ASSESSMENT ON BIOACTIVE COMPOUNDS AND THE EFFECT OF MICROWAVE ON PITAYA PEEL

Norashikin Mat Zain*, Muhd Azlan Nazeri, Nurul Aini Azman

Faculty of Chemical Engineering and Natural Resources, University Malaysia Pahang, 26300, Gambang, Pahang, Malaysia

Abstract

Over the years, a wide variety of natural colour sources have been identified. One source, the pitaya fruit is known to impart colours to products, such as food and drink. However, there have been limited studies done to determine phenolic compounds and antibacterial activity of the pitaya peel (H.polyrhizus) via Microwave Assisted Extraction (MAE) method. Both this information can escalate the potential role of pitaya fruit as a natural colour source. This study aimed to identify the types of bioactive compounds (phenolic compounds) and antibacterial activity of pitaya peel. To achieve this objective, MAE was used to extract bioactive compounds from the pitaya peel as it maintained the integrity of the compound. Based on the results, 13 types of phenolic compounds were identified from the pitaya peel extract via qualitative research using library database matching which include quinic acid, cinnamic acid, quinic acid isomer, 3,4-dihydroxyvinylbenzene, isorhamnetin 3-O-rutinoside, myricetin rhamnoglucose, 3,30-di-O-methyl ellagic acid, isorhamnetin aglycone monomer, apigenin, jasmonic acid, oxooctadecanoic acid, 2 (3,4-dihydroxyphenyl)-7-hydroxy-5-benzene propanoic acid and protocatechuic acid. The pitaya peel extract was also found to have small antibacterial effect on the Gram-positive, Staphylococcus aureus (S.aureus) and Gram-negative, Escherichia coli (E.coli). The SEM demonstrated that cell wall disruption of pitaya peel caused by microwave radiation from MAE appeared to be the main reason for rapid extraction of bioactive compounds. In conclusion, the study established that pitaya peel extract is a natural colour source with an abundance of phenolic compounds and minimal antibacterial activity, which could be used in the food and cosmetic industries.

Keywords: Microwave assisted extraction (MAE), pitaya peel, pitaya extract, phenolic compound, time-kill kinetics

Abstrak

Sepanjang tahun, banyak sumber warna semula jadi telah ditemui. Salah satu sumber adalah buah pitaya yang boleh menyumbang kepada warna untuk produk seperti makanan dan minuman. Bagaimanapun, kajian yang dilakukan terhadap untuk menentukan sebatian fenolik dan aktiviti antibakterial daripada kulit pitaya (H.polyrhizus) melalui kaedah pengekstrakan berketuhar (MAE). Kedua-dua maklumat ini berkaitan dengan sebatian fenolik dan kesan antibakteria yang boleh meningkatkan peranan berpotensi buah pitaya sebagai sumber warna semula jadi. Oleh itu, kajian ini telah dijalankan bertujuan untuk mengenal pasti sebatian bioaktif (sebatian fenolik) dan aktiviti antibakterial dari kulit pitaya. Untuk mencapai objektif ini, MAE telah digunakan untuk proses pengasingan sebatian...
1.0 INTRODUCTION

Colouring has been widely used in food and cosmetic industries to make the products more appealing and attractive. As the public become increasingly concerned about the detrimental effects of synthetic colour additives on human health and environment, there has been an increase in demand for natural colour sources. Over the years, many natural sources for product colouring have been found, one of which is pitaya fruit.

Pitaya fruit is known to contain high phenolic compounds. Phenolic compounds are secondary metabolites produced in plants. They are found mostly in fruits, vegetables, herbs, roots, leaves and seeds. As partial response to the ecological and physiological pressures, phenolic compounds are synthesized by plants [1]. These compounds are considered as natural defense substances, either for reproduction or as sensorial properties including colour, bitterness, taste, astringency and flavour. The concentration of phenolic compounds in each plant may be influenced by physiological and geographic variations, environmental conditions, and genetic factors [2, 3]. In addition, phenolic compounds have been extensively exploited due to their natural biological activities against the bacteria, carcinogens, oxidation and aging process [4].

Natural sources, such as plants and fruits have been discovered to be antibacterial, leading to numerous investigations on the potential role of plant and fruit extracts as antibacterial agents. [5]. Similarly, the by-products of plants were found to demonstrate the ability to kill bacteria [6].

Furthermore, previous studies reported that the concentration of phenolic compounds were higher in the peel than in the flesh of plants [7]. Despite this finding, agro-industrial residues, such as fruit peel remained unpopular among manufacturers. The fruit peel, generated in massive amounts from the fruit juice processing industry is normally discarded as waste material. To make the matter worse, these discarded peels often become environmental hazards which cause water pollution. [8]. The seemingly useless fruit peels could potentially be used as a source of phenolic compounds while reducing environmental pollution.

Currently, investigations of bioactive compounds of fruit peel are still lacking due to their low popularity and limited commercial application [9]. In fact, the specific bioactive compounds of fruit peels have never been reported [10]. Thus, the main objective of this study was to determine the bioactive compounds (phenolic compounds) and the antibacterial properties of fruit peels.

Pitaya fruits were chosen for this study because they contain rich polyphenols which are found to display antioxidant, antiproliferative and antibacterial activities [11]. There are three types of pitaya: Hylocereus undatus, red skin with white flesh; Hylocereus polyrhizus, red skin with red flesh and Selenicereus megalanthus, yellow skin with white flesh [12]. Hylocereus polyrhizus was selected for the experiment as it is known to contain higher phenolic compounds and antioxidant activity compared to the other pitaya species. In this study, the peel of pitaya fruits were used because 33% of the weight comes from their peel, which indicates the abundance of bioactive compounds present within the peel [13]. Besides, the fruit peel has several advantages, such as being inexpensive, can be found in abundance, and is a sustainable resource [14, 15]. In addition, the peel and pulp contain natural colorants namely betacyanins (red pigment) and betaxanthin (yellow pigment) [13].
To isolate the different types of bioactive compounds from the plants, extraction process was important. Qualitative and quantitative studies of bioactive compounds from plant materials mostly relied on the selection of a proper extraction method [16]. Methods such as solvent-based extraction and cold pressing had drawbacks including safety hazards, high energy input, low product quality, low extraction yields, long extraction time, and environment risk. Therefore, MAE was applied in this research using sample preparation techniques because it could extract bioactive compounds more rapidly and provide better recovery than the conventional extraction process [17]. In this research, MAE was employed to prepare the samples due to its ability in extracting the bioactive compounds in a rapid manner, aside from limiting the degradation of bioactive compounds. It is also feasible in extracting phenolic compounds with higher yield compared to other conventional methods, such as soxhlet. In addition, the effect of cell wall disruption in the sample due to microwave radiation from MAE can be observed via scanning electron microscopy (SEM). The use of MAE could be further justified by its environmentally friendly feature which reduced the use of organic solvents during the extraction process[18].

To obtain a higher yield of bioactive compounds via MAE, water was used as the solvent extraction in this study. This was because studies have shown a higher extraction yield with the use of water compared to ethanol and methanol. This finding was also reported by Michel et al. [19] whereby water increased the contact surface area between the plant matrix and solvent, resulting in a higher extraction yield. Besides, water is known to have high dielectric properties constant which allowed more absorption and conversion of microwave energy from MAE into the heat to facilitate partition analytes from the sample into the solvent [20]. Moreover, the non-toxicity, non-corrosiveness, non-flammability, and cost-effectiveness of water made it the greenest solvent available [13].

Following the extraction and determination of the phenolic compounds in pitaya peel, antibacterial properties of the pitaya peel extract were determined. To evaluate in vitro antibacterial activity of an extract or a pure compound, a variety of laboratory methods could be used. The most known and basic method is optical density study also known as the time kill kinetic test [21]. This test was therefore used in this study to examine the antibacterial activity of peel extract. Additionally, the microscopic structural changes of pitaya peel before and after extraction was observed to understand the characteristic of MAE methods.

In essence, this study is the first to report the phenolic compounds and antibacterial activity from the extraction of *H. Polyrhizus* using MAE and time-kill kinetics test. The findings from this study could serve as a starting point for more elaborate research on the potential use of agro-industrial residues as natural sources for various industries. Such endeavour would be essential in meeting the growing demand for natural products and more importantly, in conserving the environment.

### 2.0 METHODOLOGY

#### 2.1 Sample Collection

The pitaya peel from *H. polyrhizus* were obtained from a supermarket in Temerloh, Pahang. The peels were weighed and washed with distilled water. Subsequently, the peels were cut into 2 cm portions. As part of the freeze-drying process, the samples were first frozen overnight in the fridge freezer at a temperature of -80 ºC and then placed in the freeze dryer for 96 hours. Next, the freeze-dried samples were grounded and sifted through a 20 mesh (0.85 mm) sieve to produce the powdered samples. These samples were stored in bags and kept in a dry environment prior to conducting the experiment [22].

#### 2.2 Microwave-assisted Extraction (MAE)

Extraction of the bioactive compounds from the pitaya peels was carried out using the MAE approach with the one-factor-at-a-time method (OFAT). The liquid-solid extraction was conducted by adding the freeze-dried samples and 50 mL of distilled water into a 1000 mL extraction vessel. Subsequently, the extraction vessel was placed into MAE. The best conditions of MAE, particularly power, temperature, time, and weight of the sample were applied which are 400W, 45, 20 min and 1.2 g respectively. Immediately after the extraction process, the homogenate was centrifuged at 9000 rpm for 40 minutes at 25°C. After the centrifugation, the supernatant was collected and the same procedure was repeated twice to ensure maximum extraction of the bioactive compounds. All experiments were performed in triplicates [23, 24].

#### 2.3 Analysis of Solution Extraction

##### 2.3.1 Determination of Phenolic Compounds by UHPLC-ESI-QTRAP-MSMS

The supernatant obtained from the MAE was collected and filtered using the Whatman filter paper (size 5C). Next, the filtered solution was diluted with 5 mL of methanol (50%) UHPLC grade and filtered with 0.45 µm nylon syringe filter to inject a volume of 20 µL into LCMSMS [25]. The analysis was performed using a Flexar FX 15 ultra-high-performance liquid chromatograph (UHPLC, PerkinElmer, Inc, Massachusetts, USA) coupled with an AB SCIEX 3200 QTrap hybrid linear ion trap triple quadrupole mass spectrometer (equipped with a turbo ion spray source).
The chromatographic separation was achieved on a Phenomenex Synergi Fusion (100 mm x 2.1 mm x 3 µm) column. Mobile phase A contained water with 0.1 % (v/v) formic acid and 5 mM ammonium formate, while mobile phase B was composed of acetonitrile containing 5 mM ammonium formate. Elution was performed by means of a linear gradient from 5-95 % B (0.01-10 minutes) held for 2 minutes, then returned to 10 % B in 0.1 minutes, and finally re-equilibrated for 3 minutes before the next injection. Ionisation was achieved through electrospray ionisation on the AB Sciex Turbo V source, at a temperature of 500°C using purified nitrogen gas (99%) as the collision gas via nebulisation. Collision energy was set at 35eV for mass fragmentation purpose. A full scan with MS/MS data collection analysis was performed in a negative mode. Data analysis, processing and interpretation were carried out using the AB SCIEX Analyst 1.5 and Advance Chemistry Development, Inc (ACD/Labs, Ontario, Canada) MS Processor software. A Marker View Software (AB SCIEX, Massachusetts, USA) was used for the principal component analysis (PCA). The following parameters were used for PCA: retention time (Rt) range: 0-15 minutes, tolerance: 0.5 minutes, mass range: m/z 100-1000, mass tolerance: 0.01 Da and noise threshold: 5. [26]

### 2.3.2 Antibacterial Testing

#### 2.3.2.1 Preparation of Medium

**Preparation of Broth**

500 mL of broth was divided and placed into 2 conical flasks, each containing 250 mL of broth. 15 g of Tryptone Soya Broth (TSB) was added to 500 mL of distilled water and transferred to the Schott bottle. The solution was stirred using the magnetic stirrer and boiled on a hotplate to dissolve completely. Each flask was filled with 100 mL of broth and sterilized in the autoclave at a temperature of 121°C for 20 minutes [27].

**Preparation of Agar**

20 g of Tryptone Soya Agar (TSA) was added to 500 mL of distilled water and transferred into the Schott bottle. The mixed solution was boiled on a hotplate to dissolve completely. Petri dishes, TSB and TSA were put into the autoclave at a temperature of 121°C for 15 minutes. The cap of the Schott bottles was made loose to withstand the high pressure. Meanwhile, the laminar flow cabinet was sprayed and wiped with 70% ethanol. Then, UV rays were turned on for 15 minutes to sterilize the workspace. After the autoclave cooled down, the agar medium was poured onto the petri dishes and left to harden. Once hardened, the petri dishes containing the agar medium were sealed with parafilm to prevent any contamination. Subsequently, the petri dishes were stored upside down in the freeze[28].

#### 2.3.2.2 Bacteria Culture Preparation

**Inoculum**

S.aureus and E.coli were used as the test organisms, which were prepared from fresh colonies on TSA. One loopful of each of the bacteria was inoculated in different conical flasks containing 100 mL of TSB. The flasks were incubated in an incubator shaker at a temperature of 30 °C and at 150 rpm for 18 hours until the exponential phase. Following the incubation, 100 µL of each bacteria culture were reinoculated into new sterilized conical flasks containing 100 mL of TSB for another 22 hours to achieve 10^8 CFU/mL [29].

#### 2.3.2.3 Analysis of Antibacterial

**Time-kill Kinetics Test**

Time-kill kinetics test was the most appropriate method to reveal a time-dependent or a concentration-dependent antibacterial effect [30]. It was performed on the broth culture medium (TSB) using three conical flasks containing a bacterial suspension of S.aureus and E.coli, respectively. 50 mL of sterilized TSB were measured and placed into a conical flask. Three flasks were prepared for each bacteria suspension against the extracted sample. 1000 µL of bacterial suspension were added into each flask. Following that, 5 mL and 10 mL of pitaya peel extract were added into the first and second conical flasks of TSB respectively. The third flask was prepared by excluding the pitaya peel extract and was considered as a growth control. To obtain the initial optical density (OD), the samples were taken immediately after addition of the pitaya peel extract. Subsequently samples were taken every two-hourly up to a total of 14 hours (2, 4, 6, 8, 10, 12 and 14 hours) with the incubation being done under a suitable condition (30°C and 150 rpm) [21]. For OD reading, the samples were also diluted to an appropriate concentration at a time interval of 2 hours. After that, the absorbance of the sample was measured with a spectrophotometer at 600 nm [31]. A calibration curve to relate the absorbance with the incubation time was then generated.

### 2.4 Scanning Electron Microscopy (SEM) Observation

The SEM (FET QUANTA-450, Netherlands) was employed to observe the morphological changes of pitaya peel samples before and after MAE extraction. The samples were oven-dried at 60 °C for 15 minutes and sputtered with a thin layer of platinum. Next, the samples were observed under vacuum at an accelerating voltage of 8.0 KV and 500 magnification [32].
3.0 RESULTS AND DISCUSSION

3.1 Identification Phenolic Compounds

Table 1 shows a total of 13 phenolic compounds found in the pitaya peel extract. Besides that, 28 compounds were classified as unknown compounds because of poor match with library database and were suspected as flavonoids compounds. The compounds were identified based on library database matching and their mass fragmentation pattern showed fingerprint for each compound [26]. Based on the full chromatogram from UHPLC-ESI-QTRAP-MSMS analysis as presented in Figure 1, pitaya peel extract contained rich amounts of phenolic acid and flavonoids compounds. The analysis revealed that pitaya peel extract contained quinic acid, cinnamic acid, quinic acid isomer, 3,4-dihydroxyvinylbenzene, isorhamnetin 3-O-rutinoside, myricetin rhamno-hexoside, 3,30-di-O-methyl ellagic acid, isorhamnetin aglycone monomer, apigenin, jasmonic acid, oxoacadecanoic acid, 2 (3,4-dihydroxyphenyl)-7-hydroxy-5-benzene propanoic acid and protocatechuic hexoside conjugate.

Although phenolic contents of H.polyrhizus had previously been studied, there were no full scan studies of the phenolic compounds [10]. This study is therefore the first to have identified specific bioactive compounds in pitaya peel extract of H.polyrhizus via MAE. Based on the results, the analysis of the pitaya peel extract with UHPLC revealed two types of phenolic acids, which were hydroxycinnamic acids (quinic acid, cinnamic acid, quinic acid isomer and 2 (3,4-dihydroxyphenyl)-7-hydroxy-5-benzene propanoic acid) and hydroxybenzoic acids (3,4-dihydroxyvinylbenzene, 3,30-di-O-methyl ellagic acid and protocatechuic hexoside conjugate). Hydroxybenzoic acids had a C6-C1 chemical structure whereas hydroxycinnamic acids had a C6-C3 structure with a double bond in the side chain in cis or trans configuration [33]. The remaining compounds identified from the pitaya peel extract were flavonoids, which displayed antioxidant [11, 34-36], antiproliferative [37] and antibacterial activities [11]. The flavonoids identified in this study are isorhamnetin-3-o-rutinoside, isorhamnetin aglycone monomer, jasmonic acid, apigenin and myricetin rhamno-hexoside.

The findings in Table 1 differed from the findings reported by Lourith and Kanlayavattanakul [10] where chlorogenic acid, gallic acid and quercetin were identified from pitaya peel extract via HPLC (Waters 2695, Agilent, USA). The different findings are due to using different type of HPLC in phenolic compounds identification, which in this study used more advance HPLC. Nevertheless, the results of this research are consistent with the findings of previous work which similarly extracted phenolic compounds from other fruits and plants via MAE [20]. This indicates that MAE is an efficient method which could be used to extract different type of phenolic compounds. Additionally, Fang et al. [38] revealed that a lower consumption of solvent in MAE would give rise to a higher yield of phenolic compounds. This study established a similar finding where the use of a lower amount of solvent in MAE provided stability to the phenolic compounds, allowing them to be identified from the samples.

Table 1 Compounds identified in pitaya peel extracted based on UHPLC-ESI-QTRAP-MSMS

<table>
<thead>
<tr>
<th>Retention time (min)</th>
<th>[M-H]</th>
<th>Mass fragment, MS/MS (m/z)</th>
<th>Compound identified</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.424</td>
<td>191.40</td>
<td>191.05, 172.92, 110.98</td>
<td>Quinic acid</td>
</tr>
<tr>
<td>1.986</td>
<td>147</td>
<td>56.90, 86.95, 102.97</td>
<td>Cinnamic acid</td>
</tr>
<tr>
<td>3.866</td>
<td>191.20</td>
<td>191.04, 172.99, 110.93</td>
<td>Quinic acid isomer</td>
</tr>
<tr>
<td>3.977</td>
<td>135.19</td>
<td>135.00, 116.88, 88.80</td>
<td>3,4-Dihydroxyvinylbenzene</td>
</tr>
<tr>
<td>4.033</td>
<td>623.20</td>
<td>623.14, 605.25, 579.00, 314.04, 299.04</td>
<td>Isorhamnetin 3-O-rutinoside</td>
</tr>
<tr>
<td>5.037</td>
<td>625.32</td>
<td>625.14, 315.10, 300.20</td>
<td>Myricetin rhamno-hexoside</td>
</tr>
<tr>
<td>6.154</td>
<td>328.78</td>
<td>329.30, 311.29, 229.21, 211.16, 171.15, 139.14</td>
<td>3,30-di-O-methyl ellagic acid</td>
</tr>
<tr>
<td>6.489</td>
<td>314.80</td>
<td>315.12, 300.08, 271.06, 243.12, 164.04</td>
<td>Isorhamnetin aglycone monomer</td>
</tr>
<tr>
<td>6.600</td>
<td>269.13</td>
<td>269.15, 251.10, 225.29, 197.04</td>
<td>Apigenin</td>
</tr>
<tr>
<td>7.045</td>
<td>209.22</td>
<td>209.12, 163.15</td>
<td>Jasmonic acid</td>
</tr>
<tr>
<td>7.489</td>
<td>297.35</td>
<td>297.23, 183.00</td>
<td>Oxooctadecanoic acid</td>
</tr>
<tr>
<td>8.043</td>
<td>310.40</td>
<td>311.20, 183.03, 96.91</td>
<td>2 (3,4-Dihydroxyphenyl)-7-hydroxy-5-benzene propanoic acid</td>
</tr>
<tr>
<td>8.600</td>
<td>595.40</td>
<td>595.24, 315.03, 279.28, 241.04</td>
<td>Protocatechuic Hexoside conjugate</td>
</tr>
</tbody>
</table>
3.2 Time-kill Kinetics Test of Pitaya Peel Extract

Figures 2 and 3 show the optical density (OD) of the samples taken every 2-hourly up to 14 hours. In this study, time-kill kinetics was chosen because this method was faster, easy-to-use and flexible method.

The growth profile of the *S. aureus* and *E. coli* is shown in Figures 2 and 3 respectively. During the lag phase (0-2 hour), the growth of the *S. aureus* and *E. coli* should theoretically decline after inoculation because the cells need time to adapt to the new environment inside the conical flask. The lag phase should occur immediately after inoculation as a period of adaptation of cell to a new environment [40, 41]. During this phase, cell mass may increase a little without an increase in the cell number density. Nonetheless, the lag phase was not evident from the graph of this study. This finding could be due to the lag phase being too short to notice as a result of the two-hourly sample OD reading.

Furthermore, the length of the lag phase was known to be affected by the bacterial age and inoculum size. According to Swinnen et al. [41] the shortest lag phase and the time to the first division were obtained with the largest inoculum and the youngest inoculated parent cells. The decrease of inoculum size can be explained by the increased lag phase duration and the corresponding variation as reported by numerous researchers [42]. Thus, the short lag phase found in this experiment might be due to the age of the inoculated parent cells.

The exponential phase (log phase or growth phase) from the 2nd to 6th hour shown in Figures 2 and 3 indicate the full adaption of the cells to the new environment of the conical flask and therefore, their ability to grow and produce new cells [43]. Direct visual measurement of the different growth curve patterns was performed in this experiment. The values of the growth curve of *S. aureus* and *E. coli* for 10% and 20% pitaya peel extract were slightly lower than the control. The bioactive compounds in pitaya peel extract demonstrated an ability to inhibit *S. aureus* and *E. coli* growth even though the effect was small. The growth of the *S. aureus* and *E. coli* slowed down after the sixth hour.

Based on Figures 2 and 3, stationary phase of the growth profile started after $t=6$ hours until $t=12$ hours, while the death phase of the growth profile began after $t=12$ hours. It is unclear from this study whether inoculum size was a critical factor affecting the transition from the exponential phase to stationary phase. Hence, further studies would be needed to assess the effect of inoculum size on the late exponential and early stationary phase [42].

During the stationary phase, the death rate was equal to the growth rate of the cells. The growth of *S. aureus* and *E. coli* could be limited by nutrient depletion at the level where cell growth could not be sustained. During this phase, the growth curve of the control was expected to give a higher OD value than pitaya peel extract, with the 10% pitaya peel extract similarly giving a higher OD value than the 20% sample extracted. The result of the experiment between the 8th hour and the 10th hour however, did not show a trend as predicted. Instead, the 20% pitaya peel extract had a slightly higher OD value than that of the 10% sample extracted. This finding could be due to errors in reading the stationary phase of the growth profile graph. To avoid such errors, it is recommended that only two cuvettes be used, one for blank (fresh TSB) and the other for
samples, throughout the experiment as different cuvettes might allow different amounts of light intensity to pass through.

On further examination of the graph, it was evident that the death phase began after 12 hours. The OD of the cells from the three growth curves decreased as the cells lysed and released their intracellular metabolites into the growth medium [44, 45]. Cells might undergo apoptosis (cell suicide) process during its death phase. Nutrient depletion often initiated the apoptosis process which involved DNA cleavage, after which the cells would shrink [46].

As can be seen, the OD value between samples and control were slightly close for each incubation period. The result obtained was consistent with previous work by Majhenič et al. [47]. Their study concluded that, by using boiling water as solvent, guarana (Paulinia cupana) seed extract had little yet sufficient antibacterial activity to be used in food products as antibacterial agents. Similar findings were observed in other studies where time-kill kinetics test was reported to be able to detect differences in the rate and extent of antibacterial and antifungal over time [48, 49].

3.3 Analysis of Microscopic Changes

Figure 4 (a) presents the untreated sample that displays an intact external structure with many folded layers. Most of the external structures in pitaya peel that were analyzed via MAE had become flat and completely ruptured, as shown in Figure 4 (b). This indicates that microstructural changes in the cell walls are caused by MAE [50].

The disruption of cell wall in pitaya peel extract due to microwave radiation can promote the release of greater amounts of bioactive compounds into the solvent extraction within a short time [51, 52]. The two transport patterns pertaining to mass and heat that function in the same direction (from the inside to the outside of pitaya peel) accelerate the rates of extraction in MAE [53, 54]. Additionally, microwave radiation increased the temperature rapidly for the sample, thus causing the cell wall to rupture and facilitate the rapid release of bioactive compounds. Hu et al. [32] asserted that the effects of both cavitation and turbulence generated by microwave radiation can also destroy cell wall and promote mass transfer simultaneously, thus increasing the extraction yield.

This result is similar to that recorded by Yanik [55], who applied MAE to extract bioactive compounds from olive pomace. Hu et al. [32] also reported a similar pattern in extracting bioactive compounds from Tiger Nut (Cyperus esculentus L.), which demonstrated that cell wall disruption of the samples had been due to microwave radiation, as observed via SEM.

4.0 CONCLUSION

As evidenced by the full chromatogram from UHPLC-ESI-QTRAP-MSMS analysis, pitaya peel solution extraction was found to be high in polyphenols and flavonoids compounds. Furthermore, MAE was the optional equipment for extraction of pitaya peel as it maintained the integrity of their bioactive compounds. Additionally, time-kill kinetics test was an efficient method to determine the level of antibacterial activity in pitaya peel extract. The pitaya peel extract was proven to have antibacterial properties as the extract demonstrated small effect of antibacterial activity against both S.aureus and E.coli. In conclusion, pitaya peel is a natural colour source which could be used in food and cosmetic products. Additionally, SEM demonstrated that structural disruption of pitaya peel caused by microwave radiation was the main reason for rapid extraction. So, MAE could be a good alternative for extraction of pitaya peel and other herbs.
Acknowledgment

The authors would like to express their gratitude to University Malaysia Pahang for the financial support through grant RU150312.

References


