Air-drying temperature changes the content of the phenolic acids and flavonols in white mulberry (Morus alba L.) leaves

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ABSTRACT: The white mulberry leaves are typically available on the market in dried or encapsulated form. It was assumed in the study that appropriate drying of leaves of the white mulberry is significant for obtaining intermediate products with high content of compounds having anti-oxidative activity. The purpose of the study was to determine the influence of the temperature of mulberry leaves air drying on the content of phenolic acids and flavonols. It has been determined that the content of these compounds in the leaves depended on the drying temperature. Drying at 60 °C favored release of phenolic acids and flavonols from complexes and/or formation of new compounds. Their total content was 22% higher than in leaves dried at 30 °C. Drying at 90 °C reduced the phenolic acid and flavonol content by 24%. The most favorable drying temperature was 60 °C.

Key words: white mulberry leaves, air-drying temperature, phenolic acids, flavonols, Morus alba L.

The presence of polyphenols in the diet has considerable importance for maintaining homeostasis of the organism and in the prophylaxis of lifestyle diseases. These non-nutritive compounds include phenolic acids (PhA) and flavonols (Fla), and plants such as white mulberry constitute their source. PhA and Fla are typically reported in a low-molecular, esterified or etherified form with polymers of cellular walls (CHAN et al., 2016; KOBUS-CISOWSKA et al., 2019).

Therapeutic properties of white mulberry leaves (WML) have previously been utilized in the Far East Medicine. Presently, apart from gardening and forestry, they have been applied in food production (PRZEOR & FLACZYK, 2016; ZOU, 2015), as they are non-toxic to the human organism (KUJAWSKA et al., 2016). Positive outcomes of WML consumption have been demonstrated with regard to, among others, diabetes, hyperlipidemia, eye and skin diseases, obesity, atherosclerosis, liver cancer (BUTT et al., 2008; IQBAL et al., 2012; QIN et al., 2013).

The WML are typically available on the market in dried or encapsulated form. Adaptation of the suitable technique and drying temperature of the plant material influences their final anti-oxidative activity (LEMUS-MONDACA et al., 2018; PRZEOR & FLACZYK, 2011).

For the present study it was assumed that suitable processing of white mulberry leaves (Morus alba L.) Polish var. zolwinska wielkolistna

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(WML-P) by means of leaves drying, for the purpose of obtaining the highest possible number of bioactive compounds from them, such as PhA and Fla, is of key significance for the high health-promoting quality of the leaf intermediate products, which may be utilized in the food technology or pharmacy.

In the study white mulberry (Morus alba L.) leaves Polish var. zolwinska wielkolistna (WML-P), picked at the tree farm in Pętkowo, near Poznan, were used to prepare extracts from WML-P (WML-Pe). Unified material was dried in a convection oven (Rational CCC61/02) at the following temperatures: 30 °C (30AD), 60 °C (60AD) or 90 °C (90AD). In the experiment only 3 temperatures were tested to find a preliminary range of the best temperatures for high PhA and Fla content in WML-P. The drying process of WML-P was performed by air drying. HOSSAIN et al. (2010) demonstrated this method of drying to be the most favorable for herbs in terms of retaining anti-oxidative properties. Moreover that type of drying is the most commonly used method of drying herbs in Poland; therefore, it was used in described WML-P production process. Whole, dried leaves were disintegrated in laboratory mill (Retsch GM200, Germany). The powder (10 g) was extracted with distilled water twice (100 ml, 40 ml), each time for 5 minutes and filtered. Extraction method and conditions were based on previous own experiments with WML-P. To protect the samples for further analyzes, extracts from WML-P (WML-Pe) were dried by means of lyophilization (Christ Alpha 1-4LSC) and stored in plastic bags in the dark.

Extracts from WML-P (0.4%) (WML-Pe) were injected (10µl) into the Zorbax SB C18 (3,9x150mmx5µm, Agilent Technology, USA) chromatographic column in triplicate. Qualitative and quantitative determination of phenolic acids (PhA) and flavonoids (Fla) was determined with HPLC/DAD (Infinity 1290, Agilent Technology, USA) according to (KOBUS et al., 2009; SIGER et al., 2004). The mobile phases were H₃PO₄ (pH=2.7) and C₆H₅N (50%), and flow rate was 1,5 ml/min¹. Detection was done based on retention time and UV spectra of standard phenolic acids (Sigma Aldrich): gallic acid (GAL), protoacetucic acid (PRO), 4-hydroxybenzoic acid (HYD), vanillic acid (VAN), chlorogenic acid (CHL), caffeic acid (CAF), p-coumaric acid (CUM), ferulic acid (FER), sinapic acid (SIN), and flavonoids: rutin (RUT), isoquercitrin (ISQ), astragalin (AST), myricetin (MYR), quercetin (QUE), kaempferol (KEM), at 250nm and 310nm. Amounts of compounds were determined based on standard curves. Data were analyzed using one-way analysis of variance (ANOVA), followed by Tukey’s post-hoc test using Statistica Software, version 13 (StatSoft, Poland). Statistical differences were calculated at the significance level p<0.05 and represented by superscript letters. Analysis were made triplicate.

Benzoic and cinnamic acid derivatives were determined in WML-Pe (Table 1). Among the studied PhA, the following compounds were predominant: CHL and CAF, while among Fla: RUT, similarly to Tunisian and Spanish leaves (SÁNCHEZ-SALCEDO et al., 2015; THABTI et al., 2012). The study of HUNYADI et al. (2012) has already demonstrated that CHL and RUT are responsible for the antidiabetic properties of white mulberry leaves. In the course of the experiment, the trials were not hydrolyzed, that may have caused MYR, QUE, KEM were not detected in any samples as they remained in the bound form.

The CAF content increased by 64%, while the content of CHL by 16% as the effect of 60AD, compared to 30AD. CHL is subject to disintegration to i.a. caffeic and quinic acid as a result of technological procedures (RADOJKOVIC et al., 2016), which may explain the considerable increase of CAF observed in the study for 60AD samples. At the same time, an increase of the PhA and Fla content after 60AD application indicates the disintegration of more complex polyphenol complexes. KATSUBE et al. (2009) noted that in drying at 60°C certain polyphenols are synthesized, which influences the increase of antioxidative activity. The content of the remaining PhA was also subject to total increase by ~16% (60AD). Moreover, we have already described that the temperature of 60 °C gave the best results in antioxidants test in that kind of plant material (PRZEOR & FLACZYK, 2011). In 90AD samples, we observed a reduction of the discussed PhA by 8% (CHL) and 25% (CAF) relative to 30AD samples.

In the case of Fla only for ISQ and AST significantly higher values in 60AD samples were determined, by 17% and 41%, respectively. Drying at 90 °C resulted in reduced ISQ and AST content relative to 60AD samples. In temperature as high as 90 °C, certain PhA and Fla with antioxidative activity could have decomposed, similarly to KATSUBE et al. (2009).

To sum up, white mulberry leaf drying temperature influenced the PhA and Fla content in their aqueous extracts. The same PhA and Fla were predominant in the white mulberry var. zolwinska wielkolistna as in the leaves collected in other climatic zones. Air drying at 60 °C favored the release of PhA and Fla from compound complexes and/or formation.

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Table 1 - Phenolic Acids (PhA) and flavonols (Fla) content in WML-Pe depending on drying temperature.

| mg per g of extract | 30°C (30AD) | 60°C (60AD) | 90°C (90AD) |
|---------------------|-------------|-------------|-------------|
| Phenolic Acids (PhA) |             |             |             |
| GAL     | 0.018 ± 0.00 | 0.019 ± 0.01 | 0.018 ± 0.01 |
| PRO     | 0.156 ± 0.01 | 0.166 ± 0.01 | 0.165 ± 0.01 |
| HYD     | 0.111 ± 0.01 | 0.112 ± 0.00 | 0.135 ± 0.02 |
| VAN     | 0.761 ± 0.00 | 0.771 ± 0.01 | 0.647 ± 0.015 |
| CHL     | 4.624 ± 0.01 | 5.218 ± 0.01 | 5.270 ± 0.08 |
| CAF     | 2.540 ± 0.09 | 4.175 ± 0.01 | 1.901 ± 0.02 |
| CUM     | 0.271 ± 0.04 | 0.424 ± 0.043| 0.253 ± 0.08 |
| FER     | 0.391 ± 0.05 | 0.458 ± 0.028| 0.345 ± 0.018|
| SIN     | 0.214 ± 0.01 | 0.377 ± 0.007| 0.405 ± 0.10 |
| Flavonols (Fla) |             |             |             |
| RUT     | 2.598 ± 0.06 | 2.413 ± 0.13 | 2.377 ± 0.09 |
| ISQ     | 1.133 ± 0.08 | 1.323 ± 0.052| 1.260 ± 0.01 |
| AST     | 0.768 ± 0.15 | 1.081 ± 0.045| 0.867 ± 0.02 |
| MYR     | 0.000 ± 0.00 | 0.000 ± 0.00 | 0.000 ± 0.00 |
| QUE     | 0.000 ± 0.00 | 0.000 ± 0.00 | 0.000 ± 0.00 |
| KEM     | 0.000 ± 0.00 | 0.000 ± 0.00 | 0.000 ± 0.00 |

a, b, c – different letters show statistically significant differences in Tukey’s test, p<0.05.

WML-Pe – extracts from white mulberry (Morus alba L.) leaves Polish var. zolwinska wielkolistna; AD – air-drying temperature; GAL – gallic acid; PRO – protocatechuic acid; HYD – 4-hydroxybenzoic acid; VAN – vanillic acid; CHL – chlorogenic acid; CAF – caffeic acid; CUM – p-coumaric acid; FER – ferulic acid; SIN – sinapic acid; RUT – rutin; ISQ – isoquercitrin; AST – astragalin; MYR – myricetin; QUE – quercetin; KEM – kaempferol.

DECLARATION OF CONFLICT OF INTERESTS

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

AUTHORS’ CONTRIBUTIONS

The authors contributed equally to the manuscript.

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