Extraction of Green Acid-Base Indicators From *Acanthus Pubescens*

Flower for Use in the School Laboratory

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Abstract

The use of *Acanthus pubescens* flower extracts as natural eco-friendly substitutes for synthetic acid-base indicators is experimentally confirmed. In this study, it is indicated that ethanolic and acidified ethanol extracts of *Acanthus pubescens* flower are good replacements to phenolphthalein, bromothymol blue and methyl red in acid-base titrations involving a strong acid versus strong base and a strong acid versus weak base. They can also be used as effective substitutes to phenolphthalein in a weak acid versus strong base titrations. And also the acidified ethanol extracts can be used in place of phenolphthalein and bromothymol blue while the ethanol extracts can replace methyl red in weak acid versus weak base titrations. It is also concluded that the use of *Acanthus pubescens* flower extract as an indicator in all types of acid base titrations is beneficial because of its economy, eco-friendly nature, ease of preparation, availability, simplicity, non-carcinogenicity, precision and accuracy of results.

Keywords: *Acanthus pubescens* flower; acidified ethanol; natural indicator; anthocyanins, flavonoids

1. Introduction

Natural acid-base indicators are greener alternatives to synthetic ones as they are the demand of contemporary chemistry with regards to solving the problems of environmental pollution, cost and human health. There is an increasing need for the development of green indicators as effective substitutes for synthetic acid-base indicators used in the school chemistry laboratory.

A change in colour during acid-base titrations with variation in pH is due to presence of coloured pigments that are used in acid base titrations to show sharp end point (Mane et al., 2016). Commonly used indicators for such titrations are manmade and costly. Besides, some of them have toxic effects on users and can also cause environmental pollution. For these and other reasons, there have been increasing concerns in searching for greener sources of acid-base indicators. These greener and eco-friendly alternatives would be cheaper, more available, simple to extract, less toxic to users and environmentally viable (Abugri et al., 2012). Some highly coloured pigments obtained from plants are found to exhibit colour changes at various pH values (Mane et al., 2016). In the Present study, we prepared ethanolic and acidified ethanol *Acanthus pubescens* flower extract as a natural indicator for acid-base titration.

*Acanthus pubescens* belongs to the family of *Acanthaceae*. It is used in traditional medicine for treatment of syphilis and gonorrhea. A decoction of the leaves is used for the treatment of gastroenteritis, pneumonia and anthrax. It is also reported that, a preparation of the dried leaves is used externally as a remedy for scabies (Moshi et al., 2010). The leaves are burned to ash and licked to treat cold, cough, liver problem, chronic asthma, cancer, tonsils and flu (Seswa, 2016; Jeruto et al., 2008). The leaves of *Acanthus pubescens* contain various classes of compounds such as flavonoids, terpenes, alkaloids and saponins identified through phytochemical screening test, which might be responsible for the traditional use of the plant (Sikolia et al., 2017) where it is hypothesized that pH indicators can possibly be produced from *Acanthus pubescens* plants because of flavonoids which are pH sensitive.

Many experiments have been carried out on the use of plant pigment extracts as green substitutes to synthetic acid-base indicators. Flower extracts of *Jacaranda Acutifolia* were used as indicator in acid-base titrations (Patrakar et al., 2010) in which the equivalence point obtained by the flower extracts coincided with the equivalence point obtained by synthetic ones. In case of weak acid and weak base titration, the results obtained by the flower extract matched with the results obtained by mixed indicator. Gaurav et al (2010) studied the extracts of *Bougainvillea glabra* flowers changing colour in aqueous solutions of acids and bases. The researchers compared the results of the *Bougainvillea glabra*
flowers extracts with phenolphthalein and methyl orange in acid-base titrations as indicator. The results indicated that the extract obtained from flowers of Bougainvillea glabra could be used as indicators for strong acid vs. strong base titrations as similar results were obtained by phenolphthalein. On the other hand, methanolic extracts of the leaves of careya arborea were also used as indicators in acid base titrations (Wadkar et al., 2008). Promising results were obtained when a natural indicator prepared from methanolic extracts of the species were tested against standard synthetic indicators. The indicator was useful in all types of acid – base titrations except weak acid and weak base titration. Okonkwo et al., (2010) used Hibiscus Sabdariffa petals extract for acid – base titrations. The results were compared with those of standard end point indicators as phenolphthalein and methyl red. Prominent absorption peaks in the 500 – 550 nm wavelength region of the UV/Visible spectrum of methanolic extract confirmed the presence of anthocyanidins. In other studies, methanolic extracts of Antirrhinum majus belonging to the family of Scrophulariaceae as well as Dianthus plumaris which belong to the family of Caryophyllaceae were used which gave sharp and intense colour change as compared to phenolphthalein and methyl orange indicators (Jaspreet et al., 2011). In all these titrations the extract was found to be accurate and useful for indicating the end point of a neutralization reaction.

Colour changes of natural indicators at different pH values have been attributed to the presence of anthocyanins and flavonoids which are pH sensitive. Anthocyanins are organic compounds that are usually found in the aqueous sap of the vacuole of the epidermal plant cells. These compounds have a complex structure consisting of an aromatic three-ring molecular structure, one or more attached sugar molecules and sometimes acyl groups attached to the sugar molecules. Anthocyanins are water soluble and are usually more stable in acidic media than in alkaline solutions. A general structure of an anthocyanin is shown in Figure 1.

Almost any plants that have blue, violet, purple or red flowered colours contain organic pigments, anthocyanins that changes colour with change in pH. The colour stability of anthocyanins depends on structure of the anthocyanins, pH, temperature, oxygen, and light and water activity. They tend to be red in a more acidic solution and blue in basic solution (Nhapi, 2016).

Anthocyanins have several physiological activities which include antioxidant, anti-hepatocarcinogenic, anti-inflammatory, anti-tumour, hypolidemic, cardioprotective and cancer chemopreventive, hence they are safe to use in acid-base titration.

Figure 1. General structure of anthocyanin (Nhapi, 2016)

1.1 pH Dependant Structural Transformations of Anthocyanins

Due the high reversibility of the various forms of cyanidins in aqueous solutions, different colours are observed with respect to changes in pH. At acidic pH (1-3), as shown in Figure 2, anthocyanins exist predominantly in the form of the red or orange flavylum cation (2-phenylchromenylium cation). The colour intensity decrease as pH increases and also the concentration of the flavylum cation decreases which undergoes hydration to produce the colourless pseudo base (hemiacetal or chromenol). This is due to kinetic and thermodynamic competition between hydration reaction of the flavylum cation and proton transfer reactions related to the acidic hydroxyl groups of the aglycone. The conjugated 2-benzopyrilium system is disrupted due to a nucleophilic attack of water at position 2 of the anthocyanidin skeleton. Flavylum cation lose proton as the pH shifts higher. The equilibrium will now shift towards a purple quinoidal anyhydrobase at pH < 7 and a deep blue ionised anhydrobase at pH < 8. As the pH increases further, the carbinol form yields through opening of the central pyran ring and the light yellow chalcone will result. The anthocyanidin system undergoes a variety of molecular transformations as pH changes, thus in aqueous solutions, anthocyanidins exist as five molecular species in chemical equilibrium which are red flavylum cation, colourless carbinol pseudo base, purple quinoidal base, blue quinoidal base anion and yellowish chalcone. These transformations are shown by the mechanism
shown on Figure 2. Plant species containing anthocyanins can change colour in solution by undergoing these transformations due to change in the acidity or basicity of the solution (Nhapi, 2016).

Figure 2. Structure of Cyanidins in aqueous solution under varying pH (Nhapi, 2016)

1.2 Research Questions

In this study, we successfully prepared a natural, eco-friendly, cheaper and locally available acid-base indicator from Acanthus pubescens flower extracts. We have specifically given answers the following questions:

1. What are the characteristics of the active components that give the acid-base indicator properties of Acanthus pubescens flower extracts?
2. What are the resulting colours of Acanthus pubescens flower extracts in the pH range of 1 to 14?
3. Are there any significant differences between the results of the extracted pH indicator with phenolphthalein, bromothymol blue and methyl red pH indicators in determining the end point in different types of acid-base titration?
2. Materials and Methods

2.1 Reagents, Apparatus and Equipments

The principal raw material used for the study was the powdered *Acanthus pubescens* flower. All reagents used in the practical work were of analytical grade and used without further purification. These include hydrochloric acid (HCl, 35-36\%, UNI-CHEM Chemical Reagents, India), sodium hydroxide (NaOH, 98\%, BLULUXR Analytical Reagents), glacial acetic acid (CH₃COOH, 99.5\%, SCR-China), ammonia (NH₃, 30\%, Merck, Germany), borax (Na₂B₄O₇.10H₂O, 99.0 – 103\%, SAMIR TECH-CHEM PVT, LTD), iron (III) chloride (FeCl₃, 99.0\%, Avi Chem Industries, India), magnesium ribbon (Mg, BUCK SCIENTIFIC PURO-GRAPHIC), ethanol (CH₃CH₂OH, 97\%, Fine Chemical General Trading, Ethiopia) and distilled water. The commercial indicators were phenolphthalein (C₂₀H₁₄O₄, SAMIR TECH-CHEM PVT, LTD), methyl red (C₁₅H₁₅N₃O₂, SAMIR TECH-CHEM PVT, LTD), bromothymol blue (C₂₇H₂₈Br₂O₅S, SAMIR TECH-CHEM PVT, LTD). The instruments that were used in the study are pH meter (Elmetron CPI-501, Poland), double beam UV-Vis spectrophotometer (6705, Jenway), electronic analytical balance (AA200DS, Deriver Instrument Company, Germany) and magnetic stirrer (MS300, Germany). The following apparatus and equipments such as Erlenmeyer flasks, volumetric flasks, burette with volume size of 50 mL, micropipette (1-10 µL, 10-100 µL and 100-1000 µL), test tubes, beakers, graduated measuring cylinders with volume size of 10 mL and 25 mL, and Whatman filter paper No. 41 were used to carried out the experiment. All required reagents and volumetric solutions were prepared as per standard.

2.2 Collection and Preparation of the *Acanthus pubescens* Flower

One kilogram of fresh *Acanthus pubescens* flower were collected from around Debre Markos, Ethiopia, at the beginning of December 2019, as it is the blooming season of the plant. The fresh flower petals of *Acanthus pubescens* were separated from the whole flower by hand and washed with distilled water to remove dirt. The fresh *Acanthus pubescens* flower petals were air dried for three weeks without exposure to direct sunlight to minimize oxidative loss before pounding in to fine powder at room temperature. The dried flower petals were ground into fine powder, as shown in Figure 3 below, with coffee grinding machine and the powders were sieved and stored in a polyethylene bag before use.

![Figure 3. *Acanthus pubescens* flower (A), dried and powdered samples (B & C)](image_url)

2.3 Extraction of the Indicator

30 grams of dried powder of *Acanthus pubescens* flower was mixed with 150 mL of 97 % ethanol in 250 milliliter Erlenmeyer flask and then the mixture was stirred by using a magnetic stirrer for 2 hours to disperse the powder completely (Figure 4). The mixture was kept at room temperature for 24 hours and then triturated in mortal and pestle and the resulting solution were filtered by using Whatman filter paper to remove the remaining plant matter. Similar procedures were applied by taking the same amount of powder and solvent for all the rest extracting solutions. The resulting 97% ethanol and 0.1% HCl in ethanol extract were further used as natural pH indicator for acidimetry and alkalimetry. The extract was preserved in light closed container and stored away from direct sunlight to prevent photolysis and decomposition (Nhapi, 2016; Gupta et al., 2013).
2.4 UV-Vis Spectroscopy Analysis

The UV-Vis absorption spectra of ethanol and acidified ethanol extracts of *Acanthus pubescens* flower were determined using UV-Vis spectrophotometer (6705, Jenway) in the wavelength range of 200 nm to 800 nm. The extracted solutions were diluted by a factor of 10 with the same solvent and 5 mL of the extract was measured and placed in the quartz cuvette. The wavelength of maximum absorption of each extract was determined and the compounds present in the extracts were interpreted.

2.5 Phytochemical Screening of the Extracts

The extracts were phytochemically screened in order to determine the presence/absence of flavonoids and anthocyanins in the flowers of the *Acanthus pubescens* plant extracts. Alkaline reagent test, Shinoda’s test, hydrochloric acid test, zinc test (Loganathan et al., 2017) and ferric chloride test (Shah, 2016) were used for the presence of flavonoids. On the other hand, the presence of anthocyanins was tested by adding 2 mL of the plant extract in 2 mL of 2 M HCl. The appearance of a pink-red colour that turns purplish blue after addition of ammonia indicates the presence of anthocyanins (Obouayeba et al., 2015).

2.6 Titration Using *Acanthus Pubescens* Flower Extract, Methyl Red, Bromothymol Blue and Phenolphthalein Indicators

Obtaining equivalence point in different types of titration of the *Acanthus pubescens* flower extracts in comparison with synthetic indicators was tested. Four titrations were performed i.e. strong acid versus strong base, strong acid versus weak base, weak acid versus strong base and weak acid versus weak base. The titrations were conducted in the order HCl and NaOH; HCl and NH₄OH; CH₃COOH and NaOH and CH₃COOH and NH₄OH. A volume of 10 mL of 1 M HCl was placed in an Erlenmeyer flask and four drops of *Acanthus pubescens* flower extract indicator were added while 1 M NaOH was placed in a burette. The titrant (NaOH) was added to titrate (HCl) until a colour change was observed. Titrations were conducted in three replicate analyses. The procedure was repeated for all titrations i.e. HCl versus NH₄OH, CH₃COOH versus NaOH and CH₃COOH versus NH₄OH.

Equimolar titrations were performed using 10 mL of titrant with four drops of indicator. A set of three experiments each for all the types of acid base titrations was carried out by using 1 M solution of acid and base. The mean and standard deviation for each type of acid base titrations were calculated and recorded as mean ± standard deviation.

3. Results and Discussion

3.1 UV-Vis Spectroscopy Analysis of the *Acanthus Pubescens* Flower Extracts

UV-Vis spectrophotometric analysis was performed on ethanolic and acidified ethanol extracts of *Acanthus pubescens* flower. The wavelength regions of the spectral absorbance peaks are used to identify the the active components present in the plant species. The UV-Vis absorption spectra for the dye solution extracted using the ethanolic and acidified ethanol are shown in Figures 5 below. As shown in the figures, the extracted solutions have shown absorption peaks at 224 nm, 267 nm, 331 nm and 524 nm in ethanolic extract. These absorption peaks of the extracted dye are closely related to flavonoids and anthocyanin as reported by other investigators (Obaseki et al., 2017; Nuryanti et al., 2013; Wuletaw et al., 2016). The UV-Vis absorption spectra of flavonoids consist of two distinctive bands in a broad range of 240 – 550 nm. Band I, covering the range 300 –550 nm, while Band II, covering the range of 240 –285 nm (Obaseki et al., 2017).
Anthocyanin can be readily distinguished from other flavonoid classes by performing colour test and UV-Vis analysis ($\lambda_{\text{max}}$). Anthocyanins generally have two absorption maxima, one in the UV-region of spectrum 260 – 280 nm (band II) and the second in the visible region of spectrum 490 – 550nm (band I) (Ahmed et al., 2013). Almost all flavonoid classes give the same absorption at the region of band II, thus, anthocyanins can be distinguished from other classes by observing the absorption region wavelength of band I (Nuryanti et al., 2013).

Figures 5 (A) and 5 (B) show the UV-Vis absorption spectra of dye solutions of Acanthus pubescens flower extracted using ethanol and acidified ethanol. All the extracted solutions show a UV-Vis absorption peak in the UV-Vis region of the spectrum at 224 nm, 267 nm, 331 and 524 nm for ethanol extracts of Acanthus pubescens flower; and 224 nm, 275 nm, 331 nm and 526 nm for acidified ethanol extracts of Acanthus pubescens flower. As shown in Figures 5A and 5B below, these absorption regions are also the main characteristics of flavonoid and anthocyanin pigments (Obaseki et al., 2017; Nuryanti et al., 2013; Khoddami et al., 2013).

Figure 5. UV-Vis spectrum of ethanolic (A) and acidified ethanol (B) extract of Acanthus pubescens flower

3.2 Phytochemical Screening Test of the Acanthus Pubescens Flower Extracts

The preliminary phytochemical investigation and qualitative chemical tests of the ethanol and acidified ethanol extracts of Acanthus pubescens flower were performed using alkaline reagent test, hydrochloric acid test, ferric chloride test, zinc test, and Shinoda test which confirmed the presence of flavonoids and anthocyanin as reported in Table 1 below.

Table 1. Results of phytochemical screening of ethanol and acidified ethanol extracts of Acanthus pubescens flower

| Phytochemicals | Test               | Observation     | Ethanolic extract | Acidified ethanol extract |
|----------------|--------------------|-----------------|-------------------|--------------------------|
| Flavonoids     | Alkaline reagent test | Yellow            | +                 | +                        |
|                | Ferric chloride test | greenish-black  | +                 | +                        |
|                | Hydrochloric acid test | Red              | +                 | +                        |
|                | Shinoda test       | Red              | +                 | +                        |
|                | Zinc test          | red              | +                 | +                        |
| Anthocyanins   | Hydrochloric acid test | Purplish blue   | +                 | +                        |

+ indicates test is positive

Different studies conducted on phytochemical tests on leaves, root and flower extracts of various plant species gave similar results (Etagegnehu et al., 2016; Sikolia et al., 2017; Singh et al., 2011). The presence of pH sensitive flavonoids and anthocyanins, as confirmed by the chemical tests and UV-Vis spectra make Acanthus pubescens flower extracts ideal indicators in acid-base titrimetric analysis.

3.3 Colours of Acanthus Pubescens Flower Extracts in a Solution of pH 1 - 14

The colour change interval of Acanthus pubescens flower extracts were also determined by preparing solutions having pH in the range of 1 to 14 and adding four drops of Acanthus pubescens flower extracts to each test tube. It is shown that in the pH range of 1- 7, ethanol and acidified ethanol Acanthus pubescens imparts pink colour while in the pH
range of 8-11, it produces a colourless solution in all extracts, and at pH 12 it shows a greenish yellow colour. At pH 13 and 14, a greenish yellow colour were observed in all extracts to the solution as shown in Table 2 below.

Table 2. Colour change results after addition of ethanol and acidified ethanol Acanthus pubescens flower extract in a solutions of pH 1–14

| pH  | 1   | 2   | 3   | 4   | 5   | 6   | 7   | 8   | 9   | 10  | 11  | 12  | 13  | 14  |
|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|
| Colour | Pink | Pink | Pink | Pink | Pink | Pink | Colourless | Colourless | Colourless | Colourless | Greenish yellow | Greenish yellow | Greenish yellow |

The sensitivity of the extracts to different pH are attributed to the presence of flavonoids and anthocyanins. As described by Oswald’s ionic theory of indicators (Kolthoff, 1937; Said, 2017), the differences in colour changes brought by the Acanthus pubescens flower extracts when subjected to different pH may be due to protonation or deprotonation of the Acanthus pubescens flower extracts. The colour change might be due to intramolecular rearrangement as shown in Figure 2 that causes absorptions in the different region of the UV-Vis spectrum (Ekpo, 2007). The colour changes reported in literature for other plants extracts are from dark pink to yellow whilst in this study; it is from pink to greenish yellow for the Acanthus pubescens flower extract. This might be attributed to different plant extracts being used in the other studies (Senathirajah et al., 2017) and also the crude extract component matrix might be different between the plant extracts.

3.4 Acid-Base Titration Using Acanthus Pubescens Flower Extract, Phenolphthalein, Bromothymol Blue and Methyl Red as Indicators

In order to evaluate the potential for the use of Acanthus pubescens flower extract as indicators in acid-base titrimetry, a number of titrations using the ethanol and acidified ethanol extracts were conducted. The end points of the titrations using commercially available indicators are also reported in the table. The results in the table showed that the end points obtained with the ethanol and acidified ethanol extracts of Acanthus pubescens flower in 1M solutions of hydrochloric acid and sodium hydroxide (i.e. strong acid versus strong base) gave similar end points ranging from 9.73 ± 0.058 to 9.77 ± 0.115 mL which were close to the end points obtained using phenolphthalein (9.80 ± 0.100 mL), bromothymol blue (9.77 ± 0.115 mL) and methyl red (9.70 ± 0.100 mL) and so the Acanthus pubescens flower extract can be used in place of phenolphthalein, bromothymol blue and methyl red in acid-base titrations involving a strong acid versus strong base combination.
Table 3. Mean volume of base used (in mL) at end points and colour change for the four titrations using *Acanthus pubescens* flower extract, phenolphthalein, bromothymol blue and methyl red as indicators

| Titration          | Indicators                                    | Mean ± Sd*          | Colour change         |
|--------------------|-----------------------------------------------|---------------------|-----------------------|
| HCl vs NaOH        | Methyl red                                    | 9.70 ± 0.100        | Red to yellow         |
|                    | Bromothymol blue                              | 9.77 ± 0.115        | Yellow to blue        |
|                    | Phenolphthalein                               | 9.80 ± 0.100        | Colourless to pink    |
|                    | Ethanol *Acanthus pubescens* flower extract   | 9.73 ± 0.058        | Pink to colourless    |
|                    | Acidified ethanol *Acanthus pubescens* flower extract | 9.77 ± 0.115        | Pink to colourless    |
| CH₃COONaOH vs NaOH | Methyl red                                    | 11.20 ± 0.200       | Red to yellow         |
|                    | Bromothymol blue                              | 11.30 ± 0.100       | Yellow to blue        |
|                    | Phenolphthalein                               | 11.40 ± 0.173       | Colourless to pink    |
|                    | Ethanol *Acanthus pubescens* flower extract   | 11.33 ± 0.058       | Pink to colourless    |
|                    | Acidified ethanol *Acanthus pubescens* flower extract | 11.33 ± 0.058       | Pink to colourless    |
| CH₃COOH vs NH₄OH   | Methyl red                                    | 9.53 ± 0.058        | Red to yellow         |
|                    | Bromothymol blue                              | 9.67 ± 0.058        | Yellow to blue        |
|                    | Phenolphthalein                               | 10.40 ± 0.100       | Colourless to pink    |
|                    | Ethanol *Acanthus pubescens* flower extract   | 10.43 ± 0.058       | Pink to colourless    |
|                    | Acidified ethanol *Acanthus pubescens* flower extract | 10.40 ± 0.100       | Pink to colourless    |
| CH₃COOH vs NH₄OH   | Methyl red                                    | 8.07 ± 0.115        | Red to yellow         |
|                    | Bromothymol blue                              | 8.77 ± 0.208        | Yellow to blue        |
|                    | Phenolphthalein                               | 8.83 ± 0.058        | Colourless to pink    |
|                    | Ethanol *Acanthus pubescens* flower extract   | 8.10 ± 0.100        | Pink to colourless    |
|                    | Acidified ethanol *Acanthus pubescens* flower extract | 8.87 ± 0.115        | Pink to colourless    |

*Standard Deviation*

A comparison of the results of titration of 1M hydrochloric acid and 1M ammonium hydroxide solutions (i.e. strong acid versus weak base) using these plant extracts as indicators as presented in Table 3 indicated that the average titre using ethanolic and acidified *Acanthus pubescens* flower extracts were the same (11.33 ± 0.058 mL), while that of methyl red, bromothymol blue and phenolphthalein are 11.20 ± 0.200, 11.30 ± 0.100 and 11.40 ± 0.173 all in mL respectively. Hence the end points obtained using the extracts of *Acanthus pubescens* flower are fairly comparable to the end points obtained using the commercial indicators i.e. methyl red, bromothymol blue and phenolphthalein. The flower extracts of *Acanthus pubescens* could be used as excellent substitutes to methyl red, bromothymol blue and phenolphthalein for strong acid against weak base titrations.

For weak acid vs. strong base titration, the end point obtained using ethanol and acidified ethanol extracts of *Acanthus pubescens* flower are in the 10.40 ± 0.100 mL to 10.43 ± 0.058 mL range which is close to that obtained using phenolphthalein (10.40 ± 0.100 mL) but indicates slight statistical deviation from the end point obtained with methyl red (9.53 ± 0.058 mL) and bromothymol blue (9.67 ± 0.058 mL) in the titrations involving 1M acetic acid and 1M sodium hydroxide solutions. These slight differences are attributed to the decrease in dissociation of acetic acid in acidified and pure ethanol. The *Acanthus pubescens* flower ethanolic and acidified ethanol extracts are therefore good substitutes for phenolphthalein in this type of titration.

For the titration of weak acid with weak base (CH₃COOH vs. NH₄OH), the end point obtained with acidified ethanol extract of *Acanthus pubescens* flower was 8.87 ± 0.115 mL while ethanol extract of *Acanthus pubescens* flower changed colour at mean titre of 8.10 ± 0.100mL. The end point obtained using acidified ethanol extract of *Acanthus pubescens* flower...
flower was comparable to the end points obtained using phenolphthalein (8.83 ±0.058 mL) and bromothymol blue (8.77 ± 0.208 mL) but significantly different from the end point obtained with methyl red (8.07 ± 0.115 mL) while the end points obtained using ethanol Acanthus pubescens flower extract was closest to the end point obtained with methyl red but it shows significant deviation from the end point obtained with phenolphthalein in this medium. Hence the acidified ethanol extract of this plant could be used in place of phenolphthalein while the ethanol extract can replace methyl red in weak acid versus weak base titrations. Variations in titre volumes of the indicators for the weak acid–weak base titrations are due to lack of sharp end points for these combination and complex pH dependant protonation–deprotonation reactions, as shown in Figure 2, as well as changes in pKa values of the anthocyanin pigments. Mixed indicators are appropriate for such titrations to signal end points at the various pH values.

The results of this study are similar to the observations reported by various researchers (Jaspreet et al., 2011; Thote et al., 2015; Abbas, 2012; Senathirajah et al., 2017) that did related work on indicators using different parts of plant extracts, but there was a slight difference in the result as compared with that of Nhapi (2016) and Pimpodkar (2014).

4. Conclusion
The acid–base indicators used in the chemistry laboratory are very often synthetic which are expensive for the schools to afford for their large number of laboratory sessions, are toxic to students and teachers especially when they produce vapours in the laboratory classes, are pollutants to the school surrounding affecting animal and human health. This study focused on a natural indicator for acid-base titrations which is extracted from Acanthus pubescens flower. Four combinations of acid-base titration were studied: strong acid versus strong base, strong acid versus weak base, weak acid versus strong base, and weak acid versus weak base. The color change of the extracts, pH ranges and the average end point volumes were determined for each type of acid-base titration. These results were similar to those obtained from three selected synthetic indicators namely: methyl red, bromothymol blue and phenolphthalein. The presence of anthocyanins and flavonoids as active components of plant extract were chemically and spectrophotometrically analyzed. The authors concluded that the natural acid-base indicators prepared from pure and acidified ethanol extracts of Acanthus pubescens flower could be good replacements for synthetic indicators as they are less expensive, available, simple to manage and environmentally benign.

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