Butterfly Pea (Clitoria Ternatea L.) Extract as Indicator of Acid-Base Titration

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ABSTRACT
An anthocyanin color pigment result from butterfly pea flower maceration has been done. This study aims to determine the effect of time variation in the maceration of anthocyanin butterfly pea flower extract and the stability of butterfly pea flower extract as an indicator of acid-base titration. Color pigment levels were determined quantitatively with a UV-Visible spectrophotometer. The anthocyanin levels obtained during the duration of maceration 5 days in a row resulted from 4.3915%, 5.9869%, 7.3970%, 8.8995%, and 10.2864%. While the anthocyanin levels using a differential pH method were 14.2775 mg/L and 14.9455 mg/L with % RPD value was 4.57%. The results of the study stated that the indicators of the butterfly pea flower crown extract can be a substitute for the synthetic phenolphthalein indicator that has been used.

1. INTRODUCTION
Anthocyanin color pigments from various plants are widely used in the food and medicine industry because they can make their colors attractive and safe for health. Anthocyanin color is strongly influenced by the structure of anthocyanin and the degree of acidity (pH). Besides, anthocyanins tend to dissolve in polar solvents due to the presence of aromatic groups and glycosyl residues [1]. Anthocyanins are a type of secondary metabolite of the flavonoid family found in fruits and vegetables in large quantities. Anthocyanins are broadly divided into plant polyphenols in which flavonols, flavan-3-ol, flavones, flavanones, and flavonols are additional classes of flavonoids which differ in the oxidation of anthocyanins. Anthocyanins tend to be colorless in neutral pH areas, in solutions where the pH is very acidic (pH < 3) gives the maximum red color, whereas in an alkaline solution (pH 10.5) the color pigment of the anthocyanin agent will change color to green if in an alkaline pH atmosphere [2].

One type of plant that contains anthocyanin is the butterfly pea flower [3]. Butterfly pea flower is a plant that has the Latin name Clitoria ternatea L and Butterfly Pea in English, with anthocyanin as one of the active compounds giving a bluish-purple color to the flower crown that can function as an antibacterial as outlined in Table 1. This flower according to its name comes from Ternate region, Maluku which can grow in tropical regions such as Asia and its spread reaches South America, Africa, Brazil, North Pacific, and North America [1].
Anthocyanin compounds give purple, blue and red colors to butterfly pea flowers where the anthocyanin phytochemical content has good stability so that it can be applied to natural dyes for foodstuffs and chemical testing indicators. To obtain anthocyanin compounds, extraction methods are needed, namely the process of separating components using certain solvents. There are 2 methods of anthocyanin extraction in testing namely cold and hot. Ways of extraction without heating include maceration and percolation, while methods that require heat include reflux, digestion, infusion, and soxhletation. Factors that can affect the extraction rate are the type of sample preparation, extraction time, a quantity of solvent, temperature of the solvent, and type of solvent.

The best solvent used for anthocyanin extraction in ethanol solvent [5]. The extraction process used is simpler maceration and is one of the most widely used methods and can avoid the destruction of thermolabile compounds found in butterfly pea flowers [1]. This research wants to study the use of solvents and effectiveness for the extraction of butterfly pea flowers (Clitoria ternatea L) and their application as an alternative indicator of acid-base. The structure of anthocyanin can be seen in Scheme 1.

Organic compounds that can be used as indicators in acid-base titration have the characteristics of compounds that can provide color changes to the atmosphere of a pH solution. Color changes can occur due to the process of balancing the molecular and ionic shapes of an indicator compound. For example, phenolphthalein compounds are indicators of strong-weak acids and bases, undergoing a change in ion equilibrium followed by a change in color from colorless under acidic conditions to red under alkaline conditions. From the equilibrium reaction, it is known that the indicator compound is in the form of ions which can produce red color changes [6]. Color changes due to what happens because the compounds in phenol in the form of ions undergo delocalization to form quinoids. This
research will study the process of anthocyanin extraction using the maceration method with ethanol solvent for five (5) days and the quality of the extract is tested as an alternative indicator of acid-base titration. Similar studies have been carried out for the titration indicator of standard NaOH solutions and standard solutions of CH$_3$COOH using azo derivatives from eugenol which is 4-allyl-2-methoxy-6-hydroxiazobenzene [6].

2. METHOD

2.1 Tools and Materials
The tools used in this study include a set of glassware, analytical balance (Ohaus), pH meter (Ohaus), hotplate, magnetic stirrer, a set of support devices, single beam UV-Visible Thermo scientific genesis 20 spectrophotometers.

The materials used are butterfly pea flowers, methanol (Merck), concentrated HCl, ethanol solution 96%, HCl solution 1%, NaOH solution 10%, NaOH solution 0.1N, billberry standard, KCl solution 0.0125M, solution CH$_3$COONa.3H$_2$O 0.2M, phenolphthalein indicator, anhydrous oxalic acid, distilled water, filter paper, and aluminum foil.

2.2 Making a standard solution of bilberry
The standard solid of bilberry 20.20 mg was dissolved with 60 mL of ethanol and then put into a 100 mL volumetric flask, ethanol was added to approach the etching boundary mark then shaken to homogenize the solution.

2.3 Sample Preparation
The cleaned butterfly pea flowers were put into a mortar. The fine butterfly pea flowers were weighed 25 g using a beaker glass, added a solvent in the ratio of 1: 4 100 mL 96% ethanol and 100 mL distilled water then stirred using a magnetic stirrer at a temperature of 30 °C. Samples were allowed to stand for 24 hours. Every day 20 mL extract was taken and then filtered to get the filtrate, the same thing was delivered for 5 days.

2.4 Qualitative identification test
The filtrate from the butterfly pea flower was taken for 7 mL, 2-3 drops of 10% NaOH were added, the color changes to green, then 2-3 concentrated HCl was added until the color turns red.

2.5 Determination of anthocyanin levels in butterfly pea flower
A 12 mL extract solution was added with methanol containing 1% HCl to a 100 mL volumetric flask. The absorbance of anthocyanin extract was measured at a maximum wavelength of 528 nm then the absorbance was recorded at that wavelength range. Anthocyanin levels were determined using equation 1.

\[
\text{Anthocyanin level} = \frac{\text{Sample absorbance}}{\text{Standard absorbance}} \times \text{concentration of billberry standard} \times \text{DF} \times \text{Extract volume} \times 0.418
\]  

For testing of anthocyanin level using a differential pH, for 1 mL of butterfly pea flower extract, added 9 mL of KCl solution pH 1, put into 10 mL volumetric flask and then homogenized. The same was done for the CH$_3$COONa.3H$_2$O solution pH of 4.5. The absorbance of anthocyanin extract was measured at a maximum wavelength of 500 nm and 700 nm then recorded at that wavelength range. The test was carried out two replicates where the anthocyanin level using the differential pH method is determined using equation 2.

\[
A = (A_{520} - A_{700})pH_1 - (A_{520} - A_{700})pH_{4.5}
\]

\[
\text{Anthocyanin level} = \frac{A \times \text{MW} \times \text{DF} \times 1000}{\text{Ex} \times \text{I}}
\]

Keterangan: A for absorbance
MW for the molecular weight of cyaniding-3-glucoside (449,2 g/mol)
DF for the dilution factor
ε for molar absorptivity of cyaniding-3-glucoside (26900 L/mol)
I for cuvette wide (cm)
3. RESULT

3.1 Results of identification of color pigments

This research was carried out with a maceration time of 5 days with a solvent ratio of 1: 4 and a flower weight of 25 g flower at room temperature with a wavelength of 528 nm. The results of tests in the form of qualitative identification are presented in Table 2.

| Maceration time | Basic color | After adding NaOH 10% | After adding HCl glacial |
|-----------------|-------------|-----------------------|-------------------------|
| Day 1           | Purple      | Green                 | Red                     |
| Day 2           | Purple      | Green                 | Red                     |
| Day 3           | Purple      | Green                 | Red                     |
| Day 4           | Purple      | Green                 | Red                     |
| Day 5           | Purple      | Green                 | Red                     |

To identify anthocyanin, the extract obtained from maceration is filtered and then put into a test tube as much as 1 mL. The color of the anthocyanin compound is purple with a pH of 5, then the extract is dropped with 10% NaOH by 2 drops, the anthocyanin compound changes to green at pH 10, after that the extract is dropped later with concentrated HCl 2 drops then the anthocyanin compound turns red with a pH of 3.5. The butterfly pea flower has a blue base color where the blue color appears due to the color degradation process of anthocyanin which is in the form of a red cavity flavilium to a blue quinoidal base. In a liquid medium, anthocyanin undergoes structural changes because the instability of anthocyanin is influenced by pH. Anthocyanin compounds in very acidic conditions (pH <2) is dominated by flavilium cations that are red, whereas in conditions of weak acidity, neutral, or alkaline the carbinol (colorless) and quinoidal base (blue) dominate the cavity flavilium so that the color fades (colorless) and changes from red to blue. Increasing pH value makes carbinol and chalcone base compounds more and more formed which causes colorless [7]. Changes in color at different pH values indicate that flower extracts of butterfly pea have the potential to be used as an alternative indicator of acid-base titration.

The event of anthocyanin color changes at different pHs due to the intermolecular copigmentation process between anthocyanin color pigments and copigment compounds characterized by a shift between bathochromic and hyperchromic. Bathochromic shift (redshift or bathochromic effect) is a shift at the peak of absorption in the direction of a larger wavelength. This happened because of the substitution of the glycone or aglycone group or the effect of the solvent. Hyperchromic effects are effects that can be caused by functional groups, causing an increase in the value of maximum absorption intensity. The addition of hydroxyl groups results in a shift toward green (pelargonidin → cyanidin → delphinidin), where glycoside formation and methylation produce a shift toward red (pelargonidin → pelargonidin-3-glucoside; cyanidin → peonidin) [7].
3.2 Result of Anthocyanin Concentration

The study of anthocyanin levels was carried out with maceration time variables, namely 1, 2, 3, 4 and 5 days and the weight of 25 g flowers at 30 °C and a wavelength of 528 nm. The results of tests in the form of anthocyanin levels are presented in Table 3.

TABLE III. Results of Anthocyanin Levels

| No | Maceration time | Dilution factor | Anthocyanin Concentration (mg/L) |
|----|-----------------|-----------------|----------------------------------|
| 1  | Day 1           | 5 kali          | 4,391                            |
| 2  | Day 2           | 5 kali          | 5,987                            |
| 3  | Day 3           | 5 kali          | 7,397                            |
| 4  | Day 4           | 5 kali          | 8,899                            |
| 5  | Day 5           | 5 kali          | 10,286                           |

The anthocyanin extract obtained from maceration was then diluted again with 1% HCl in methanol as much as 500 mL, after which it was inserted into a UV-Vis spectrophotometer. The first blank that added 1% HCl solvent in methanol obtained 0,000 absorbances. The next blank is a bilberry extract that has known concentration. The absorbance yield of bilberry extract was 0.526. The diluted sample was entered into a UV-Vis spectrophotometer for the results compared to a standard comparison of bilberry extract. Anthocyanin levels were calculated using equation 1, while the relationship between anthocyanin levels and maceration time is shown in Figure 3.
The results of the stem diagram of the sample levels of the butterfly pea flower extract show that the color pigment levels obtained are increased where \( y \) is the color pigment level (mg/L) and \( x \) is the maceration time (day). From the results of the study note that the longer the maceration time used, the greater the level of color pigment obtained is also proportional to the concentration of the anthocyanin compound. The higher concentration of anthocyanin pigments causes increased chroma, which can be heard by higher extract levels [9]. In addition to the duration of time, the type of solvent extraction process affects the levels of color pigments because anthocyanins dissolve in polar solvents such as ethanol which are easily absorbed by cell membranes so that membrane breakdown occurs on the surface of tissue particles in flower petals [5].

In this study, the extraction process depends entirely on the ethanol solvent and the duration of time alone because there is no addition of acid solutions and the use of high temperatures (heating). As is known, acidic solution and heating will further optimize the yield of anthocyanin but the potential for damage (degradation) of anthocyanin compounds also arises due to heating through the hydrolysis of glycosidic bonds to produce aglycones and the opening of the aglycone ring to form carcinol groups and colorless chalcones [10].

![Figure 4. Mechanism of anthocyanin degradation at temperatures over 70 ° C](image)

### 3.3 Total Concentration of Anthocyanin using pH Differential

Color pigments that have been dissolved in pH 1 and pH 4.5 are measured for their absorbance with wavelengths of 520 nm and 700 nm. The wavelength of 520 nm is the maximum wavelength to determine cyaniding-3-glucoside while the wavelength of 700 nm is to correct deposits or impurities that are still present in the sample. It is known that anthocyanin compounds have 4-5 different structures in aqueous solutions with different pH as outlined in Table 4.

| pH value | Molecular Form                  |
|----------|---------------------------------|
| 1-3      | Red flavylum cation             |
| 4-5      | Colorless carbinol pseudo base  |
| 6-7      | Purple quinonoidal base         |
| 7-8      | Blue anionic quinonoidal base    |
| 8-9      | Pale yellow chalcone            |

For anthocyanin compound extracts of butterfly pea flower which were dissolved at pH 1 and pH 4.5 showed a typical spectrum pattern shown in Figure 5 as reported by [11] who conducted a study of color stability and spectrum characteristics. At pH 4, the butterfly pea flower extract shows two wave peaks (\( \lambda_{\text{peak}} \)) representing quinonoidal base and anionic quinonoidal base while one supporting wave peak (\( \lambda_{\text{shoulder}} \)) represents the species flavylum cation.
If the sample is really clear then the absorbance at 700 nm is 0 [12]. The absorbance of anthocyanin that has been dissolved at pH 1 and 4.5 can be seen in Tables 5 and 6.

**TABLE V. The absorbance results using a differential pH of 1.0**

| Wavelength (nm) | Absorbance | Average |
|-----------------|------------|---------|
| 520             | 0.095      | 0.0945  | 0.095   |
|                 | 0.095      | 0.095   | 0.094   |
| 700             | 0.005      | 0.006   | 0.005   |
|                 | 0.006      | 0.005   | 0.005   |

**TABLE VI. The absorbance results using a differential pH of 4.5**

| Wavelength (nm) | Absorbance | Average |
|-----------------|------------|---------|
| 520             | 0.08       | 0.08    | 0.08    |
|                 | 0.079      | 0.079   | 0.079   |
| 700             | 0.008      | 0.008   | 0.008   |
|                 | 0.008      | 0.008   | 0.008   |

The results of the study can be calculated using equation 2. Total anthocyanin level in the first butterfly pea flower is 14.2775 mg/L, while the second anthocyanin level is 14.9455 mg/L with a % RPD value of 4.57%, the value of % RPD produced is good because it is less than the specified threshold of > 5%. The results show that the higher the absorbance obtained, the more anthocyanin levels. Determination of anthocyanin concentration by the pH differential method because at pH 1.0 anthocyanin forms a colored oxonium (cation flavilium) compound and at pH 4.5 it forms a colorless carbinol/hemiketal [13].

Measurement of total anthocyanin levels using this method is a calculation through visible light at different pHs. Research on anthocyanin content which is mostly found in plants is cyaniding-3-
glucoside with molar absorptivity ($\epsilon$) of 26,900. Generally, cyanidin-3-glucoside is used as a reference compound of anthocyanin [14].

The indicator results of the butterfly pea flower extract show a change in color i.e. in a red acid solution and a green base [15]. Anthocyanin in its structure contains cation flavilium, color changes can occur due to changes in the shape of the structure caused by the influence of pH. The results of the analysis of butterfly pea flower extracts are the color of the solution in pH is bluish-green where the shape changes in the color pigment structure due to the influence of the pH value. The anthocyanin structure in Scheme 1, under red acidic conditions, if the pH is increased (pH < 4) a colorless carbinol base (3) is formed and further as shown in Figure 6.

![Figure 6](image)

Figure 6. The equilibrium form of the flavilium cation in anthocyanins in various pH ranges ([16][17])

**3.4 Control Chart of Butterfly Pea Flower Extract as An Indicator of Titration**

Butterfly Pea flower extract obtained from maceration for 5 days will be used as an indicator in the simulation of acid-base titration. In this test, the performance indicator of butterfly pea flower extract will be assessed for its performance with the phenolphthalein indicator as a dick or reference. The results of titration testing are shown in Table 7.

| TABLE VI. The titration results using the phenolphthalein indicator and the butterfly pea flower extract indicator |
|---------------------------------------------------------------|
| **Day** | **Phenolphthalein (mL)** | **Butterfly Pea (mL)** |
| 1 | 8,47 | 8,60 |
| 2 | 8,70 | 9,80 |
| 3 | 8,85 | 8,70 |
| 4 | 8,60 | 9,70 |
| 5 | 8,85 | 8,85 |
| 6 | 8,53 | 9,85 |

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The volume of NaOH produced from acid-base titration using the phenolphthalein indicator and the butterfly pea flower extract indicator has an average difference of 0.59 mL so that the butterfly pea flower extract indicator can be used as an alternative indicator of phenolphthalein. To find out the stability of butterfly pea flower extract as an indicator, a control chart is needed as shown in Figures 7 and 8. Acid-base titration was used in this study because anthocyanin compounds in the flower had a change in color at a certain pH value which also corresponds to the pH range of the phenolphthalein indicator.

The control chart of phenolphthalein indicator control

![Figure 7. The control chart of phenolphthalein indicator control](image)

Figure 7. The control chart of phenolphthalein indicator control

The phenolphthalein indicator is an acid-base titration indicator which has a pH range of 8.0-9.6 used as a comparison indicator. The graph shows that for six (6) days, the phenolphthalein indicator and the banana flower extract are still between the Upper Control Limit (UCL) and Lower Control Limit (LCL), meaning that both indicators are at the control limit or stable test results. These results indicate that the phenolphthalein indicator and butterfly pea flower extract can be used for 6 days even though there is a difference in the volume of NaOH needed at titration but is not significant. The volume of NaOH produced by the butterfly pea flower extract indicator is greater than the phenolphthalein indicator due to differences in pH values. The control chart itself is used to see the consistency of the analyst, the stability of the equipment, the difficulty level of the method used, and to know the durability of the butterfly pea flower extract as a test sample.

4. CONCLUSION

Color pigments can be obtained by extracting butterfly pea flowers using a maceration method
that can produce concentrated purple extracts. Anthocyanin from butterfly pea flowers can be used as an indicator of the acid-base because the color pigment changes its color when it drops acid or base and natural dyes. When the maceration was obtained anthocyanin levels for 5 consecutive days namely 4.3915%, 5.9869%, 7.3970%, 8.8995%, and 10.2864%.

The total concentration of color pigments from butterfly pea flowers can be determined by the pH differential method by spectrophotometry. The results showed a total concentration of 14.2775 mg/L and 14.9455 mg/L with a% RPD value of 4.57%. Color changes occur in butterfly pea flower extracts due to the presence of anthocyanin compounds, which in their structure contain cations of flavilium to form anhydrobase due to changes in pH. Butterfly pea flower extract indicator has similarities with phenolphthalein, so it can be a substitute for acid-base titration indicators.

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