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A multifunctional organophosphonic ligand-templated poly(polyoxotungstate) with potential in building a genosensor for thalassemia gene detection†

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A novel P^V-Sb^{III} heteroatom-inserted poly(polyoxotungstate) (poly(POT)) [H₂N(CH₂)₂]₅Na₁₁H₉[(HEDATMPA)SbW₁₅O₅₀]{[PrW₄O₁₀][B-α-SbW₉O₃₃]}₂·38H₂O (**1**) was synthesized and is composed of an attractive [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ hybrid fragment and one sandwich-like Pr^{III}-inserted [PrW₄O₁₀(B-α-SbW₉O₃₃)₂]¹¹⁻ entity by using ethylenediamine tetramethylphosphoric acid (H₈EDATMPA) with potential heteroatom sites and functional organic groups. Considering the advantages of poly(POT) clusters in storing and transferring electrons, we fabricated a **1**-PEDOT (PEDOT = poly(3,4-ethylenedioxythiophene)) hybrid film, which was used as an electrode-modified material to construct a **1**-based genosensor (GS) for detecting the β-thalassemia gene. Research results show that this **1**-based GS demonstrates high specificity, outstanding antijamming ability and low detection limit for β-thalassemia gene detection. This work not only provides a novel and feasible preparative strategy to construct poly(POT) clusters, but also promisingly promotes the development of poly(POT)-based GSs.

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Introduction

Metal–oxygen clusters have been widely acknowledged as star candidates in diverse fields (*e.g.* catalysis, energy storage, biomedicine and photoelectric materials), which mainly benefits from their abundant structures, skeleton size and functional components.^{1–3} Among metal–oxygen cluster materials, polyoxotungstates (POTs), mainly formed from tungsten centers and oxygen linkers, have been extensively studied since more and more applications of POTs are being discovered in different fields.^{4–8}

It is worth mentioning that the introduction of heteroatoms (HAs) in the tungsten-oxo reaction system can produce lacunary POT units.^{7,9} It has been revealed that due to abundant surface reactive oxygen sites, lacunary POT units can work as rigid multidentate inorganic ligands to coordinate with func-

tional components such as rare-earth (RE) ions, transition-metal (TM) ions and organic ligands.¹⁰ Because of diverse coordination sites and flexible combination modes between POT units and functional components, a great deal of attractive poly(POT) clusters have been obtained.^{11–13} Particularly, organic–inorganic hybrid POTs have been considered as one important branch in POT chemistry because various organic ligands can provide a great opportunity for creating novel poly (POT) hybrids.^{8,14,15} For example, with the assistance of fumaric acid, our group prepared a fascinating RE-encapsulated poly(POT) hybrid {[RE₃(H₂O)₇]₂[W₄O₈(C₄H₂O₄)(C₄H₃O₄)]₂[SeW₆O₂₅]₂[B-α-SeW₉O₃₃]₄}²⁴⁻ (RE = Ce³⁺, La³⁺, Fig. S1a†), in which two fumaric acid ligands bridge two {[RE₃(H₂O)₇][W₄O₈(C₄H₂O₄)(C₄H₃O₄)]}[SeW₆O₂₅][B-α-SeW₉O₃₃]₂}¹²⁻ moieties.¹⁵ In 2019, Ding and his colleagues utilized citric acid (H₄CIT) as the organic ligand and isolated a strip-shaped octamer [Eu₁₆Co₇Se₁₆W₁₂₈O₄₄₈(CIT)₁₀ (HCIT)₂(NO₃)₄(OH)₄(H₂O)₅₂]⁴⁸⁻ (Fig. S1b†), which represents the longest poly(POT) macromolecule at that time.¹³ This macromolecule consists of two tetrameric {Eu₈Se₈W₆₄Co₃O₂₂₄(OH)₂(NO₃)₂(CIT)₆(H₂O)₂₆} segments joined together by H₄CIT ligands and TM ions. It is to be noted that not only the introduction of organic ligands tremendously makes for the formation of poly(POT) hybrids, but also the addition of RE ions helps in gathering adjacent POT units together.^{6,16}

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Until now, although many innovative poly(POT) hybrids have been greatly developed, oriented design and precise synthesis of poly(POT) clusters remain a challenge. It has been found that POT precursors, HAs or organic ligands can function as the template-guiding role in the formation of poly(POT) clusters.^{17–19} The well-defined template can effectively induce the assembly of multiple components. HAs have been often used as the template-guiding agent to induce the aggregation of POT units. For instance, by employing SeO_3^{2-} as the template, Cronin's group acquired a battery of high-nuclear POTs $[\text{H}_2(\text{SeO}_3)_3\text{W}_{43}\text{O}_{139}]^{24-}$, $[\text{H}_4(\text{SeO}_3)_5\text{W}_{77}\text{O}_{250}]^{44-}$, $[\text{H}_6(\text{SeO}_3)_6\text{W}_{63}\text{O}_{203}]^{34-}$ and $[\text{H}_8(\text{SeO}_3)_{16}\text{W}_{101}\text{O}_{317}]^{52-}$ (Fig. 1a), highlighting that the HA template (SeO_3^{2-}) effectively can give rise to lacunary $\{\text{SeW}_9\text{O}_{33}\}$ units instead of the "closed" saturated Keggin-type $\{\text{SeW}_{12}\text{O}_{40}\}$ cluster.²⁰ In recent years, researchers have no longer been satisfied with the studies of single HA-inserted POTs and have begun to introduce different types of HAs into a one POT system. Cronin's group and Zhao's group have demonstrated the feasibility of using dual-HA templates to synthesize novel poly(POT) clusters with novel layered POT building blocks $\{\text{XX}'\text{W}_{15}\text{O}_{54}\}$ ($\text{X} = \text{HP}^{\text{III}}, \text{Bi}^{\text{III}}, \text{X}' = \text{Se}^{\text{III}}, \text{Te}^{\text{VI}}, \text{As}^{\text{III}}, \text{Sb}^{\text{III}}$), providing a promising approach for continually discovering POT architectures (Fig. 1b–d).^{19,21–23} Beyond that, the critical template-guiding role of organic ligands can't be ignored. A typical example is the group of multi-Ni^{II} substituted silicotungstate clusters $\{\text{SiW}_9\text{Ni}_4\}_{2n}\text{L}_{3n}$ ($n = 2, 3$); $\{\text{SiW}_9\text{Ni}_4\} = \{[(\text{A}-\alpha\text{-SiW}_9\text{O}_{34})\text{Ni}_4(\text{OH})_3]^{5-}, \text{L} = \text{ligand}\}$ with regulable interior voids (Fig. 1e).²⁴ In this work, by utilizing the $[\text{A}-\alpha\text{-SiW}_9\text{O}_{34}]^{10-}$ precursors as inorganic nodes and rigid ligands with different bending angles as linkers, Fang *et al.* successfully realized the regulation of the skeletons and interior voids of $\{\text{SiW}_9\text{Ni}_4\}_{2n}\text{L}_{3n}$.²⁴

In previous research studies, it has been proved that some elemento-organic compounds can be used as template

linkers or provide HA templates to construct uncommon poly(POT) hybrids.^{6,14,25} As early as in 2000, Pope's group introduced phenyltin trichloride $(\text{PhSn})\text{Cl}_3$ into the POT system and obtained sandwich-like POT clusters $[(\text{PhSn})_3\text{Na}_3(\alpha\text{-SbW}_9\text{O}_{33})_2]^{6-}$ and $\{[(\text{PhSn})_2\text{O}]_2\text{H}(\alpha\text{-AsW}_9\text{O}_{33})_2\}^{9-}$, in which organotin ligands serve as a template linker to join two $[\alpha\text{-XW}_9\text{O}_{33}]^{9-}$ ($\text{X} = \text{Sb}^{\text{III}}$ and As^{III}) building blocks (Fig. 1f).²⁶ In 2020, organoboron ligands (BBLs) were first introduced into the POT system and led to tetrameric $[\{\text{BBL}\}_4(\text{M}_3\text{P}_2\text{W}_{15}\text{O}_{62})_4]^{28-}$ and dodecameric $[\{\text{BBL}\}_{15}(\text{M}_3\text{P}_2\text{W}_{15}\text{O}_{62})_{12}]^{86-}$ ($\text{M} = \text{Ta}^{\text{V}}, \text{Nb}^{\text{V}}$), in which each BBL covalently links to three Dawson-like lacunary POT blocks (Fig. 1g).²⁷ More interestingly, in the organophosphonic ligand modified poly(POT) cluster $\{[(\text{AsW}_9\text{O}_{33})\text{Zn}(\text{H}_2\text{O})\text{W}_5\text{O}_{13}(\text{N}(\text{CH}_2\text{PO}_3)_3)_2]^{14-}$, aminotrimethylene phosphonic ligands serve as the HA-supplier other than the organic linker (Fig. 1h).²⁸ Based on these results, given that both HAs and organic ligands can simultaneously act as templates for the assembly of poly(POT) clusters, it is possible that special organic ligands containing both potential HA sites and functional coordination groups will play a dual-template-guiding role. That is to say, these special organic ligands not only work as template linkers to combine with POT different blocks, but also can provide HA templates to induce novel building blocks. Thus, we began to use elemento-organic ligands with potential HA sites and functional organic groups to react with the POT precursors in the participation of RE ions for searching for innovative poly(POT) clusters.

In this work, we chose ethylenediamine tetramethylphosphonic acid ($\text{H}_8\text{EDATMPA}$) (Fig. S2†) with potential HA sites and functional organic groups as the multifunctional ligand and successfully obtained a $\text{P}^{\text{V}}\text{-Sb}^{\text{III}}$ double-HA intercalated poly(POT) cluster $[\text{H}_2\text{N}(\text{CH}_2)_2]_{15}\text{Na}_{11}\text{H}_9\{(\text{HEDATMPA})\text{SbW}_{15}\text{O}_{50}\}[\{\text{PrW}_4\text{O}_{10}\}[\text{B}-\alpha\text{-SbW}_9\text{O}_{33}]_2\} \cdot 38\text{H}_2\text{O}$ (1). In the poly-

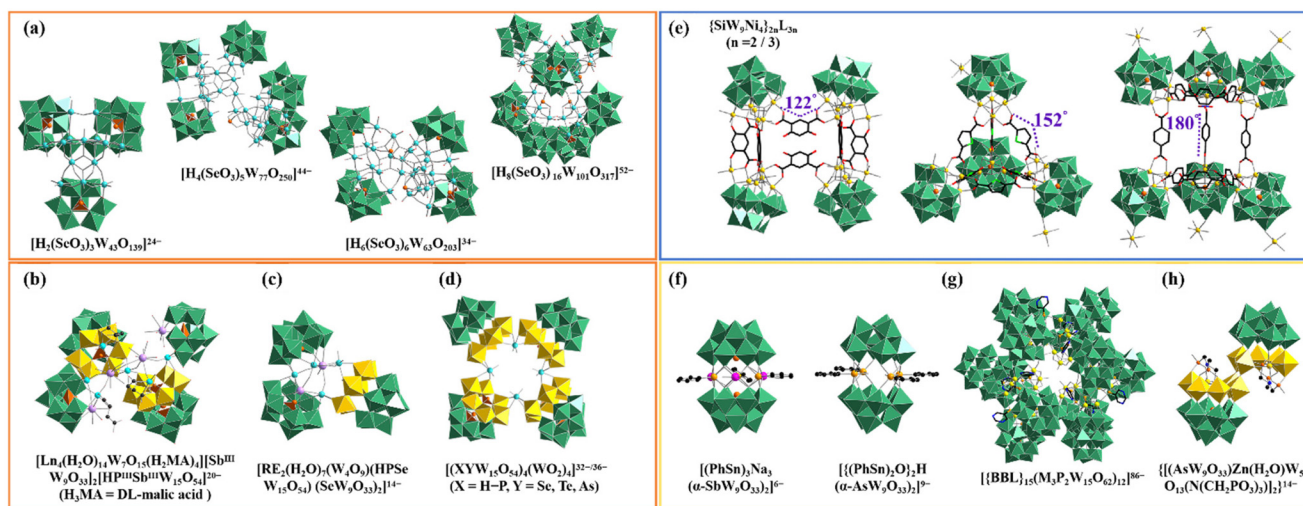


Fig. 1 (a) Some poly(POT) clusters using single-HA templates. (b–d) Some poly(POT) clusters with double-HA templates. (e) Some organic ligands regulated poly(POT) clusters. (f) Two poly(POT) clusters based on organotin linkers. (g) An organoboron ligand-modified poly(POT) cluster. (h) A organophosphonic modified poly(POT) cluster ($\{\text{WO}_6\}$: green and yellow octahedra).

nion of **1**, as expected, HEDATMPA⁷⁻ not only offers the P^V center as the HA template contributing to a layered [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ building block, but also works as a linker to bridge [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ and {PrW₄O₁₀[B- α -SbW₉O₃₃]₂}¹¹⁻ entities. Moreover, **1** was composited with poly(3,4-ethylenedioxythiophene) (PEDOT) to prepare the **1**-PEDOT hybrid film using the electropolymerization strategy. Furthermore, the **1**-PEDOT film was utilized as the electrode-modified material to construct the **1**-based genosensor (GS) for the β -thalassemia gene detection. The constructed **1**-based GS shows high sensitivity as well as strong anti-interference ability in determining the β -thalassemia gene. Undoubtedly, this work provides a new idea for the design and synthesis of novel poly(POT) aggregates and enriches the application of poly(POT)-based materials in the detection of genes.

Results and discussion

Structure description

The agreement of experimental and simulated powder X-ray diffraction curves of **1** indicates the good phase purity of the collected crystals (Fig. S3[†]). Subsequently, IR spectroscopy (Fig. S4[†]) and thermogravimetric analysis (Fig. S5[†]) were conducted to characterize **1**. The crystallographic data (Table S1[†]) show that **1** belongs to the *P* $\bar{1}$ space group. The polyanion [((HEDATMPA)SbW₁₅O₅₀)[{PrW₄O₁₀[B- α -SbW₉O₃₃]₂}]²⁵⁻ (**1a**) features an HEDATMPA⁷⁻ connecting trimeric hybrid architecture (Fig. 2a). Specifically, **1a** can be deemed as an aggregate of a Dawson-like [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ entity and a sandwich-like Pr^{III}-inserted {PrW₄O₁₀[B- α -SbW₉O₃₃]₂}¹¹⁻ segment bridged by a HEDATMPA⁷⁻ linker (Fig. 2b). It should be highlighted that the organic-inorganic hybrid Dawson-like [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ entity can be imagined as a second HA (P^V) embedded [(HEDATMPA)W₆O₂₇]²⁵⁻ subunit grafting onto a trivalent Keggin [B- α -SbW₉O₃₃]⁹⁻ fragment (Fig. 2c). More significantly, the P^V HA comes from the organophosphonic ligand. In other words, due to the template-guiding effect of the P^V1 atom in the organophosphonic ligand, six {WO₆} octahedra were successfully induced to form a {C₆N₂P₄W₆} ring around the P^V1 HA, and further connect the {Sb1W₉} block by sharing apical oxygen atoms. Meanwhile, another phosphate group in the organophosphonic ligand provides the P^V2 atom to share an oxygen atom with the W13 atom, leading to a chelate ring, which will potentially stabilize the skeleton (Fig. 2d). More than being an HA provider, the organophosphonic ligand also functions as the linker between adjacent POT fragments. To be specific, the left two phosphate groups on the other side of the organophosphonic ligand further joined two [W₂O₁₁]¹⁰⁻ units to form the [(POCH₂O)₃(POCH₃O)₂C₂H₄N₂W₄O₁₈]¹⁹⁻ {C₆N₂P₄W₄} segment (Fig. 2e and f), which is encapsulated by the [PrO₄(B- α -SbW₉O₃₃)₂]²³⁻ ({PrSb₂W₁₈}) entity. As for the sandwich-like {PrSb₂W₁₈} unit (Fig. 2g), it can be observed that the eight-coordinated Pr³⁺ ion adopts a distorted bicapped triangular prismatic geometry and connects two {SbW₉} subunits

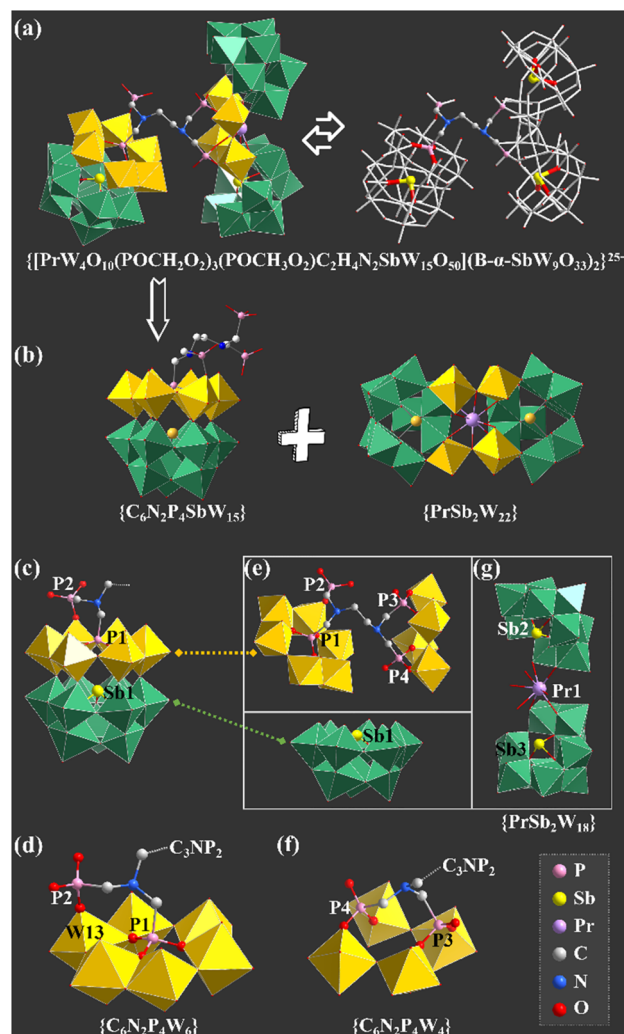


Fig. 2 (a) The **1a** polyanion. (b) The Dawson-like [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ entity and the sandwich-type {PrW₄O₁₀[B- α -SbW₉O₃₃]₂}¹¹⁻ segment. (c) The Dawson-like [(HEDATMPA)SbW₁₅O₅₀]¹⁴⁻ entity. (d) The {C₆N₂P₄W₆} ring with the P^V1 atom as the HA. (e) The bridging role of the organophosphonic ligand. (f) The {C₆N₂P₄W₄} segment. (g) The sandwich-like {PrSb₂W₁₈} unit.

through Pr–O–W bonds (Fig. S6 and S7[†]). Interestingly, in the whole structure, four P^V atoms in the organophosphonic ligand exhibit three different coordination modes. Three oxygen atoms in the –P1O₃H₂ group are all coordinated with W atoms, while the P2 atom has two uncoordinated oxygen atoms (Fig. S7a and b[†]). On the other side of the ligand, both P3 and P4 atoms provide two oxygen atoms to connect additional W atoms (Fig. S7c and d[†]).

So as to demonstrate the structural novelty of **1**, the polyanion **1a** is here compared with the reported selenotungstate anion [Ce₂(H₂O)₇(W₄O₉)(HP^{III}SeW₁₅O₅₄)(SeW₉O₃₃)₂]¹⁴⁻ (**2a**), which is similar to **1a** and contains a pure inorganic Dawson-like [HP^{III}SeW₁₅O₅₄]¹¹⁻ ({HP^{III}SeW₁₅}) fragment (Fig. 3a).²² This {HP^{III}SeW₁₅} entity in **2a** consists of a firstly reported [HP^{III}W₆O₂₇]¹⁴⁻ ({HP^{III}W₆}) subunit with P^{III} as the second HA

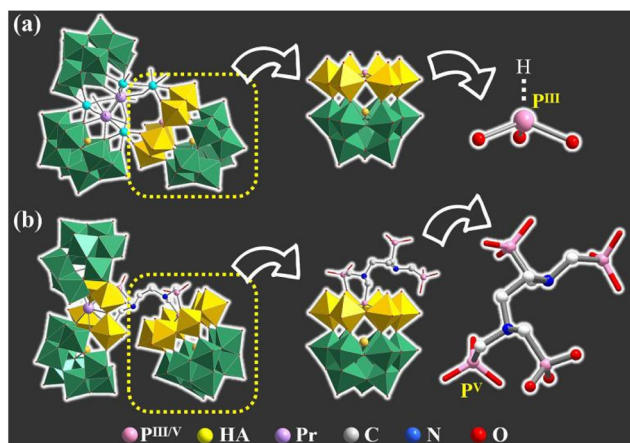


Fig. 3 Comparison of the inorganic HP^{III}-inserted Dawson-like (HP^{III}SeW₁₅) unit in **2a** (a) and the organo-inorganic hybrid (C₆N₂P^VSbW₁₅) unit in **1a** (b).

located on the top of the classical {SeW₉} fragment. As we know, the coordination number of the P^{III} atom from inorganic phosphorous acid is four. Thus, the P^{III} atom in the {HP^{III}SeW₁₅} entity can't further coordinate with other functional components such as organic ligands or metal ions. Therefore, structural expansion of the {HP^{III}SeW₁₅} entity can be only achieved through the active oxygen sites in the {WO₆} octahedra. In comparison, the {C₆N₂P^VSbW₁₅} block with the P^V atoms from the organophosphonic ligand in **1a** has exposed oxygen sites from tungsten-oxygen clusters and coordination-active oxygen sites from organic groups, which offers more opportunity to bridge more POT blocks together (Fig. 3b). This result indicates that such organophosphonic ligand is supposed to provide more chances for achieving novel poly(POT) aggregates.

In the crystal structure, some lattice water molecules, [H₂N(CH₃)₂]⁺, Na⁺ and H⁺ cations can interact with **1a** polyanions through hydrogen bonding interactions and electrostatic attraction.²⁹ Moreover, **1a** polyanions, counter cations and lattice water molecules are arranged orderly in three axes. In order to more clearly describe the three-dimensional stacking mode of **1a** polyanions, we omit those charge compensation ions here and further leave only Sb1, Sb2 and Sb3 atoms in the structure, simplifying the **1a** polyanion into a triangle. It is clearly observed that the **1a** polyanions exhibit an “-ABAB-” stacking pattern along the *a*, *b*, and *c* axes, respectively (Fig. S8†).

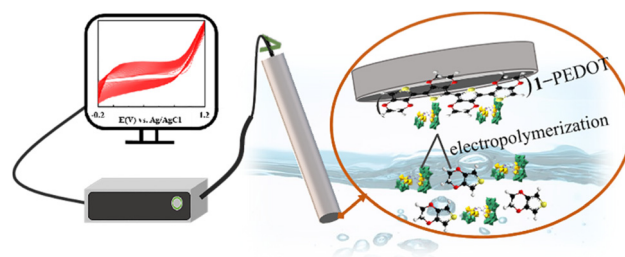
Characterization of the 1-based hybrid film

Recently, the electrochemistry of poly(POT) clusters has been explosively studied, especially in electrochemical catalysts/sensors, energy-related devices and so on.^{8,21,30} Owing to their reversible electron transmittability between stable skeletons, poly(POT) clusters have been considered as promising candidates in electrochemical applications.³¹ Except that they can

be used as electron conductors, poly(POT) clusters have also been demonstrated to be stability-enhancers, conductivity-facilitators, or even electronic repositories.^{32–34} Nevertheless, the advancements of poly(POT) clusters in electrochemistry have been limited by their water-solubility. To break this bottleneck, more and more preparative methods for poly(POT)-based composite materials have been put forward, for instance, physical adsorption,²¹ chemical/electro-deposition,³⁵ layer-by-layer assembly,³⁶ and electropolymerization.³⁷ As the electropolymerization technology can provide uniform polymeric films with high chemical/electrochemical stability and remarkable conductivity, it has become a favored route to modify electrodes in constructing electrochemical sensors or devices.³⁸ It is well-known that PEDOT has been widely used as a conductive polymer, which was first reported in the early 1990s, and has characteristics of excellent conductivity, good film-forming performance and a long cycle life.^{39,40} Herein, we successfully prepared a hybrid film **1**-PEDOT *via* an electropolymerization process in the mixed solution of **1** and 3,4-ethylenedioxythiophene (EDOT). Therefore, the prepared **1**-PEDOT hybrid film was also supposed to show outstanding electrochemical properties.

The homogeneous **1**-PEDOT film was constructed based on a typical three-electrode platform, including a glassy carbon electrode (GCE) as the working electrode, an auxiliary platinum electrode, and the reference silver electrode. As shown in Scheme 1, the electropolymerization process was realized through a cyclic voltammetry (CV) method under -0.2–1.2 V potential in a mixed solution containing **1**, EDOT and NaClO₄. During the electropolymerization, **1a** polyanions and protonated EDOT migrate to the surface of GCE under the electric field. The polymerization results in a uniform **1**-PEDOT film firmly attached to the GCE surface.

Since the reaction took place in aqueous solution, the stability of **1** in water was first tested by electrospray ionization mass spectrometry (ESI-MS). In the ESI-MS spectrum, three obvious ESI-MS signals at *m/z* 1424.22 (7⁻), 1665.31 (6⁻) and 2012.26 (5⁻) were observed and can be respectively assigned to {Na₉H₉[PrW₄O₁₀(POCH₂O₂)₃(POCH₃O₂)₂C₂H₄N₂SbW₁₅O₅₀][B-α-SbW₉O₃₃]₂}⁷⁻, {Na₁₀H₉[PrW₄O₁₀(POCH₂O₂)₃(POCH₃O₂)₂C₂H₄N₂SbW₁₅O₅₀][B-α-SbW₉O₃₃]₂}⁶⁻ and {[H₂N(CH₃)₂]⁺Na₉H₉[PrW₄O₁₀(POCH₂O₂)₃(POCH₃O₂)₂C₂H₄N₂SbW₁₅O₅₀][B-α-SbW₉



Scheme 1 Schematic diagram of preparing the **1**-PEDOT modified electrode.

$O_{3.3}S_{1.2}P_{0.5}^{5-}$ segments, which implies that the basic skeleton of the **1a** polyanion is stable in aqueous solution (Fig. S9†). Moreover, we monitored the structure variation of the **1a** polyanion after dissolving it in an aqueous solution for different times by ultraviolet–visible (UV–vis) absorption spectroscopy. As the UV–vis spectra of **1** in the aqueous solution almost remained constant over time (Fig. S10†), the stability of the **1a** polyanion in water is demonstrated, laying the foundation for its further application exploration.

Besides, in order to confirm the successful preparation of the **1**-based film, CV curves of the naked electrode and different modified electrodes were collected using a 0.2 mol L⁻¹ CH₃COOH–CH₃COONa buffer solution (pH = 4.5, Fig. 4a). It can be observed that there are no paired redox peaks of PEDOT in the CH₃COOH–CH₃COONa buffer solution, while a couple of redox peaks can be observed in the CV curve of **1**-PEDOT after electropolymerization, which may imply the successful recombination of **1** and PEDOT. To further verify the combination of **1** and PEDOT in the hybrid film, IR spectra (Fig. S11†), X-ray photoelectron spectroscopy (XPS, Fig. 4b and Fig. S12, S13†) and energy dispersive spectroscopy (EDS) elemental mapping (Fig. S14†) measurements of **1**, PEDOT and the **1**-PEDOT film were carried out. It can be seen from the comparison of IR spectra of **1**, PEDOT and the **1**-PEDOT film (Fig. S11†) that the characteristic bands of both **1** [1065 (P^V–O), 948 (W–O_t), 899 (W–O_b), 780 (W–O_c) and 718 cm⁻¹ (Sb^{III}–O_a)] and PEDOT [1521 cm⁻¹ (C=C), 686 cm⁻¹ (C–S)]⁴⁰ can be observed in the IR spectrum of **1**-PEDOT, stating the co-existence of **1** and PEDOT in the **1**-PEDOT film. In the mean-

time, comparing the XPS spectrum of **1** and that of the **1**-PEDOT film, we can see that both the characteristic peaks of **1** (W 4f_{7/2}: 35.9 eV, W 4f_{5/2}: 38.0 eV) and those corresponding to PEDOT (S 2p_{3/2}: 163.4 eV, S 2p_{1/2}: 164.6 eV) appear simultaneously in the XPS pattern of the **1**-PEDOT film (Fig. 4c and Fig. S12, S13†).^{30,41,42} Furthermore, the EDS mapping images exhibit the uniform distribution of W, P, Sb, Pr and S elements in the **1**-PEDOT film (Fig. S14†). All the above results synthetically evidence the combination of **1** and PEDOT in the **1**-PEDOT hybrid film.

It is well known that electrical conductivity is the primary criterion to determine whether the **1**-PEDOT hybrid film can be used as the electrode modification material. So, experimental conditions for preparing the **1**-PEDOT film as well as follow-up measurements were optimized by taking electrical conductivity (through CV and DPV signals) as the evaluation index. First of all, the electropolymerization cycles that mainly affect the conductivity of the **1**-PEDOT film were optimized by varying the number of sweep cycles (10, 20, 30, 40, 50 and 60 cycles) in the CV preparation procedure. The **1**-PEDOT modified GCE (**1**-PEDOT/GCE) was used as the working electrode and the corresponding CV and DPV curves were collected in 0.10 mol L⁻¹ phosphate buffered saline (PBS, pH = 7.4) containing 0.01 mol L⁻¹ 1 : 1 [Fe(CN)₆]^{3-/4-} (marked as Fe-PBS). As shown in Fig. S15,† various redox peaks appear under different electropolymerization cycles, which are caused by the faster electron transfer rate on the PEDOT-modified GCE. This performance improvement can be attributed to the larger effective surface area of **1**-PEDOT/GCE.⁴³ The electroactive area

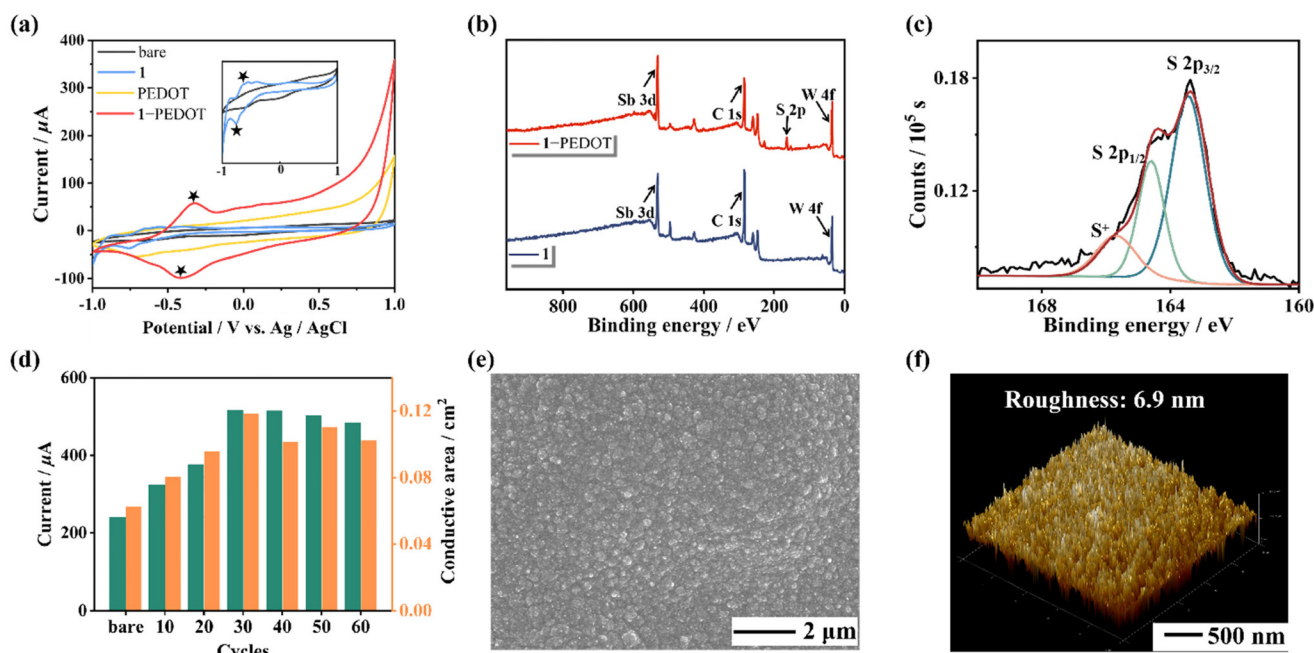


Fig. 4 (a) CV curves of bare GCE and GCEs modified using **1**, **1**-PEDOT and PEDOT in 0.2 mol L⁻¹ CH₃COOH–CH₃COONa buffer solution. Inset: CV curves of bare GCE and **1**-PEDOT/GCE. (b) Comparison of XPS spectra between **1** and the **1**-PEDOT film. (c) XPS spectra of S 2p in the **1**-PEDOT film. (d) Comparison of the conductive area and DPV peak of **1**-PEDOT/GCEs with different electropolymerization cycles (0/10/20/30/40/50/60) in the Fe-PBS solution (pH = 7.4). (e and f) SEM and AFM images of the **1**-PEDOT film on the ITO substrate (30 electropolymerization cycles).

under different sweep cycles in the CV method was calculated according to the Randles–Sevcik equation shown below:

$$I_p = 2.69 \times 10^5 n^{3/2} A D^{1/2} \nu^{1/2} c$$

Here, I_p refers to the redox peak of the CV curve measured in Fe-PBS (pH = 7.4) solution, A (cm^2) is the electrode conductive area, n is the number of transferred electrons ($n = 1$), D represents the diffusion coefficient of $\text{K}_3[\text{Fe}(\text{CN})_6]$ ($D = 7.6 \times 10^{-6} \text{ cm}^2 \text{ s}^{-1}$), ν represents the scanning rate ($\nu = 0.1 \text{ V s}^{-1}$) and c is the concentration of $\text{K}_3[\text{Fe}(\text{CN})_6]$ ($c = 0.01 \text{ mol L}^{-1}$).³¹ While compared with the bare GCE ($A = 0.06 \text{ cm}^2$), the modified 1-PEDOT/GCE exhibits a larger electrode conductive area ($A = 0.12 \text{ cm}^2$), which might be explained by the fact that the 1-PEDOT hybrid film on the GCE surface can provide more exposed redox sites after electropolymerization.³¹ It can be seen that on increasing the electropolymerization cycles, the conductive area and the intensity of the DPV current at 0.252 V (Fig. 4d and Fig. S15, S16[†]) also increase and reach the maximum when the electropolymerization cycle is 30. In other words, the optimized electropolymerization cycle for preparing the 1-PEDOT film is 30. Moreover, it is found from both CV and DPV measurements (Fig. S17[†]) that the signal strength of CV and DPV peaks and the integral area of the CV curve (that is the electrode conductive area A) of 1-PEDOT/GCE are obviously enhanced compared with those of bare GCE, which indicates that the 1-PEDOT film has good electrical conductivity. In order to investigate the influence of the pH value of phosphate buffered saline (PBS) solution on the sensitivity of 1-PEDOT/GCE, a series of 0.10 mol L^{-1} PBS solutions containing 0.01 mol L^{-1} 1:1 $[\text{Fe}(\text{CN})_6]^{3-/4-}$ with different pH values (pH = 5.4/6.4/7.4/8.4/9.4) were prepared. Then, the DPV curves of 1-PEDOT/GCE were tested in these solutions (Fig. S18[†]). It can be observed that the detected DPV signal of 1-PEDOT/GCE is the strongest when the pH value is 7.4. Therefore, 7.4 was selected as the optimized pH value and the subsequent experiments were carried out in the pH = 7.4 Fe-PBS solution. At the same time, the electrochemical double-layer capacitance (C_{dl}) test is an important method for obtaining the electrochemically active surface area (ECSA) involved in the electrochemical reaction, which can be achieved from CV curves with different scan rates under voltages in the non-Faraday interval.^{44,45} Therefore, the modified GCEs were tested by CV technology with different scanning rates (0.02–0.20 V s^{-1}) in the voltage range of 0.32–0.42 V. Thus, the C_{dl} of 1-PEDOT/GCE was calculated as 16.56 mF cm^{-2} while that of PEDOT/GCE was calculated as 9.30 mF cm^{-2} , indicating the larger ECSA of 1-PEDOT/GCE. That is to say, the participation of **1** significantly improves the ECSA of the film-modified GCE (Fig. S19[†]).

Subsequently, to investigate the morphology of the 1-PEDOT hybrid film, the 1-PEDOT film was fabricated on the indium-tin oxide (ITO) substrate under optimized conditions and was characterized by scanning electron microscopy (SEM) and atomic force microscopy (AFM). The SEM image clearly demonstrates that the obtained 1-PEDOT film features outstanding flatness and uniformity (Fig. 4e). The AFM image

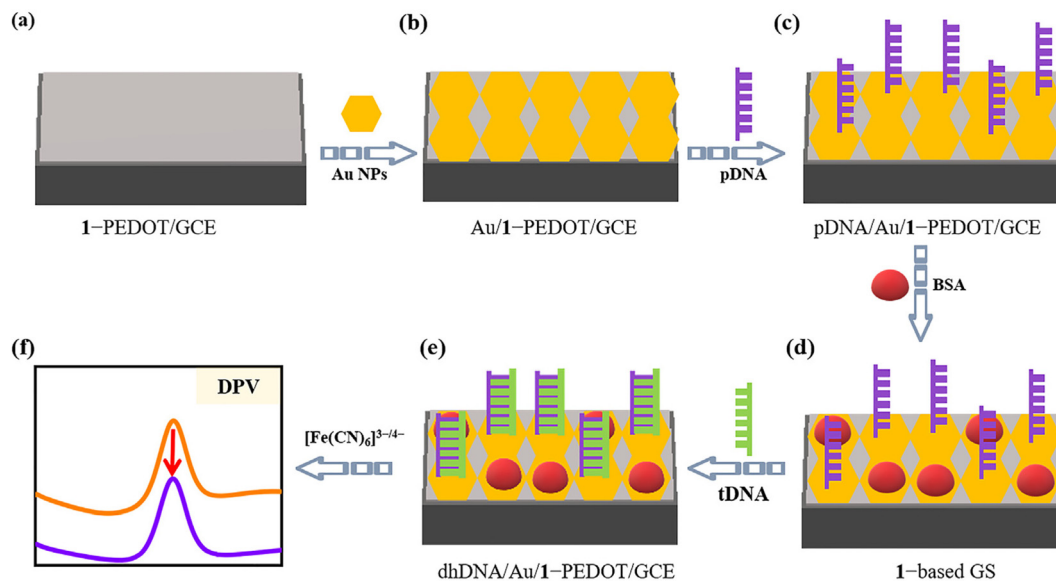
indicates that the roughness of the 1-PEDOT film is about 6.9 nm (Fig. 4f).

Long-term stability is another critical evaluation criterion for further practical application as electrode materials, so the stability evaluation of the 1-PEDOT hybrid film was conducted. Specifically, the conductivity variation of bare GCEs and 1-PEDOT/GCEs during the natural storage at room temperature for a period of time was monitored by recording their DPV curves. As illustrated in Fig. S20,[†] the DPV signal intensity of 1-PEDOT/GCE goes down slightly during the storage for 30 days while the bare GCE is fast passivated and the conductivity decreases significantly in two hours. Thus, the 1-PEDOT hybrid film shows good stability, enabling it to be a promising electrode modification material.

Fabrication of the 1-based GS for β -thalassemia gene detection

Thalassemia is a common genetic defect disease that is chronic hemolytic anemia caused by a disorder of globin peptide chain synthesis, and has been a serious threat to human health.⁴⁶ Thus, the growing demand for simple, fast and inexpensive testing measures, as opposed to time-consuming and expensive detection of nucleic acids with specific sequences, has largely encouraged researchers to study GSs for the detection of the thalassemia gene. Until now, diverse response signals (*e.g.* electrochemistry,^{47,48} fluorescence,⁴⁹ surface plasmon resonance optics⁵⁰) have been applied in thalassemia GSs. Among them, electrochemical detection has attracted wide attention due to its advantages of low cost, high sensitivity and fast response.

The electrochemical detection of genes usually requires effective modification of the electrode surface to access enhanced signals along with improved stability of the working electrode. Herein, considering that the 1-PEDOT film has good conductivity and stability, we designed a novel GS using the 1-PEDOT/GCE as the working electrode for electrochemical identification of the β -thalassemia gene. The basic principle of this GS is presented in Scheme 2. Firstly, to provide fixed sites for the pDNA chains, Au nanoparticles (NPs) are electrochemically deposited on the surface of 1-PEDOT/GCE (Scheme 2a) through an electrodeposition procedure in AuCl_4^- (0.10 g L^{-1}) solution, leading to the Au NP modified electrode (Au/1-PEDOT/GCE, Scheme 2b). Then, 5 μL of a solution containing a sulfhydryl-modified pDNA fragment at one end (5'-SH-ACTTCAGGATG AGTCTATGG-3') as pDNA is dropped down onto the Au/1-PEDOT/GCE surface to acquire the pDNA graft on the exposed Au sites *via* Au-S bonds, leading to pDNA/Au/1-PEDOT/GCE (Scheme 2c). Afterwards, the unoccupied Au sites on the electrode surface are closed using 20 μL of 2.0 wt% bovine albumin (BSA) solution to avoid the possible nonspecific adsorption of DNA chains in the subsequent steps (Scheme 2d). Next, 10 μL antisense DNA strand of the human β -thalassemia gene (5'-CCATAGACTCATCCTGAAGT-3') as the target DNA (tDNA) was dropped onto the surface of BSA-treated pDNA/Au/1-PEDOT/GCE (1-based GS). After a period of time for hybridization, pDNA will capture the tDNA strand forming double helix DNA (dhDNA) on the electrode (marked



Scheme 2 Mechanism of detecting the β -thalassemia gene by the 1-based GS: (a) the 1-PEDOT film modified GCE (1-PEDOT/GCE); (b) Au deposited 1-PEDOT/GCE (Au/1-PEDOT/GCE); (c) pDNA grafted Au/1-PEDOT/GCE (pDNA/Au/1-PEDOT/GCE); (d) BSA-treated pDNA/Au/1-PEDOT/GCE (1-based GS); (e) 1-based GS capturing tDNA (dhDNA/Au/1-PEDOT/GCE); (f) DPV response of dhDNA/Au/1-PEDOT/GCE.

as dhDNA/Au/1-PEDOT/GCE, Scheme 2e) based on the principle of complementary base pairing. At this point, the electrochemical signals will be markedly changed after detecting tDNA, giving out the obviously decreased DPV signal (Scheme 2f).

The DPV response can be explained as follows. As we all know, because of the contained negatively-charged phosphate residues, the whole DNA chain is negative. For dhDNA/Au/1-PEDOT/GCE, the dhDNA formed by the hybridization of pDNA and tDNA will possess more negative charges than single pDNA strands, leading to larger electrostatic repulsion between negatively charged electrode surfaces and $[\text{Fe}(\text{CN})_6]^{3-/4-}$ ions. That is to say, the dhDNA chain formed by the hybridization of pDNA and tDNA may work as an “electronic shield layer”, which will largely hinder interfacial electron transfer on the electrode surface.^{51,52} In contrast, the electrostatic repulsion between $[\text{Fe}(\text{CN})_6]^{3-/4-}$ and single strand pDNA is relatively weaker. Consequently, once tDNA is detected, the dhDNA/Au/1-PEDOT/GCE will show reduced conductivity and decreased DPV signal compared with pDNA/Au/1-PEDOT/GCE.

In order to obtain the most reliable results for tDNA detection, some key parameters in the process of constructing GS were optimized with DPV signals being the criterion. First of all, to provide enough active sites for grafting pDNA, the electrochemical deposition time of Au NPs was optimized. As can be seen from Fig. 5 and Fig. S21,[†] the conductivity of Au/1-PEDOT/GCEs gradually increases with electrochemical deposition time changing from 20 s to 80 s, whereas the conductivity begins to decrease with prolonging electrochemical deposition time. As a result, 80 s was chosen as the most suitable electrochemical deposition time of Au NPs. For the characterization of the Au/1-PEDOT film, XPS, AFM and SEM-EDS were

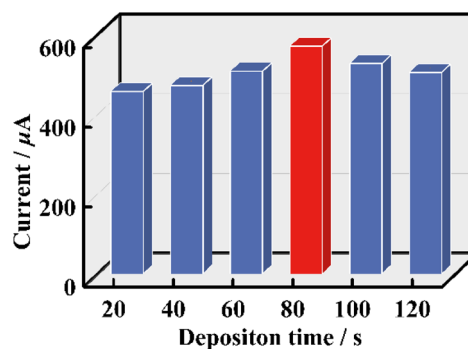


Fig. 5 DPV signals of Au/1-PEDOT/GCEs with different electrodeposition times of Au NPs in Fe-PBS solution (pH = 7.4).

conducted. In addition to the contained elements in the 1-PEDOT film, two more characteristic peaks belonging to Au (Au 4f5/2: 84.8 eV, Au 4f7/2: 88.5 eV) are observed in the XPS spectrum of the Au/1-PEDOT film,⁵³ testifying the deposition of Au NPs in the 1-PEDOT film (Fig. S22[†]). Moreover, both AFM and SEM images indicate the flatness as well as the integrity of the 1-PEDOT film after Au deposition (Fig. S23 and S24[†]), while the SEM-EDS elemental mapping images further confirm the homogeneous distribution of Au NPs on the film (Fig. S24[†]).

Secondly, to ensure the grafting of pDNA on Au sites, the pDNA incubation time and the concentration of pDNA were optimized. As shown in Fig. 6, when the same concentration of pDNA ($1 \times 10^{-8} \text{ mol L}^{-1}$) is applied for the grafting of pDNA on Au/1-PEDOT/GCE, the read out DPV signals increases obviously when the incubation time is less than 2 h.

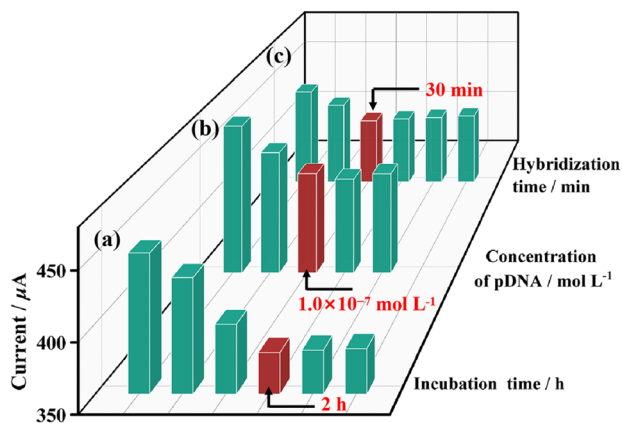


Fig. 6 (a and b) DPV signal variation of Au/1-PEDOT/GCE with different pDNA incubation time and different pDNA concentration. (c) DPV signal variation of dhDNA/Au/1-PEDOT/GCE with different hybridization time. [DPV curves were measured in Fe-PBS (pH = 7.4)].

Whereafter, the DPV signal intensity remains stable, which manifests that the incubation reaction is accomplished in 2 h (Fig. S25[†]). So, a period of 2 h was chosen as the best pDNA incubation time. Then, DPV signals of pDNA/Au/1-PEDOT/GCEs with different pDNA concentrations were measured. It can be seen that the intensity of DPV peaks reduces when the concentration of pDNA ranges from 1×10^{-9} mol L⁻¹ to 1×10^{-7} mol L⁻¹, and then keeps almost unchanged on increasing the pDNA concentration (Fig. 6 and Fig. S26[†]). Accordingly, the optimized conditions for the graft pDNA were 2 h and 1×10^{-7} mol L⁻¹ pDNA solution. To evaluate the best reaction time for the hybridization of pDNA and tDNA, 10 μ L tDNA solution (5×10^{-10} mol L⁻¹) was applied on the pDNA/Au/1-PEDOT/GCE surface, and the obtained dhDNA/Au/1-PEDOT/GCEs with different hybridization times (10–60 min) were tested by DPV. According to the experimental results, the current signal response shows a decreasing tendency at first and then reaches a stable platform after 30 min, indicating that the hybridization process is basically completed in 30 min (Fig. 6 and S27[†]). Therefore, 30 min was supposed to be sufficient for the hybridization procedure.

To verify the successful modification of the working electrode in each step during the process of the 1-based GS construction, the CV method was employed to examine the variation of the electrical conductivity of the working electrode in each step. As shown in Fig. S28,[†] all the modified GCEs exhibit enhanced peak currents and electroactive areas compared with the bare GCE, demonstrating that the electrical conductivities of the modified electrodes increase. Moreover, the electrical conductivity of Au/1-PEDOT/GCE is much higher than that of 1-PEDOT/GCE owing to the electrochemical deposition of Au on the electrode surface. After the electronegative pDNA is fixed on the surface of Au/1-PEDOT/GCE, the peak current and electroactive area of pDNA/Au/1-PEDOT/GCE decrease. Finally, the formation of double helix DNA by combining tDNA with pDNA leads to a further decrease of the peak current and electroactive area.

Under these optimal conditions, the sensibility of the 1-based GS platform (pDNA/Au/1-PEDOT/GCE) detecting tDNA was evaluated using DPV and EIS techniques. As can be seen from Fig. 7a and b, the read-out DPV signal of the GS detecting tDNA goes down linearly with the concentration of tDNA ranging from 1×10^{-12} mol L⁻¹ to 5×10^{-9} mol L⁻¹. This is because more tDNA strands in the solution result in more dhDNA (“electron shield layer”) on the surface of the GS, which hinders the conduction of electrons to some extent. Moreover, within this concentration range, the current intensity (I) has a linear relationship with the logarithm of the tDNA concentration, and the regression equation is $I = -26.67 \lg c + 82.91$ with a correlation coefficient (R^2) of 0.9866. On this basis, the limit of detection (LOD) for tDNA identification was 2.6×10^{-13} mol L⁻¹ (Fig. 7c). Excitingly, compared with some previously reported sensors for the β -thalassemia gene, the 1-based GS has a relatively wider linear range and a lower LOD (Table S2[†]).

Beyond that, repeatability, selectivity and anti-interference ability are also important criteria for evaluating whether a newly-developed GS is qualified. For this purpose, five pDNA/Au/1-PEDOT/GCEs were prepared under the same conditions to detect 5×10^{-11} mol L⁻¹ tDNA. As no significant changes in DPV signals (the relative standard deviation was about 2.9%) can be observed, it proves the outstanding repeatability of

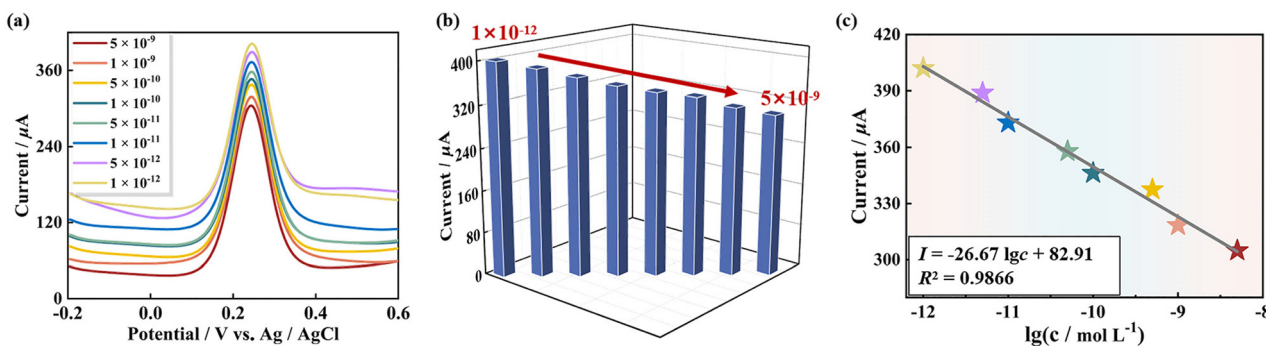


Fig. 7 (a and b) DPV curves and signal intensity of pDNA/Au/1-PEDOT/GCEs capturing tDNA with different concentrations. (c) The linear relationship between the DPV signal intensity and $\lg c(\text{tDNA})$. All signals were measured in solution Fe-PBS (pH = 7.4).

such a 1-based GS (Fig. S29†). Additionally, according to the principle of complementary base pairing, the specific binding of pDNA and tDNA should be irreplaceable. To test the specificity of this GS, tDNA, and DNA with non-complementary sequences (ncsDNA: 5'-AATCTCATGGCCGATTTCG TT-3'), double-base mismatched DNA (dbmDNA: 5'-CGATTGACTCATCCT GAAGT-3') and three-base mismatched DNA (tbmDNA: 5'-CGATTGACA CATCCTGAAGT-3') were respectively dropped on the surface of pDNA/Au/1-PEDOT/GCEs (the concentrations of all the DNAs were 5×10^{-11} mol L⁻¹). Apparently, only the 1-based GS capturing tDNA (dhDNA/Au/1-PEDOT/GCE) exhibits a significant decrease of the peak current, while none of the other DNA chains (ncsDNA, dbmDNA or tbmDNA) can be identified by the 1-based GS (Fig. S30†). This is because the hybridization process occurs between completely complementary tDNA and pDNA. In addition, the anti-interference ability of this GS was evaluated by detecting tDNA accompanied by one or several interferents (ncsDNA, dbmDNA and tbmDNA). It can be observed that the existence of interferents hardly influences the DPV response for tDNA, demonstrating a favorable anti-interference ability (Fig. S31†). The above-mentioned results confirm the great potential of 1-based GS in future applications.

Conclusions

In conclusion, an innovative P^V-Sb^{III} double-HA intercalated poly(POT) **1** was synthesized using ethylenediamine tetramethylphosphoric acid with potential HA sites and functional organic groups. Intriguingly, the organic-inorganic hybrid Dawson-like {C₆N₂P₄SbW₁₅} building block is present in the architecture of **1**, which is quite different from the previously-reported either the {(XW₆)(XW₉)} containing single-HA or the emerging dual-HA inserted {(XW₆)(X'W₉)}. Whereafter, a 1-PEDOT film with prominent conductivity was prepared by the electropolymerization method, which exhibits great potential in electrochemical applications. Furthermore, a thalassemia GS was fabricated using 1-PEDOT modified GCE as the working electrode, producing a sensitive DPV response. It shows a low detection limit and wide response range, providing a feasible strategy for trace detection of the β-thalassemia gene. As we know, such a poly(POT)-based thalassemia GS is the first example of a poly(POT)-based thalassemia electrochemical sensor. Applying elemento-organic compounds as dual-functional ligands in this work provides a new idea for building more giant and innovative poly(POT) nanoclusters. Meanwhile, the performance exploration for **1** being used in constructing β-thalassemia GS in some degree promotes the integration of chemistry and biology.

Author contributions

Wenshu Zhang and Dan Wang performed syntheses, characterization, electrochemical properties and wrote the manuscript.

Zixu Wang and Mengya Cao participated in PXRD, IR and TG studies. Lijuan Chen and Junwei Zhao provided research ideas, determined crystal structures and revised all over the manuscript.

Conflicts of interest

There are no conflicts to declare.

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