Biopolymer Test Kit for Colorimetric Detection of Chlorine in Water

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Abstract. The objective of the study was to fabricate the colorimetric sensor of biodegradable material for free chlorine determination. The colorimetric reagent of N,N-diethyl-p-phenylenediamine sulfate (DPD) was entrapped in the hybrid biopolymer film of agar (AG) and tapioca starch (TAS) and it was coated on the plastic micro-PCR tube. The pink product obtained from the reaction between DPD reagent and chlorine could indicate the presence of residual chlorine in the water. The condition for the sensor film synthesis was optimized by the digital image analytical technique with mobile phone application. The results were showed that Red-Green-Blue (RGB) intensity of reaction product was not changed, even through the DPD reagent was added over 0.2 g/mL. The addition of 16 g/L EDTA in the buffer solution could reduce the interference effect from some metals, especially Fe³⁺, contaminated in water sample. The water pH could be maintained for best analysis at the volume ratio between buffer and DPD solution of 0.5:1. The incubation of colorimetric film at 60 °C and 60 minutes provided the best sensor performance with fast analysis of 1 min reaction time. In conjunction with the digital image colorimetry (DIC), the developed test kit did not provided only the qualitative information, but the rapid quantitative analysis could be also fulfilled. A wide linear range of 0.3 to 15 mg/L chlorine concentration with good linearity (R² > 0.99) was achieved by this coupled technique. The application of biopolymer film to various kind of real water samples showed the good performances, which were comparable with the standard spectrophotometry (no significantly different results at 95% confidence level). These could promote the use of biopolymer test kit as the environmentally-friendly analytical method for chlorine in water.

Keywords: Biopolymer, colorimetric sensor, free chlorine, RGB intensity, digital image analysis.
1. Introduction

Chlorine (Cl₂) is a typical oxidizing agent widely used for water disinfection in tap water, drinking water, swimming pools and recycled water from wastewater [1, 2]. When a concentration of liquid solid or gas chlorine are added into water, they are transformed into hypochlorous acid (HClO) and hypochlorite ion (ClO⁻) which generally defined as free residual chlorine [3]. Too low chlorine concentration in water cannot kill pathogenic bacteria and causes many hazards of an insufficient disinfection [4]. On the other hand, free residual chlorine with too high level might produce many undesirable byproducts such as trihalomethanes (THMs) [5, 6], which have been proved to be carcinogenic and harmful to human and animals [7]. The World Health Organization (WHO) has recommended that acceptable chlorine level in drinking water should not exceed 5 mg/L [8]. The United States Environmental Protection Agency (EPA) has established the maximum free chlorine concentration level of 4 mg/L for both swimming pools, when it is opened to the public, and drinking water [9, 10]. In Thailand, the concentration of free chlorine residual in natural waters allowed by legislation should not exceed 1 mg/L [11].

A large number of analytical methods have been developed for determination of free chlorine in water including ion chromatography [12], chemiluminescence [13], fluorescence [14], amperometry [15], and spectrophotometry [16]. Of all methods, the most commonly used one is a colorimetric method, which is based on the reaction of N,N-diethyl-p-phenylenediamine (DPD) reagent [17, 18] and analysis of free chlorine by spectrophotometer. However, the application of conventionally spectrophotometric technique is suffered from some drawbacks, such as non-portability of laboratory instrument and low capability for real-time or on-line monitoring. In recent years, the digital image-based methods have frequently been used as an alternative to develop fast and low-cost colorimetric detections for quantitative chemical analysis [19-21]. The principle of these methods is based on the analysis of RGB (red-green-blue) color system from the digital images of colorimetric product. Various color analysis program, e.g. Adobe Photoshop and MATLAB-based program, could be used to provide the recorded RGB values [22, 23]. Also, the digital image colorimetry was successfully developed for the detection by using other commercial applicationor customized-built programs on the smartphone [24, 25].

The colorimetric sensor showed the advantage of rapid and on-site measurement. It is based on the entrainment of color reagent in support materials, e.g. sol-gel, hydrogel and synthetic polymer. Even through, most substrates trend to be recalcitrant to naturally biodegraded process and some can be dissolved in water or widespread to the environment. For this reason, the natural polymers, especially a group of polysaccharides such as starch, become considerably interested for the development of biodegradable sensors [26-29]. Several studies were performed to analyze the properties of starch-based films [30-32] and they reported that films possessed the poor mechanical properties and high moisture sensitivity. However, the addition of agar was able to enhance the tensile strength of starch-based film to obviously reduction on elongation [33, 34], and also showed the good compatibility with the starch texture [35, 36]. Thus, agar-strach blending film is an interesting alternative as a biopolymer for the development of chemical sensor due to a low cost polymer and eco-friendly material.

The objectives of this study was focused on the development of the chemical sensor from hybrid biopolymer of starch and agar, which could be applied with digital image technique for the determination of free residual chlorine in water. The testkit of film sensor was fabricated in the small plastic tube to promote the portability for the field measurement, less reagent consumption and environmentally friendly method.

2. Experiment

2.1. Chemicals and Materials

Sodium hypochlorite (NaOCl with available chlorine 7% w/w) was purchased from Carlo Erba (Milan). N,N-diethyl-p-phenylenediamine sulfate (DPD) was obtained from Sigma-Aldrich (USA), and the other reagents were obtained from Ajax Finechem (Australia). NaOCl solution was used as the free chlorine with concentration of approximately 5 mg as Cl₂/L for all tests, if not mentioned others and it was freshly prepared for daily use. The plastic micro-PCR tube 1.5 mL was used as a reaction container, where a coated film was on its lid (as showed in Fig. 1.)

![Fig. 1. Testkit with a thin sensor film coated on the lid (A) of the plastic micro-PCR tube (B).](image)

2.2. Preparation of Sensing Polymers

2.2.1. Colorimetric reagents

There were 2 reagents related to the colorimetric reaction of free chlorine, i.e. N,N-diethyl-p-phenylenediamine (DPD) sulfate reagent and phosphate buffer. The DPD
reagent as chromogenic species was daily prepared by dissolving appropriate amount of DPD sulphate in 10 mL ultrapure water containing 80 µL of 3M H₂SO₄ solution (pH 2) before stored in an amber bottle to avoid any interferences from a light. Phosphate buffer at pH 6 was prepared by dissolving 0.24 g Na₂HPO₄ and 0.46 g KH₂PO₄ in 10 mL of ultrapure water with proper disodium ethylenediaminetetraacetic acid (EDTA) and it was stored in a refrigerator at 4°C before used.

2.2.2. Agar-starch hybrid polymer films

Tapioca starch (TAS) (Erawan Brand, Nakhon Pathom, Thailand) and agar (AG) (Nang Kwak Brand, Bangkok Thailand), which were used as the film forming components, could be easily purchased from a local market in Phuket, Thailand. A mass series of agar (0.025, 0.05, 0.1, 0.2 g) and starch (0.05, 0.1, 0.2, 0.3, 0.4 g) solution were prepared in 10 mL ultrapure water purified by a water purification system (Merek, Darmstadt, Germany). Each solution was simultaneously heated at 90 °C for 15 min before cooled down to ambient temperature for 5 min. Then, both solutions were mixed well at a 2:1 volume ratio of AG: TAS solution under a magnetic stirring condition and the clear solution (AG-TAS) was obtained in a few seconds.

The sensing film was fabricated by blending the appropriate proportion of colorimetric reagent and buffer mixture into 3.5 mL AG-TAS solution. 100 µL of the resulting mixture was then dropped on the inside lid of a micro-PCR tube and dried in hot air oven at 60 °C for 60 minutes. After cooled to room temperature, the sensor testkit was in a ready-to-use condition with 1 mL water sample of free chlorine.

2.3. RGB Imaging System and Colorimetric Quantification of Chlorine

Since the principle of digital image analysis was based on the measurement of light reflection from the reaction product, the custom-built analytical box was set up to limit the detection interference from the environmental light, such as sunlight. A photographic box with 15.5 cm width x 15.5 cm length x 19.5 cm height (Fig. 2) was made of opaque black corrugated plastic board with a white interior. Three light-emitting diode (LED) rows were placed on the walls inside the box as light sources. Front side of the box was perforated for an installation of camera built in the mobile phone (iPhone 7 with 12MP and backside illuminated CMOS), which was operated in the flash-off mode (f/1.8 aperture and ISO 50) throughout experiments.

A series of chlorine standard solution (0.1-50 mg/L) was freshly prepared to prevent any errors from unstability of chlorine. Colorimetric analysis was performed when 1 mL of water sample was poured into a testkit tube with the biodegradable hybrid film on the lid and mixed by shaking for 1 min. The resultant pink products were appeared in the tube, due to the reaction between free chlorine and DPD dissolved from the film. In a process of digital image colorimetry (DIC), three sensor tubes were hung on the top of the box and they was photographed for 6 images (4032 × 3024 pixels, 1.33 MB). The average intensities of the red, green and blue colors (RGB values) across six images from three sensors were investigated using an in-house RGB analysis program. Those average RGB data of each standard concentration were used as a single data point to establish calibration curve by Microsoft Excel 2010 for further chlorine quantification.

Fig. 2. A custom built photographic box.

2.4. Analysis of Real Sample

Real samples were analyzed by using the biopolymer test kit and DIC for measurement of chlorine concentration. These grab samples were randomly collected from different kinds of water possibly contaminated by chlorine, i.e. the swimming pool, salt water pool, tap water and sea water at Patong beach, Phuket. Each sampling point was done with three replicates of samples. The concentration of chlorine in all water samples were analyzed by the developed test kit coupled with DIC within an hour after collection. Those average concentration were compared with the results from standard spectrophotometric method, or named as DPD colorimetric method [37].

3. Results and Discussion

3.1. Effect of Film Content on Reaction Product

3.1.1. Concentration of DPD and sulfuric acid in chromogenic reagent

A colorimetric test of chlorine was based on the reaction between free chlorine and DPD chromogenic species, i.e. a pink product of Würster dye was formed when DPD reacts with small amounts of chlorine [16]. Thus, the determination of DPD concentration in the hybrid film plays an important role to chlorine analysis. Moreover, the chemical cost was one of reasons prohibited someone to use any methods to detect the
concerned pollutant. Because of these, the effect of DPD concentration on the color intensity of reaction product was studied in the range of 0.0011-0.6 g/mL.

As showed in Fig. 3(a), the red intensity was higher than the blue and green, respectively. This was because the reaction product was pink. The reflection of red color was highest among others. In constrast, the green intensity was lowest. This was attributed to its complimentary color of red. The results in Fig. 3(a) were also revealed that the color intensity was decreased with the increasing DPD concentration in the range of below 0.2 g/mL. This caused by more reaction product formation leading to the strengthen color and lower light reflection. Even through, the intensity became constant after 0.2 g/mL DPD concentration. It might be due to the chlorine was used up. Therefore, a concentration of 0.2 g/mL of the DPD reagent contant was applied to form the colorimetric film.

The DPD was unstable to the oxidation from either atmospheric oxygen or dissolved oxygen presented in the prepared water. It was reported that the oxidation of DPD by oxygen was pH dependent [38]. Therefore, the solution of 3M sulfuric acid was recommended to add into the DPD reagent so as to lower the pH and retard the oxidation of DPD species. The effect of concentrated sulfuric acid volume in DPD reagent was therefore investigated from 0 to 400 µL/mL. However, the results in Fig. 3(b) suggested that the sensor response was decreased with the increasing volume of acid solution. In fact, many kinds of waters might often contain the dissolved oxygen, thus the sensors with/without acid were tested with the real water sample to monitor the effect of DPD loss from oxidation. The experiment was designed by the application of the 0.8 µL of 3M sulfuric acid into 1 mL DPD reagent used for the film fabrication. The sensor film was tested with samples of the tap water and swimming pool and compared the results from the sensor of no acid addition. Since the digital image colorimetry was based on the measurement of object color, the color of background sample could interfere to RGB light intensity. In order to eliminate the interference from background color of water sample, the RGB intensity of blank was subtracted from those of sample. According to Fig. 3(c), it was clear that RGB intensity difference between the reaction product and

![Fig. 3. (a) Effect of DPD concentration in film on RGB intensities, (b) Effect of sulfuric acid concentration in film on RGB intensities and (c) the application of film sensor with/without the sulfuric acid to real water samples.](image-url)
blank obtained from the sensor film with 0.8 µL/mL sulfuric acid was better than the sensor without acid for both kinds of real samples. Thus, the addition of 0.8 µL of 3M sulfuric acid was recommended as the good proportion of 1 mL DPD reagent used for the film fabrication.

3.1.2. Concentration of EDTA in a phosphate buffer solution

EDTA was well-known as a powerful chelating agent, which could form the highly stable complex with a metal ligand. It was suggested to insert into the buffer solution so as to eliminate the metal interference in water sample. In this study, Fe³⁺ was selected as nominal metal, because it was often found in many kinds of water, e.g. surface water, groundwater and even tap water for some cases. According to Fig. 4(a), various Fe³⁺ concentration was added into 5 mg/L chlorine solution and it was found that the presence of Fe³⁺ would be not affected to the measurement of free chlorine, if its concentration was less than 50 mg/L. This indicated that Fe³⁺ could competitively react with DPD to form the pink dye reaction product. Therefore, it was necessary to add EDTA into the buffer solution to reduce the interference effect of Fe³⁺. The appropriate EDTA concentration in the range 0-28 g/L was investigated for the chlorine sample containing 50 mg/L Fe³⁺ ion and the results were displayed in Fig. 4(b). It was noticed that the increasing EDTA concentration increased in the RGB intensity ratio between the chlorine sample with Fe³⁺ and those without Fe³⁺. The ratio was risen up closely to 1.0, when the amount of EDTA was more than 16 g/L. This implied that the optimum concentration of EDTA in buffer solution, which could mitigate the problem of metal interference, was 16 g/L.

3.1.3. Volume ratio of the phosphate buffer to DPD reagent

The phosphate buffer played an important role to maintain the solution pH and stabilize the reaction product, leading to enhance the detection sensitivity [16, 18]. In this study, the sensor film was fabricated from three portion of solution, i.e. DPD reagent, phosphate buffer and hybrid biopolymer (AG and TAS). To monitor the effect of buffer solution on the colorimetric reaction, the proportional ratio of phosphate buffer to DPD reagent volume was varied at 0.5:1, 1:1, 2:1 and 3:1 mL. The volume of biopolymer was adjusted correspondingly with those ratio to keep the total volume of mixed solution to be 5 mL, as showed in Table 1.

In general, the free chlorine was employed in the disinfecting process for the swimming pool and tap water. The standard pH of swimming pool and tap water was recommended at 7.2-7.6 and 6.5-8.5, respectively. Thus, the effect of buffer and DPD ratio was studied in the range of 0:1 to 3:1. As presented in Fig. 5, there was insignificant difference in RGB intensity of the ratio between 0.5-2. To save the chemical cost and reduce the chemical waste after analysis, the ratio of phosphate buffer and DPD reagent was suggested at 0.5: 1 for testkit production.

### Table 1. Combination ratio of phosphate buffer to DPD reagent in the biopolymer blends for 5 mL of total solution.

| Ratio of reagents | Phosphate buffer (mL) | DPD reagent (mL) | AG+CAS (mL) (2:1 ratio) |
|-------------------|-----------------------|------------------|-------------------------|
| 0:1               | 0                     | 1.0              | 4                       |
| 0.5:1             | 0.5                   | 1.0              | 3.50                    |
| 1:1               | 1.0                   | 1.0              | 3.0                     |
| 2:1               | 2.0                   | 1.0              | 2.0                     |
| 3:1               | 3.0                   | 1.0              | 1.0                     |
3.1.4. Effect of hybrid biopolymer component

It has been reported that AG was able to provide a very good cohesive matrix, which contributed to enhance the mechanical properties of polysaccharide based film. Besides, the blending AG with TAS increased the surface wettability, inducing an improvement in elongation and tensile strength [36]. In this research, the biopolymer was generated from the blending of AG and TAS solution at the ratio of 2:1 (v/v), before mixing with DPD and buffer solution at above proportion. The optimum concentration used to prepare the AG solution was studied in the range of 2.5-20 g/L, while those for TAS solution was 5-40 g/L. Their results was presented in Fig. 6(a) and (b), respectively.

According to Fig. 6(a), the increasing in agar concentration decreased the RGB intensity, or built up the magnitude of reaction product. It was attributed that the structure of biopolymer with more agar concentration would foster the characteristics of wettability [36], which could possibly promote the diffusion of colorimetric reagent from sensor film, leading to accelerate the reaction kinetic. However, the excess of agar concentration caused to reduce the flexibility of sensor film and affected to the difficulty of fabricated process. The results in Fig. 6(b) showed the same phenomenon as the agar section. The existence of high starch concentration reduced the formation of color reaction product. It was because the increasing in the starch molecule would patronize the denser structure of biopolymer. This could impede the diffusion of chromogenic species from the polymer film to water layer, leading to lower reaction rate. That's why the concentration of 2.5 g/L of AG and 5 g/L of TAS was chosen as good optimum condition for film forming.
3.2. Effect of Incubated Temperature and Time

After 100 μL mixture of the buffer-containing DPD reagent and polymer solution were dropped on the testkit lid, the polymerization was operated at high temperature. The effect of incubated temperature and time was studied in the range of 50-110 °C and 20-90 min, respectively. As exhibited in Fig. 7(a), the best performance of sensor kit was obtained by the highest subtracted RGB intensity at 70 °C. The film polymerization at higher temperature was resulted in the regression of subtracted intensity. This implied that the excess energy delivered to the polymer could deteriorate the film structure and homogeneity. Some parts of film were turned brown as noticed by naked eye. Since RGB intensity from 60 °C and 70 °C incubated films was different less than 4%, the the polymerization at 60 °C was selected as the optimum incubated temperature to save the energy for the film production.

For the effect of incubated time as presented in Fig. 7(b), there was insignificantly difference in the subtracted RGB intensity over the studied range, even at short incubated time where the film did not formulated. This indicated the trapping of chromogenic species by polymer did not obstruct to its diffusion to the water phase. That’s why the subtracted RGB intensity from DPD in reagent condition was not different with those of film condition. However, the incubated time was recommended at 60 min (for 60 °C), because of the film thoroughly formulated without any liquid textures.

3.3. Effect of Reaction Time on the Chlorine Measurement

The effect of time on the reaction between chlorine and DPD from the thin film under the synthetic condition optimized as above was studied in the range of 0 – 30 min. The experimental results revealed that the color product was completely generated within 1 min, as shown in terms of the maximum subtracted RGB intensity in Fig. 8. However, the subtracted intensity was trended to decrease when the reaction time was longer than 4 min. There were 2 possible reasons. First, the increasing in dissolution of color reagent from the blank film could be detected and it was therefore resulted in the subtracted RGB value. Second, the color product could be destabilized at longer reaction time. To prevent the discolor of subtracted RGB, the optimum time for detection should be applied at 1 min.

3.4. Quantification of Chlorine by Using the Biopolymer Test Kit with DIC

After the optimum condition of film fabrication was determined, the test kit was ready-to-use for qualitative analysis. However, the rapid quantitative information could be also achieved by the developed film in conjunction with DIC. The pink complexes obtained from testing the film with the chlorine standard solution at different concentration were photographed and analyzed for RGB intensities. The relationship between chlorine concentrations and RGB values was plotted as the calibration curve (as shown in Fig. 9(a)) and the linear equation was then determined as a key element for quantitative analysis. In constrast with the green (I_G) and blue intensities (I_B), the red intensity (I_R) was hardly changed with the chlorine concentration. It was lower than I_G and I_B at low concentration. However, I_R became the highest one, when other two intensities was decreased, especially at higher concentration than 7 mg/L. This was due to the darkening of pink complex formed, as noticed by naked eye (Fig. 10), at initial concentration level of 7 mg/L.
Fig. 9. Relationships between free chlorine concentrations and (a) RGB intensities and (b) calculated RGB absorbances.

In the standard method based on spectrophotometry, the absorbance of the colorimetric product was a key parameter playing an important role in its measurement. For DIC technique, the absorbance could be estimated from each color band intensity by using Eq. (1) [21, 39-40]:

\[ A_X = -\log \left( \frac{(I_X - I_{X,B})}{(I_{X,W} - I_{X,B})} \right) = -\log \left( \frac{(I_X)c}{(I_{X,W})c} \right) \]

where for each X color (R, G, B), \( A_X \) was the absorbance of X, \( I_X \) was the intensity of X, \( I_{X,B} \) was the intensity of black color or 0, \( I_{X,W} \) was the intensity of white color or 255, and C was the concentration of X.

According to Fig. 9(b), an inverse relationship was obtained by calculated absorbance profile and the green channel (500-580 nm) dominated in absorbance over the blue (430-470 nm) and red channels (660 – 760 nm). These results indicated that the pink complexes could remarkably absorb the visible light in the green and blue ranges, while they reflected red light. This was in good agreement with spectrophotometric results that the pink complex provided the maximum absorbance at 515 nm [37], standing in the wavelength of green region.

Both intensities and calculated absorbances exhibited a linear relationship with the chlorine concentration which offered a convenient means of quantifying chlorine. As summarized in Table 2, the linear portions for \( I_G, I_B, A_G \) and \( A_B \) were in the same range of 0.3 – 15 mg/L chlorine with good linearity (R\(^2\) > 0.99). However, the sensitivity provided by the intensity value seems better than those by absorbance. The \( I_G \) possessed the highest sensitivity of 4,5449 a.u. L/mg, while the lowest one of 0.0085 a.u. L/mg was fed by \( A_B \). In addition to the greatest linearity (R\(^2\) = 0.9984), the \( I_G \) was obtained directly from the analysis of a digital image, unlike the molecular absorption calculated from the intensity. Thus, the \( I_G \) was recommended for quantification of chlorine.
Table 2. Calibration equation and linear range for chlorine detection by using the colorimetric film and DIC.

| Parameter | Calibration equation \( y = a \cdot x + b \) | Linear range (mg/L) | Linearity \( (R^2) \) |
|-----------|----------------------------------|----------------------|------------------|
| \( I_C \) | \( y = 0.39x + 0.15 \)          | 0.0 - 0.2            | 0.9952           |
| \( I_B \) | \( y = 0.20x + 0.10 \)          | 0.0 - 0.2            | 0.9952           |
| \( A_C \) | \( y = 0.39x + 0.15 \)          | 0.0 - 0.2            | 0.9952           |
| \( A_B \) | \( y = 0.20x + 0.10 \)          | 0.0 - 0.2            | 0.9952           |

3.5. Analysis of Real Water Sample

Four water samples were randomly collected from each water source possibly contaminated by chlorine, i.e., the swimming pool, salt water pool, tap water and sea water at Patong beach, Phuket. The average concentrations were analyzed by using the biopolymer film in conjunction with DIC, and spectrophotometry for comparison. The results were shown in Table 3.

Except sea water, the free chlorine was found in all samples. In the traditional swimming pool, the chlorine concentration quantified by the proposed method was 1.73 mg/L, comparable with 1.87 mg/L from the standard spectrophotometric technique. The level of chlorine in the conventional pool was higher than the modern salt pool, where they were found at 0.56 and 0.62 for the test kit and spectrophotometric detection, respectively. This was because the free chlorine was added directly to the water in the traditional pool, while hypochlorous acid (free chlorine) was turned from the sodium chloride via electrolysis process in the salt pool. The chlorine concentrations found in both pool samples did not exceed the level of 3.5 mg/L, suggested by World Health Organization (WHO) for safe swimming pool from the harmful disinfection by-product [41]. For the tap water sample, the chlorine concentration was detected at 1.34 mg/L by the developed test kit, which was in a good agreement with 1.39 mg/L chlorine from spectrophotometric method (only 3.60% relative error). Noted that the amount of chlorine in the tap water sample was also under the concentration degree of 1.0 – 1.5 mg/L controlled by Thai Provincial Waterworks Authority. Statistically, all results from the proposed method were not significantly different with the standard spectrophotometric technique at 95% confidence level (calculated \( t \)-value=0.94, and critical \( t \)-value=2.45). This indicated the excellent performance of the biopolymer hybrid film combined with DIC for the analysis of chlorine in water samples.

Table 3. Concentration of free chlorine in various water sample.

| Sample          | Developed test kit (mg/L) | Spectrophotometry (mg/L) | %Relative Error |
|-----------------|---------------------------|-------------------------|----------------|
| Swimming pool   | 1.73                      | 1.87                    | 7.49           |
| Salt water pool | 0.56                      | 0.62                    | 9.68           |
| Tap water       | 1.34                      | 1.39                    | 3.60           |
| Sea water       | n.d.*                     | n.d.                    | -              |

* n.d. = not detected

4. Conclusion

A novel film of hybrid biopolymer was successfully developed from the crosslinkage of two natural products, i.e., the tapioca starch and agar. It was employed to entrap N,N-diethyl-p-phenylenediamine (DPD) colorimetric indicator for free chlorine analysis. The analytical film was in-situ coated on the lid of plastic micro-PCR tube so that 1 mL water sample could directly added into the tube, acting as the portable color reactor. The hybrid film would be well prepared from the mixture of 3 solutions, i.e., 0.5 mL phosphate buffer containing 16 g/L EDTA, 1 mL of 0.2 g/mL DPD reagent and 3.5 mL of 2:1 (v/v) 2.5 g/L agar : 5 g/L starch solution. One hundred microliter of those mixture dropped on the plastic lid was polymerized at 60°C for 60 min to obtain the best performance film. The test kit showed very fast response of 1 min for the development of pink product color, after 1 mL of chlorine sample added into the tube. When the developed film sensor was used in combination with digital image colorimetry (DIC), it provided an ideal novel platform for the rapid quantitative analysis of free chlorine. The good linear response with wide range of 0.3-15 mg/L at greatest linearity \( (R^2 = 0.9984) \) was achieved. This indicated the possibility of application for the portable, rapid, cost-effective and environmentally friendly testkit.
Acknowledgements

Authors would like to thank for the funding and facility supporting by Faculty of Technology and Environment, Prince of Songkla University, Phuket Campus. It was also appreciate to the Office of Higher Education Commission (OHEC), the S&T Postgraduate Education and Research Development Office (PERDO) and Center of Excellence on Hazardous Substance Management (HSM), Chulalongkorn University for their invaluable supports in terms of financial seeding.

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