ABSTRACT

Developing countries are still counting on herbal medicine which aims to promote primary health care with better cultural, human acceptability and fewer side effects. Fragrea acuminatissima is one of underutilized herb consumed by Temiar tribe in Malaysia which able to give the similar benefits. The most important attribute in developing herbal based product is the extraction condition and selection of the technique used will potentially affect the expression of antioxidant compound in herbal plant. Technical problem arising from the extraction phase also urge the needs to elucidate the best techniques for respective herbal product. Therefore, this study was aimed to investigate the best extraction techniques (maceration, orbital shaker assisted, ultrasonic assisted and microwave assisted) on Total Phenolic Content (TPC), Total Flavonoid Content (TFC) and antioxidant activity of F. acuminatissima stem and root parts by using methanol solvent. Folin-Ciocalteu method was used for determination of TPC, aluminium chloride colorimetric method for TFC, while antioxidant activity was determined by DPPH radical scavenging assay. Overall, the best extraction method of F. acuminatissima was ultrasonic assisted extraction which exhibited the highest percentage yield, TPC and TFC values for both plant parts. Based on the DPPH assay, F. acuminatissima have procured the excellent IC50 of 29.954 µg/ml by ultrasonic assisted extraction for stem while maceration techniques showed the best IC50 for root at 76.996 µg/ml. Positive moderate and strong correlation between TPC, TFC and DPPH was detected which support the result obtained. In brief, ultrasonic assisted technique was suggested to be used in future research of this plant for phytochemical profiling, bioactivity assays and herbal product development.

Keywords: Fragrea acuminatissima, extraction techniques, Total Phenolic Content (TPC); Total Flavonoid Content (TFC), DPPH radical scavenging effect.

Introduction

Plant extracts from fruits, herbs, vegetables and cereals which had been scientifically proven to contain abundance secondary metabolites are currently becoming a major interest in the food industry because of their ability to impede the oxidative degradation of macro nutrients and thus improving the quality and nutritional value of food (Žugić et al., 2014; Abdul Mutalib et al., 2013). Herbs and spices have been widely used not only as food preservatives and flavoring, but also as traditional medicines for thousand years (Yi & Wetzstein, 2011). These herbal medicine are being used extensively because they can be considered as safe since they are from natural resources (Aziz et al., 2014). The usage of herbs in processed food as a source of alternative antioxidant to substitute synthetic antioxidant is well-known in food industry as they are cheaper, easily consume and available locally either in raw or conventional preparations method (Nićiforović et al., 2010).

The plant species Fragrea acuminatissima (Loganiaceae) or locally known as Tengkuk Biawak grows wild in Peninsula region of Malaysia. Their best habitats are at inland forest and lowland area. This plant is an epileptic climber and falls under the shrub tree category. Fragrea acuminatissima are being widely used by Temiar tribe, one of the local indigenous population in Kelantan for treatment of fever and body ache.
Currently, there are no other studies had been done on this plant especially on the antioxidant and phytochemical aspects. However, there were studies had been conducted for the plant under Loganiaceae family, for example, *Anthocephalista nobilis* was claimed to effectively cure fever, diarrhea and stomach ache in West Africa (Ngwoke et al., 2015). According to the previous finding, antioxidant activity of stem bark extracted by acetone and methanol solvents as well as by using maceration technique showed good antioxidant properties. These findings provided evidence that *F. acuminatissima* might be able to exhibit the same potential as it falls under the same family of Loganiaceae.

The extraction yield of active secondary metabolites from plant materials is mainly influenced by the process of extraction with aqueous and/or organic solvent (Cacace & Mazza, 2003). Effective separation of antioxidants with high extraction yield and concentration of bioactive compound from a complex plant matrix is a difficult procedure due to co-extraction of other compound, which are undesirable in antioxidant extract (Bimakr et al., 2010).

Maceration technique had been conventionally used to extract the bioactive compounds from plant material. This technique does not require complicated instrumentation and only involve a few simple procedures. However, the main drawback of this technique is that it requires longer time for extraction process to complete (Azmir et al., 2013). In order to increase time efficiency and extraction yield, shaking technique had been incorporated in the conventional maceration method and some studies showed better results (Amorim-Carrilho et al., 2014; Abdul Mutalib et al., 2013). Other than that, ultrasound has been recognized for potential industry application in the phyto-pharmaseutical extraction for a wide range of herbal extract as it creates forces that destruct cell walls mechanically and enhance material transfer (Truta et al., 2010). Simple, inexpensive and efficient ultrasonic assisted extraction have major benefits over conventional techniques as it has faster kinetic and able to increase the extraction yield (Wang & Weller, 2006). Microwave assisted extraction also can be considered as a potential alternative for solid-liquid extraction of the metabolites from plants. This method is applied to extract nutraceutical which able to improve extraction yield, reduce solvent usage and extraction time (Wang & Weller, 2006). Dipole rotation of the solvent in microwave field generate heat effect which enhance the product recovery contributed to solvent temperature rise, thus increase the solubility of the compound present (Zaheer, 2011).

Various extraction techniques had been innovated mainly to increase the efficiency and ease the extract production. However, different extraction techniques might exhibit different antioxidant compounds in respective plant. Since there are insufficient information regarding this underutilized plant, thus, this preliminary research aimed to study the antioxidant potential of root and stem of *F. acuminatissima* through different extraction techniques. Total phenolic and flavonoid content were assessed as they represent the most abundant antioxidant content in natural resources and DPPH scavenging activity assay was evaluated since this method is established well and involve only simple procedures.

Materials and Methods

**Materials**

Chemical used in this study were Folin-ciocalteu reagent, NaH2PO4, Na2HPO4, sodium carbonate, aluminium trichloride, gallic acid, trichloroacetic acid, potassium persulfate ascorbic acid and methanol as supplied by Merck, Germany. Whereas quercetin, gallic acid and 2,2-diphenyl-picrylhydrazyl (DPPH) were obtained from Sigma Aldrich, USA. Plant samples were collected at Kuala Betis, Gua Musang in Kelantan, Malaysia.

**Sample preparation**

The samples were separated into stem and root parts, thoroughly washed and blot dried. Then, samples were course ground into flakes and dried in the electric oven (Lab Companion, Korea) for 3 days at 50 °C. The dried samples were further ground to fine powder using electrical grinder (Panasonic, Japan) and kept in 4 °C chiller prior extraction.

**Sample extraction**

Sample extraction was done by using simple maceration technique, and assisted with orbital shaker (Lab Companion, Korea), ultrasonic bath (Lab Companion, Korea) and microwave oven (Sharp, Japan) according to the methods described by Savita & Prakashchandra (2011) with a slight modification. Overall, 50 g of powdered samples were mixed with 250 ml of 95 % methanol. Then, mixtures were proceeded for incubation by using incubation protocols stated in Table 1. After the incubation period, the mixtures were filtered by Whitman no. 4 filter paper and samples were extracted again using the same protocols. Then, the solutions obtained were pooled and evaporated to obtain the semi-solid crude sample, respectively.

**Table 1: Incubation protocols of different extraction techniques**

<table>
<thead>
<tr>
<th>Techniques</th>
<th>Temperature</th>
<th>Period</th>
<th>Condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maceration</td>
<td>Room temperature</td>
<td>24 hours</td>
<td>-</td>
</tr>
<tr>
<td>Orbital Shaker Assisted (OAE)</td>
<td>Room temperature</td>
<td>3 hours</td>
<td>Shaking frequency at 150 rpm</td>
</tr>
<tr>
<td>Ultrasonic Assisted (UAE)</td>
<td>Room temperature</td>
<td>1 hour</td>
<td>Medium sonication (17 kHz sound wave)</td>
</tr>
<tr>
<td>Microwave Assisted (MAE)</td>
<td>Heat by dipole rotation of solvent molecules</td>
<td>15 minutes</td>
<td>Power of 90 W</td>
</tr>
</tbody>
</table>
Total Phenolic Content

Total phenolic content was determined by using Folin-Ciocalteu assay based on method described by Singleton & Rossi (1965) with slight modification by Ghazi et al. (2012). Folin-Ciocalteu reagent was prepared by diluting the stock reagent in distilled water with ratio of 1:10. Then, 50 µl of sample extract at concentration of 1 mg/ml were mixed with 250 µl of 10 % Folin-Ciocalteu reagent in the test tubes. The mixtures were incubated for 5 minutes at room temperature and 750 µl sodium carbonate were added into the test tube. The mixtures were shaken and incubated in the dark for 2 hours at room temperature. Then, absorbance values were recorded at 765 nm using UV-Vis spectrophotometer (Thermoscientific, USA). Gallic acid was used as a standard following the similar procedures and graph was constructed in range of 6.25 to 400 µg/ml. Linear equation obtained was used to calculate the total phenolic content and values were expressed as µg gallic acid equivalent (GAE)/mg extract.

Total Flavonoid Content

Total flavonoid content was evaluated by using aluminium chloride colorimetric method described by Christ & Muller (1960) with slight modification by Ahmed et al. (2014). Briefly, 0.3 ml of sample extracts at concentration of 1 mg/ml were mixed with 3.4 ml of 30 % aqueous methanol, 150 µl sodium nitrite solution (0.5 M) and 150 µl aluminium chloride solution (0.3 M). After 5 minutes, 1 ml of NaOH solution (1 M) was added. Then, absorbance values were measured at 506 nm against blank. Standard graph of quercetin was constructed in range of 50 to 1600 µg/ml following the similar procedures and linear equation was obtained. Total flavonoid content of plant extracts were calculated and expressed as µg quercetin equivalent (QE)/mg extract.

Free Radical Scavenging activity

Free radical scavenging activity was done by using 2,2-diphenyl-picrylhydrazyl (DPPH) reagent following the method described by Brand-Williams et al. (1995) with slight modification by Gan et al. (2013). DPPH solution was prepared by diluting in 95 % methanol at different concentrations ranged from 6.25 µl/ml to 400 µl/ml. Amount of 2 ml of diluted extracts were added with 2 ml DPPH solution and incubated in the dark for 30 minutes. Then, absorbance values were measured at 517 nm using UV-Vis spectrophotometer against blank. Control containing 2 ml of 95 % methanol and 2 ml DPPH was prepared following the similar procedures. Scavenging activity was calculated according to following equation:

\[ \text{DPPH radical scavenging activity (\%) = \frac{(Ac - As)}{Ac} \times 100, } \]

where Ac is the absorbance of the control and As is the absorbance of the sample extracts. Graph of scavenging activity was plotted and inhibition concentration at 50 % (IC50) was estimated. Ascorbic acid was used as standard comparison.

Statistical Analysis

Data were reported as mean ± standard deviation using triplicate measurement values. Statistical analysis was done using Statistical Packages for Social Science (SPSS) version 20.0 (IBM Corp., USA). One-way analysis of variance (ANOVA) and Duncan’s test were performed to compare the mean between samples. Pearson correlation coefficient was done to correlate between antioxidant activity and content. Data were considered significant differences at p ≤ 0.05.

Result and Discussion

Sample Extraction

The ability of different extraction techniques in exhibiting the antioxidant potential were investigated. In brief, the time of extraction was ranged from 15 minutes to 24 hours, where MAE had the lowest extraction time while maceration technique had the highest extraction time. MAE, UAE and OAE are the instrumental techniques while maceration extraction is only using gravitational force. After solvent removal, the colour produced by all crude extracts was dark brown in colour and the extracted condition were in highly viscous or semi solid state.

Table 2 shows the extraction yield of F. acuminatissima stem and root by using different extraction techniques. From the result, the UAE for both stem and root of F. acuminatissima showed the highest yield, meanwhile the lowest yield for stem extract was obtained through maceration and MAE technique for the root extract.

<table>
<thead>
<tr>
<th>Extraction Techniques</th>
<th>Stem</th>
<th>Root</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Raw (g)</td>
<td>Yield (g)</td>
</tr>
<tr>
<td>Maceration</td>
<td>50</td>
<td>2.26</td>
</tr>
<tr>
<td>Orbital Shaker Assisted (OAE)</td>
<td>50</td>
<td>3.20</td>
</tr>
<tr>
<td>Ultrasonic Assisted (UAE)</td>
<td>50</td>
<td>3.93</td>
</tr>
<tr>
<td>Microwave Assisted (MAE)</td>
<td>50</td>
<td>2.93</td>
</tr>
</tbody>
</table>
Ultrasonic assisted extraction decreases the inner and external mass transfer limitation, thus gaining the yield percentage (Gimbut et al., 2014). A comparison study on different extraction techniques in yellow tea had suggested that ultrasound assisted extraction could have been used successfully for extraction of polyphenol and methylxanthines, thus proving that this technique provide more energy efficient (Horžič et al., 2012). Although microwave assisted extraction shown higher percentage of recovery effect of anthraquinoines extraction, but after 60 minutes of extraction, the yield become plateau and constant (Hemwimon et al., 2007).

Total Phenolic Content (TPC)

TPC of stem and root of F. acuminatissima at different extraction techniques were estimated by Folin-Ciocalteu’s reagent and using gallic acid as standard. The graph of gallic acid concentration ranged from 6.25 µg/ml to 400 µg/ml at 750 nm was plotted with a regression co-efficient (R²) = 0.9967 and linear equation of y = 0.008x + 0.078. The absorbance was steadily increased with the incremental of concentration.

Table 3 shows TPC content of plant parts at different extraction techniques. The highest TPC for both stem and root parts were obtained from UAE and the lowest TPC for both stem and root parts were acquired from MAE. Statistical analysis showed that there were significant differences between the TPC values of entire sample (p < 0.05).

<table>
<thead>
<tr>
<th>Extraction Techniques</th>
<th>Total Phenolic Content (µg GA/mg extract)</th>
<th>Stem</th>
<th>Root</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maceration</td>
<td></td>
<td>205.33 ± 0.78a</td>
<td>137.99 ± 0.33b</td>
</tr>
<tr>
<td>Orbital Shaker Assisted (OAE)</td>
<td></td>
<td>188.63 ± 0.13c</td>
<td>156.13 ± 0.63c</td>
</tr>
<tr>
<td>Ultrasonic Assisted (UAE)</td>
<td></td>
<td>206.88 ± 0.13b</td>
<td>184.33 ± 0.20c</td>
</tr>
<tr>
<td>Microwave Assisted (MAE)</td>
<td></td>
<td>171.00 ± 0.13d</td>
<td>83.46 ± 0.20c</td>
</tr>
</tbody>
</table>

* The values are means ± standard deviation (n = 3). Values with different superscripts were significantly different by Duncan test at level of p < 0.05.

The findings were supported by other study where ultrasonic assisted extraction technique also showed higher yield of phenolic compounds in Nigella sativa crude extract (Gimbut et al., 2014). Cavitation phenomena in ultrasonic was possibly occurred by propagation of ultrasound pressure waves through the solvent and the plant materials (Shirsath et al., 2012). Phenolic component reacts differently depending on the extraction condition and composition (Ksouri et al., 2008). Therefore, antioxidant capabilities are influenced by several factors that cannot be evaluated by single condition where other criteria such as solvent extracting power, duration of the extraction and their interaction are also important (Falleh, 2012). Many researchers also noted that TPC assay not only specific to phenols but sometimes also oxidized another components (Escarpa & Gonzalez, 2001).

Total Flavonoid Content (TFC)

TFC for stem and root of F. acuminatissima at different extraction techniques were measured with the aluminium chloride colorimetric assay using quercetin as standard. The graph of quercetin concentration ranged from 50 µg/ml to 1600 µg/ml at 510 nm was plotted with a regression co-efficient (R²) = 0.9802 and linear equation of y = 0.0003x + 0.0379.

Table 4 shows TFC of stem and root of F. acuminatissima at different extraction techniques. The highest TFC for the extraction of F. acuminatissima stem was obtained by using UAE technique and the lowest TFC was gained through extraction using maceration technique. Meanwhile, for the root part, the highest TFC was obtained from ultrasonic UAE technique and OAE technique showed the lowest TFC value. Overall, there were significant differences between the TFC values of entire sample (p < 0.05). However, further analysis using Duncan test showed no significant different between TFC in UAE and MAE of root part.

<table>
<thead>
<tr>
<th>Extraction Techniques</th>
<th>Total Flavonoid Content (µg QE/mg extract)</th>
<th>Stem</th>
<th>Root</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maceration</td>
<td></td>
<td>479.67 ± 3.33c</td>
<td>449.67 ± 3.33b</td>
</tr>
<tr>
<td>Orbital Shaker Assisted (OAE)</td>
<td></td>
<td>581.89 ± 5.09c</td>
<td>443.00 ± 3.33c</td>
</tr>
<tr>
<td>Ultrasonic Assisted (UAE)</td>
<td></td>
<td>873.00 ± 3.33b</td>
<td>566.33 ± 1.92d</td>
</tr>
<tr>
<td>Microwave Assisted (MAE)</td>
<td></td>
<td>609.81 ± 3.33d</td>
<td>563.00 ± 3.33a</td>
</tr>
</tbody>
</table>

* The values are means ± standard deviation (n = 3). Values with different superscripts were significantly different by Duncan test at level of p < 0.05.

Higher efficiency of UAE in exhibiting the flavonoid compounds could be attributed to acting sonic vibration, which produce particle size reduction and cell disruption. Moreover, the ultrasonic vibration towards solid’s surface lead to greater contact area between solid and liquid phase (Chen et al., 2007). Ultrasonic assisted extraction gives the highest extraction yield of some...
flavonoid components such as tectoridin, iristectorin A, iristectorin B, tectorigenin, iris-tectorigenin A and total isoflavones in shorter time next to soxhlet extraction and maceration extraction (Sun et al., 2011). However, study done by Gimbun et al. (2014), showed otherwise where, maceration techniques revealed higher yield of TFC when compared to UAE. This results support the previous statement by Falleh (2012) which stated that there are other extraction conditions that need to be considered in order to exhibit the antioxidant capabilities.

**DPPH Radical Scavenging Assay**

Figure 1 shows the graph of DPPH scavenging effect of ascorbic acid and different parts of *F. acuminatissima* at different extraction techniques. Generally, standard and samples showed a plateau pattern starting from concentration of 200 µg/ml except for ultrasonic assisted root which only reached the plateau stage at 400 µg/ml.

Figure 1: DPPH scavenging activity of ascorbic acid and samples at different plant parts and extraction techniques

This study revealed that all samples had exhibited IC_{50} value of DPPH radical scavenging activity as shows in Table 5. The value of IC_{50} was ranging from 29.33 µg/ml to 82.00 µg/ml for stem extracts where, UAE was found to be the best and OAE was the weakest extract to scavenged 50 % of DPPH free radicals. Other than that, IC_{50} of root extract ranged from 74.17 µg/ml to 82.51 µg/ml where, extract from maceration technique need lowest amount of concentration to obtain 50 % scavenging effect and MAE need much higher concentration. Meanwhile, the standard ascorbic acid showed the IC_{50} value of 70.483 µg/ml. Generally, all the samples were statistically different at p < 0.05. Despite of that, further Duncan analysis showed there were no significant differences in IC_{50} value between OAE and UAE of root extracts as well as between OAE of stem extract and MAE of root extract.

**Table 5: IC_{50} of *F. acuminatissima* stem and root**

<table>
<thead>
<tr>
<th>Extraction Techniques</th>
<th>IC_{50} (µg/ml) Stem</th>
<th>IC_{50} (µg/ml) Root</th>
</tr>
</thead>
<tbody>
<tr>
<td>Maceration</td>
<td>62.83 ± 0.577^{c}</td>
<td>74.17 ± 0.577^{a}</td>
</tr>
<tr>
<td>Orbital Shaker Assisted (OAE)</td>
<td>82.00 ± 0.500^{d}</td>
<td>81.00 ± 0.500^{c}</td>
</tr>
<tr>
<td>Ultrasonic Assisted (UAE)</td>
<td>29.33 ± 0.289^{a}</td>
<td>80.17 ± 0.289^{d}</td>
</tr>
<tr>
<td>Microwave Assisted (MAE)</td>
<td>50.67 ± 0.577^{b}</td>
<td>82.51 ± 0.500^{f}</td>
</tr>
</tbody>
</table>

* The values are means ± standard deviation (n = 3). Values with different superscripts were significantly different by Duncan test at level of p< 0.05.

DPPH assay is being widely used in the determination of free radical scavenging activity of natural antioxidants, mainly due its conventional reproducibility and high sensitivity (Wijekoon et al., 2011). Ultrasonic assisted extraction does not have long extraction time compared to other parameters, as ultra-sonication could induce free radical’s formation within the liquid medium thus causing degradation and oxidation of the sample (Hemwimon et al., 2007).

The lower the value of IC_{50} indicated more strength of the antioxidant potential. According to the current and previous findings, the antioxidant activity of the extracts also varies from one to another (Alothman et al., 2009). There are two factors that affect the result collected by the antioxidant assays such, some phytochemicals available in the extracts may contain molecular weight
antioxidants or antioxidants bound to complex molecules and some phenolic compound might not possess the antioxidant strength (Gursoy et al., 2009).

**Correlation between TPC, TFC and DPPH Radical Scavenging assay**

Table 6 shows the correlation of TPC, TFC and DPPH radical scavenging activity assay analysed by using Pearson’s correlation coefficient test. Statistically, the correlations between all antioxidant content and activity were significantly different. From the table, TPC r-value indicated a positive and moderate correlation with DPPH, meanwhile, TFC showed a positive and strongly correlated with DPPH activity.

<table>
<thead>
<tr>
<th></th>
<th>DPPH Radical Scavenging Activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total Phenolic Content</td>
<td>0.478*</td>
</tr>
<tr>
<td>Total Flavonoid Content</td>
<td>0.841**</td>
</tr>
</tbody>
</table>

* Correlation is significant at the level p < 0.05.
** Correlation is significant at p < 0.01.

The medium correlation between TPC and DPPH assay was possibly caused by the present of non-phenolic compound, where Folin-Ciocalteu reagent also react with simple phenol which is not compelling antioxidant. Moreover, distinctive phenolic component may also show different antioxidant activity depending on their structure, with the present of other compound that included in the crude extract (Nićiforović et al., 2010).

The correlation coefficient between total phenolic and flavonoid content with DPPH assay showed moderate and slightly strong coefficient, respectively, which might be due to the instability of DPPH free radical. Previous research by Molyneux (2004) described that the stability of DPPH solution can last for 3 days. Nevertheless, this hypothesis was considered inaccurate due to the inconsistency of the results. Therefore, preparation of fresh DPPH-methanol stock solution before an experiment is recommended.

In brief, the ultrasonic assisted extraction process proved to produce higher amount of TPC and TFC for both stem and root part of *F. acuminatissima*. This is probably because during sonication, production of sound waves caused the generation of bubbles (pit), close to the tissue of the plant cell, cleaved and disrupt the cell walls, thus caused the solvent to penetrate and thereby releasing the contents of cells. Besides, sonication also accelerates the movement of the molecules, thus bringing together the molecules of solvent with those of the sample (Roidaki et al., 2015).

Microwave assisted extraction method is not a good option for extracting antioxidant compounds in *F. acuminatissima*, where high temperature may have caused enzymatic oxidation and led to the denaturation, or destruction of plant secondary metabolites (Khoddami et al., 2013). Besides, other operational conditions (e.g., high extraction pressure) of microwave assisted extraction may modify the chemical structures of the target compounds (Zhang et al., 2011). Acceleration of chemical reactions or changes of some target metabolites caused by microwave irradiation (Ghani et al., 2008).

**Conclusion**

This study demonstrated that different extraction techniques were able to reveal significant amount of antioxidant compounds and activities in *Fragrea acuminatissima*. Surprisingly, all findings on TPC and TFC showed that UAE had the highest antioxidant content. Antioxidant activities of stem sample also conveyed similar result. Hence, it can be concluded that UAE is a highly efficient technique not only good in extraction yield but also excellent in revealing the antioxidant capacities of the sample. In addition, recent findings also support the principles of this technique which capable to minimize the extraction time and solvent usage as compared to the conventional techniques. In contrast, despite of shorter extraction time and solvent usage, MAE was proved to be a poor method for crude extraction of *F. acuminatissima* stem and root, with respect to extraction efficiency and antioxidant capacity.

Since there are other factors that may influence the potential of antioxidant activity, further research on the other extraction condition of UAE extract such as solvent extracting power, duration of the extraction and compounds interaction need to be carried out. Moreover, natural product developers had shown increasing interest in water-based extract due to the safety reason in commercializing the plant. Furthermore, future investigation is needed to delve into the major or individual polyphenolic group and other bioactive compound present in *Fragrea acuminatissima* stem and root.

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