SPECTROPHOTOMETRIC EVALUATION OF FLAVONOIDS, PHENOLCARBOXYLIC ACIDS AND TOTAL PHENOLIC CONTENTS OF SEVERAL INDIGENOUS HERBAL PRODUCTS WITH POTENTIAL HEPATOPROTECTIVE EFFECT

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Abstract
Chronic liver diseases (CLD) are leading causes of morbidity worldwide. Current evidence highlights the beneficial properties associated to polyphenols in CLD, due to their antioxidant activity. Therefore, the aim of our study was the evaluation of phenolic content of several indigenous herbal products (agrimony, cichory and dandelion aerial parts; artichoke and rosemary leaves), in view of obtaining a phytomedicine with hepatoprotective effect. Flavonoids, phenolcarboxylic acids and total phenolic contents were assessed by means of spectrophotometric methods using different solvents (70% ethanol, 50% ethanol, 20% ethanol and water). Agrimony aerial parts and rosemary leaves showed the highest flavonoids (1.17 g rutin/100 g dried herbal product and 1.27 g ru tin/100 g dried herbal product) and phenolcarboxylic acids contents (6.55 g chlorogenic acid/100 g dried herbal product and 5.67 g chlorogenic acid/100 g dried herbal product). Moreover, according to our results, 50% ethanol provided the optimal extraction of phenolic compounds for all analysed herbal products, except for dandelion aerial parts, for which 20% ethanol was the best extraction solvent. Analysed herbal products are a source of phenolic compounds. Nevertheless, the solvent is a key factor that greatly influences the phenolic content.

Rezumat
Afeziunile hepatice de natură cronică reprezintă o cauză majoră de deces la nivel mondial. Cercetări recente au evidențiat rolul benefic al polifenolilor în tratamentul acestor afecțiuni, datorită activității lor antioxidant. Scopul lucrării a constat în evaluarea conținutului de polifenoli a unor materii prime vegetale (părți aeriene de cicoare, păpădie, turță mare, frunze de anghinare, de rozmarin), în vederea obținerii unui fitopreparat asociat în tratamentul hepatopatiilor. Pentru evaluarea conținutului de polifenoli totali, acizi fenolcarboxilici și flavone s-au utilizat metode spectrofotometrice. Determinările au fost realizate utilizând solvenți diferiți utilizând etanolul de concentrație 50% (cu excepția părților aeriene de turță mare și frunzele de rozmarin) avut cel mai mare conținut de flavone (1,17 g rutin/100 g produs vegetal uscat și 1,27 g rutin/100 g produs vegetal uscat) și de acizi fenolcarboxilici (6,55 g acid clorogenic/100 g produs vegetal uscat și 5,67 g acid clorogenic/100 g produs vegetal uscat). Cea mai mare cantitate de principii active s-a obținut utilizând ca solvent de extracție etanolul de concentrație 50% (cu excepția părților aeriene de turță mare). Produsele analizate reprezintă un sursă importantă de polifenoli. Totuși, solventul reprezintă un factor cheie care influențează modul de extracție a principiilor active de interes.

Keywords: agrimony, total phenolic content, liver diseases, oxidative stress

Introduction
Chronic liver diseases (CLD) such as non-alcoholic fatty liver disease, hepatitis, cirrhosis or hepatocellular carcinoma are leading causes of morbidity worldwide [13, 17]. The main aetiologies of liver diseases are alcohol abuse, hepatitis virus infections and metabolic syndrome. The most important pathological processes of liver diseases are inflammation, immune response disruption, alteration of gut-microbiota axis, alteration of gut-liver axis and oxidative stress [5, 8, 13, 17, 21, 31]. It is well known that oxidative stress represents a hallmark of liver injury [5, 8, 13, 15, 31], since it is involved in: (i) inflammation and apoptosis [17], (ii) alteration of the mitochondrial membrane permeability and transition potential which initiates the release of pro-apoptotic factors [17], (iii) alteration of lipids, proteins and DNA contents, which further leads to steatosis/fibrosis [17]. Moreover oxidative stress is involved in modulation of several pathways that regulate gene transcription, protein expression and the immune response [5, 17].

Current evidence highlights the beneficial properties associated to polyphenols for chronic liver diseases treatment [2, 3, 22, 27]. Several mechanisms are involved in the hepatoprotective effect of polyphenols:
(i) scavenger activity of free radicals; (ii) inhibiting the formation and expression of inflammatory cytokine [17, 25]; (iii) regulation of gene-expression or signalling pathways - up-regulation of Bcl-2 (B-cell lymphoma 2) apoptotic pathway, activation of Nrf2 - nuclear factor erythroid 2-related factor 2 mediated pathway, suppressing of canonical NF-kB and PI3K/AKT (phosphatidylinositol-3-kinase/protein kinase B) signalling pathways etc.; (iv) inhibition of cyclooxygenase and iNOS (nitric oxide synthase); (v) down-regulation of lipogenic genes (fatty acid synthase, acetyl-CoA carboxylase); (vi) up-regulation of PPARα (peroxisome proliferator-activated) receptors and (vii) reduced serum fibrosis markers [11, 17, 25].

Considering the scientific data, the aim of our paper was the evaluation of phenolic content of several indigenous herbal products - agrimony aerial parts (Agrimoniae herba), common cichory aerial parts (Cichorii herba), rosemary leaves (Rosmarini folium), dandelion aerial parts (Taraxaci herba) and artichoke leaves (Cynarae folium), in view of obtaining a phytomedicine with hepatoprotective effect. Indigenous herbal products were selected based on their chemical composition, rich in phenolic compounds [7, 9, 10, 12, 14].

### Materials and Methods

#### Plant Material

Herbal products (agrimony aerial parts - encoded A, common cichory aerial parts - encoded CH, dandelion aerial parts - encoded T, artichoke leaves - encoded CY and rosemary leaves - encoded R) were acquired in 2019 (as medicinal tea), from indigenous manufacturers.

#### Reagents and solvents

All chemicals (aluminium chloride, chlorogenic acid, ethanol, hydrochloric acid, rutin, sodium hydroxide, sodium nitrite, tannic acid, sodium carbonate) were purchased from Sigma-Aldrich (Germany).

#### Obtaining the extractive solutions

For evaluation of the total phenolic, flavonoids and phenolcarboxylic acids contents, 1 g of each herbal product was heated twice with 25 mL solvent (70% alcohol, 50% alcohol, 20% alcohol and water) on a heated condenser for 30 min. After cooling, the solutions were filtered in a 50 mL volumetric flask and filled to mark with the same solvent. Analysed solutions were encoded as follows (Table I).

#### Table I

| Herbal product       | Solvent 70% ethanol | Solvent 50% ethanol | Solvent 20% ethanol | Solvent water |
|----------------------|---------------------|---------------------|---------------------|--------------|
| Agrimony aerial parts| SA70                | SA50                | SA20                | SAA          |
| Common cichory aerial parts | SCHA               | SCHA                | SCHA                | SCHA         |
| Artichoke leaves     | SCT70               | SCT50               | SCT20               | SCYA         |
| Rosemary leaves      | SRT70               | SRT50               | SRT20               | SRA          |

These analyses were necessary, thereby to choose the solvent, that provides the best extraction yield of active substances. The optimum solvent will be further used for obtaining freeze-dried extracts, rich in phenolic compounds.

#### Methods

**Spectrophotometric assays.** Total phenolic content (TPC) was determined with the Folin-Ciocalteu reagent as previously described [14, 19, 30]. Results were expressed as g tannic acid/100 g dried herbal product, based on a calibration curve (2.04 - 9.18 µg/mL, \( R^2 = 0.9994 \), \( n = 8 \)). The flavonoids content (FL) was determined based on the chelating reaction with aluminium chloride and results were expressed as g rutin/100 g dried herbal product using the following calibration curve (5.0 - 35.0 µg/mL, \( R^2 = 0.9998 \), \( n = 11 \)) [14, 19]. Phenolcarboxylic acids (PCAs) were evaluated based on formation of oximes in the presence of sodium nitrite/hydrochloric acid and sodium hydroxide. Results were expressed as g chlorogenic acid/100 g dried herbal product (chlorogenic acid calibration curve - 11.3-52.7 µg/mL, \( R^2 = 0.9998 \), \( n = 6 \)) [14, 19]. For all spectrophotometric assays, a Jasco V-530 spectrophotometer (Jasco, Japan) was used.

### Statistical analysis

For each herbal product, three samples were analysed and all assays were carried out in triplicate (\( n = 3 \)). The results were expressed as mean ± SD (standard deviation). Standard deviation was determined using Microsoft Office programme (Excell, 2010). Statistical analysis was performed using the open source software R [24]. Since the samples related to our study were too small for a classical approach, we elaborated robust measures for mean and standard deviation and we applied a robust ANOVA version for comparing our datasets [20]. We used a similar parametric framework, but with a bootstrap approach, without worrying about basic violation of normality, homoscedasticity and sphericity [4]. Statistical significance was accepted for alpha-level 0.05 and post hoc analysis for a Bonferroni adjusted alpha level. The factors that affected the continuous variable concentration were a focal variable denoted with Type (with five levels: Agrimoniae herba, Cichorii herba, Cynarae folium, Rosmarini folium and Taraxaci herba) and other two moderator variables: the solvent (with four levels: ethanol 20%, ethanol 50%, ethanol 70% and water) and the active substance (with three levels: PCAs, FL and TPC). We computed
a three-way robust ANOVA to evaluate how each independent factor (herbal product, solvent and active substance) interact with a continuous variable denoted by concentration. There was a statistically significant three-way interaction between the herbal product, the solvent and the active substance (p = 0.001) and a simple two-way post hoc interaction for all pairwise of factors implied (p = 0.001).

Results and Discussion

The results for our spectrophotometric assays are presented in Table II.

Table II

Spectrophotometric results for analysed herbal products

| FL (g rutin/100 g dried herbal product) | SA70 | SA50 | SA20 | SAA |
|----------------------------------------|------|------|------|-----|
| 1.1745 ± 0.0293                        | 1.0332 ± 0.0836 | 0.9447 ± 0.1368 | 0.6453 ± 0.0496 |
| SCH70                                  | 0.4771 ± 0.0348 | 0.5596 ± 0.0482 | 0.4350 ± 0.0116 | 0.3814 ± 0.0312 |
| SCY70                                  | 0.7382 ± 0.0590 | 0.7497 ± 0.0244 | 0.4361 ± 0.0476 | 0.4121 ± 0.0277 |
| ST70                                   | 1.0271 ± 0.1130 | 1.2771 ± 0.0684 | 0.7991 ± 0.1469 | 0.1156 ± 0.0099 |
| PCAs                                   | 0.6737 ± 0.0192 | 0.6497 ± 0.1479 | 0.9154 ± 0.1138 | 0.7808 ± 0.1655 |

| TPC (g tannic acid/100 g dried herbal product) | SA70 | SA50 | SA20 | SAA |
|-----------------------------------------------|------|------|------|-----|
| 4.1655 ± 0.2108                               | 4.7469 ± 0.1683 | 6.5580 ± 0.4671 | 3.1842 ± 0.1978 |
| SCH70                                         | 1.8201 ± 0.0503 | 1.4614 ± 0.0766 | 1.4840 ± 0.0273 | 1.4974 ± 0.1054 |
| SCY70                                         | 0.6572 ± 0.1201 | 0.5643 ± 0.0401 | 0.6170 ± 0.0697 | 0.3802 ± 0.0428 |
| ST70                                          | 5.6123 ± 0.2365 | 5.6733 ± 0.7740 | 4.9732 ± 0.4402 | 0.8151 ± 0.0582 |
| TPC                                           | 2.0383 ± 0.1333 | 2.1674 ± 0.1824 | 2.6529 ± 0.0438 | 2.8350 ± 0.0867 |

Legend: FL - flavonoids, PCAs - phenolcarboxylic acids, TPC - total phenolic content, SA - agrimony aerial parts extractive solutions (SA70 - using 70% ethanol, SA50 - using 50% ethanol, SA20 - using 20% ethanol, SAA - using water); SCH - chicory aerial parts extractive solutions (SCH70 - using 70% ethanol, SCH50 - using 50% ethanol, SCH20 - using 20% ethanol, SCHA - using water); SCY - artichoke leaves extractive solutions (SCY70 - using 70% ethanol, SCY50 - using 50% ethanol, SCY20 - using 20% ethanol, SCYA - using water); SR - rosemary leaves extractive solutions (SR70 - using 70% ethanol, SR50 - using 50% ethanol, SR20 - using 20% ethanol, SRA - using water); ST - dandelion aerial parts extractive solutions (ST70 - using 70% ethanol, ST50 - using 50% ethanol, ST20 - using 20% ethanol, STA - using water).

According to our results (Table II), all analysed herbal products are a source of flavonoids and phenolcarboxylic acids. Agrimony aerial parts and rosemary leaves showed the highest flavonoids (1.17 g rutin/100 g dried herbal product and 1.27 g rutin/100 g dried herbal product) and phenolcarboxylic acids contents (6.55 g chlorogenic acid/100 g dried herbal product and 5.67 g chlorogenic acid/100 g dried herbal product). According to previous published papers, agrimony aerial parts are a rich source of flavonoids and hydroxycinnamic acids [29]. Our results regarding the flavonoids content of Agrimoniae herba are much lower compared to Ciobanu N et al. [9] that found 3.72 g flavonoids (expressed as rutin equivalents)/100 g dried herbal product, using 70% ethanol as extraction solvent [9]. For Agrimoniae herba, there were no statistically significant mean differences between all solvents in case of flavonoids, however regarding the total phenolic content, water showed a different behaviour. Still, there was a statistically significant simple main effect of all active substances for agrimony aerial parts (p < 0.001) (Figure 1a).

According to the scientific literature, chicory aerial parts are also an important source of phenolcarboxylic acids and flavonoids [1, 12]. Regarding chicory aerial parts total phenolic content, it was difficult to compare...
our results with other reports, that found 14 - 23 g polyphenols/100 g dried herbal product (expressed as tannic acid equivalents), since these results were obtained using a 70% hydroalcoholic dry extract [12]. Still, our results are much lower compared to Sahan Y et al. that found 3.3 - 8.8 g polyphenols/100 g dried herbal product (expressed as tannic acid equivalents) [26]. Regarding Cichorii herba, there was a statistically significant simple main effect of all active substances (p < 0.001) and solvents (p < 0.001). A different behaviour was observed for 70% ethanol regarding the PCAs content (Figure 1b).

For artichoke leaves (Cynarae folium) we did not find a statistically significant mean difference between PCAs and flavonoids (p = 0.23). Moreover, according to our results, we have not found a statistically significant mean difference between 50% ethanol and 70% ethanol (p = 0.21) (Figure 2a). Still, our results regarding the flavonoids (0.41 g rutin/100 g dried herbal product) content (using water as extraction solvent) are much lower compared to other authors, that found 1.99 g flavonoids/100 g dried herbal product (expressed as rutin equivalents) [7]. We assume that these differences are the consequence of the herbal product origin, pedoclimatic conditions and extraction methods.

Figure 1.
Statistical analysis (a two-way interaction boxplot) for a – Agrimoniae herba (agrimony aerial parts), b – Cichorii herba (common cichory aerial parts)

Figure 2.
Statistical analysis (a two- way interaction boxplot) for a – Cynarae folium, b – Rosmarini folium, c – Taraxaci herba
Our results regarding the total phenolic and flavonoids contents of rosemary leaves (using 50% ethanol as extraction solvent) (8.10 g tannic acid/100 g dried herbal product and 1.27 g rutin/100 g dried herbal product) are similar to our previous published paper (7.88 g tannic acid/100 g dried herbal product and 0.90 g rutin/100 g dried herbal product respectively) [14]. In case of rosemary leaves, there was a statistically significant simple main effect for all active substances (p < 0.001) and all solvents (p < 0.001). According to our results statistically significant mean differences were found regarding PCAs and flavonoids contents, when using hydroalcoholic solutions compared to water (p < 0.001). However, regarding the total phenolic content, we did not find statistically significant results, when using 70% hydroalcoholic and aequous solutions (Figure 2b).

In case of dandelion aerial parts, there was a statistically significant interaction between active substances and solvent variables (p < 0.002). Significant differences were found regarding the flavonoids and total phenolic contents (except for 20% ethanol in case of TPC) (Figure 2c). Our results (Table II) have shown that dandelion aerial parts are a rich source of flavonoids and phenolcarboxylic acids, which is in agreement with previous published reports [10, 16, 18]. Taking into account our results (Table II), 50% ethanol provided the optimal extraction of phenolic compounds for all analysed herbal products, except for dandelion aerial parts, for which 20% ethanol was the best extraction solvent.

It is well known that different extraction solvents (methanol, ethanol, mixtures of ethanol/methanol with water, mixture of acetone with water etc.) are used for evaluation of phenolic compounds in herbal products [23]. The efficacy of a certain solvent depends on its ability to dissolve specific phenolic groups; in addition the solvent also influences the plants cell permeability [23]. All-over, flavonoids present a high solubility in alcohol and a higher extraction yield as the content of ethanol in water increases from 50% ethanol and upward [6, 23]. Ethanol - in - water solutions (50%) are more effective than pure water or alcohol for hydroxyccinnamic acids extraction [6, 23].

Interesting results were obtained in case of dandelion aerial parts, since 20% ethanol provided the highest extraction yield of phenolcarboxylic acids and flavonoids. We assume that these results are probably a consequence of the high amount of polyphenols, mainly found as glycosidic forms (that are more soluble in water) [28].

Conclusions

All analysed herbal products are a source of flavonoids and phenolcarboxylic acids; still the solvent represents a key factor that greatly influences the phenolic content. According to our results, 50% ethanol provided the optimal extraction of phenolic compounds for all analysed herbal products, except for dandelion aerial parts (for which 20% ethanol was the best extraction solvent). Going forwards, our research will look on obtaining freeze-dried extracts rich in phenolic compounds. Furthermore, the extracts will be characterized by means of spectrophotometric/HPLC assays and we shall assess their antioxidant activity and hepatoprotective effects (animal studies).

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Conflict of interest

The authors declare no conflict of interest.

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