Changes in the content of bioactive substances among Hypericum montbretii populations from Turkey

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In the present study, we investigated the variation in the content of seventeen secondary metabolites among Hypericum montbretii Spach., Hypericaceae, populations from five different growing zones in Turkey for the first time. The plants were collected at full flowering, and after they were dried at room temperature, they were assayed for chemical contents by HPLC. Chemical constituents of plants varied significantly among populations except for 2,4-dihydroxybenzoic acid which was accumulated at similar levels. Plants from population - 1 yielded the highest amount of hypericin and pseudohypericin (1.27 and 2.97 mg/g, respectively) while hyperforin and adhyperforin accumulations were the highest in plants from population - 2 (6.64 and 1.24 mg/g, respectively). (+)-Catechin and (-)-epicatechin were accumulated at significantly higher levels by plants of population - 4 (1.54 and 4.35 mg/g, respectively). The highest accumulation level of the rest compounds namely, chlorogenic and neochlorogenic acids, amentoflavone, hyperoside, isoquercitrin, quercitrin, quercetin, avicularin and rutin was reached in plants from population - 5 (2.64, 4.37, 2.35, 10.26, 3.52, 4.37, 1.55, 1.56 and 20.54 mg/g, respectively). The pronounced chemical diversity between populations is discussed to possibly be the result of different environmental, morphological and genetic factors.

Original article

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present study we investigated the content variation of the above-mentioned compounds, as well as other newly identified bioactive substances such as, adhyperforin (4), neochlorogenic (6) and 2,4-dihydroxybenzoic (7) acids, amentoflavone (8), isoquercitrin (10), avicularin (13), (+)-catechin (15) and (-)-epicatechin (16) among five H. montbretii populations from different growing sites in Turkey for the first time.

Materials and methods

Plant material

The plant materials were described in our previous studies (Cirak & Radusiene, 2007; Cirak et al., 2008). The samples were authenticated by Dr. Samim Kayikci, Mustafa Kemal University, Faculty of Arts and Sciences, Department of Biology, Turkey. Voucher specimens were deposited in the herbarium of Ondokuz Mayis University Agricultural Faculty and voucher numbers of the collections are shown in Table 1.

Experimental procedures

The aerial parts of Hypericum montbretii Spach, Hypericaceae, plants representing a total of 30 shoots were collected at full flowering from five populations of Turkey (Table 1). The tops of 2/3 plants were harvested between 11 am and 13 pm and dried at room temperature (20° ± 2°C).

Preparation of plant extracts

Air-dried plant material was mechanically grounded using a laboratory mill until a homogenous powder was obtained. Samples of about 0.1 g (weighed with 0.0001 g precision) were extracted in 10 ml of 100% methanol by ultrasonication at 40°C for 30 min. The prepared extracts were filtered and kept in a refrigerator until analysis. The extracts for naphthodianthrones analysis were exposed to light from a xenon lamp (765 W/m²) for 8 min to allow photoconversion of protohypericins into hypericins.

Table 1
Geographical data and seasonal climatic conditions of Hypericum montbretii growing localities in Turkey.

<table>
<thead>
<tr>
<th>Populations</th>
<th>Collection date</th>
<th>Voucher nº</th>
<th>Latitude (n)</th>
<th>Longitude (E)</th>
<th>Elevation (m)</th>
<th>Mean temperature (°C)</th>
<th>Precipitation (mm)</th>
<th>Habitat</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>June 16, 2011</td>
<td>BMYO nº 4/1</td>
<td>41° 16’</td>
<td>41° 33’</td>
<td>500</td>
<td>10.01</td>
<td>850</td>
<td>Rocky and open slopes</td>
</tr>
<tr>
<td>2</td>
<td>June 19, 2011</td>
<td>BMYO nº 4/2</td>
<td>40° 54’</td>
<td>40° 28’</td>
<td>800</td>
<td>14.10</td>
<td>535</td>
<td>Arid pasturelands</td>
</tr>
<tr>
<td>3</td>
<td>June 20, 2012</td>
<td>BMYO nº 4/3</td>
<td>41° 52’</td>
<td>39° 14’</td>
<td>1720</td>
<td>11.00</td>
<td>825</td>
<td>Igneous slopes and rock ledges</td>
</tr>
<tr>
<td>4</td>
<td>June 15, 2011</td>
<td>BMYO nº 4/4</td>
<td>39° 54’</td>
<td>37° 28’</td>
<td>1500</td>
<td>15.17</td>
<td>525</td>
<td>Igneous slopes and rock ledges</td>
</tr>
<tr>
<td>5</td>
<td>June 13, 2011</td>
<td>BMYO nº 4/5</td>
<td>41° 04’</td>
<td>38° 06’</td>
<td>2750</td>
<td>11.12</td>
<td>819</td>
<td>Rocky and open slopes</td>
</tr>
</tbody>
</table>
**HPLC analysis and identification**

A Waters Alliance 2695 (Waters, Milford, USA) separation module system, equipped with Waters 2487 UV/Vis and Waters 996 PDA diode-array detectors, was used for HPLC analysis. Separation of flavonoids, epicatechin and hyperforins was carried out using a SunFire C18 column (3.5 µm, 150 mm x 3.0 mm i.d.; Waters, Milford, USA) with 10 mm guard-precolumn. The binary gradient elution was used for the detection of corresponding compounds. The mobile phase consisted of eluent A (water acidified with 0.3% phosphoric acid), and eluent B (acetonitrile containing 0.3% phosphoric acid). The elution profile was adjusted as follows: 0-12 min 16% B, 12-18 min (B 53→97%), 30-36 min (B 97→16%). Flow rate was 0.6 ml.min⁻¹ at a constant 25°C column temperature and the injected volume was 10 µl. Detection was monitored at 270-360 nm.

The ACE C18 column (5.0 µm, 250 x 4.6 mm i.d.; MAC-MOD Analytical, Inc) with guard-precolumn was used for separation of phenolic acids, catechin and hypericins. The mobile phase of gradient elution of phenolic acids and catechin was composed of eluent A (water acidified with 0.5% glacial acetic acid), and eluent B (acetonitrile). The separation was performed using the following program: 0-30 min (B 5→35%), 30-36 min (B 35→90%), and 36-37 min (B 90→5%). The flow rate was 1.0 ml.min⁻¹ at 25°C column temperature, injected volume 10 µl. Detection was performed at 277-324 nm.

Naphthodianthrones were analysed according to the modified pharmacopeial method (Pharm. Eur., 2010). The mobile phase of isocratic elution consisted of ethyl acetate, aqueous 0.1 M sodium dihydrogen phosphate solution, adjusted to pH 2.0 using phosphoric acid and methanol (16:17:67% v/v). The flow rate was 1.0 ml.min⁻¹ at 40°C column temperature and volume of extract injected 20 µl. Detection was performed at 590 nm.

Quantification of compounds was carried out by the external standard method. Standard stock solutions were prepared freshly in methanol and diluted in six different concentrations to obtain a set of corresponding concentration ranges for the linearity study. Chromatogram peak areas of analytes on their absorption maxima: 270 nm for hyperforins; 277 nm for epicatechin and catechin; 290 nm for 2,4-dihydroxybenzoic acid; 324 nm for chlorogenic and neochlorogenic acids; 360 nm for flavonoids; and 590 nm for hypericins, were plotted against the known concentrations of the standard solutions to establish the calibration equations. The regression coefficients of all calibration curves were $r^2 \geq 0.999$ confirming linearity of concentration ranges. The precision of the method was demonstrated for all analytes, since all relative standard deviations (RSD) values were lower than 5.0%.

Chromatographic peaks were assigned based on the retention time, UV spectra of the standard compounds using HPLC-DAD.

Solvents used were HPLC grade and purchased from Roth GmbH (Karlsruhe, Germany). Water was filtered through the Millipore HPLC grade water cartridge. Reference substances were purchased from ChromaDex (Santa Ana, USA), Sigma-Aldrich (Saint Louis, USA) and HWI Analytik GmbH (Germany).

**Data analysis**

Data for the content of each chemical were analysed using an ANOVA, and significant differences between mean values tested with the Duncan Multiple Range Test ($p \leq 0.01$), using MSTAT-C statistical software.

**Results and discussion**

The tested chemical contents in plants varied significantly among populations ($p < 0.01$) except for, 2,4-dihydroxybenzoic acid (7) which was accumulated at similar levels between populations. Plants from population-1 yielded the highest amount of hypericin (1) and pseudohypericin (2) (1.27 and 2.97 mg/g DM, respectively) while hyperforin (3) and adhyperforin (4) accumulations were the highest in plants from population-2 (6.64 and 1.24 mg/g DM, respectively). (+)-Catechin (15) and (-)-epicatechin (16) were accumulated at significantly higher levels by plants of population-4 (1.54 and 4.35 mg/g DM, respectively). The highest accumulation level of the rest of the compounds: chlorogenic (5) and neochlorogenic (6) acids, amentoflavone (8), hyperoside (9), isoquercitrin (10), quercitrin (11), quercetin (12), avicularin (13) and rutin (14), was reached in plants from population-5 (2.64, 4.37, 2.35, 10.26, 3.52, 4.37, 1.55, 1.56 and 20.54 mg/g DM, respectively) (Fig. 1). Similarly, significant changes in hypericin (1), pseudohypericin (2), hyperforin (3), quercitrin (11), quercetin (12), rutin (14), hyperoside (9) and chlorogenic acid (5) contents were reported among wild populations of Hypericum perforatum L.; the most abundant and well-known species (Sirvent et al., 2002; Cirak et al., 2007a; Bagdonaite et al., 2012), as well as in Hypericum triquetrifolium Turra (Cirak et al., 2011) and Hypericum orientale L. (Cirak et al., 2012). The present findings and the above mentioned reports indicate different geographic origin as an evident source for the chemical diversity observed among wild Hypericum populations. The huge variation in chemical content among the investigated populations could be partially related to the plant’s phenotypic plasticity at different environmental conditions for each population as seen in Table 1. Population-5, producing the highest level of chlorogenic acid (5), neochlorogenic (6) acid, amentoflavone (8), hyperoside (9), isoquercitrin (10), quercitrin (11), quercetin (12), avicularin (13) and rutin (14), was located at the highest altitude (2750 m). It can be assumed that the altitude may have a favorable impact on chemical accumulation in H. monbretii. On the other hand, the genetic diversity among plants may have an impact on the differences in chemical content as well. For example, the populations produced similar amount of 2,4-dihydroxybenzoic acid (7), while 3.14 and 5.64 fold differences were detected among them in hyperforin (3) and adhyperforin (4) content. However, it should be noted that detailed biochemical and molecular studies of the genetic diversity of wild populations are necessary to identify the exact influence of genetic and environmental factors on the evident phytochemical diversity among populations. As an example, Afef et al. (2012) assessed the genetic diversity within and among seven wild populations of Hypericum
humifusum L., from different geographic regions of Tunisia by using eleven isozymic polymorphic loci and 166 RAPD markers. Combined chemical, molecular and morphological studies on the relationships between genetic and chemical diversities within and among species of Hypericum are currently ongoing (Nürk & Crockett 2011).

_H. montbretii_ is a member of the section Drosocarpium Spach. To our knowledge, the chemical profiles of two other species from the section; _Hypericum bithynicum_ Boiss. and _Hypericum perfoliatum_ L., have been previously described (Cirak et al., 2007b; Smelcerovic et al., 2008). The comparison of the present results with previously published reports reveals that the three members of section Drosocarpium have a similar chemical profile, and include hypericin (1), pseudohypericin (2), hyperforin (3), rutin (14), chlorogenic acid (5), hyperoside (9), quercitrin (11) and quercetin (12). Among the chemicals, hypericins were reported to have an apparent taxonomic worth for the infrageneric classification of the genus _Hypericum_ (Crockett & Robson, 2011). Because hypericin and pseudohypericin were not detected in members of the primitive sections and seem to be specific only for the taxa of phylogenetically more advanced sections (Kitanov, 2001). Thus, detection of hypericins as well as the other chemicals tested in _H. montbretii_ in the present study solidifies the taxonomic standing of the section Drosocarpium within the genus Hypericum.

**Conclusions**

The present results indicate a considerable chemical variation among Turkish populations of _H. montbretii_. The chemical instability of the populations is a possibly due to different geographic origins and should be taken into account while processing the plant material for medicinal aims. Further multiple studies on the genetic and environmental reasons of the observed chemical diversity among wild populations are needed to make more substantial inferences. The accumulation of adhyperforin (4), neochlorogenic acid (6), 2,4-dihydroxybenzoic acid (7), amentoflavone (8), isoquercitrin (10), avicularin (13), (+)-catechin (15) and (-)-epicatechin (16) in _H. montbretii_ was documented by us for the first time. The present data could also be helpful to explain the chemotaxonomical significance of the corresponding compounds as well as to serve as a phytochemical evaluation of _H. montbretii_.

**Authors’ contributions**

CC provided the idea. CC and NC performed the field studies and sampled the plant populations in their wild habitats. JR, LI and VJ performed the chemical analyses. CC performed the statistical data evaluation and JR prepared the figures. All authors participated in the writing of the manuscript and approved its content.

![Figure 1](image-url) - Chromatograms of Hypericum montbretii whole flowering shoots extract detected by HPLC-DAD at 270-360 nm wavelength on SunFire C18 column for flavonoids, epicatechin and hyperforin, and on ACE C18 column for phenolic acids and (+)-catechin. Peaks identified: A: 1. (-)-epicatechin (16); 2. rutin (14); 3. hyperoside (9); 4. isoquercitrin (10); 5. avicularin (13); 6. quercitrin (11); 7. quercetin (12); 8. amentoflavone (8); 9. hyperforin (3); 10. adhyperforin (4); B: 11. 2,4-dihydroxybenzoic acid (7); 12. (+)-catechin (15); C: 13. neochlorogenic acid (6); 14. chlorogenic acid (5); 15. caffeic acid (17); D: 16. pseudohypericin (2); 17. hypericin (1).
Conflicts of interest

The authors declare no conflicts interest.

REFERENCES


