MORPHOGENETIC AND DIURNAL VARIATION OF TOTAL PHENOLS IN SOME HYPERICUM SPECIES FROM TURKEY DURING THEIR PHENOLOGICAL CYCLES

ALI KEMAL AYAN, OGUZHAN YANAR, CUNEYT CIRAK* AND MAHMUT BİLGENER

The High School of Profession of Bafra, University of Ondokuz Mayıs, Samsun, Turkey

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Abstract

The genus Hypericum has attracted scientific interest worldwide in recent years, since it is a source of a variety of biologically active compounds including the phenolics. The present study was to determine ontogenetic, morphogenetic and diurnal variation of the total phenol contents in some Hypericum species growing in Turkey namely, H. nummularioides, H. hyssopifolium, H. pruinatum and H. scabrum. Plants were separated into stem, leaf and reproductive tissues, which were dried separately, and subsequently assayed for total phenol contents. Among different tissues, leaves had higher phenol contents and the highest content reached at noon in all species tested. But, ontogenetic change in the total phenol contents of whole plants varied among the species. It reached the highest level at floral budding in H. hyssopifolium and H. scabrum, at full flowering in H. pruinatum and at fresh fruiting in H. nummularioides.

Introduction

The genus Hypericum contains approximately 400 different species of annuals, perennials, shrubs and small trees, ranging from very small perennials to trees. The species of this genus have been used as healing agent due to their various medicinal properties for the last two hundred years (Dias et al. 1998). Hypericum species are also used as sedatives, antiseptics, and antispasmodics in Turkish folk medicine under the names “kantaron, peygamber çiçeği, kılıçotu, kanotu, kuzukıran and binbirdelik otu” (Baytop 1999). The Hypericum genus is represented in Turkey by 89 species of which 43 are endemic. The most widespread and abundant species is H. perforatum L. (Davis 1988).

Morphologically, Hypericum plants are characterised by the presence of different kinds of secretory tissues including light glands, dark glands and secretory canals. These secretory structures are sites of synthesis and/or accumulation of biologically active substances and their localizations are different depending on plant tissue (Cicracelli et al. 2001). Therefore, organ-dependence of phenolic compounds has an important role to the understanding of the underlying sources of variation in phenolic contents of Hypericum plants.

Methanolic extract from the aerial parts of Hypericum plants typically contains hypericins, hyperforins and phenolic compounds (Barnes et al. 2001). It is especially rich in phenolics, caffeic acid, chlorogenic acid, proanthocyanidin (dimers and oligomers of catechin and epicatechin), prenylated derivatives of chloroglucinol and flavonoids, hyperin, rutin, quercitrin, isoquercitrin and bis-apigenins. (Mojca et al. 2005). Phenolic compounds are important for their contribution to the colour, sensory attributes and nutritional and antioxidant properties of plants (Christel et al. 2000). Phenolic compounds are reported to have multiple biological effects, including antioxidant activity, antitumor, antimutagenic and antibacterial properties (Shui and Leong 2002).

Diurnal fluctuations in the concentration of plant secondary metabolites such as saponins in Phytolacca dodecandra (Ndamba et al. 1993), alkaloids in Papaver somniferum (Itenov et al. 1999), essential oils in Laurus nobilis (Kevseroğlu et al. 2003) and hypericins in H. perforatum,
H. pruinatum and H. aviculariifolium (Çırak et al. 2006) have been reported. However, there is no information about diurnal variation of phenols in species of Hypericum.

Increased market demand for Hypericum plants has led to several investigations of secondary metabolite levels in plants from different areas of the world. These previous investigations were carried out on H. perforatum and did not give homogenous results. Furthermore, they are fully related with hypericins (Sirvent et al. 2002), hyperforin (Kirakosyan et al. 2002) or essential oils (Schwob et al. 2004) and ontogenetic stages were not established in most of them. To date, little effort has been given to the study of variation of phenolic compounds such as quercitrine, isoquercitrine (Marfonti and Repcak 1994) and quercetin (Tekel’ova et al. 2000). To our knowledge, no study has been done on the variation of total phenols in H. nummularioides Trautv., H. hyssopifolium Chaix, H. pruinatum Boiss.& Bal. and H. scabrum L.

In the present study, morphogenetic and diurnal variations of total phenols were investigated in the four Turkish species of Hypericum at different stages of plant phenology. The aim was to determine if there was a link between phenol content of plant materials and development stages during diurnal and phenological cycles.

Materials and Methods

H. nummularioides: Stems 10-20 cm, erect or ascending from a branching base, not rooting, glabrous. Leaves 7-15 mm, ovate to oblong-elliptic, plane, glabrous with conspicuous intramarginal glands. Inflorescence corymbose, 1-5 flowered. Sepals ovate to oblong-elliptic, black-glandular-denticulate. Petals 10-16 mm. Capsule unknown.

H. hyssopifolium: Stems 15-80 cm, erect, glabrous or with minute red or amber glands. Leaves on main stem 10-30 mm, narrowly elliptic-oblong to linear, often revolute. Inflorescence elongate, narrowly cylindric to pyramidal, many-flowered. Sepals unequal or subequal, ovate to oblong or rarely lanceolate, entire or with sessile glands. Petals 8-16 mm, sometimes red-veined. Capsule 5-15 mm, ovoid to subglobose.

H. scabrum: Stems 15-45 cm, erect or decumbent at the base, glabrous, scabrid with unbranched red-gland-tipped emergences. Leaves on main stems 7-20 mm, oblong or oblong-elliptic to lanceolate or linear. Inflorescence corymbose, many-flowered. Sepals oblong, subacute to rounded. Petals 5-8 mm. Capsule 5-8 mm, ovoid or ovoid-rigorous, not or scarcely rostrate.

H. pruinatum: Stems 15-35 cm, erect or ascending from a rooting and branching base, pruinose. Leaves on main stems 10-35 mm, oblong to elliptic, pruinose. Inflorescence pyramidal to cylindric. Sepals broadly oblong to broadly elliptic, rounded, entire or minutely black-glandular-denticulate. Petals 9-14 mm. Capsule 7-10 mm, ovoid (Davis 1988). The plant materials were identified and Voucher specimens were deposited in the Herbarium of Agricultural Faculty, Ondokuz Mayis University.

The Hypericum plants were collected from mountain pasture of Kuşmer in Aydintepe district of Bayburt province (40°15’ N Lat., 40°14’ E Long., and 1884 m elevation), Turkey between May and September of 2004 at different stages of plant development. The sampling sites were not grazed or mown during the sampling periods. Collections were done three times a day (06.00, 12.00 and 18.00 h) for each development stage. Sampling was done in these wild populations by a randomized collection of 30 individuals for each collection time within a day and each phenological stage.

Shoots with leaves were harvested at the vegetative stage. For the floral budding stage, only shoots with floral buds were selected. At the full flowering stage, only shoots with full opened flowers were harvested. At the fresh fruiting stage, the shoots which had green capsules were harvested. At the mature fruiting stage, the shoots which had dark brown capsules were harvested. After collection, ten individuals were kept as whole plants and the rest were separated into floral,
leaf and stem tissues. The plant materials were dried at room temperature (20 ± 2°C) separately. The dried materials were bulked and subsequently assayed for total phenols.

The procedure of Swain and Hills (1959) was used to determine the total phenolic contents of the plant extract. The total phenolics were measured in a spectrophotometer at 725 nm. A reagent blank and tannic acid standard solutions (0.01 to 0.06 mg/ml) were tested in triplicate and the mean value was calculated.

Phenol content of plant materials was given for each species. Data for phenol content of whole plant were objected to ANOVA and significant differences among phenol contents were tested with the DMRT ($p < 0.01$). However, because of the absence of generative tissues at vegetative stage as an experimental factor, no statistical analysis was performed for data belonging to the phenol content of different plant tissues (Steel and Torrie 1980).

**Results and Discussion**

Total phenolic contents of plant tissues varied with species greatly. *H. nummularioides* had the highest phenolic content and approximately eightfold difference between total phenol contents of this species and that of *H. scabrum*, supplying the lowest value.

Change in phenol content of whole plant within a day during the course of ontogenesis did not follow the same trend in the four species of *Hypericum*. In *H. hyssopifolium* and *H. scabrum*, phenol content in whole plant increased during flower ontogenesis and reached their highest level at floral budding (1.92 and 0.24% DW, respectively), then it decreased. The difference among phenol contents in whole plant at different development stages was found to be significant ($p < 0.01$). Diurnal fluctuation in phenol content of whole plant was also observed for *H. hyssopifolium* and it was highest (1.68% DW) at noon (Table 1). Similarly, phenol contents in whole plant were statistically different among different developmental stages in *H. pruinatum* and *H. nummularioides* ($p < 0.01$) but the highest concentration reached at full flowering for *H. pruinatum* (0.71% DW) and fresh fruiting for *H. nummularioides*. Similar to *H. hyssopifolium*, a significant difference was detected in phenol content in whole plant among diurnal harvesting and it was highest (0.63 and 1.73% DW, respectively) at noon in both species (Table 1).

Ontogenetic and diurnal fluctuations in phenol content of stems, leaves and reproductive parts varied with plant tissues and species. Leaves had higher phenolic content when compared to other tissues and total phenols content of all tissues was the highest at noon for all species. Leaf samples collected at noon gave the highest values depending on phenological stage in *H. nummularioides*, *H. hyssopifolium* and *H. pruinatum*, 2.36, 2.09 and 0.89% DW, respectively (Figs. 1, 2 and 3). Similarly, total phenols content of leaves was generally higher than that of other tissues in *H. scabrum* during its phenological cycle, although the highest value (0.32% DW) for this species was obtained from floral buds collected at noon (Fig. 4).

Investigations of ontogenetic variation of secondary metabolites have been made over several decades, e.g. alkaloid changes during fruit development in *Papaver somniferum* (Miriam and Pfeifer 1959) and *Conium maculatum* (Fairbairn and Challen 1959), essential oil changes during the course of ontogenesis in *Hypericum perforatum* (Schwob et al. 2004), artemisinin changes during phenological cycle of *Artemisia annua* (Gupta et al. 2002) and foliar monoterpenoid variation in *Umbellularia californica* in seedlings, saplings and adult trees stages (Raymond et al. 1996). Chemical concentrations vary considerably during the course of ontogenesis in a medicinal plant, not only do the concentrations of plant chemicals fluctuate through the season, but they can also be short-lived and experience rapid turnover. (Smith et al. 1996). Likewise, in the present study, total phenol contents of whole plants at different phenological stages were significantly different for each *Hypericum* species and the highest levels reached at different stages of plant
phenology depending on species (Kazlauskas and Bagdonaite 2004). Similarly, Osińska and Weglarz (2003). Ontogenetic fluctuations in phenolic content were also reported for other plant species such as *Malus domestica* (Treutter 2001), *Morus alba* and *Morus nigra* (Sıvacı and Sökmen 2004).

Table 1. Variation of total phenol contents of *Hypericum hyssopifolium*, *H. scabrum*, *H. pruinatum* and *H. nummularioides* within a day during the course of ontogenesis (% DW).

<table>
<thead>
<tr>
<th>Developmental stages</th>
<th>Diurnal collecting times</th>
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<tbody>
<tr>
<td></td>
<td>06.00 h</td>
</tr>
<tr>
<td><em>H. hyssopifolium</em></td>
<td></td>
</tr>
<tr>
<td>Vegetative</td>
<td>1.11 o*</td>
</tr>
<tr>
<td>Floral budding</td>
<td>1.85 ab</td>
</tr>
<tr>
<td>Full flowering</td>
<td>1.17 nm</td>
</tr>
<tr>
<td>Fresh fruiting</td>
<td>1.70 bcd</td>
</tr>
<tr>
<td>Mature fruiting</td>
<td>1.38 ik</td>
</tr>
<tr>
<td>Mean</td>
<td>1.44 B</td>
</tr>
</tbody>
</table>

| *H. scabrum*         |          |          |          |           |
| Vegetative           | 0.12 d*  | 0.18 abc | 0.18 abc | 0.15 B**  |
| Floral budding       | 0.22 ab  | 0.28 a   | 0.23 ab  | 0.24 A    |
| Full flowering       | 0.17 bc  | 0.21 ab  | 0.20 ab  | 0.19 AB   |
| Fresh fruiting       | 0.15 bc  | 0.19 abc | 0.17 bc  | 0.17 B    |
| Mature fruiting      | 0.14 bc  | 0.15 bc  | 0.12 d   | 0.14 B    |
| Mean                 | 0.16     | 0.20     | 0.18     |           |

| *H. pruinatum*       |          |          |          |           |
| Vegetative           | 0.62 bc* | 0.67 b   | 0.53 cd  | 0.61 B**  |
| Floral budding       | 0.55 cd  | 0.61 bc  | 0.54 cd  | 0.56 AB   |
| Full flowering       | 0.72 ab  | 0.78 a   | 0.61 bc  | 0.71 A    |
| Fresh fruiting       | 0.66 b   | 0.67 b   | 0.60 bc  | 0.64 B    |
| Mature fruiting      | 0.27 f   | 0.41 e   | 0.33 f   | 0.34 C    |
| Mean                 | 0.56 AB  | 0.63 A   | 0.52 B   |           |

| *H. nummularioides*   |          |          |          |           |
| Vegetative           | 1.19 ghi*| 1.39 fg  | 1.02 i   | 1.20 D**  |
| Floral budding       | 1.30 gh  | 1.92 b   | 1.71 d   | 1.64 B    |
| Full flowering       | 1.62 de  | 1.71 d   | 0.89 i   | 1.41 C    |
| Fresh fruiting       | 1.91 ab  | 2.10 a   | 1.92 ab  | 1.98 A    |
| Mature fruiting      | 1.51 ef  | 1.54 ef  | 1.44 efg | 1.49 C    |
| Mean                 | 1.51 B   | 1.73 A   | 1.40 C   |           |

Values followed by same small letters and capital letters do not differ significantly at 1% level. Plant material representing a total of ten individuals includes leaves and stems for vegetative stage; reproductive parts, leaves and stems for floral budding; full flowering; fresh fruiting stages; stems, brown capsules with seeds for mature fruiting stages.

Similar to primary metabolites, tissue-dependence of secondary metabolites is very common among medicinal plants. Leaves and flowers generally contain greater levels of phenolic acids and terpenoids than stems and roots (Hakulinen and Julkunen-Tiitto 2000). In the present study, leaves had the highest level of total phenols in all species tested and similar findings were reported by Radusiene *et al.* (2004) and Valentao *et al.* (2003).
Although, the general role of phenolic compounds in plant physiology and allelopathy has been known for many years, a less well reported aspect is their activity as defence factors against various types of stresses caused by pathogens or adverse environmental conditions (Treutter 2001). Plants can accumulate phenolic compounds in their different tissues in response to challenge by various stress factors (Pasqualini et al. 2003). During the course of ontogenesis, plants are subjected to different kinds of stress factors, such as drought, heat, herbivore/pathogen attack and air pollution (Paliyath et al. 1997). Most plants suffer from physiological and biochemical damage due to the exposure to temperatures higher or lower than optimal for growth (Grace et al. 1998). Climatic changes like high temperature stress promote production of phenolic compounds (Christie et al. 1994, Dixon and Paiva 1995, Sıvacı and Sökmen 2004). The increase
observed here in phenolic content of plant tissues of the Turkish *Hypericum* species collected at noon may be attributed to higher temperatures of midday.

**Fig. 3.** Diurnal and ontogenetic changes in total phenol contents of reproductive parts, leaf and stem tissues in *H. pruinatum*.

**Fig. 4.** Diurnal and ontogenetic changes in total phenol contents of reproductive parts, leaf and stem tissues in *H. scabrum*.

**Conclusion**

It can be concluded that there is a close relationship between phenolic content of plant tissues and growth stages during diurnal and phenological cycles in *Hypericum* species. Considering the pharmacological significance of phenolics, their possible use in therapeutics and the growing interest in analytical data on natural phenols in plants, it is important to find the different sources of these compounds. At this point, the high phenolic content of *H. nummularioides* and *H. hyssopifolium* encourages the cultivation, and biological evaluation of these species in Turkey.
The results also indicated that the quantitative variation of total phenol contents in four Turkish species of *Hypericum* allows the selection of the best plant samples for their cultivation and conservation in field collections. Therefore, wild populations of these *Hypericum* species are potentially important sources for breeding and improvement of the cultivated varieties.

References


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