Potentiometric Phosphate Ion Sensor Based on Electrochemically Modified All-Solid-State Copper Electrode for Phosphate Ions’ Detection in Real Water

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Abstract: The importance of phosphates has sparked researchers’ considerable interest in the electrochemical detection of phosphates within aqueous solutions in recent years. In this study, we present a novel all-solid-state phosphate ion-selective electrode (ISE) that integrates copper, copper nanoparticles, and copper phosphate. By modifying the copper substrate of the electrode with a copper nanoparticle film and creating a lamellar copper phosphate film through electrochemical treatment, we significantly enhanced the electrode’s electron transfer efficiency. This microstructure with large specific surface area markedly improved the electrode’s responsiveness to the targeted ions by accelerating the achievement of chemical equilibrium on the electrode surface, thereby boosting its sensitivity and stability. The newly developed electrode was capable of detecting phosphate ions in solutions with a pH range from 6 to 11 and performed optimally in neutral solutions at pH 7, following Nernst principle, with a detection limit of 1 × 10^-6 M. The electrode exhibited a short response time of less than 10 s with significant reproducibility, stability, longevity—maintaining functionality for more than two months. It also displayed good selectivity as the electrochemical equilibrium was not influenced by up to 1 mM of potential competing species like HCO3-, NO3-, Cl- and SO42-. We compared the detection results of current phosphate ion sensor and conventional determination methods for phosphate content in natural lake and aquaculture water samples, with a detection discrepancy of about 10% (RSD). Considering all feasible performance characteristics combined with its low cost, simple manufacture and portability, the sensor provides a new possibility for rapid, reliable, and long-term real-time in situ detection of phosphates.

Keywords: ion-selective electrode; electrochemical sensor; phosphate determination; copper phosphate; nano-modification; electrodeposition

1. Introduction

Phosphorus is widely distributed in the biological ecosystem and in environmental systems and is of great significance to the biogeochemistry cycle and healthy habitat. Phosphorus usually occurs in phosphate and phosphate is the key substance in inorganic chemistry, biochemistry and biogeochemistry. In recent years, the rising global population and substantial growth of industrial and agricultural activities resulted in the release of a significant amount of phosphorus-containing pollutants into nature waters. An excessive accumulation of phosphate in lakes, oceans and other natural water bodies induces eutrophication—a phenomenon followed by algal blooms, anoxia, acidification, and a biodiversity crisis. Eventually, a series of water quality and environmental problems are triggered [1–3]. Therefore, phosphates act as both a crucial nutrient and a growth inhibitor in aquatic ecosystems; real-time detection of phosphates is of significant importance as
it serves to reflect the water quality and health status of natural water bodies and water used in hydroponic and aquaculture systems [1,3]. In addition, phosphorus is one of the important constituent elements of living cells, and phosphate excessiveness or deficiency in the human body will cause a series of physiological problems, such as hyperphosphatemia, which will harm the kidneys, thyroid gland and so on [4–6]. Additionally, phosphates are widely used in food processing as well as food additives and ingredients. Therefore, effective determination of phosphate concentration is important in various fields such as chemistry, biomedicine, agriculture, industry and environmental pollution prevention and control [7–9]. In a word, it is of great value to design and develop a low-cost, portable, efficient, accurate and real-time on-line sensor for phosphate ion detection.

Over the past few decades, environmental agencies in many countries have specified recommended range values for total phosphorus or phosphate concentrations and standard procedures for their detection [10]. The phosphor molybdenum blue spectrophotometric method is currently the most widely used phosphate detection method [11,12]. With the advancement of instrumentation, ion chromatography has gradually become a common method for determining phosphates [13,14]. The methods mentioned above possess good sensitivity and reliability. However, they necessitate sample pretreatment, are complex, time-consuming, and costly. Additionally, they cannot meet the demands of long-term real-time online monitoring. These limitations to some extent restrict the application of these methods [15,16]. In recent years, electrochemical analysis techniques, especially ion-selective electrodes (ISEs), have been widely studied and applied due to their capabilities of rapid and efficient measurement, easy operation, high sensitivity, and strong anti-interference ability [17,18].

At present, there is no commercial phosphate ion sensor in the electrochemical ion sensor market. The development of ion-selective sensors for phosphates is more challenging compared to other ions. The large tetrahedral structure of the phosphate ions results in high hydration energy and a tendency to remain in aqueous solutions, meaning that it is not easy for the ion-selective electrodes to exhibit excellent selectivity for phosphates [19]. Furthermore, phosphates in aqueous solutions undergo three levels of dissociation, and the valence state and relative distribution ratio of phosphates vary under different pH conditions, which brings great difficulty to the quantitative detection of phosphates. In recent years, some researchers have reported a lot of research progress on phosphate ion sensors. Many have developed a series of spectrophotometric phosphate sensors based on the spectral characteristics and fluorescent response of phosphates [20–28]. In terms of electrochemical sensors, cobalt metal is most commonly used in the research of phosphate sensors due to its unique responsiveness and selectivity to phosphate ions. Xiao et al. proposed a novel all-solid-state cobalt-based phosphate ion sensor, which presented a selective response to $\text{H}_2\text{PO}_4^-$ by modifying the surface of metallic cobalt with a cobalt phosphate-sensitive membrane [29]. Meruva et al. confirmed that the selective response of this cobalt-based phosphate electrode to $\text{H}_2\text{PO}_4^-$ was a mixed potential response mechanism and the response potential was affected by dissolved oxygen [30]. Xu et al. made further improvements to this cobalt-based electrode by coating cobalt with cobalt phosphate on the surface and conducted a further analysis of the electrode’s mechanism and performance [31]. To improve the sensitivity and stability of cobalt-based phosphate sensors, polymers and electron transfer media were used to fix cobalt powder and improve the electron transport efficiency. Zhao et al. proposed a new type of phosphate ISE composed of polypyrrole, cobalt, and ordered mesoporous carbon, which had good sensitivity and stability in response to $\text{H}_2\text{PO}_4^-$ and performed well in actual water sample detection [32]. Since cobalt-based sensors can only respond to $\text{H}_2\text{PO}_4^-$ under weakly acidic conditions, Xu et al. used the method of surface doping with phosphate complexes to develop a new type of tungsten-based phosphate ISE. This electrode showed a selective response to $\text{H}_2\text{PO}_4^-$ under neutral to weakly alkaline conditions, with a detection limit of $10^{-6}$ M under neutral conditions [33].
In addition, other metals, such as silver [34] and nickel [35], played a role in the research of phosphate ion sensors. While there has been extensive research on inorganic metals and their phosphates, some researchers have embarked on exploring the construction of polymer-sensitive membranes on an all-solid-state phosphate ISE. Jeong et al. prepared an effective phosphate ionophore using the special affinity of niacinamide and phosphate through hydrogen bonding and charge interactions; the sensor modified by this ionophore can determine phosphate with a relatively lower limit of $0.85 \times 10^{-6}$ M compared with most reported electrodes [36]. Ben et al. reported a novel PE-based phosphate sensor in which they embedded octamolybdate anions with periodic mesoporous organosilica nanosphere, which was suitable for detecting extremely low concentrations of phosphates with a detection limit of 0.16 nM [37].

In this study, we proposed a novel all-solid-state phosphate ISE that is efficient, accurate, low cost, easy to operate, and has a long lifetime. Our approach involved enhancing the surface electrochemical activity of the copper-based electrode through the modification of a copper nanoparticle film. Subsequently, the copper nanoparticle film created a mesh-like film structure with a significantly larger specific surface area by electrochemical modification. This microstructure not only enhanced electron transfer efficiency within the electrode but also improved the electrode surface membrane's response to target ions, with excellent sensitivity and stability. We analyzed and studied the response mechanism of the electrode by cyclic voltammetry, scanning electron microscopy, and energy spectrum analysis. Our systematic characterization of the prepared phosphate ISE covered aspects such as its detection limit, response time, selectivity, reproducibility, stability, and lifespan. Experimental results evidenced that the current phosphate ISE was capable of detecting phosphate ions across a broad pH range, from neutral to weakly alkaline.

2. Materials and Methods

The preparation process of the electrochemically modified all-solid-state copper electrode, designated as the copper phosphate electrode, is briefly described as follows. Firstly, we prepared a uniform copper nanoparticle film on the surface of copper wire through electrochemical deposition. Then, we fabricated a uniform copper phosphate-sensitive membrane on the surface of the above electrochemically treated copper wire through electrochemical synthesis. The schematic diagram is shown in Figure 1.

![Figure 1. Schematic diagram of electrode fabrication process.](image)

2.1. Reagents and Apparatus

The copper sulfate ($\text{Cu}_2\text{SO}_4$), sodium chloride ($\text{NaCl}$), sodium hydroxide ($\text{NaOH}$), nitric acid ($\text{HNO}_3$), disodium hydrogen phosphate ($\text{Na}_2\text{HPO}_4$), sodium dihydrogen phosphate ($\text{NaH}_2\text{PO}_4$), sodium sulfate ($\text{Na}_2\text{SO}_4$), sodium bicarbonate ($\text{NaHCO}_3$), sodium acetate ($\text{CH}_3\text{COONa}$), sodium silicate ($\text{Na}_2\text{SiO}_3$), sodium nitrate ($\text{NaNO}_3$), and copper wire (99.9%, 0.5 mm in diameter) were obtained from Sinopharm Chemical Reagent Co., Ltd.
(Shanghai, China). All chemical reagents were of analytical grade and used as received without further purification. All solutions used in the experiment were prepared with ultrapure water with a resistivity of 18.2 MΩ·cm.

We applied a SG1020A function generator (Rhett, Huaian, China) to carry out the nanoparticles’ electrodeposition process by connecting a 1N5819 Schottky diode (OnSemi, Scottsdale, AZ, USA). The electrochemical synthesis and the electrochemical performance testing of the electrode were implemented using a CHI760D electrochemical workstation (CHI, Shanghai, China). We purchased the Ag/AgCl reference electrode (3.8 mm) from Jinhong Electronic Technology Development Co., Ltd. (Shanghai, China). We acquired the platinum electrode (3.8 mm) from Gaoss Union Electronic Technology Co., Ltd. (Chengdu, China).

The SEM analysis and the EDS measurements were performed with a Zeiss Sigma 500 field emission scanning electron microscope (Carl Zeiss, Oberkochen, Germany). The pH of the solutions was measured with a E-201F composite pH electrode (INESA, Shanghai, China). The phosphomolybdenum blue spectrophotometric method was performed with a CleverChem 380 Plus auto discrete analyzer (DeChem-Tech, Hamburg, Germany).

2.2. Electrode Pretreatment and Fabrication

We polished a copper wire (φ0.5 mm × 50 mm) by using abrasive paper of #1000, #3000, #5000 and #7000 in turns for 2 min. It was then ultrasonically washed with 0.1 M nitric acid, acetone and deionized water in sequence for 15 min, and finally dried for later use.

In order to form a copper nanoparticle film, we applied a SG1020A function generator with sine wave parameters of an amplitude of 1.5 V and frequency of 50 Hz. The cathode was connected to the pretreated copper wire, and the anode was attached to a Schottky diode and contributing copper wire. Two copper wires were simultaneously immersed into a 0.05 M Cu₂SO₄ solution and were electroplated for 60 s. In the process of electrochemical deposition, the metallic luster on the surface of the copper electrode gradually disappeared, and the color became darker. Ultimately, a layer of copper nanoparticles was uniformly fabricated on the surface of the cathodic copper wire.

We carried out the electrochemical deposition process of copper phosphate by using a three-electrode system on the CHI760D workstation. A saturated Ag/AgCl electrode served as a reference electrode, a copper electrode modified with a copper nanoparticle film functioned as a working electrode, and a platinum electrode acted as a counter electrode. We placed the abovementioned electrodes simultaneously in 0.1 M Na₂HPO₄ solution at pH 9.0 and the electrochemical synthesis process of the copper phosphate-sensitive membrane was accomplished through a chrononamperometry method for 300 s at a potential of 0.8 V. In the process of constant potential electroplating, the surface of the copper electrode was gradually covered with a blue crystal film. After 300 s of electroplating, the electroplating was stopped when the surface morphology of the electrode was no longer changed. Then, the copper phosphate electrode dried naturally for the characterization and performance evaluation.

2.3. Electrode Performance Characterization

The SEM analysis was carried out to observe the surface morphology of the electrode and its elemental distribution was determined by the EDS measurement. All potential measurements were carried out with a two-electrode system on a CHI760D electrochemical workstation at room temperature using an Ag/AgCl electrode as a reference electrode and the prepared copper phosphate electrode as a working electrode. We recorded the potential values instantaneously until it was stable and evaluated the performance of the electrode according to the measurement results.
3. Results and Discussion
3.1. Electrochemical Characteristics of the Copper Electrode Modified with a Copper Nanoparticle Film in a Na₂HPO₄ Solution

We compared the electrochemical properties of a copper electrode modified with a copper nanoparticle film, designated as Cu-MCN, in systems with and without phosphate ions using cyclic voltammetry. The aim of this experiment was to ascertain whether Cu-MCN could undergo a specific electrochemical reaction with phosphate ions in a phosphoric acid-containing system, leading to the formation of an insoluble phosphate film. We analyzed the cyclic voltammetry (CV) curve to identify the appropriate reaction potential of Cu-MCN with phosphate ions. As depicted in Figure 2a, in a sodium chloride solution at pH 9, Cu-MCN exhibited an anodic peak around 0.3 V, and two distinct anodic peaks at 0.3 V and 0.75 V in a hydrogen phosphate disodium solution at the same pH. Comparing systems with and without phosphate ions revealed that the anodic wave at approximately 0.3 V occurred in both, attributable to the oxidation of Cu nanoparticles. This suggested that the dual peaks at 0.3 V and 0.75 V may stem from the electrochemical interaction between Cu-MCN and phosphate ions.

![Cyclic voltammetry curves of the Cu-MCN in different solutions at pH 9.0: (a) in 0.1 M Na₂HPO₄ and 0.1 M NaCl; (b) in Na₂HPO₄ of various concentrations.](image)

To delve deeper into the characteristics of these two anodic peaks, we assessed the concentration dependency of Cu-MCN in various phosphate solutions using cyclic voltammetry. There is a corresponding increase in the CV peak intensity as the concentration of Na₂HPO₄ solution increased from 0.01 M to 0.1 M (see Figure 2b), which confirms that the electrochemical activity at 0.75 V is indeed due to the reaction between Cu-MCN and phosphate ions.

We proceeded with the electrodeposition of the copper phosphate-sensitive membrane on the surface of the Cu-MCN, following the cyclic voltammetry experiments, which established the optimal reaction conditions for Cu-MCN with phosphate ions. This process utilized Cu-MCN and employed a chronoamperometry method at a potential of 0.8 V in a 0.1 M Na₂HPO₄ solution at pH 9. As illustrated in Figure 3a, the initial electrolytic current was around 2 mA. The current then surged as the copper nanoparticles dissociated from the solid metal to form copper ions. The high electrochemical activity of the copper nanoparticles on the copper wire’s surface caused the current to quickly peak at approximately 3 mA after around 30 s. Subsequently, the current gradually decreased due to the formation of copper phosphate compounds through the reaction of copper ions with phosphate ions on the electrode’s surface. As the electrodeposition continued, these phosphate compounds accumulated, covering the copper electrode’s surface. Approximately 5 min later, the current stabilized near zero and remained constant.
This ratio is consistent with the molecular structure of phosphates, indicating copper phosphate adheres uniformly on the electrode surface after the electrochemical electrodeposition, the copper particles with tens of nanometers in size uniformly coated on the surface of copper wire (Figure 4b), enhancing its specific surface area and electrochemical activity. In addition, this nanoparticle film tightly attached to the substrate surface can also strengthen the bond between the substrate and the sensitive membrane. Utilizing nanomaterials—recognized for their large specific surface area and exceptional electronic transport capabilities in microstructures—to modify electrodes can significantly boost electrochemical performance metrics such as sensitivity and response range, as cited in references [38,39]. As shown in Figure 4c, dense copper phosphate-sensitive film coated uniformly on the electrode surface after the electrochemical modification of the copper nanoparticle film, which indicated the electroplating was perfect. When enlarged amplification times, it can be observed that the copper phosphate adheres closely to the surface of the electrode substrate, forming a sheet-like network interwoven across the surface (Figure 4d). This thin film structure with a large specific surface area formed on the basis of copper nanoparticles not only enhances its responsiveness to target ions, but also has a low internal resistance [38].

Figure 3b illustrates the elemental composition and relative distribution ratios within the copper phosphate film, revealing an atomic ratio of phosphorus to oxygen nearing 1:4. This ratio is consistent with the molecular structure of phosphates, indicating copper nanoparticles successfully react with phosphate ions to form insoluble phosphate solid films. The copper phosphate electrode is a crystalline membrane electrode with metal insoluble salt as ion-sensitive membrane, and this ISE produces equilibrium potential and specific response to the anions of membrane components by precipitation dissolution equilibrium. CuHPO$_4$ and Cu$_3$(PO$_4$)$_2$ are the two main forms of insoluble phosphate salts of phosphate ion and copper ion (Cu(H$_2$PO$_4$)$_2$ is a water-soluble phosphate). For comparison, the copper to phosphorus atomic ratios in CuHPO$_4$ and Cu$_3$(PO$_4$)$_2$ are 1:1 and 3:2, respectively. The copper to phosphorus atomic ratio in the copper phosphate film is approximately 4:3, which falls between the ratios of the two referenced compounds. This indicates that the combined film is possibly composed mainly of Cu$_3$(PO$_4$)$_2$, with a smaller proportion of CuHPO$_4$. Therefore, the reaction described below Equations (1)–(7) [40] seems to occur on the electrode surface. Copper phosphate-crystalline film selectively respond to HPO$_4^{2-}$ in solution by achieving chemical equilibrium with it.

\[ \text{Cu + HPO}_4^{2-} \rightleftharpoons \text{CuHPO}_4 + 2e^- \] (1)
3Cu + 2PO$_4^{3-}$ $\rightleftharpoons$ Cu$_3$(PO$_4$)$_2$ + 6e$^-$  \hspace{1cm} (2)

2Cu + H$_2$O $\rightarrow$ Cu$_2$O + 2H$^+$ + 2e$^-$  \hspace{1cm} (3)

Cu$_2$O + H$_2$O $\rightarrow$ 2CuO + 2H$^+$ + 2e$^-$  \hspace{1cm} (4)

Cu$_2$O + 2HPO$_4^{2-}$ + H$_2$O $\rightleftharpoons$ 2CuHPO$_4$ + 2HO$^-$ + 2e$^-$  \hspace{1cm} (5)

CuO + HPO$_4^{2-}$ + H$_2$O $\rightleftharpoons$ CuHPO$_4$ + 2HO$^-$  \hspace{1cm} (6)

3CuHPO$_4$ + 3HO$^-$ $\rightleftharpoons$ Cu$_3$(PO$_4$)$_2$ + PO$_4^{3-}$ + 3H$_2$O  \hspace{1cm} (7)

Figure 4. SEM images of the surface of pure Cu wire (a), Cu nanoparticle layer (b), copper phosphate membrane layer (c), viewed at 3000× magnification; (d), viewed at 30,000× magnification.

3.3. Response Characteristics of the Copper Phosphate Electrode

In order to maintain a constant ionic strength in the standard calibration solution, we chose 1 mM NaNO$_3$ (nitrate is an inert electrolyte for the electrode) solution as the supporting electrolyte, which can maintain stable background currents and ionic strength. Prior to measurement, we must activate the electrode by immersing it into $10^{-3}$ M Na$_2$HPO$_4$ solution for 1 h at room temperature.

3.3.1. Response Slope, Detection Linear Range, Detection Limit and Reproducibility

The response potential of the electrode was measured in Na$_2$HPO$_4$ solutions (pH = 8) of varying concentrations, from $10^{-8}$ M to $10^{-1}$ M, and plotted to construct a calibration curve as depicted in Figure 5. This figure delineates two fitted lines across the concentration span. According to IUPAC guidelines [41], the segment with the steeper slope, corresponding to higher concentrations, is considered the detection range. The detection limit is defined by the intersection point on the x-axis of the two lines. The calibration curve showed that the electrode responded linearly to phosphate ion concentrations ranging from...
10⁻⁵ to 10⁻¹ M, with a response slope of −27.8 mV·dec⁻¹ and an R² of 0.9985, characteristic of Nernst behavior [41]. The detection limit was calculated to be 4 × 10⁻⁶ M.

![Graph showing response characteristics](image)

**Figure 5.** Response characteristics of the copper phosphate electrode in 10⁻⁸–10⁻¹ M Na₂HPO₄ standard solution at pH 8.

We test the repeatability of the electrode through five repetitive measurements of a 10⁻³ M Na₂HPO₄ solution at pH 8, yielding response potentials of 14.67 mV, 14.28 mV, 13.95 mV, 14.42 mV, and 15.03 mV. The standard deviation of these measurements was under 1 mV. For further testing of the reproducibility of the electrode, we measured five simultaneously fabricated phosphate ISEs by using 10⁻⁶ to 10⁻¹ M Na₂HPO₄ solutions at pH 8 and recorded the results. The response slopes of the five electrodes were −27.80, −27.45, −27.93, −28.12, and −27.68 mV·dec⁻¹, respectively. The average value was −27.8 (±0.4) mV·dec⁻¹, with a relative standard deviation (RSD) under 0.5%, which affirmed the electrodes’ great reproducibility.

### 3.3.2. Response Time

The response time is a crucial characteristic for analytical applications, impacting the accuracy of real-time monitoring and the sensor’s throughput. We evaluate the electrode’s response time by t₁/₂, which is the time required for the potential value to reach more than 95% of its final value according to the definition by IUPAC [41]. We immersed the prepared electrode into Na₂HPO₄ solutions ranging from 10⁻⁵–10⁻¹ M at pH 8, with continuous and repeated testing, and each measurement lasted for 300 s. Figure 6 shows that the electrode’s response time is less than 3 s from higher to lower Na₂HPO₄ concentration, while the average response time is approximately 10 s when measured in the reverse order. The swift response time of the prepared phosphate ISE can be ascribed to the efficient ion-to-electron transfer facilitated by the copper nanoparticle solid contact layer, which hastens the attainment of chemical equilibrium on the electrode surface.

Figure 6 also presents the final potential values at each stage as 69.59, 44.84, 14.67, −13.96, −40.52, −13.41, 14.60, 45.42, 69.97, −40.07, and 70.02 mV, respectively. Notably, the potential difference did not exceed 1 mV at the same concentration, indicating that the response behavior of the developed phosphate ISE remained consistent when the potential values were measured across concentrations in either direction. Furthermore, this experiment corroborated the absence of any memory effect in the electrode, as the results displayed no evidence of such an effect.
We evaluated the lifespan of the copper phosphate electrode by storing it in deionized water for 90 days, conducting periodic performance tests in a series of Na$_2$HPO$_4$ solutions at pH 8. Table 1 shows that for over 75 days, the electrode’s slopes remained stable between

![Figure 6](image-url)

**Figure 6.** Dynamic response time curve of the electrode in 10$^{-5}$–10$^{-1}$ M Na$_2$HPO$_4$ solutions at pH 8 using an open circuit technique.

3.3.3. Electrode Stability and Lifetime

The stability of the ISE is another concern for its practical application. To assess the copper phosphate electrode’s capability for long-term real-time monitoring of low-concentration phosphate ions, we put the prepared electrode into a Na$_2$HPO$_4$ solution of various concentration (10$^{-5}$ M, 10$^{-4}$ M, and 10$^{-3}$ M) for real-time monitoring for 5 h and continuously recorded its response potential. As shown in Figure 7, the average potential fluctuation was less than ±2 mV. Notably, the potential fluctuated significantly at a 10$^{-5}$ M concentration, possibly because of the solution instability at such low concentrations or that the concentration approached the electrode’s lower detection limit. At a 10$^{-3}$ M concentration, the potential gradually increased which may attribute to temperature changes.

![Figure 7](image-url)

**Figure 7.** Response potential measurements of the electrode over 5 h in 10$^{-5}$–10$^{-3}$ M Na$_2$HPO$_4$ solutions at pH 8.

The lifetime of an electrode is crucial for its application in long-term in-situ monitoring. We evaluated the lifespan of the copper phosphate electrode by storing it in deionized water for 90 days, conducting periodic performance tests in a series of Na$_2$HPO$_4$ solutions at pH 8. Table 1 shows that for over 75 days, the electrode’s slopes remained stable between
−27.01 and −29.42 mV·dec⁻¹, with determination coefficient R² exceeding 0.99, and the linear response range consistently maintained at 10⁻⁵−10⁻¹ M. On the 90th day, there was a significant alteration in response potential with marked declines in both the response slope and linear range, indicating a substantial loss in the electrode’s ability to detect phosphate ions. Thus, the copper phosphate electrode is viable for continuous use in an aqueous environment for up to 75 days, after which it requires modification.

Table 1. Performance of the electrode in 10⁻⁵–10⁻¹ M Na₂HPO₄ solutions at pH 8 over 90 days.

<table>
<thead>
<tr>
<th>Days</th>
<th>Potential (mV) of Different Concentrations</th>
<th>Linear Range</th>
<th>Slope</th>
<th>R²</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>10⁻⁵ M</td>
<td>10⁻⁴ M</td>
<td>10⁻³ M</td>
<td>10⁻² M</td>
</tr>
<tr>
<td>1</td>
<td>70.60</td>
<td>45.24</td>
<td>14.71</td>
<td>−10.36</td>
</tr>
<tr>
<td>3</td>
<td>69.32</td>
<td>44.51</td>
<td>16.27</td>
<td>−8.55</td>
</tr>
<tr>
<td>7</td>
<td>72.38</td>
<td>47.62</td>
<td>18.14</td>
<td>−11.54</td>
</tr>
<tr>
<td>15</td>
<td>68.19</td>
<td>42.25</td>
<td>13.68</td>
<td>−13.36</td>
</tr>
<tr>
<td>30</td>
<td>73.77</td>
<td>46.35</td>
<td>19.81</td>
<td>−7.65</td>
</tr>
<tr>
<td>45</td>
<td>71.42</td>
<td>41.63</td>
<td>15.58</td>
<td>−10.34</td>
</tr>
<tr>
<td>60</td>
<td>70.95</td>
<td>40.16</td>
<td>12.35</td>
<td>−14.74</td>
</tr>
<tr>
<td>75</td>
<td>75.63</td>
<td>47.81</td>
<td>16.48</td>
<td>−11.26</td>
</tr>
<tr>
<td>90</td>
<td>40.25</td>
<td>28.56</td>
<td>10.73</td>
<td>−8.61</td>
</tr>
</tbody>
</table>

3.3.4. Electrode Selectivity

The specific selectivity of ISE towards target ion is relative, and it can also respond to coexisting ions in the solution to varying extents. Therefore, the specific response of the electrode to the target ion can be disturbed by other ions with a similar charge in aqueous solution. Given the significantly complex ion environment in real water bodies, the selectivity of an electrode for the common ions is crucial for reliable determination in actual water bodies. In the presence of interfering ions, the Nikolskii–Eisenman equation [42] (Equation (8), the generalized Nernst equation) determines the response potential of the electrode to the target ion. In Equation (8), the selectivity coefficient Kᵢ⁾j quantifies the influence of interfering ion j on the response potential of target ion i, indicating the electrode’s preferential ability to selectively respond to the ion i in the presence of ion j.

\[
E = E^0 + \frac{2.303RT}{z_iF} \log(a_i + \sum_j K_i^{pot} a_j) \tag{8}
\]

Here, ion i and ion j are the target ion and the interfering ion, respectively; E is the measured response potential; E⁰ is the reference potential under standard conditions, which can be regarded as a constant under certain measurement conditions; R is the gas constant (8.314 J/K·mol), T is the absolute temperature (273.15 + t °C), F is the Faraday constant (9.648 × 10⁴ C/mol); zᵢ and zⱼ are the charge numbers of the ions i and j, respectively; Kᵢ⁾j is the selectivity coefficient of the target ion i toward interfering ion j, and the logarithm of selectivity coefficient logKᵢ⁾j < 0 means that the electrode displayed excellent selectivity for ion i over the ion j; the smaller the value of logKᵢ⁾j, the better the selectivity of the electrode to the target ion [43]; aᵢ and aⱼ are the activities of ions i and j, respectively. There is the following equation (Equation (9)) between ion activity and ion concentration in solution:

\[
a_i = y_i c_i \tag{9}
\]

Here, aᵢ is the activity of ion i; yᵢ is the activity coefficient of ion i, which is related to the ionic strength in the solution; cᵢ is the ion concentration. In general, in a solution with a relatively simple composition and constant ionic strength, the ion activity in the Nernst equation can be replaced by the ion concentration.
The separated solution method (SSM) and the fixed interference method (FIM) are two common analytical methods recommended by IUPAC [44] for determining selectivity coefficient \( K_{ij}^{pot} \). We analyzed and calculated the selectivity coefficients \( K_{ij}^{pot} \) of the electrode for different interfering ions using the FIM based on Equation (8). In this method, within mixed solutions containing both interfering ion \( j \) and target ion \( i \), we maintained the concentration of ion \( j \) constant (10\(^{-3}\) M) while varying the concentration of ion \( i \) (10\(^{-6}\)–10\(^{-1}\) M). We measured the response potential of the electrode in a series of mixed solutions with concentration gradients of ion \( i \), and calculated the selectivity coefficient \( K_{ij}^{pot} \) by the simplified equation (Equation (10)) based on Equation (8):

\[
K_{ij}^{pot} = \frac{a_i a_{z,j}}{a_j}
\] (10)

As shown in Table 2, we established the following selectivity sequence: \( \text{HPO}_4^{2-} > \text{OH}^- > \text{Cl}^- > \text{SiO}_3^{2-} > \text{HCO}_3^- > \text{CH}_3\text{COO}^- > \text{SO}_4^{2-} > \text{NO}_3^- \). The \( \log K_{ij}^{pot} \) values for common anions, aside from the hydroxide ion, are less than \(-2\), indicating that the electrode’s response to the target phosphate ion is unaffected by these interfering ions when their concentration is less than 100 times that of the phosphate ion. However, the hydroxide ion exhibits strong interference compared to other interfering ions, likely because the solubility product of \( \text{Cu(OH)}_2 \) is quite low (\( K_{sp}(\text{Cu(OH)}_2) = 2.2 \times 10^{-20} \)). At relatively high concentrations of hydroxide ions, a chemical equilibrium is established between copper ions and hydroxide ions, which can significantly interfere with the copper phosphate electrode—an crystalline membrane electrode that relies on chemical equilibrium for a Nernst response.

Table 2. Selectivity coefficients of common anions measured by FIM.

<table>
<thead>
<tr>
<th>Interfering Ions</th>
<th>Reagent</th>
<th>( \log K_{ij}^{pot} )</th>
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<tbody>
<tr>
<td>( \text{SO}_4^{2-} )</td>
<td>( \text{Na}_2\text{SO}_4 )</td>
<td>-4.13</td>
</tr>
<tr>
<td>( \text{HCO}_3^- )</td>
<td>( \text{NaHCO}_3 )</td>
<td>-3.28</td>
</tr>
<tr>
<td>( \text{NO}_3^- )</td>
<td>( \text{NaNO}_3 )</td>
<td>-4.76</td>
</tr>
<tr>
<td>( \text{CH}_3\text{COO}^- )</td>
<td>( \text{CH}_3\text{COONa} )</td>
<td>-3.57</td>
</tr>
<tr>
<td>( \text{Cl}^- )</td>
<td>( \text{NaCl} )</td>
<td>-2.04</td>
</tr>
<tr>
<td>( \text{OH}^- )</td>
<td>( \text{NaOH} )</td>
<td>-0.85</td>
</tr>
<tr>
<td>( \text{SiO}_3^{2-} )</td>
<td>( \text{Na}_2\text{SiO}_3 )</td>
<td>-3.46</td>
</tr>
</tbody>
</table>

3.3.5. pH Interference on the Phosphate Ions’ Measurement

As phosphoric acid belongs to ternary weak acid, which has a three-stage dissociation in aqueous solution (see Equations (11)–(13)), Equations (14)–(16) (in the equation: \([\text{H}_3\text{O}^+] = [\text{H}^+] = 10^{-p\text{H}}\) show the dissociation equilibrium constant of phosphoric acid at different stages at room temperature. Under different pH conditions, the existing forms of phosphate hydroxyl groups and the distribution proportions of various valence phosphate ions in different dissociation states are different. The morphological distribution diagram (Figure 8) of phosphate ions was plotted through the calculation of Equations (11)–(16). Figure 8 shows that the dominant species of phosphate ions is \( \text{H}_2\text{PO}_4^- \) in the solution from pH 5 to pH 6, and \( \text{HPO}_4^{2-} \) predominates in the solution pH range of 8 to 11.

\[
\begin{align*}
\text{H}_3\text{PO}_4 + \text{H}_2\text{O} & \rightleftharpoons \text{H}_2\text{PO}_4^- + \text{H}_3\text{O}^+ \quad (11) \\
\text{H}_2\text{PO}_4^- + \text{H}_2\text{O} & \rightleftharpoons \text{HPO}_4^{2-} + \text{H}_3\text{O}^+ \quad (12) \\
\text{HPO}_4^{2-} + \text{H}_2\text{O} & \rightleftharpoons \text{PO}_4^{3-} + \text{H}_3\text{O}^+ \quad (13) \\
K_{a1} & = \frac{[\text{H}_2\text{PO}_4^-][\text{H}_3\text{O}^+]}{[\text{H}_3\text{PO}_4]} = 7.5 \times 10^{-5} \quad (14)
\end{align*}
\]
We studied the effect of solution pH in the range of 5 to 12 on the electrode’s potential response of phosphate ions. The phosphate solutions with different pH values were adjusted by using different concentrations of sodium hydroxide and nitric acid based on Na$_2$HPO$_4$ and NaH$_2$PO$_4$. As shown in Figure 9, the electrode exhibited optimal performance at pH 7 with a detection limit of 10$^{-6}$ M when the dominant species of phosphate ions gradually transitioned to HPO$_4^{2-}$ in the solution pH range of 6 to 11, while the detection limit decreased to about 10$^{-4}$ M at pH 6 and pH 11 as well as about 10$^{-5}$ M in the pH range of 8 to 10. The response slopes from pH 6 to pH 11 were $-29.4$, $-29.31$, $-27.9$, $-26.7$, $-25.5$ and $-22.32$ mV·dec$^{-1}$, respectively. In addition, the electrode’s response to phosphate ions completely deviated from the Nernst behavior at pH 5. We infer that the performance loss and gradual deviation from Nernst response of the electrode in alkaline solutions arise from the interference of a relatively high concentration of OH$^-$, which leads to the precipitation–dissolution equilibrium of insoluble Cu(OH)$_2$ formed by the reaction of hydroxide ions, and copper ions gradually dominate the electrode surface.

As a result, the prepared electrode exhibited an excellent selectivity to HPO$_4^{2-}$ and can be used for detecting solutions with a pH range from neutral to weakly alkaline. In the practical applications, it is crucial to consider the actual pH of the solutions to ensure accurate phosphate ions detection.
phosphate was detected in these samples (either these samples may not contain phosphate or the phosphate concentration is less than $10^{-5}$ M, and we only need to detect them when their phosphate content exceeds the normal standard or higher), after adding a known quantity of phosphate, the recovery values derived from the electrode’s detection results affirmed the method’s accuracy. Table 4 presents that the total phosphate content detected by the electrode in lake water and aquaculture water samples are close to the results measured by the CleverChem 380 Plus, with a relative standard deviation (RSD) of about 10%. The results showed that the prepared phosphate ISE was capable of application for phosphate detection in real water bodies.

**Table 3.** Recovery test in different solutions.

<table>
<thead>
<tr>
<th>Sample</th>
<th>pH</th>
<th>Adjusted pH</th>
<th>Added (mM)</th>
<th>Found (mM)</th>
<th>Recovery Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tap water</td>
<td>7.20</td>
<td>8.03</td>
<td>0.1</td>
<td>0.097 ± 0.002</td>
<td>97</td>
</tr>
</tbody>
</table>
Table 3. Cont.

<table>
<thead>
<tr>
<th>Sample</th>
<th>pH</th>
<th>Adjusted pH</th>
<th>Added (mM)</th>
<th>Found (mM)</th>
<th>Recovery Rate (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drinking natural water</td>
<td>7.63</td>
<td>9.02</td>
<td>1</td>
<td>1.02 ± 0.01</td>
<td>102</td>
</tr>
<tr>
<td>Laboratory wastewater</td>
<td>5.28</td>
<td>7.02</td>
<td>10</td>
<td>9.50 ± 0.4</td>
<td>95</td>
</tr>
</tbody>
</table>

Table 4. Comparison of the phosphate concentrations detected by the phosphate ISE and CleverChem 380 Plus in real aqueous samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>pH</th>
<th>Adjusted pH</th>
<th>Determined by CleverChem 380 Plus (µM)</th>
<th>Determined by ISE (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lake water</td>
<td>8.24</td>
<td>7.01</td>
<td>5.33 ± 0.08</td>
<td>6.04 ± 0.21</td>
</tr>
<tr>
<td>Aquatic water</td>
<td>8.05</td>
<td>6.98</td>
<td>10.76 ± 0.06</td>
<td>11.35 ± 0.13</td>
</tr>
</tbody>
</table>

3.3.7. Comparison of the Copper Phosphate Electrode and Previously Reported Electrodes

In summary, Table 5 presents a comparative analysis with the relevant literature, showcasing the competitive edge of our copper phosphate ISE. The detection range and the limit of our phosphate ISE are on par with those of similar ISEs reported in prior research. Notably, our phosphate ISE exhibits a wider pH detection range for phosphate ions, with a detection limit in neutral solutions of $1 \times 10^{-6}$ M. Additionally, it features the fastest response time recorded at 10 s and an impressive service life of 75 days, outperforming most recently reported phosphate ISEs.

Table 5. Performance comparison of references and current research.

<table>
<thead>
<tr>
<th>Phosphate ISE Type</th>
<th>Detection Range/M</th>
<th>Detection Limit/M</th>
<th>Slope (mV·dec$^{-1}$)</th>
<th>Response Time (s)</th>
<th>pH Range</th>
<th>Lifetime</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Co-PPy-OMC/GCE</td>
<td>$10^{-5}$–$10^{-2}$</td>
<td>$6.81 \times 10^{-6}$</td>
<td>$-31.6$</td>
<td>9</td>
<td>3–5</td>
<td>NG</td>
<td>[32]</td>
</tr>
<tr>
<td>Nickel</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>$10^{-5}$</td>
<td>$-81$</td>
<td>20</td>
<td>4–7</td>
<td>4 weeks</td>
<td>[35]</td>
</tr>
<tr>
<td>Tungsten</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>$1 \times 10^{-6}$</td>
<td>$-24.6$</td>
<td>60</td>
<td>7–10</td>
<td>4 weeks</td>
<td>[33]</td>
</tr>
<tr>
<td>Cobalt wire</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>NG</td>
<td>$-39$</td>
<td>30</td>
<td>4.0–6.5</td>
<td>4 weeks</td>
<td>[31]</td>
</tr>
<tr>
<td>Molybdenum</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>$10^{-5}$</td>
<td>$-27.8$</td>
<td>300</td>
<td>8–11</td>
<td>1 month</td>
<td>[47]</td>
</tr>
<tr>
<td>llP</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>$7.6 \times 10^{-6}$</td>
<td>$-3.7$</td>
<td>1800</td>
<td>2–7</td>
<td>NG</td>
<td>[48]</td>
</tr>
<tr>
<td>Copper</td>
<td>$10^{-5}$–$10^{-1}$</td>
<td>$1 \times 10^{-6}$</td>
<td>$-27.8$</td>
<td>10</td>
<td>6–11</td>
<td>75 days</td>
<td>This work</td>
</tr>
</tbody>
</table>

NG not given.

4. Conclusions

In this study, we proposed an innovative all-solid-state ISE for phosphate ions, utilizing a copper/nano-copper/copper phosphate matrix. This sensor is not only economical to produce but also simple in its manufacturing process. We characterized the electrode’s surface morphology and elemental composition with a scanning electron microscope and an energy dispersive spectrometer, respectively. Nanoscale modifications to the electrode have significantly expanded its specific surface area, boosting its efficiency in responding to target ions and enhancing both its sensitivity and stability. The electrode performed remarkable sensitivity and stability, evidenced by a swift response time of less than 10 s and fewer potential fluctuations, staying within ±2 mV over a 5 h continuous testing period. The interference tests indicated the electrode’s excellent selectivity against common anions found in natural water sources. The detection range and limits of the phosphate ISE developed through this research are on par with the existing literature. The advantage of this electrode is its easy fabrication, capacity to detect accurately across a wide pH spectrum from 6 to 11, and a quick response time, combined with a service lifetime of two months. And these qualities hold the potential to mark it as a viable candidate for commercial test instrumentation. Furthermore, the results of this study provide feasible
reference information for the design of new chemical sensors tailored for environmental monitoring purposes.


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