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Abstract
A new method of microwave assisted extraction (MAE) was used to extract antioxidant compounds from Plantago major L. Ultrasound-assisted extraction (UAE) and kinetic maceration were studied as comparison process. The effect of part of plant, duration and temperature on the free radical scavenger activity, total phenol and flavonoid content of the extract were determined. MAE was found to be the most effective method capable of yielding 1.72 %GAE total phenol, 1.94 %CE total flavonoid and giving EC50 equivalent to 474.11 ppm of crude drug with 70°C and 20 min of extraction temperature and time, respectively. The percent recovery of antioxidant compounds was found to increase with increasing MAE time and temperature. It was also found that antioxidant compounds of the leaf was much greater than that of root and petiole.

Keywords: microwave assisted extraction, total phenol, total flavonoid, antioxidant, Plantago major

INTRODUCTION
Plantago major L. (daun sendok) has been used by people in various countries for the treatment of skin diseases, infections, problems related to the digestive organs, respiratory, reproductive, circulatory, against tumors, eliminate pain and reduce fever. This herb is also reported to have antioxidant activity and free radical scavenger, although lower than in tea. In addition to 15 kinds of flavonoids, this plant also contains terpenoids, alkaloids, derivatives of caffeic acid, iridoid glycosides, polysaccharides, fats, vitamins and organic acids (Samuelsen, 2000). Flavonoids is estimated as the main antioxidant components from these plants. Consequently, it is important that effective methods for extracting these components from P. major be developed.

The availability of standardized plant extracts is important to commercial production. For extracts preparation, selection of appropriate extraction method is a key consideration. Presently, a number of extraction methods including kinetic maceration, UAE and MAE are available (Zhang et al., 2005). However, comparative studies of these methods, especially for the extraction of antioxidant components from P. major have not been attempted previously.

The purpose of the present study is to examine different extraction methods for P. major and compare them relative merits. The experiments and conclusion are given in the following section.

METHODS
Material, chemical and instrumentation
Fresh herbs of P. major were purchased from Purwodadi Botanical Garden, Pasuruan, Indonesia. Standards for gallic acid and (+)-catechin hydrate were obtained from Sigma-Aldrich. Other chemicals including DPPH (Sigma-Aldrich), Folin Ciocalteu (Merck),
\[ \text{AlCl}_3 \text{ (Merck), NaOH (Merck), NaNO}_2 \text{ (Merck), Na}_2\text{CO}_3 \text{ (Merck) and demineralized water (FF, UBAYA). Instruments including microwave oven (2450 MHz, 700 W, from Sharp), ultrasonic bath (Branson 2000), magnetic stirrer (Ceramig Midi IKA® Works), spectrophotometer (Cintra), analytical balance (Sartorius BL210s), blender (Hitachi), stopwatch (Alba), sieve (30 mesh), vortex mixer (Thermolyte), and laboratory glassware equipment.} \]

**Preparation of plant materials and kinetic maceration**

Fresh herbs of *P. major* were harvested, washed, and separated into roots, leaves and petioles. These three parts of plant then called as “sample”. These samples then dried under indirect sunlight. Dried materials were ground to an average size of 30 mesh in diameter. The ground samples were kept in a dry place until use. A conventional solvent extraction method was used in this study: kinetic maceration. In this method, two grams of sample was extracted in 70 ml of water in a 250 ml beaker glass at room temperature using magnetic stirrer (360 rpm, 20 minutes). The extract was then filtered into a 100 ml volumetric flask. The sample residue after extraction was washed in approximately 30 ml of water, and lastly fixed with water until 100.0 ml. The antioxidant activity, concentration of total phenol and flavonoid were measured by a spectrophotometer.

**UAE**

For the UAE experiments, an ultrasonic bath was used as an ultrasound source. The bath was basically a rectangular container (15 cm x 15 cm x 15 cm), to which 47 kHz ± 6% transducer was annealed at the bottom, and the bath power rating was 270W. The extraction of antioxidant compounds were performed by adding two gram of sample into 70 ml of water in a 250 ml beaker glass. The beaker glass was then partially immersed into the ultrasonic bath, which contains 2.2 litre of water. The bottom of the beaker glass was approximately five cm above the bottom of the bath. The solvent surface in the beaker glass was kept at the level of the water in the ultrasonic bath. Extraction was conducted for up to 20 min at room temperature. The extract was then filtered into a 100 ml volumetric flask. The sample residue after extraction was washed in approximately 30 ml of water, and lastly fixed with water until 100.0 ml. The antioxidant activity, concentration of total phenol and flavonoid were measured by a spectrophotometer.

**MAE**

MAE experiments were performed on a microwave oven. The extraction of antioxidant compounds was conducted by adding two gram of sample into 70 ml of water in beaker glass. The beaker glass were placed symmetrically in the microwave field. Experiments were carried out to determine the effect of irradiation time (5, 10, 20, 30 min) and temperature (40, 70, 100°C) on MAE efficiency. After microwave irradiation, the solution was filtered into a 100 ml volumetric flask. The sample residue after extraction was washed in approximately 30 ml of water, and lastly fixed with water until 100.0 ml. The antioxidant activity, concentration of total phenol and flavonoid were measured by a spectrophotometer.

**Measurement of total phenol concentration**

The total phenol content of extracts were determined by using the Folin-Ciocalteu assay (Kahkonen et al., 1999). An aliquot (one ml) of extracts or standard solution of gallic acid (60, 80, 90, 100 and 110 mg/l) was added to 10 ml volumetric flask, containing four ml of demineralized water. 0.2 ml of Folin-Ciocalteu phenol reagent was added to the mixture and shaken. After five min, 0.2 ml of 15%
Na$_2$CO$_3$ solution was added to the mixture and fixed with addition of water until 10.0 ml. After incubation for 15 minutes at room temperature, the absorbance against prepared reagent blank was determined at 706 nm with an UV-Vis spectrophotometer. The total phenol content of extract was expressed as gram gallic acid equivalent (GAE)/100 gram of dry weight. All samples were analysed in five times replication, each in duplicates.

Measurement of total flavonoid concentration
Total flavonoid content measured by the aluminum chloride colorimetric assay (Marinova et al., 2005). An aliquot (one ml) of extracts or standard solution of catechin (12, 15, 18, 21 and 24 mg/l) was added to 10 ml volumetric flask containing four ml of water. To the flask was added 0.3 ml 5% NaNO2. After five min, 0.3 ml 10% AlCl$_3$ was added. At 6$^{th}$ min, two ml 1 M NaOH was added and the total volume was made up to 10.0 ml with water. The solution was mixed well and absorbance was measured against prepared reagent blank at 503 nm. Total flavonoid content of extract was expressed as gram catechin equivalent (CE)/100 gram of dry mass. All samples were analysed in five times replication, each in duplicates.

Antioxidant activity measurement
Antioxidant activities of $P$. major extracts obtained using UAE, MAE, and kinetic maceration were tested and compared by measuring the ability of the extracts to scavenge the free radical DPPH (1,1-diphenyl-2-picrylhydrazyl) in vitro. The assay method was modified from that described in Wu and Ng (2008). For the purpose of comparing the antioxidant activity in various extracts, concentration of sample producing 50% reduction of the radical absorbance (EC$_{50}$) was used as an index. To find this value, the extract was diluted in series with water and 1.5 ml of each diluted extract was added to 3.0 ml of 0.004% DDPH solution. The solutions were mixed using a vortex mixer and the mixture was then incubated for 10 minutes in darkness at room temperature, after which the absorbance was measured at the wavelength of 520 nm using water and ethanol as a reference. EC$_{50}$ can be found from a plot of percent inhibition (PI) versus the concentration of extract. The values of PI can be calculated using the following equation:

$$\text{PI} (%) = (1 – \frac{A_t}{A_r}) \times 100$$

in which $A_t$ and $A_r$ are absorbance of test sample and absorbance of the DPPH reference, respectively.

RESULTS AND DISCUSSION
Comparison of extraction methods
Fig.1 shows the extraction profiles for MAE in comparison with other extraction methods such as kinetic maceration and UAE. The yield of total phenol and flavonoid also antioxidant activity from MAE is higher than UAE and kinetic maceration. When compared with kinetic maceration conducted at room temperature, MAE gave considerably higher yields due to the heating effect. More importantly, microwave heating occurred at much faster rate. Heating is known to affect the morphological changes in the plant sample matrix and thus enhances product mass transfer, faster heating in MAE should therefore be responsible for increased mass transfer, and thus antioxidant compounds release rate observed. Thus, in MAE, the effects of higher heating and mass transfer rates synergistically increase the rate of antioxidant compounds extraction. When MAE was compared with UAE at room temperature in which mass transfer was enhanced by cavitation effect, MAE yet resulted in higher of extraction. Although UAE did not require a long extraction time, it is commonly known that ultrasonic could
induce free radicals formation within the liquid medium, causing oxidation and degradation of the phenol, including that of flavonoid (Hemwimon, 2007).

![Figure 1. EC₅₀, total phenol and total flavonoid content in *P. major* extract resulted from Kinetic Maceration (MK), UAE and MAE](image1)

**Comparison of parts of plant**

A certain part of the plant has higher levels of chemical compounds than those of the others. Among others, depending on how and where the metabolism of these compounds. Flavonoids are found in all part of plant including leaves, roots, wood, bark, pollen, nectar, flowers, fruit and seed. Higher concentrations are in the leaves and barks compared with other parts. There is estimated about 2% of all carbon photosynthesized by plant or approximately 1x10⁹ tons/year converted into flavonoids (Markham, 1988). Mostly photosynthetic process occurs in green (contain chlorophyll) leaves. Therefore, this experiment has showed the water extract of *P. major* leaves has higher level of total flavonoids, total phenols, and antioxidant activity than the roots and petioles (Fig. 2).

![Figure 2. EC₅₀, total phenol and total flavonoid content in *P. major* extract resulted from leaves, roots and petioles by MAE](image2)
**Effect of extraction temperatures**

Generally, the higher extracting temperature is profitable for extraction due to the increased solubility. In a closed microwave vessel used in this study, the temperature of the solvent could be increased above the boiling point temperature. As a result, the solubility of phenol and flavonoid could greatly be enhanced. The effect of extraction temperature is shown in Fig. 3 which clearly demonstrates that increasing the temperature of the solvent from 40 to 100°C significantly increases the extraction efficiency. This is because higher temperature causes intermolecular interactions within the solvent to decrease, giving rise to higher molecular motion, and causing the solubility to increase. The increasing temperature may also cause opening cell matrix, and as a result, phenol and flavonoid availability for extraction increases. Moreover, at high temperature, solvent viscosity decreases and the diffusivity increases, thus the efficiency of extraction increases (Pan et al., 2000; Camel, 2000).

![Figure 3. Effect of extraction temperature of MAE](image)

**Effect of duration of extraction**

Generally, with increasing duration of extraction, the number of dissolved compounds will increase and moreover, pharmacological activity of the extract will increase too. If the compound is thermostable or does not evaportate, with increasing duration of heat the number of compounds dissolved will increase and more until reach a state where the compounds cannot be extracted again. In such conditions, if the extraction time increased, the amount of dissolved compounds would not be increased, thus pharmacological activities will remain and this position is the optimal conditions. The effect of duration of extraction is shown in Fig. 4 which clearly demonstrates that increasing the duration of extraction from five min to 30 min significantly increases the extraction efficiency, except on phenol content. This is probably caused presence thermolabile phenolic compounds (besides flavonoid) in the extract.
CONCLUSIONS
MAE of *P. major* leaves gives the highest yields of antioxidant compounds. The appropriate condition for maximum antioxidant components with MAE was at the extraction temperature of 100°C and extraction times of 30 min. These results demonstrate the new MAE potential to extract the antioxidant compounds from the leaves of *P. major*.

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