm). Fluorescence images were obtained digitally on a Nikon-80i upright fluorescence microscope. GC analysis was performed with a FULI 9790 instrument equipped with FID detector, using benzyl acetate as internal standard. The chromatographic conditions were as follows: Column temperature, 170; injection temperature and detector temperature, 300; Carrier gas, nitrogen; N\(_2\) stream, 30 mL min\(^{-1}\); split ration, 1:50. The peak retention times were as follows: l-menthyl acetate, 2.89 min; l-menthol, 3.24 min; benzyl acetate, 3.81 min. The yield was calculated from the amount of the easter peak. Substrate and product concentrations were calculated from calibration curves using stock solutions of pure compounds.

Synthesis of GQD

GQD was prepared by pyrolyzing the mixture of citric acid and L-cysteine following the previous report, with some modifications [28]. Briefly, a 4 g of citric acid and 2 g of L-cysteine were dissolved in 10 mL of ultrapure water. Followed by heated at 200 for 3 h with evaporation until dry. The resulting black product was dissolved by dropwise addition of 1 M NaOH until pH of the solution was neutral. The solution was subsequently purified by dialyzing and a powdery product was obtained by lyophilisation.

Modification of PCL

First, GQD (500 μg) was dispersed in water to form a concentrated GQD dispersion. The dispersion was subsequently transferred to a sprayer and sprayed onto the surface of dry PCL (150 mg) under vigorously vortex shaking. Finally, the hybrid was dried by freeze drying to enhance long-term stability of GQD-PCL.

General procedure for the synthesis of l-menthyl acetate

A 0.5 g of l-menthol (3.2 mmol), 300 μL of acetic anhydride, 1 ml of ionic liquid and 30 mg lipase PCL were added into a 25 mL conical flask with stopper in sequence, followed by incubation the above mixture at 40 in an air bath constant temperature oscillator with a stirring speed of 180 r min\(^{-1}\). At different times, 10 μL aliquots were withdrawn and dispersed in 400 μL of hexane. After being precipitated with centrifugation, 10 μL of benzyl acetate (internal standard) was added to the supernatant (100 μL) and then the solution was
analyzed by GC. All experiments were repeated for five times, and mean values were reported.

**Half-lifetime measurement**

Mixtures containing 1.0 mL of ionic liquid and 30 mg of lipase PCL or modified PCL were incubated at 40. After a certain time, 0.5 g of l-menthol and 300 μL of acetic anhydride were added and the initial activity of biocatalyst was measured. Then the specific activity $A$ was calculated based on the measured results. Specific activity is defined as the enzyme amount needed for synthesis of 1 μmol of products under optimal conditions. All the experiments were carried out in duplicate. Half-lives were calculated from a first-order exponential decay of the activity:

$$A = A_0 \times e^{-kt}$$

where $k$ is the first-order deactivation rate constant, and $A_0$ and $A$ are the initial and residual enzyme activity, respectively. The half-life $t_{1/2}$ is defined as the $t$ at $A = A_0/2$ or $t_{1/2} = \ln 2/k$.

**Results**

**Materials characterization**

SEM image of GO and TEM image of GQD as well as their FTIR spectra are shown in Figure 1. It can be seen that GQD offers a narrow particle size distribution with the average size of 2.3 ± 0.4 nm. While GO is composed of relatively large graphene sheets of a few micrometer. On the FTIR spectrum in Figure 1D there are strong absorption peaks at 3420 cm$^{-1}$, 1701 cm$^{-1}$ and 1400 cm$^{-1}$. They could be attributed to stretching vibration of $-\text{OH}$, $-\text{C}=\text{O}$ and $-\text{COOH}$, respectively. The results also show that GQD contains the rich of oxygen-containing groups, which renders GQD strongly hydrophilic and water soluble. The presence of peak at 1296 cm$^{-1}$ suggests the existence of C-N and C-S groups.

**Optimization of reaction conditions**

The GQD-modified lipase (GQD-PCL) was investigated as biocatalyst for the model reaction of esterification of l-menthol with acetic anhydride in [i-C$_4$min][PF$_6$] ionic liquid medium (schemed in Figure 3). Herein, the conversion of l-menthol can well present the catalytic activity of the biocatalyst. To optimize the experimental conditions for this esterification reaction, effects of amounts of GQD, reaction time, reaction temperature and alcohol to anhydride molar ratio on activity of modified lipase were tested in the laboratory.

An appropriate amount of GQD should be considered to obtain high catalytic efficiency. Figure 4A shows that the amounts of GQD obviously effects on the conversion of l-menthol. The conversion increased with increasing the GQD content, and reaches the maximum value when the amounts of GQD are about 100 μg. However, the l-menthol conversion will rapidly decrease with the amount of GQD is more than 100 μg. This is because the excessive GQD for the modification...
of lipase could cause conformational hindrance and structural damage of PCL. This may reduce the interfacial area, leading to a low conversion. Thus, the GQD amount of 100 μg was selected for the modification of PCL.

The effect of reaction time on the l-menthol conversion was investigated. Figure 4B shows that the conversion was rapidly increased with the increase of the reaction time from 0 to 1 h. However, the conversion doesn’t significantly increase for the reaction times in excess of 8 h. This indicates that the reaction reaches the equilibrium within 8 h. Thus, the reaction time of 8 h was selected for the esterification reaction of l-menthol.

Influence of the molar ratio of l-menthol to acetic anhydride on the synthesis of l-menthyl acetate catalyzed by GQD-PCL was examined, in which the amounts of GQD was fixed at 100 μg. As shown in Figure 5, the l-menthol conversion increased with the increase of acetic anhydride content. The highest conversion were observed when the alcohol-to anhydride molar ratio was 1:1. However, the conversion of l-menthol decreased with the addition of anhydride. The result is similar to that of a well-established method [29]. This may be because a low concentration of anhydride leads to a low reaction rate. Meanwhile, the progressively produced acetic acid may be acted as main acyl donors along with the decrease of anhydride. However, the lower ability of acetic acid acylation leads to a decreased conversion. Besides, too much acetic anhydride will dehydrate into acetic acid, decreasing the activity of biocatalyst. Therefore, a best alcohol-to anhydride molar ratio of 1:1 was selected in the subsequent studies.

Figure 4: Effect of GQD amount (A) and reaction time (B) on l-menthol conversions.

Figure 5: Effect of the molar ratio of alcohol to anhydride on conversion of l-menthol.

The temperature mostly has the significant influence on the activity of lipase. High temperatures offer an important advantage of accelerating the reaction rates due to the reduced medium viscosity and increased substrate diffusion coefficient. However, high temperatures lead to promote protein denaturation [4]. Thus, it is of great importance to investigate the thermo-stability of lipase, especially for industrial applications. The effects of temperature in the range of 10 to 70 on the activity of the bare lipase and the modified one were investigated. With GQD amount being kept constant at 100 μg, pre-treatment of GQD-PCL in different temperatures and each l-menthol conversion was measured, respectively. It can be seen from Figure 6, the conversion of l-menthol has the same change trend when the GQD-PCL and bare PCL were used as the biocatalyst. At the temperatures of lower than 40, the conversion increased rapidly with the increase of reaction temperature. However, a gradual decrease in the conversion was observed with the further increase in temperature due to the gradual denaturation of lipase at high temperature. The optimum temperature was 40 for both bare PCL and GQD-PCL. More importantly, the conversion of l-menthol achieved by GQD-PCL is much higher than that by bare lipase under the same temperature. For the temperature at 50, the modified PCL retained nearly 85% of its relatively activity for an incubation of 8 h, whereas that of the bare PCL only left 50%. However, at 60, the bare nearly lost all its initial activity, whereas the modified PCL still retained 50% of its activity. Such significant resistance to heat of GQD-PCL could be due to attachment of GQD on PCL, which may cause robust conformational transitions at high temperature.

Figure 6: Effect of reaction temperature on l-menthol conversion.

According to the above studies, the optimal conditions for the esterification reaction of l-menthol catalyzed by the GQD-PCL were as follows: A GQD amount of 100 μg, reaction temperature of 40 and alcohol-to anhydride molar ratio of 1:1. Under the above conditions, initial reaction rate, the conversion of l-menthol was 97.3%, which was much higher than that of bare PCL (90.2%).

Possible Mechanism of the Improved Activity

To further understand the improved catalysis efficiency of the GQD-PCL, another additive, GO was used to replace GQD for the modification of PCL. shows the change of l-menthol conversion with increasing reaction time, in which bare PCL, GO-PCL or GQD-PCL was employed as the catalyst. It can be seen the initial reaction rates and l-menthol conversions for
GO-PCL and GQD-PCL were higher than that of bare PCL, indicating both GO and GQD can accelerate esterification reaction. This may be due to the hydrophilic groups of both GO and GQD provide a benign microenvironment for enzyme’s biological function. However, an obvious difference was unexpectedly observed when using these two biocatalysts. The reaction catalyzed by GQD-PCL reaches equilibrium within 8 h with a high conversion (97.3%), which is 1.04-fold greater than that of the GO-PCL and 1.08-fold greater than that of bare lipase, respectively.

![Image](https://via.placeholder.com/150)

*Figure 7: Time-courses of l-menthol conversion of PCL, GQD-PCL and GO-PCL.*

The key criterion for selecting an appropriate additive for the enzyme modification is the stability within the reaction medium. In this experiment, the stability of GO-PCL, GQD-PCL as well as bare lipase were investigated by incubating bare lipase and modified lipases in ionic liquids without substrates, followed by measurement of the residual activity using the method described in the general procedure.

![Image](https://via.placeholder.com/150)

*Figure 8: Deactivation curves of bare PCL, GO-PCL and GQD-PCL.*

Figure 8 shows the deactivation curves of the different biocatalysts. The half-life time of bare lipase, GO-PCL and GQD-PCL were found to be 108, 135 and 224 h, respectively, which were calculated using the formula \( t_{1/2}=\frac{ln2}{k} \). The data confirm that the stability of GQD-PCL is obviously higher than that of the other two catalysts. Its half-life time is more than 2.07-fold that of bare lipase and 1.66-fold that of GO-PCL. The result demonstrates that the addition of GQD improve the stability of lipase. On the one hand, ionic liquid \([i-C_4min][PF_6]\) provides a benign microenvironment for the lipase, which helps to retain the activity [5]. The large number of carboxylic groups existing at the GQD sheets’ periphery provide a hydrophilic microenvironment that enables lipase to maintain the ‘essential water’ for its biological function [30,31].

The above two factors improve the enzyme stability, which is necessary for the enzyme to display its biological function.

**Enzyme recycling**

To evaluate the possibility of the reuse of GQD-PCL, it was applied as a biocatalyst for esterification reaction of l-menthol repeatedly. The mixed solution was completely removed after l-menthol reacts with acetic anhydride for 8 h, and the collected GQD-PCL was directly used for the next esterification reaction with new raw materials under the same reaction conditions. After each reaction was completed, the conversion of l-menthol was measured by GC and the results are shown in . It can be seen that the conversion of l-menthol decreased slightly after 10 cycles from 97.3% to 94.3%, indicating no substantial diminution of enzyme activity. Figure 9 also indicates that the cycling stability of GQD-PCL is much better than that of the bare PCL. This demonstrates that the introduction of GQD can improve the stability of lipase.

![Image](https://via.placeholder.com/150)

*Figure 9: Conversions for the esterification reaction of l-menthol based on reuse of GQD-PCL (black) and bare PCL (green).*

**Discussion**

The study demonstrates the preparation of lipases-nitrogen and sulfur co-doped GQD composite and its application as the biocatalyst for esterification reaction of l-menthol with acetic anhydride in \([i-C_4min][PF_6]\) ionic liquid. The result shows that the GQD are composed of tiny graphene sheets with the rich of oxygen-containing groups. The small size of graphene sheets reduces the steric hindrance for the substrates entrance. The oxygen-containing groups increase with the polarity of lipase. The doped nitrogen and sulfur groups further improve the affinity of lipase with the substrate due to their relatively big electronegativity. The association of GQD with the lipase causes a special structural change that leads to exposure of the active site and results in an improvement in the enzyme activity. The lipase modified with the GQD exhibits an obviously enhanced activity, stability and thermo-stability. The study also provides a promising approach for enzyme modification in industrial catalysis to improve the enzyme activity and stability using GQD with good biocompatibility.
References


