Enhancing the specificity of polymerase chain reaction by graphene oxide through surface modification: zwitterionic polymer is superior to other polymers with different charges

Yong Zhong1,*
Lihong Huang2,*
Zhisen Zhang3
Yunjing Xiong4
Liping Sun1
Jian Weng1

1Department of Biomaterials, College of Materials, 2State Key Laboratory of Cellular Stress Biology, School of Life Sciences, 3Department of Physics, Research Institute for Biomimetics and Soft Matter, Xiamen University, Xiamen, People’s Republic of China
4These authors contributed equally to this work

Abstract: Graphene oxides (GOs) with different surface characteristics, such as size, reduction degree and charge, are prepared, and their effects on the specificity of polymerase chain reaction (PCR) are investigated. In this study, we demonstrate that GO with a large size and high reduction degree is superior to small and nonreduced GO in enhancing the specificity of PCR. Negatively charged polyacrylic acid (PAA), positively charged polyacrylamide (PAM), neutral polyethylene glycol (PEG) and zwitterionic polymer poly(sulfobetaine) (pSB) are used to modify GO. The PCR specificity-enhancing ability increases in the following order: GO-PAA < GO-PAM < GO-PEG < GO-pSB. Thus, zwitterionic polymer-modified GO is superior to other GO derivatives with different charges in enhancing the specificity of PCR. GO derivatives are also successfully used to enhance the specificity of PCR for the amplification of human mitochondrial DNA using blood genomic DNA as template. Molecular dynamics simulations and molecular docking are performed to elucidate the interaction between the polymers and Pfu DNA polymerase. Our data demonstrate that the size, reduction degree and surface charge of GO affect the specificity of PCR. Based on our results, zwitterionic polymer-modified GO may be used as an efficient additive for enhancing the specificity of PCR.

Keywords: PCR additive, charge, pSB, Pfu DNA polymerase

Introduction
Polymerase chain reaction (PCR) is one of the most ubiquitous and well-developed tools in molecular biology.1,2 Due to its ability to produce billions of DNA copies by rapid and selective amplification of a specific region in the DNA chain, it has a wide range of applications in gene amplification, molecular cloning and disease diagnosis.3 A basic PCR usually requires the following five components: 1) DNA template containing the target DNA region, 2) two primers to initiate DNA synthesis, 3) a thermostable DNA polymerase to catalyze DNA synthesis, 4) deoxyribonucleoside triphosphates (dNTPs, the building blocks of new DNA strand) and 5) buffer including bivalent cations (usually Mg2+).4 Often a time-consuming optimization process may be required in order to obtain a successful PCR, primarily when amplification from genomic DNA is required.

Many factors will affect the specificity of PCR, such as the primer purity and sequence, purity of the template DNA, Mg2+ concentration, annealing temperature and other additives such as dimethyl sulfoxide (DMSO), which are frequently included
in the PCR mixture.\textsuperscript{5,6} Usually, when the concentration of template DNA is very low, or the structure of DNA template is very complicated, such as GC-rich gene or mammalian genomic DNA, the specificity of PCR might be very low. Different strategies have been developed to improve the specificity of PCR, such as nested PCR using two separate sets of primers, touchdown PCR with gradually decreased annealing temperatures and hot start PCR withholding the DNA polymerase or the primers until the reaction mixture has reached a temperature above the threshold for nonspecific binding of primer to template.\textsuperscript{4} However, optimizing the primer design, PCR cycling conditions and the annealing temperatures and adjusting the concentrations of PCR components are lengthy and frustrating processes. To enhance the specificity of PCR, several small molecules have been used as additives, such as DMSO, glycerin, betaine, formamide and bovine serum albumin (BSA).\textsuperscript{7,8} Using them either singly or in combination does not guarantee enhancing the specificity of PCR. Thus, the effect of these additives is unpredictable. Therefore, it remains a challenge to find new additives to enhance the specificity of PCR.

In recent years, nanomaterials have been used in PCR system, including metal nanoparticles, semiconductor quantum dots, carbon nanomaterials and polymer nanoparticles.\textsuperscript{9-19} Nanomaterials often possess unique physical and chemical properties, eg, surface effect arising from their large surface-to-volume ratio, which differ greatly from macroscopic materials. The effect of nanomaterials on PCR mainly depends on the strong interaction between PCR components and nanomaterials.\textsuperscript{20} However, conflicting results were obtained from different research groups.\textsuperscript{21-25} Therefore, the effect of nanomaterials with different surface characteristics on PCR is still unclear and much more systemic and deep investigations are necessary to understand the corresponding mechanism.

Our previous study has demonstrated that reduced graphene oxide (RGO) can improve the specificity of PCR significantly.\textsuperscript{19} However, RGO has poor solubility in water due to its shortage of functional groups on its surface, which inhibits the further application of RGO. In comparison with RGO, there are many hydroxyl, epoxy and carboxyl groups on graphene oxide (GO) sheet, which result in high polarity and hydrophilicity of GO. Thus, GO is hydrophilic and can be well dispersed in water. Further surface modification of GO can provide GO with some new surface properties, such as size, reduction degree and surface modification, on the specificity of PCR. We found that large and RGO was superior to small and non-RGO in enhancing the specificity of PCR. The surface charge of GO also affected the specificity of PCR in the following order: GO-PAA < GO-PAM < GO-PEG < GO-pSB. For polymer-modified GO, the zwitterionic pSB was found to be superior in enhancing the specificity of PCR compared to neutral and negatively and positively charged polymers. These GO derivatives with different surface charges have been used successfully as additives in amplification of the mitochondrial DNA using blood genomic DNA as template. Based on our results, we suggest that GO derivatives can also enhance the specificity of PCR of low abundance DNA obtained from clinical samples. Molecular dynamics simulations and molecular docking are further used to investigate the interaction between the polymers and \textit{Pfu} DNA polymerase for understanding the mechanism of specificity enhancing. Our research results may ultimately progress the application of GO in biomedical fields.

Materials and methods

Materials

Natural graphite powder (320 mesh, Alfa, 99%) was purchased from Tianjin Guangfu Chemical Agent Co., Ltd. (Tianjin, People’s Republic of China). NaNO\textsubscript{3}, KMnO\textsubscript{4}, K\textsubscript{2}PO\textsubscript{4} (99%), L-lysine (98%), ninhydrin (97%) and H\textsubscript{2}O\textsubscript{2} were purchased from Guoyao Co., Ltd. (Shanghai, People’s Republic of China). Acrylic acid (AA), acrylamide (AM), H\textsubscript{2}SO\textsubscript{4}, HCl, hydrazine hydrate (80%), ethanol, NaHCO\textsubscript{3}, NaOH, Mg\textsubscript{2}SO\textsubscript{4} (99%), methyl-red and sodium borate were purchased from Xilong Chemical (Guangzhou, People’s Republic of China). Potassium peroxosulfate was purchased from Dahao Nanhong Industrial Co., Ltd. (Shantou, People’s Republic of China). mPEG-NH\textsubscript{2} was purchased from Kailzheng Biotech Development Co., Ltd. (Beijing, People’s Republic of China). 1-Ethyl-3-[3-dimethylaminopropyl] carbodiimide hydrochloride (EDC, 98%) was purchased from Aladdin Industrial Corporation (Shanghai, People’s Republic of China). BSA (98%) and [2-(methacryloyloxy) ethyl] dimethyl-(3-sulfopropyl) ammonium hydroxide (SB, 98%) were purchased from Sigma-Aldrich. All reagents used were of analytical-reagent grade except specially stated. The dialysis tubing (8,000–14,000 MWCO) was purchased from Solarbio Life Sciences (Beijing, People’s Republic of China). The pET-32a plasmid DNA was purchased from Novogen, Shanghai, People’s Republic of China. The P\textit{fu} polymerase, dNTP, 10× buffer, 0.22 μm filter, agarose, glycerol, Luria-Bertani broth, ethidium bromide (EB), DNA
marker, DNA-loading buffer, anhydrous CaCl₂, PCR tubes, 50× TAE buffer and ampicillin were purchased from Sangon Biotech Co., Ltd. (Shanghai, People’s Republic of China). The primers were synthesized by Sangon Biotech Co., Ltd. (Table S1). The PCR products were sequenced by Sangon Biotech Co., Ltd. Mini Plasmid Kit was purchased from TIANGEN Biotech Co., Ltd. (Beijing, People’s Republic of China). High Pure PCR Template Preparation Kit was purchased from Roche Ltd. (Germany). Deionized water was used in all experiments.

**Synthesis of GO and its derivatives**

GO was synthesized according to our previous method. First, 2.0 g of graphite powder and 1.0 g of NaNO₃ were added to 46 mL of 98% H₂SO₄ in an ice bath, and then 6.0 g of KMnO₄ was added slowly into the mixture with vigorous agitation. The mixture was stirred continually for 2 hours in the ice bath and placed in a water bath at 35°C for 30 minutes. Then 92 mL of water was gradually added, and the temperature was raised to 95°C. The solution was stirred for 3 hours. Finally, 400 mL of water was added, and 6 mL of 30% H₂O₂ was dropped into the reaction. After cooling to room temperature, the mixture was filtered and washed with 1:10 HCl (volume ratio), followed by repeated washing with water to remove the acid. The filter cake was then dispersed in water to obtain graphite oxide. Exfoliation of graphite oxide to GO was achieved by ultrasonication for 3 hours at 500 W. The aggregate was removed by centrifugation at 4,000 rpm for 10 minutes. The supernatant was collected and dialyzed in water for 1 week to remove residual salts.

Small GO (S-GO) and large GO (L-GO) were prepared according to the literature. The dialyzed GO solution was sonicated for 10 hours, followed by centrifugation at 12,000 rpm for 20 minutes. The supernatant was collected as S-GO. The aggregate was dispersed in water again and centrifuged at 2,000 rpm for 10 minutes. The supernatant was collected as L-GO. RGO was prepared from GO, which was reduced by hydrazine hydrate in ammonia. A total of 50 mL of 1 mg mL⁻¹ GO solution was added to 4 mL of 80% hydrazine hydrate and 100 μL of 28% ammonium hydroxide were added. After thorough mixing, the solution was heated in an oil bath at 95°C for 1 hour. The supernatant was collected and dialyzed for a week in water. Another sample of RGO with a higher reduction degree was prepared by adding 42 μL of 80% hydrazine hydrate and 700 μL of 28% ammonium hydroxide.

GO-PEG was prepared according to the literature. First, 10 mL of 3 M NaOH was mixed with 20 mL of 2 mg mL⁻¹ GO solution and sonicated for 3 hours. Then 10% HCl was added to neutralize the solution to pH 7. The mixture was filtered and rinsed with water. The filter cake was then dispersed in 40 mL of water. A total of 80 mg mPEG-NH₂ was added in the solution and sonicated for 5 minutes. Then 20 mg of EDC was added and sonicated for another 30 minutes, followed by the addition of 60 mg of EDC and overnight stirring. The supernatant was collected and dialyzed for a week in water to remove excess mPEG-NH₂. After 30 minutes of centrifugation at 4,000 rpm, the supernatant was collected as GO-PEG.

**Characterization of GO and its derivatives**

Atomic force microscopy (AFM) images were obtained in the tapping mode in air using Nanoscope Multimode 8 (Veeco Instruments Inc., USA). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The Fourier transform infrared (FTIR) spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China). The UV–vis absorption spectra were obtained using a TU-1901 UV–Vis spectrometer (Beijing Purkinje General Instrument, People’s Republic of China).
PCR and agarose gel electrophoresis

pET-32a plasmid was used as the template of PCR. The following components were mixed in the first round of PCR: 1× PCR buffer containing 2 mM Mg²⁺, dNTP (each 0.2 mM), primer F1 (0.4 μM), primer R1 (0.4 μM) (Table S1), plasmid DNA (0.5 ng μL⁻¹) and Pfu polymerase (0.1 U μL⁻¹). GO or its derivatives with different concentrations were added in the PCR system. Each sample was made up to the final volume of 25 μL with water. In the second-round PCR, the same reaction condition was set, except that 0.5 μL of PCR product was used to substitute the plasmid DNA as the template. The PCR conditions are described in Table S2.

The study was approved by the Ethical Committee of the Medical College, Xiamen University. Blood samples were collected from healthy volunteers who provided written informed consent. Human genomic DNA was isolated using High Pure PCR Template Preparation Kit. A total of 20 ng μL⁻¹ DNA was added instead of plasmid DNA. Primers F2 and R2 (Table S1) were used to amplify a fragment of human mitochondrial DNA. The PCR components were the same as above, except that plasmid DNA was replaced with 2.5 μL of genomic DNA. In the second- or third-round PCR, 2.5 μL of last-round PCR product was used as the template instead of genomic DNA. The PCR conditions are listed in Table S2. PCRs were performed in a T3 thermocycler (Biometra, Germany). The PCR products were electrophoresed in 1.5% agarose gel for 30 minutes, stained with 0.5 μg mL⁻¹ EB and analyzed using a Gel Doc XR system (Bio-Rad, USA). The fluorescence intensity of the band of target was measured by Image Lab software.

Results and discussion

Preparation and characterization of GO and its derivatives

Figure 1 demonstrates the preparation of S-GO, L-GO, GO-PAA, GO-PEG, GO-PAM and GO-pSB. GO was synthesized from the natural graphite using modified Hummers method. S-GO and L-GO were prepared with high-speed centrifugation technology. The AFM images showed that the average size of S-GO (50 nm) was much smaller than that of L-GO (500 nm), while the thickness was ~0.9 nm for both samples (Figure S2). The result shows that both S-GO and L-GO are monolayer sheets with same thickness. Thus, we can exclude the effect of GO thickness on the specificity of PCR in the latter experiments. Grafting polymer onto the surface of GO resulted in the thickness increasing to 1–3 nm (Figure S2C–F). The zeta potential of S-GO (~45.1±0.6 mV) was lower.
than that of L-GO (−35.6±0.8 mV), which means that the content of carboxyl groups in S-GO is higher than that of L-GO, and the result is consistent with the results from Boehm titrations.

GO-PAA, GO-PAM and GO-pSB were prepared by free-radical polymerization of AA, AM and SB monomers on GO surface, respectively. On the other hand, mPEG-NH₂ was conjugated to the carboxyl groups on GO via carbodiimide-catalyzed amide formation. Ultraviolet–visible (UV–vis) spectra (Figure S2), FTIR spectra (Figure S4), XRD spectra (Figure S5) and XPS spectra (Figure S6) were used to characterize all samples and listed in the Supplementary materials. All these characterizations indicated that PAA, PAM, PEG and pSB were successfully grafted on GO. TGA of GO, GO-PAA, GO-PAM, GO-PEG and GO-pSB is shown in Figure S7. TGA of GO with 8.39% carboxyl content
exhibits two regions of mass loss. The first weight loss at temperature \(<100^\circ\text{C}\) is attributed to the loss of adsorbed water. The second weight loss between 180°C and 230°C is attributed to the decomposition of the oxygen-containing functional groups, such as carboxyl, epoxide and hydroxyl groups, in GO. The weight loss of graphene decreases with decreasing the content of carboxyl (Figure S7A). Compared to GO, the second weight loss of GO derivatives happens in a higher temperature, which mainly results from the decomposition of polymeric chain. By calculating the weight loss percentage, the calculated contents of PAA, PAM, PEG and pSB were 43.48\%, 40.96\%, 42.53\% and 38.70\%, respectively (Table S3). These polymer-modified GOs have similar content of polymer. Thus, we can exclude the effects of polymer content on the specificity of PCR in the latter experiments.

The zeta potentials of GO and RGO were \(-39.1\pm0.5\) mV and \(-15.6\pm0.3\) mV, respectively, indicating that GO was more stable than RGO. After grafting polymers onto the surface of GO, the zeta potential of GO-PAA \((-46.9\pm0.9\) mV) became more negative due to the negative charge of COO\(^-\) in PAA. The zeta potential of GO-PAM increased to \(-24.1\pm1.2\) mV due to positively charged NH\(_2\)+ in PAM. The negative potential of GO-PEG \((-35.7\pm0.8\) mV) and GO-pSB \((-20.4\pm0.9\) mV) resulted from the COO\(^-\) groups on GO surface, even though PEG was a neutral polymer and pSB was a zwitterionic polymer (Tables S4 and S5).

The effect of GO size on PCR

A common process of PCR includes 25–30 cycles of repeated temperature change. A typical PCR cycle consists of the following three steps: 1) denaturation of the template DNA, 2) annealing of two synthetic primers to the template DNA and 3) extension of the bound primers catalyzed by a thermostable DNA polymerase. Although PCR can multiply the target gene exponentially, nonspecific amplification remains a problem. Especially in the multiple-round PCR, the unreacted initial DNA template and primers disturb the reaction and cause unspecific products accumulation. In the initial studies, we used pET-32a plasmid as the template of PCR amplification. The optimal template concentration was determined by serial dilution experiments (Figure S8). A total of 0.5 ng \(\mu\text{L}^{-1}\) plasmid was considered as the optimal template concentration in the first-round PCR, and 0.5 \(\mu\text{L}\) of the first-round PCR product was the optimal template concentration in the second-round PCR. The effect of GO with different sizes (S-GO and L-GO) on the two rounds of PCR was investigated. In the first-round PCR, the intensity of the 689 bp target band decreases with increasing the GO concentration, indicating that the reaction is inhibited by both S-GO and L-GO at high concentration (4.8 \(\mu\text{g mL}^{-1}\) for S-GO and 6.4 \(\mu\text{g mL}^{-1}\) for L-GO; Figure 2A–C). In the second-round PCR, the first-round PCR product was used as the template. Since the PCR product contains not only the target product but also the unreacted initial DNA template, primers and unspecific products, these substances would disturb the second-round PCR and cause the accumulation of nonspecific products, leading to obvious smear bands without GO (Figure 2D and E). However, after adding S-GO or L-GO, the smear bands disappeared. The nonspecific products were decreased with increasing the concentration of S-GO and L-GO. L-GO had a better effect than S-GO, because it had a wider efficient concentration range to enhance the specificity of PCR (1.6–3.2 \(\mu\text{g mL}^{-1}\) for S-GO versus 0.8–4.8 \(\mu\text{g mL}^{-1}\) for L-GO; Table S6).

Our working hypothesis for the observed GO concentration effect is as follows. When the concentration of GO is low, the adsorbed amount of positively charged \(Pfu\) polymerase and Mg\(^{2+}\) on negatively charged GO surface was relatively high. Then negatively charged molecules, such as DNA template, primers and dNTPs, are attracted by positively charged \(Pfu\) polymerase on GO surface. Thus, the probability of mismatch is decreased and the specificity of PCR is improved. However, when further increasing the GO concentration, the amount of negatively charged GO increases in the PCR system, so the adsorbed amount of positively charged \(Pfu\) polymerase and Mg\(^{2+}\) on single GO surface decreases. Moreover, the excess negative charges of GO will repel the negatively charged primers and dNTPs, leading to suppression of PCR. We measured the zeta potentials of GO and its mixture with \(Pfu\) polymerase in water (Table S4). The zeta potential of S-GO increased from \(-45.1\pm0.6\) mV to \(-33.2\pm0.9\) mV, while the zeta potential of L-GO increased from \(-35.6\pm0.8\) mV to \(-29.8\pm1.1\) mV, confirming a stronger electrostatic interaction between S-GO and \(Pfu\) polymerase than that of L-GO and \(Pfu\) polymerase. The similar trend is obtained in PCR buffer (Table S5). The result is attributed to higher content of negatively charged carboxyl groups in S-GO (9.01\%) than that of L-GO (8.17\%). Therefore, S-GO inhibits PCR at a lower concentration (1.6 \(\mu\text{g mL}^{-1}\)) than L-GO (6.4 \(\mu\text{g mL}^{-1}\)) and L-GO has a wider concentration range to enhance the specificity of PCR.

The effect of GO reduction degree on PCR

We prepared RGO using hydrazine as the reductant in ammonia. Boehm titrations and TGA (Figure S7A) showed that the carboxyl content was reduced from 8.39\% to 5.43\%
Enhancing the specificity of PCR by GO through surface modification

following reduction. The zeta potentials were increased from $-39.1\pm0.5$ mV to $-15.6\pm0.3$ mV after reduction. In the first-round PCR, the bands intensity decreased with increasing the concentrations of GO and RGO (Figure 3A–D). At the same time, the inhibition effect was decreased with reducing the content of carboxyl groups on GO. In the second-round PCR, obvious smear bands appeared and decreased with increasing the concentrations of GO and RGO (Figure 3A–D). The efficient concentration range for optimal specificity of PCR of GO (8.39%), RGO (7.28%) and RGO (5.43%) is $0.8–3.2 \mu g m L^{-1}$, $3.2–4.8 \mu g m L^{-1}$ and $0.8–6.4 \mu g m L^{-1}$, respectively. The specificity of PCR increases and the inhibition effect was decreased with increasing the concentrations of GO and RGO (Figure 3A–D). The efficient concentration range for optimal specificity of PCR decreases with decreasing carboxyl content in GO, demonstrating that the carboxyl groups on GO have significant effect on PCR. RGO has fewer negative charges than GO, resulting in less electrostatic repulsion with the negatively charged PCR components. In addition, a stronger $\pi$-$\pi$ stacking interaction between the ring structures of the nucleobases and the polyaromatic hydrocarbon structure of RGO leads to greater affinity of RGO with DNA strands than that of GO with DNA. RGO has more perfect hexagonal lattice than GO; thus, it has stronger hydrophobic and $\pi$-$\pi$ stacking interactions with Pfu polymerase than GO with Pfu polymerase. The Pfu polymerase is composed of 775 amino acids, including 26 phenylalanines, 47 tyrosines and 11 tryptophans, that all these molecules contain benzene rings. The rich nonpolar benzenes of these aromatic amino acids can bind on the graphene sheet strongly via $\pi$-$\pi$ interaction. Therefore, RGO is superior to GO in enhancing the specificity of PCR due to less electrostatic repulsion with the negatively charged PCR components and higher affinity with DNA template and Pfu polymerase. The electrostatic absorption between positively charged Pfu polymerase and RGO is smaller than that between Pfu polymerase and GO with more negative charges, and hydrophobic interaction is the driving force for enzyme adsorption onto RGO. Thus, RGO has stronger affinity with Pfu polymerase than GO. This conclusion was further supported by the isothermal titration calorimetry (ITC) analysis (Table S7 and Figure S9). The dissociation constants ($K_d$) for graphene–Pfu polymerase interaction were $12.3 \text{ mol}^{-1}$, $10.5 \text{ mol}^{-1}$ and $3.96 \text{ mol}^{-1}$ for GO (8.39% carboxyl), RGO (7.28% carboxyl) and RGO (5.43% carboxyl), respectively. Negative values of Gibbs free energy change ($\Delta G$) confirmed the spontaneous binding of graphene

Figure 2. Effect of GO with different sizes on the first- and second-round PCRs.

Notes: M: DNA marker. (A) S-GO in the first-round PCR, (B) L-GO in the first-round PCR, (C) PCR band intensity at different concentrations of GO in the first-round PCR, (D) S-GO in the second-round PCR, (E) L-GO in the second-round PCR, (F) PCR band intensity at different concentrations of GO in the second-round PCR. The GO concentration in lanes 1–7 is $0 \mu g m L^{-1}$, $0.8 \mu g m L^{-1}$, $1.6 \mu g m L^{-1}$, $3.2 \mu g m L^{-1}$, $4.8 \mu g m L^{-1}$, $6.4 \mu g m L^{-1}$ and $8.0 \mu g m L^{-1}$, respectively.

Abbreviations: GOs, graphene oxides; L-GO, large GO; PCRs, polymerase chain reactions; S-GO, small GO.
with *Pfu* polymerase.\(^{36,47}\) The more RGO (5.43%) had the lowest $K_a$ and more negative $\Delta G$ values, indicating higher affinity between RGO and *Pfu* polymerase than between GO and *Pfu* polymerase.

### The effect of GO surface modification on PCR

To further investigate the effect of GO surface modification on the specificity of PCR, we modified GO with negatively charged PAA, positively charged PAM, neutral PEG and zwitterionic pSB. In the first-round PCR, no smear bands were found for all samples, indicating good specificity of PCR (Figure 4A–E). In the second-round PCR, nonspecific smear appeared in the control experiments without additives, but the smears decreased with increasing the concentrations of GO derivatives, indicating that the specificity of PCR is enhanced in all GO derivative-assisted PCR systems. Compared to GO-PAA, GO-PAM and GO-PEG, GO-pSB has lesser smears decreased with increasing the concentrations of GO and RGO in lanes 1–7 is 0 $\mu$g mL\(^{-1}\), 0.8 $\mu$g mL\(^{-1}\), 1.6 $\mu$g mL\(^{-1}\), 3.2 $\mu$g mL\(^{-1}\), 4.8 $\mu$g mL\(^{-1}\), 6.4 $\mu$g mL\(^{-1}\) and 8.0 $\mu$g mL\(^{-1}\), respectively. Abbreviations: GO, graphene oxide; RGO, reduced graphene oxide; PCR, polymerase chain reaction; TGA, thermal gravimetric analysis.

In order to confirm that enhancing the specificity of PCR results from the polymer-modified GOS and not from the polymers, we investigated the effect of bare polymers on the specificity of PCR without GO. The calculated amounts of polymers according to the TGA data of GO derivatives were added into PCR system. The results show that there is no change in the band (Figure S10). Only when the concentration of polymer was increased >1,000-fold from microgram per milliliter to milligram per milliliter, the specificity of PCR could be improved (Figure S11). These results indicate that enhancing the specificity of PCR results from the polymer-modified GO and not from the bare polymers. Based on previous results, GO is not a good enhancer in the specificity of PCR. Thus, enhancing the specificity of PCR by GO derivative is attributed to the combination or synergistic effect of GO and polymers.
Figure 4  Effect of GO derivatives on the first- and second-round PCRs.

Notes: M: DNA marker. The concentration of GO derivative in lanes 1–8 in (A), (B), (F) and (G) is 0 μg mL⁻¹, 0.8 μg mL⁻¹, 1.6 μg mL⁻¹, 3.2 μg mL⁻¹, 4.8 μg mL⁻¹, 6.4 μg mL⁻¹, 8.0 μg mL⁻¹ and 9.6 μg mL⁻¹, respectively. The concentration of GO derivative in lanes 1–6 in (C), (D), (H) and (I), is 0 μg mL⁻¹, 4.8 μg mL⁻¹, 9.6 μg mL⁻¹, 14.4 μg mL⁻¹, 19.2 μg mL⁻¹, and 24.0 μg mL⁻¹, respectively. The contents of PAA, PAM, PEG and pSB in GO derivatives are 43.48%, 40.96%, 42.53% and 38.70%, respectively. (E) PCR band intensity at different concentrations of GO derivatives in 1st round PCR. (J) PCR band intensity at different concentrations of GO derivatives in 2nd round PCR.

Abbreviations: GO, graphene oxide; PAA, polyacrylic acid; PAM, polyacrylamide; PCRs, polymerase chain reactions; PEG, polyethylene glycol; pSB, poly(sulfobetaine).
Figure 5 shows that at high concentrations of GO or GO derivatives, the control bands (lane 1) are inhibited. In order to test if *Pfu* polymerase is being adsorbed onto the surfaces of GO and its derivatives, BSA was added into the PCR system inhibited by excess GO and GO derivatives. After the addition of BSA, all PCRs are recovered and the intensities of band increase with increasing concentration of BSA. The result is attributed to the competitive adsorption of BSA on the surface of GO and its derivatives. Thus, the adsorbed *Pfu* polymerase was released into the solution, resulting in the recovery of PCR. The results indicate that there is strong interaction between GO or GO derivatives and *Pfu* polymerase. We also incubated GO and its derivatives with *Pfu* polymerase for 1 hour in ice to allow sufficient absorbance of *Pfu* polymerase with GO or its derivatives. Then we added other PCR components and performed PCRs.

Figure 5 confirms efficient PCR following adsorption of *Pfu* polymerase on GO or its derivatives.

PCR was inhibited when the concentration of GO derivative was further increased (Figure 4). Each GO sheet can adsorb positively charged *Pfu* polymerase or Mg\(^{2+}\) due to its large surface area and many negatively charged carboxyl groups. Increasing the concentration of GO derivative will decrease the adsorption amount of *Pfu* polymerase and Mg\(^{2+}\) on single GO sheet, resulting in less PCR efficiency. When excess Mg\(^{2+}\) is added into the PCR system, more Mg\(^{2+}\) will be further adsorbed on negatively charged GO derivatives with *Pfu* polymerase to recover PCR. Figure 6 shows that the inhibited bands are recovered after the addition of 1–2 mM Mg\(^{2+}\). Therefore, the results further indicate that PCR will be inhibited if the *Pfu* polymerase and Mg\(^{2+}\) adsorbed on single GO derivative decrease.

Figure 6 confirms efficient PCR assisted by GO and its derivatives.

Notes: M: DNA marker. (A) GO (9.6 μg mL\(^{-1}\)). (B) GO-PAA (4.8 μg mL\(^{-1}\)). (C) GO-PAM (8.0 μg mL\(^{-1}\)). (D) GO-PEG (24.0 μg mL\(^{-1}\)) and (E) GO-pSB (24.0 μg mL\(^{-1}\)). The concentration of added Mg\(^{2+}\) in lanes 1–4 is 0 mM, 0.4 mM, 1.0 mM and 2.0 mM, respectively (the original 2 mM Mg\(^{2+}\) was not included).

Abbreviations: GO, graphene oxide; PAA, polyacrylic acid; PAM, polyacrylamide; PCR, polymerase chain reaction; PEG, polyethylene glycol; pSB, poly(sulfobetaine).
Pfu polymerase possesses a proofreading activity and can be used instead of Taq polymerase to perform higher fidelity of DNA synthesis. To confirm whether GO or its derivatives would affect the fidelity of PCR, all PCR products in the absence or presence of GO and its derivatives were sequenced. The data confirm that GO and its derivatives have no effect on the fidelity of Pfu polymerase (Figure S13). The results further indicate that the specificity of PCR is enhanced and fidelity of Pfu polymerase is guaranteed in the presence of GO derivatives.

The application in clinical samples

Finally, GO derivatives were also used to enhance the specificity of PCR for amplifying the complicated clinical samples. Human blood genomic DNA was used as the template. A 324 bp fragment of the mitochondrial DNA was amplified. In the first- and second-round PCRs, no smear bands are observed for all samples, but the target band is weaker for the second-round PCR without GO derivative than that of the sample with GO derivatives (Figure 7). In the third-round PCR, there is a strong smear band for PCR without GO derivative and the target band becomes negligible. However, the target bands are still remarkable with low unspecific products in the presence of GO derivatives. Therefore, GO derivatives enhance the specificity of PCR for amplifying the complicated clinical samples. The results indicate that GO derivatives may have potential applications in clinical molecular diagnosis.

Simulation of the interaction between polymerase and polymer

It is very difficult to directly simulate the interaction between polymerase and polymer-modified GO. Thus, molecular dynamics simulation and molecular docking were used to study the possible interaction between the polymer and Pfu polymerase when GO was neglected for simplification. Pfu polymerase contains the following five domains: the N-terminal (residues 1–130 and 327–368), exonuclease (131–326), palm (369–450 and 501–588), fingers (451–500) and thumb domains (589–775) (Figure S14). The proofreading and polymerization mechanism are coordinated from the interactions between the loop (144–158) of the exonuclease domain and the positively charged edge of the thumb domain. Molecular docking showed that the binding sites of polymers were located at the exonuclease, thumb, palm and fingers domain by docking calculations (Figure S14). The binding affinities decreased in the following order: PAA > PAM > PEG > pSB (Table S8). The Coul Short Range energies fluctuated widely over the course of the 10 ns equilibration, reaching steady after ~4 ns (Figure S15A and B). Coul Short Range energy analysis indicates that there is a strong electrostatic interaction between PAA and Pfu (−2,248±203 kJ mol⁻¹). The Coul Short Range energy of Pfu-PAM is −959±107 kJ mol⁻¹, whereas PEG and pSB have nearly no electrostatic interaction with Pfu (Table S9). This order is consistent with the PCR results (Figure S11).

The solvent-accessible surface area (SASA) is the surface area of a biomolecule that is accessible to a solvent. SASAs of Pfu-PEG and Pfu-pSB are almost equivalent in 10 ns (Figure S15C), indicating that PEG and pSB have similar influence on the enzyme–substrate binding. The minimum distance of PEG and Pfu fluctuates in a wide range, indicating higher flexibility of PEG than pSB (Figure S15D). Cumulative solvent distribution shows that the dipole area of the solvent molecule around PEG is much larger than that of pSB (Figure S15E and F), indicating that PEG and pSB affect solvent and protein in different levels. Molecular dynamics simulations showed that the Pfu–PAA complex (PAA: the brown strip below Pfu) had the strongest interaction followed by Pfu–PAM complex (PAM: the blue strip below Pfu) (Figure 8A and B). The interaction would be electrostatic and hydrophobic interactions and van der Waals’ force. Obviously, there is no direct interaction between Pfu polymerase and PEG, and Pfu polymerase and pSB (Figure 8C and D). These results are consistent with the Coul Short Range energy analysis. The results of molecular dynamics simulation and molecular docking are reasonable because PEG can reduce...
protein adsorption, and zwitterionic polymer has better ability to reduce protein adsorption than PEG. Binding affinities of polymers with Pfu polymerase (Table S8) also support the result.

Besides the interaction between Pfu polymerase and polymer, negatively charged GO and positively charged Pfu polymerase also have strong electrostatic, hydrophobic and π–π stacking interactions. PEG and pSB will decrease these interactions due to reducing the adsorption of Pfu polymerase. Thus, GO-PEG and GO-pSB do not inhibit PCR in a wider range of concentration than GO-PAA and GO-PAM, which is consistent with experimental results (Figure 4). A little amount of positively charged Pfu polymerase still is adsorbed by negatively charged GO in GO-PEG and GO-pSB, even though PEG and pSB will reduce its adsorption. Thus, the specificity of PCR is also enhanced by GO derivative-assisted PCR. Therefore, these results further confirm that these GO derivatives can enhance the specificity of PCR.

**Conclusion**

We investigated the effect of GO with different sizes, reduction degrees and surface modifications on the specificity of PCR. The results indicate that L-GO and RGO is superior to S-GO and non-RGO in enhancing the specificity of PCR. For GO derivatives, the ability to enhance the specificity of PCR increases in the following order: GO-PAA < GO-PAM < GO-PEG < GO-pSB. Our study demonstrates that both GO and its surface properties affect the specificity of PCR. The GO derivatives may be used as efficient additives to enhance the specificity of PCR in clinical molecular diagnosis.

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**Author contributions**

All authors contributed toward data analysis, drafting and critically revising the paper and agree to be accountable for all aspects of the work.

**Disclosure**

The authors report no conflicts of interest in this work.

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