Detection and difference analysis of the enzyme activity of colloidal gold nanoparticles with negatively charged surfaces prepared by different reducing agents

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Abstract: Research on the activity of nanoenzymes has always been a focus of nanomaterials. In this study, several reducing agents with different structures were used to prepare colloidal gold with a negative charge and similar size by controlling the temperature and pH. The affinity analysis of the substrate H2O2 and TMB showed that the activity of colloidal gold nanoenzymes prepared by different reducing agents was in the order of Cc, Hq, Rs, Vc, Ga, Sc, Sm, St. The rule is that the enzyme activity of colloidal gold reduced by benzene ring is higher than that of colloidal gold enzyme reduced by linear chain. Finally, we discussed the activity of colloidal gold peroxidase from the number and position of isomers and functional groups, and demonstrated that the nanoenzyme activity affected the surface activity of colloidal gold, the elimination of hydroxyl radical and TMB binding efficiency.

Keywords: colloidal gold; negative charge; nanoenzyme;

Introduction

Natural enzymes, such as proteins, RNAs, or their complexes, are widely used in the fields of disease diagnosis, clinical treatment, biosensing, and environmental protection due to their high catalytic efficiency and selectivity [1-4]. However, under extreme conditions (strong acid, strong alkali, high temperature, etc.), these enzymes are easily inactivated and have high extraction costs and low yields, making them difficult to use in large-scale industrial production [5]. Therefore, it is desirable to find natural enzyme substitutes with strong catalytic activities and stabilities [6,7]. In 2007, Yan et al. discovered for the first time that ferrous oxide magnetic nanoparticles (Fe3O4 MNPs) possess intrinsic peroxidase-like activity [8]. The enzymatic properties of nanomaterials were systematically studied from the perspective of enzymology, and corresponding determination standards were established, which were used as substitutes for enzymes [9]. In 2013, Wei et al defined such nanomaterial mimetic enzymes as nanoenzymes. Nanoenzymes are artificial mimetic enzymes with catalytic functions and unique physicochemical properties [10]. At present, most metal oxides, noble metals, and carbon-based nanomaterials are found to have enzymatic activity, and can simultaneously exhibit the activity of one or more enzymes, such as oxidase, superoxide enzyme, catalase [11-15]. Their catalytic reaction is similar to that of natural enzymes, which not only conforms to the kinetic curve of Michaelis-Menten equation but also is affected by temperature, substrate concentration, and pH [16].
The catalytic reaction of nanoenzymes occurs on the surface of nanomaterials, which is essentially the process of surface electron transfer; the size effects and surface active sites of these materials are key factors that affect the catalytic activity [17-20]. The kinetic analysis of enzyme reaction shows that the surface modification of nano-mimic enzyme can change its affinity to the substrate, thus affecting its catalytic performance [21,22]. Therefore, the size and surface modification of nanoenzymes have become an important mean of regulate the activity of nanoenzymes and to provide a new route for the further research and application of nanoenzymes.

As a type of nanomaterials, colloidal gold have also been widely studied in the field of nanoenzyme due to its unique optical properties. Li et al. found that colloidal gold can exhibit the activity of oxidase [23]. Luo et al. speculated that the reaction followed the Eley-Rideal mechanism [24]. In addition, JV et al. explored whether colloidal gold have the characteristics of intrinsic mimic peroxidase and applied them to glucose detection and studied the difference in peroxidase activity between gold nanoparticles modified by different surface charges and bare gold nanoparticles, and finally found that the peroxidase activity of bare gold nanoparticles was the highest through a series of characterization data [25].

In this study, reductants with obvious structural differences were used to prepare colloidal gold with similar particle size and the enzyme activity was detected. Through the optimization of the morphology and particle size of the colloidal gold, the characterization of a series of data, and the detection of enzyme activity, it was found that the enzyme activity of the enzyme activity of colloidal gold reduced by benzene ring was higher than that of colloidal gold enzyme reduced by linear chain. The above experimental results verify the differences in nanoenzyme activities with different surface modifications and provide a theoretical basis for the in-depth study and application of nanoenzyme surface modification.

1.2 Experiment methods:

1.2.1 Preparation of colloidal gold:

To avoid affecting the accuracy of the experimental results, it is necessary to prepare colloidal gold by reduction without any stabilizer. Certain concentrations of the reducing agent, pH regulator and chloroauric acid solution were added in a certain proportion and order, and nanoparticles with similar particle size and morphology were prepared by controlling the reaction conditions [26]. The specific proportioning methods are shown in Table1.

Table1: Proportion of reagents for preparing colloidal gold

| Reducing agent concentration | dDW$_2$O | HAucl$_4$ (1%) | NaOH(0.1M) | K$_2$CO$_3$(0.1M) | HCL(0.1M) | Reaction conditions |
|----------------------------|---------|---------------|------------|-----------------|---------|-------------------|
| ▲ 72.7 µL Hq (30mM)        | ● 9.696 mL | ■ 81.6 µL    | ★ 150 µL  | 50°C Stir       |         |                   |
1.2.2 Nano-enzyme activity verification

First, the activity of peroxidase-mimicking enzymes was preliminarily verified. It is generally believed that peroxidase-mimicking enzymes can catalyse the decomposition of H₂O₂ to produce free radicals and chemically react with a certain chromogenic substrate. In this study, TMB was used as the chromogenic substrate, and its oxidation product has an absorption peak at 652 nm. The activity of the peroxidase-mimicking enzyme was preliminarily verified according to the change of absorbance. The enzyme activity was measure after confirming that it has peroxidase activity, and a (IU·mg⁻¹) was calculated.

Table 2: Nano-enzyme activity determination verification

| Vial number | Reactant | NaAc-HAc buffer (mL) | Nanozyme (mL) | TMB (µL) | H₂O₂ (µL) |
|-------------|----------|----------------------|---------------|----------|-----------|
| Vial 1      |          | 2                    | 1.2           | 40       | 363.7     |
| Vial 2      |          | 2.364                | 1.2           | 40       | 0         |
| Vial 3      |          | 3.2                  | 0             | 40       | 363.7     |

1.2.3 Nanoenzyme activity determination and catalytic kinetics research

Different nanoenzyme solutions (1.2 ml) were added into vials containing 2 ml of a 0.2 M NaAc-HAc buffer (pH=3.6); Forty microliters of TMB solution (10 mg·mL⁻¹) was added into the vial and mixed. In addition to the above samples prepared, prepare a sample without adding H₂O₂ as a blank; the reaction is carried out in the dark at 35°C, and the absorbance at 652 nm is measured every 20 s; the relationship between the absorbance at 652 nm and the reaction time is plotted to obtain the reaction -Time curve.

The following formula was used to calculate a nanozyme (IU·mg⁻¹) [27]:

\[
b_{\text{nanozyne}} = \frac{V}{(\varepsilon \times l) \times \Delta A / \Delta t}
\]

\[
a_{\text{nanozyme}} = b_{\text{nanozyne}} / [m]
\]
By changing the concentration of the substrate \( \text{H}_2\text{O}_2 \) within a certain concentration range, the enzyme kinetics of the nanozyme was evaluated by the steady-state kinetic method. TMB was oxidized via the oxygen produced by catalytic decomposition to undergo a colour chromogenic reaction. As the concentration of the substrate increases, the reaction rate increases linearly and then tends to be saturated at a high concentration. This finding conforms to a typical Michaelis-Menten. The Michaelis curve was fitted with substrate concentration and initial catalytic velocity as the horizontal and vertical coordinates, respectively. Then, using the double reciprocal method, the Michaelis equation

\[
\frac{V}{V_{\text{max}}} = \frac{[S]}{(K_m + [S])}
\]

was constructed to calculate the reaction kinetic parameters. \( K_m \) is the Michaelis constant, \( V \) is the initial rate of the reaction, and \( [S] \) is the concentration of substrate. The Michaelis constant is a measure of the binding ability of the enzyme to the substrate. The smaller the \( K_m \) value is, the stronger the binding capacity to substrate, the greater the \( K_m \) value and the weaker the affinity.

2 Results and discussion

2.1 Optimization of nanoparticles

Due to the requirements of experimental research, spherical gold nanoparticles with identical size were synthetized. The characterization outcomes demonstrated that the colloidal gold, produced with the methods in the relevant literature [28-32], have different size and various morphologies, except for some linear chain organic acids (tartaric acid, malic acid, citric acid) and Catechol. Resorcinol cannot be rapid oxidative self-conversion into their quinone forms because of the lack of electronic resonance in aromatic nucleus; therefore, its slow reduction process, unlike other Hydroxylphenols, causes the colloidal gold to exhibit irregular diameters. The negative and positive charges of Hydroxylphenols, whose magnitudes also affect the ability to gain and lose electrons, are mainly distributed on the oxygen and aromatic ring. Compared with other hydroquinones, 1,4-hydroquinone has the strongest electron gain and loss ability. The colloidal gold fabricated using pyrogallic acid by reducing the corresponding metal ions (\( \text{Au}^{3+} \)) are affected by the quantity and position of their Hydroxyphenols, which makes their charge distribution uneven and ultimately leads to the uneven particle size. The formation of colloidal golds mediated by ascorbic acid occurs through releasing protons and electrons in an acidic medium, and the freed electrons eventually reduce \( \text{Au}(\text{III}) \) ions into colloidal golds, which are then covered by dehydroascorbic acid (DHAA) [33]. Clearly, the four alcoholic hydroxyl groups contained by nanoparticles are extremely reductive, as observed by the structure of the ascorbic acid, which ultimately results in the inhomogeneity of the colloidal gold sizes caused by the rapid reaction. In summary, there are two feasible main options to acquire colloidal golds with uniform particle size and spherical shape. The first is to add a stabilizer during the preparation process, and the second is to adjust the temperature and pH by altering the forms of the reducing agent and reaction rate. We
eventually chose the second option owing to realities. The results shown in Table 1 are the specific optimization methods.

2.2 Characterization of Nanoenzymes

It is generally accepted that both the temperature and pH or enzyme concentration have a serious influence on enzyme activity. On the basis of the experiment carried out at room temperature, with a pH 3.6, it can easily be discovered that the OD values of all kinds of colloidal gold are similar UV-vis spectroscopy (Figure 1), the results show that the concentrations of colloidal gold are extremely similar, which eliminates the impact of enzyme concentration.

![Figure 1: Visible light (400–700 nm) absorption spectra of colloidal gold (λ/ nm)](image)

On the other hand, the size and surface modification of colloidal gold have a significant impact on their activity from the perspective of nanomaterials, illustrating that it is vital to fabricate colloidal gold with analogical particle sizes. As shown in Table 1, the concentration and dosage of various reducing agents are different in that the agents demonstrate a large structural difference. First, it can be concluded from the maximum absorption peak of the UV-vis spectroscopy finding that there is little difference between nanoparticle size prepared by different reducing agents (Figure 1). The zeta potential shows that the surface charge of all colloidal gold is negative. Transmission electron microscopy (TEM) images also intuitively reflect this result (Figure 2).
2.2 Nanoenzymes activity verification

The activity of mimic enzyme was initially verified by virtue of the colour development reaction, and its catalytic ability was determined, as shown in Figure 3.

![Figure 3](image)

Figure 3: (a) Mechanism of peroxidase activity of the colloid gold (b) UV spectrogram and photo of nanoenzyme activity verification

The results of the activities of eight prepared nanoenzymes indicated that their catalytic effects were similar. Taking hydroquinone as a reducing agent as an example, as shown in Figure 3b, the clear solution in the No.1 bottle containing both nanoenzymes, TMB and H₂O₂, gradually turns blue, and the product has a strong absorption peak at 652 nm which is determined to be the absorption peak of TMB oxidation. This is due to H₂O₂ decomposed after being catalyzed by nanoenzymes to produce a large amount of O₂⁻ in the system, which oxidizes TMB to cause a colour development reaction so that it appears blue (Figure 3a). A slightly blue colour appeared in the solution of the second bottle containing nanoenzymes and TMB, and product also has a faint peak at 652 nm which is judged to be absorption peak of partial TMB oxidized by a small quantity of oxygen-containing substances adsorbed on the surface of nanoenzymes. On the basis of no-change in bottle 3 containing TMB and H₂O₂, H₂O₂ was not decomposed, and TMB was not oxidized when nanoenzymes were added to system. Conclusions: Nanoenzymes possess peroxidase catalytic activity. For the sake of unambiguous differences between
nanoenzymes fabricated by various reducing agents, the enzyme activity and enzymatic reaction kinetics were explored.

2.3 Study on activity determination and catalytic kinetics of nanoenzyme

2.3.1 Research on the most suitable conditions for nanoenzyme activity

Taking the preparation of colloidal gold by Sc reduction as an example, Through orthogonal design on different temperature and pH conditions, the optimal temperature and pH conditions are shown in the following Figure 4: Clearly, when pH is 3.6 and temperature is 40°C, the OD value (2.91) is the highest, which reflects the highest activity of nanoenzyme.

![Figure 4: Orthogonal analysis of pH and temperature of colloidal gold prepared by Sc reduction method](image)

2.3.2 Determination of nanoenzyme activity

It is well known that the peroxidase-like activity of colloidal gold results from their own. Before use, the prepared colloidal gold was centrifuged and then redispersed with ultrapure water to confirm the accuracy of the experiment and eliminate the influence of other impurities. H₂O₂ is decomposed into double HO⁺ radicals, which are adsorbed on the surface of AuNPs and stabilized by AuNPs through the interaction of partial electron exchange [34]. Finally, the catalytic performance of AuNPs was improved. In this experiment, the effects of reductants with different chemical structures on the properties of the prepared mimic enzymes were explored by investigating the enzyme activity and catalytic reaction kinetics of the mimic enzymes. According to the standard method provided by yan [27], the initial reaction rate under the condition of substrate saturation was selected as the standard to evaluate the enzyme activity. The change rate of absorbance with time is converted into that of concentration with time according to lambert-beer law. It is obvious that the activities of these eight
enzymes are in the following sequence, namely, Cc, Hq, Rs, Vc, Ga, Sc, Sm and St from high to low (Figure 7). The control variable method was used to analyse whether the difference of colloidal gold surface modification leads to the difference in the affinity with the substrate. The affinity of the enzyme to the two substrates, \( \text{H}_2\text{O}_2 \) and TMB, was compared. From the \( \text{Km} \) value, it is not difficult to find that when the substrate is \( \text{H}_2\text{O}_2 \), the \( \text{Km} \) value is in the order of Sm, St, Sc, Vc, Rs, Hq, Cc and Ga (Figure 5). It is not difficult to find that the affinity of the mimic enzyme for the substrate has an almost a linearly positive correlation with the reducibility of the reductant, and the stronger the reducibility of the simulated enzyme is, the greater the affinity of the mimic enzyme for the substrate. It can be seen from table 3 that the affinity of the mimic enzyme modified by benzene ring is strongest, and the affinity of the mimic enzyme modified by Vc is stronger than that of the mimic enzyme modified by linear chain. The surface properties of AuNPs will change as well. By affecting the absorption of \( \text{H}_2\text{O}_2 \), the electron transfer processes mediated by particles are different. The stronger the reducibility of the reductants is, the easier it is for the reductants to bind to the colloidal gold surface and the easier it is for charge transfer to occur, making the colloidal gold surface more active and facilitating \( \text{H}_2\text{O}_2 \) adsorption on the colloidal gold surface to form \( \text{HO}^- \) radicals [35]. The \( \text{Km} \) value magnitudes, when the substrate is TMB are in the following order: Hq, Rs, Cc, Ga, Vc, Sc, St, Sm (Table 4). The research shows that the activity of colloidal gold peroxidase was positively correlated with the affinity of the substrate hydrogen peroxide [1]. From the Table 5, the colloidal gold peroxidase activity modified by eight different reducing agents is in order: Cc, Hq, Rs, Vc, Ga, Sc, Sm, St(from high to low).

Figure 5: Michaelis–Menten curves for nanozymes prepared with eight different reducing agents, such as Cc, Hq, Rs, Vc, Ga, Sc, Sm and St. The concentration of TMB used was 1%, and the \( \text{H}_2\text{O}_2 \) concentration varied from 0 to nearly 30%. Error bars shown represent the s.e. derived from three independent experiments.
Table 3: $K_m$ and $V_{max}$ with $H_2O_2$ as substrate

| Type of reducing agent | Cc    | Hq    | Rs    | Ga    | Vc    | Sc    | Sm    | St    |
|------------------------|-------|-------|-------|-------|-------|-------|-------|-------|
| $K_m$(mM)              | 0.249 | 0.497 | 0.378 | 0.226 | 0.849 | 1.634 | 2.75  | 2.382 |
| $V_{max}$              | 6.596 | 6.057 | 6.12  | 5.385 | 6.897 | 4.228 | 3.785 | 6.369 |

Note: $K_m$ is the Michaelis constant, $V_{max}$ is the maximal reaction velocity.

Figure 6: Michaelis–Menten curves for nanozymes prepared with eight different reducing agents, such as Cc, Hq, Rs, Vc, Ga, Sc, Sm and St. The concentration of $H_2O_2$ used was 30 %, and the TMB concentration varied from 0 to nearly 1%. Error bars shown represent the s.e. derived from three independent experiments.

Table 4: $K_m$ and $V_{max}$ with TMB as substrate

| Type of reducing agent | Cc    | Hq    | Rs    | Vc    | Ga    | Sc    | Sm    | St    |
|------------------------|-------|-------|-------|-------|-------|-------|-------|-------|
| $K_m$(mg/ml)           | 0.0074| 0.0153| 0.0105| 0.003 | 0.0059| 0.0026| 0.0008| 0.0016|
| $V_{max}$              | 6.689 | 4.496 | 5.01  | 6.039 | 3.495 | 1.866 | 1.306 | 2.729 |

Note: $K_m$ is the Michaelis constant, $V_{max}$ is the maximal reaction velocity.

Table 5: The results of enzyme activity measurement

| Type of reducing agent | Cc    | Hq    | Rs    | Vc    | Ga    | Sc    | Sm    | St    |
|------------------------|-------|-------|-------|-------|-------|-------|-------|-------|
| Total                  |       |       |       |       |       |       |       | 4ml   |
| The amount nanozyme    |       |       |       |       |       |       |       | 500 µL|
| nanozyme concentration |       |       |       |       |       |       |       |       |
| (mg·ml$^{-1}$)         | 0.0816| 0.1   | 0.1   | 0.1   | 0.1   | 0.1   | 0.1   | 0.1   |
| b(IU)                  | 0.058 | 0.023 | 0.018 | 0.012 | 0.011 | 0.004 | 0.0023| 0.0022|
| a(IU·mg$^{-1}$)        | 1.415 | 0.464 | 0.378 | 0.242 | 0.219 | 0.084 | 0.047 | 0.044 |
However, why is there a difference in the end? (Figure 6) First, due to the resonance between the nonbonding PZ orbital of the phenolic hydroxyl oxygen atom and the \( \pi \)-bonding orbital of the benzene ring, the phenolic hydroxyl oxygen has a higher electron density [36]. And it was bound to the surface of colloidal gold, which greatly increased its reactivity. Second, as a radical scavenger, Vc competed with TMB, which resulted in the radicals HO’ not completely reacting with TMB. Finally, colloidal gold capped with carboxyl groups (citrate, etc.) tend to electrostatically attract the amino groups of TMB, moreover, the carboxyl moieties of linear chain organic acids are more easily adsorbed on the surface of colloidal gold, making the colloidal gold surface exposed moieties relatively less, leading to the relative difficulty of \( \text{H}_2\text{O}_2 \) adsorption on the colloidal gold surface. Compared with nanoparticles modified by other structures, its affinity with TMB is greater. The results showed that the affinity of the nanoenzyme prepared by reduction of phenolic compounds was higher than that of the nanoenzyme reduced by linear chain organic acids.

Figure 7: The initial linear part of the reaction–time curves of TMB colorimetric reaction catalyzed by colloidal gold nanoenzyme prepared with eight different reducing agents, such as Cc, Hq, Rs, Vc, Ga, Sc, Sm and St.

The IR spectra of the colloidal gold prepared by Vc, Sc, Sm and St reduction were analysed (Figure 8b), and the results showed that there was a \(-\text{OH}\) stretching vibration peak at 3449 cm\(^{-1}\). The results indicate that the peak intensity of Sc is the lowest, but the peak intensity of Sm is very strong. The reason is that it may contain water molecules. The wavenumbers 2984 cm\(^{-1}\) and 2980 cm\(^{-1}\) belong to C-H stretching vibration peak on saturated carbon chain, 1750-1680 cm\(^{-1}\) belong to carbonyl stretching vibration peak, 1422 cm\(^{-1}\),1314 cm\(^{-1}\) and 1039 cm\(^{-1}\) mainly belong to C-H in-plane bending vibration and C-C single skeleton vibration. However, in the process of preparing colloidal gold, the ring of Vc is opened and
the double bond is reduced, resulting in the two kinds of infrared characteristic peaks not appearing. To further study the effect of the amount and position of carboxyl and hydroxyl groups in linear chain organic acids on the enzyme activity, three linear chain organic acids with insignificant structural differences (tartrate, citrate, malate) were used as reductants to prepare colloidal gold, and the enzymatic activity of colloidal gold was examined. The detection results are shown in Figure 8c, with the order of Sodium citrate > Sodium malate > Sodium tartrate, Citric acid is tricarboxylic acid, while malic acid and tartaric acid are dicarboxylic acids. Malic acid has one fewer alcohol hydroxyl group than tartaric acid. Under conditions with the same carbon skeleton, tartaric acid has one more hydroxyl group than malic acid, which eventually leads to the hydroxyl radicals generated from hydrogen peroxide adsorbed on the surface of colloidal gold being more easily eliminated. Malic acid has one fewer carboxyl group than citric acid. Although they are both @ - hydroxyl group, due to bond angle differences, citric acid is less active in hydroxyl groups than malic acid, ultimately leading to the generation of hydroxyl radicals that better bind to TMB.

Figure 8: (a) The molecular structure of Vc, Sc, Sm and St and (b) the infrared spectrum of the reduction to prepare colloidal gold (c) Part of Figure 7.

The colloidal gold prepared by four kinds of phenol reduction method were analyzed by FTIR (Figure 9b), It is not difficult to find that there is a broad peak at 3417 cm⁻¹, which can be attributed to the stretching vibration peak of water molecules, C-H stretching vibration peak at 2991 cm⁻¹ and 2900 cm⁻¹, the quinone carbonyl stretching vibration peak at 1667 cm⁻¹ and benzene absorption peak at 1426 cm⁻¹ and 1314 cm⁻¹. However, no characteristic peaks of four phenols were found in the fingerprint area, indicating that all phenols participated in the reduction and stabilization of colloidal gold. Experimental studies on Cc, Rs, Hq were carried out mainly to comparatively analyse the differences in the nanzyme activity of prepared colloidal gold from the perspective of isomers. Due to their different molecular structures, the ease of their redox reactions varies. The hydroxyl group on the benzene ring belongs to electron withdrawing group, but because its conjugation effect is greater than its electron withdrawing ability, the hydroxyl group will eventually act as electron donating group. The two
hydroxyl groups on the phenyl ring differ in position and therefore in the distribution of charge density, which leads to differences in their degree of redox. Among the three isomers, when the two hydroxyl positions are in the para position of benzene ring, the charge density is the highest, followed by the ortho position and the meta position. However, because the phenolic hydroxyl group of catechol is in the ortho position, their electron cloud densities influence each other, which greatly enhances the charge effect. Finally, the colloidal gold prepared by catechol reduction method had the highest surface activity, followed by hydroquinone and resorcinol.

Figure 9: (a) The molecular structure of Cc, Hq, Rs, Ga and (b) the infrared spectrum of the reduction to prepare colloidal gold (c) Part of Figure 7.

Conclusion

Colloidal gold with similar particle size was prepared by optimization processes, and the enzyme activity of eight different surface-modified colloidal gold was verified. The results showed that the enzyme activity of colloidal gold reduced by benzene ring is higher than that of colloidal gold enzyme reduced by linear chain. The effects of isomers, number, and location of functional groups on the enzymatic activity of colloidal gold prepared with three different hydroquinones and three kinds of linear chain organic acids as reductants are discussed separately. Hydroquinone affects the reactivity of the colloidal gold surface, resulting in a different number of hydroxyl radicals produced, while linear chain organic acids affect the reaction efficiency of hydroxyl radical and TMB, resulting in the difference in enzyme activity.
• Ethics approval and consent to participate
Not applicable
• Consent for publication
Not applicable
• Availability of data and materials
The data and materials that support the findings of this study are available from the corresponding author upon reasonable request.
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Feng Shi provided the experimental platform, Junjun Cao carried out the Conception and design of the research, Jie Zhang participated in the Acquisition of data. Ning Yuan carried out the Analysis and interpretation of data. Junjun Cao and Yixiao Sun participated in the design of the study and performed the statistical analysis. Junjun Cao and Zhihua Xu conceived of the study, and participated in its design and coordination and helped to draft the manuscript and revision of manuscript for important intellectual content. All authors read and approved the final manuscript.
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