Chemical composition and bioactive properties of the lichen, *Pleurosticta acetabulum*

Jovica Tomović1, Marijana Kosanić2, Svetlana Ristić2, Branislav Ranković2, Tatjana Stanojković3, Nedeljko Manojlović1*

1University of Kragujevac, Serbia, Faculty of Medical Sciences, Department of Pharmacy, 2University of Kragujevac, Faculty of Science, Department of Biology, 3University of Kragujevac, Faculty of Oncology and Radiology of Serbia, 11000 Belgrade, Serbia

*For correspondence: Email: mtndeljko@gmail.com; Tel: +381-691137150; Fax: +381-34306800

Sent for review: 18 August 2017
Revised accepted: 22 November 2017

**Abstract**

**Purpose:** To investigate the chemical composition and bioactivity of the acetone extract of *Pleurosticta acetabulum* lichen.

**Methods:** Phytochemical analysis of the acetone extract of the lichen (*Pleurosticta acetabulum*) was carried out by high-performance liquid chromatography (HPLC). The antioxidant activity of the lichen extract was evaluated by determining the radical scavenging capacity on 2,2-diphenyl-1-picrylhydrazyl (DPPH) radicals and reducing power. To determine total phenolics and flavonoids, we used spectrophotometric methods. Antimicrobial activity was estimated by determining the minimal inhibitory concentration using broth microdilution method. Anticancer activity of the lichen extract was tested using 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT).

**Results:** Salazinic, norstictic, protocetraric, evernic acid and atranorin were identified as compounds of lichen. *P. acetabulum* extract exhibited moderate free radical scavenging activity (half-maximum inhibitory concentration, IC50 of 151.7301 µg/mL). The spectrophotometric absorbance of the extract for reducing power varied from 0.035 to 0.127, while the total phenolics and flavonoids in the extract were 35.39 µg PE/mg and 12.74 µg RE/mg, respectively. Minimum inhibitory concentration (MIC) was in the range of 1.25 to 20 mg/mL while cytotoxic activity (based on IC50 values) ranged from 24.09 to 45.94 µg/mL.

**Conclusion:** The results confirm that lichen extract contains secondary metabolites that possess antioxidant, antimicrobial and anticancer activities, which opens up some possibilities for the extract to be developed as food supplements and pharmaceutical raw materials.

**Keywords:** Pleurosticta acetabulum, Antioxidant activity, Antimicrobial activity, Cytotoxic activity, Lichen

This is an Open Access article that uses a funding model which does not charge readers or their institutions for access and distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0) and the Budapest Open Access Initiative (http://www.budapestopenaccessinitiative.org/read), which permit unrestricted use, distribution, and reproduction in any medium, provided the original work is properly credited.

Tropical Journal of Pharmaceutical Research is indexed by Science Citation Index (SciSearch), Scopus, International Pharmaceutical Abstract, Chemical Abstracts, EMBASE, Index Copernicus, EBSCO, African Index Medicus, JournalSeek, Journal Citation Reports/Science Edition, Directory of Open Access Journals (DOAJ), African Journal Online, Bioline International, Open-J-Gate and Pharmacy Abstracts

**INTRODUCTION**

A great diversity of living organisms can be used for the improvement of human health, such as biologically active components extracted from the plants, mushrooms and lichens [1-3]. In searching for new therapeutic alternatives, many different lichen species were investigated, due to significant antiviral, antimicrobial, anticancer, antihyperglycemic, cardioprotective, antiparasitic,
anti-inflammatory and antibiotic effects that they have exhibited so far [3-5].

Many lichen species exert interesting biological and pharmacological activities. However, in the literature, only a few studies have investigated the potential of Pleurosticta acutabulum lichen.

Thus, the aim of this study is to present the results of the phytochemical analysis of the acetone extract of P. acutabulum lichen and its antioxidant, antimicrobial and anticancer activities in order to search for available natural antioxidant, antibiotic and anticancer agents that can be used as possible food supplements, raw materials for the pharmaceutical industry and remedies for the treatment of various diseases.

EXPERIMENTAL

Collection and identification of lichen sample

The samples of lichen P. acutabulum (Neck.) Elix & Lumbsch were collected from the mountain Kopaonik, Serbia, during May 2013. Identification was done using the relevant key and monographs by Ranković [6,7]. The samples of collected species of lichen have been herbarium-stored at the Department of Biology and Ecology of Kragujevac, Faculty of Science, (voucher no. 109) for future reference.

Preparation of lichen extract

The samples of Lichen Parmelia acutabulum (100 g) were extracted with a Soxhlet extractor using acetone. The extract was filtered and then concentrated under reduced pressure in a rotary evaporator. The dried extract was dissolved in DMSO (5 % dimethyl sulfoxide) for further experiments.

High-performance liquid chromatography (HPLC)

The dry lichen extract was dissolved in 500 µL of acetone and carried out on a 1200 Series HPLC (Agilent Technologies) instrument with C18 column (C18; 25 cm 4.6 mm, 10 µm). A UV spectrophotometric detector was used with methanol-water-formic acid (70:30:0.8, v/v/v) as a solvent. The detection wavelength was 254 nm and the injection volume was 5 µL, with a flow rate of 1 ml/min. Deionized water was purified using a Milli-Q academic water purification system (Milford, MA, USA). Formic acid was used as an analytical-grade reagent. HPLC-grade methanol was purchased from Merck (Darmstadt, Germany). These lichen substances are not available commercially.

Secondary metabolites were isolated manually from the lichen species using thin-layer chromatography (TLC) and column chromatography. The purity of the isolated metabolites was checked by HPLC. The standards used were obtained from the following sources: salazinic acid (tR = 1.56 ± 0.20 min) isolated from the lichen Lobaria pulmonaria, norstictic acid (tR = 2.70 ± 0.10 min) from the lichen Ramalina farinacea and protocetraric acid (tR = 3.24 ± 0.20 min) from the lichen Tonia transc. Evernic acid (tR = 5.08 ± 0.10 min) and atranol (tR = 14.88 ± 0.10 min) were isolated from the lichen Evernia prunastri.

Determination of antioxidant activity

Scavenging DPPH radicals

DPPH (2,2-diphenyl-1-picryl-hydrazyl-hydrate) radical was used to evaluate the free radical scavenging activity of the lichen extract [8]. Two millilitres of 0.05 mg/mL methanol solution of DPPH radical and 1 mL of the lichen extract (1 mg/mL) were placed in cuvettes. The mixture was stored at room temperature for 30 min. Then, the absorbance was measured at 517 nm in a spectrophotometer (Jenway, UK). Ascorbic acid was used as a positive control. DPPH radical scavenging activity (D) was calculated using Eq 1.

\[ D(\%) = \frac{A_0 - A_1}{A_0} \times 100 \]  

where \( A_0 \) is the absorbance of the negative control (DPPH solution) and \( A_1 \) is the absorbance of the reaction mixture or standard. All the measurements were repeated three times. The inhibitory concentration (IC50) was the parameter used to compare the radical scavenging activity.

Reducing power

The reducing power of the extract was determined according to the method of Oyaizu [9]. One millilitre of extract (1 mg/mL) was mixed with 2.5 mL of phosphate buffer (2.5 mL, 0.2 M, pH 6.6) and potassium ferricyanide (2.5 mL, 1 %). The mixtures were incubated at 50 °C for 20 min. Trichloroacetic acid (10 %, 2.5 mL) was added to the mixture and centrifuged. The upper layer was mixed with distilled water (2.5 mL) and ferric chloride (0.5 mL, 0.1%). The absorbance of the solution was measured at 700 nm in a spectrophotometer (Jenway UK). Ascorbic acid was used as a positive control. The greater the absorbance of the reaction mixture the higher the reducing power of the samples.
Determination of total phenolic compounds

Total soluble phenolic compounds in the lichen extract were determined with Folin-Ciocalteu method [10], using pyrocatechol as a standard phenolic compound. One millilitre of the lichen extract (1 mg/mL) was diluted with distilled water (46 mL), and the content was mixed in a volumetric flask after adding one millilitre of Folin-Ciocalteu reagent. After 3 min, 3 mL of 2 % sodium carbonate was added and left for 2 h with intermittent shaking. The reaction mixture absorbance was measured at 760 nm in a spectrophotometer (Jenway UK). The total concentration of phenolic compounds in the extract was expressed as microgram of pyrocatechol equivalent (PE) per milligram of dried extract. The total phenolics content was determined as the pyrocatechol equivalent using an equation obtained from a standard pyrocatechol graph (y = 0.0057 x total phenols (μg PE/mg of dry extracts) - 0.1646, R²=0.9934).

Evaluation of total flavonoid content

The total flavonoids content in the lichen extract was determined with the spectrophotometric method using aluminium trichloride based on flavonoid-aluminium complex formation [11]. Two millilitres of 2 % aluminium trichloride in methanol was mixed with the same volume of the extract solution (1 mg/mL). The mixture was incubated at room temperature for 10 min, and the absorbance was measured at 415 nm in a spectrophotometer (Jenway UK) against a reagent blank (consisting of all the reagents except the extract or rutin standard solution being substituted with methanol). The total flavonoid content was determined as microgram of rutin equivalent (RE) per milligram of dried extract. The total amount of flavonoid compounds was determined as the rutin equivalent using an equation obtained from a standard rutin graph (y = 0.0296 x total flavonoid [μgRE/mg of dry extracts] + 0.0204, R²=0.9992).

Assessment of antimicrobial activity

Antimicrobial activity of the tested sample was evaluated against 15 microorganisms, including five strains of bacteria: *Bacillus cereus* (ATCC 11778), *B. subtilis* (ATCC 6633), *Staphylococcus aureus* (ATCC 25923), *Escherichia coli* (ATCC 25922), *Proteus mirabilis* (ATCC 12453) and 10 species of fungi: *Aspergillus flavus* (ATCC 9170), *A. niger* (ATCC 16888), *Candida albicans* (ATCC 10231), *Mucor mucedo* (ATCC 20094), *Trichoderma viride* (ATCC 13233), *Cladosporium cladosporioides* (ATCC 11275), *Fusarium oxysporum* (ATCC 62506), *Alternaria alternata* (ATCC 11680), *Penicillium expansum* (ATCC 20466), *P. chrysogenum* (ATCC 10106) obtained from the American Type Culture Collection (ATCC).

The bacterial isolates were isolated from overnight cultures using Müller-Hinton agar and the suspensions were prepared in sterile distilled water by adjusting the turbidity to match that of a 0.5 McFarland standard (approximately 10⁶ CFU/mL). Fungal suspensions were prepared from 3-7-day-old cultures that grew on a potato dextrose agar, except for *C. albicans* which was maintained on Sabouraud dextrose (SD) agar. Sterile distilled water was used to rinse the spores, the turbidity was measured spectrophotometrically at 530 nm, and then further diluted to a concentration of approximately 10⁶ CFU/mL according to NCCLS recommendations [12].

In order to determine the minimum inhibitory concentration (MIC) of the active extract, the 96-well microtitre assay using resazurin as the indicator of cell growth was employed [13]. The starting solutions of the tested extract were obtained by dissolving it in 5 % dimethyl-sulfoxide. Serial twofold dilutions of the extract were made within a concentration range from 0.04 to 40 mg/mL in sterile 96-well plates containing Mueller–Hinton broth for bacterial cultures and Sabouraud Dextrose SD broth for fungal cultures. Resazurin solution was added as an indicator to each well. The MIC was determined visually and defined as the lowest concentration of the tested extract that prevented resazurin colour change from blue to pink. Streptomycin and ketoconazole were used as positive controls, while 5 % DMSO was used as a negative control.

Determination of cytotoxic activity

Human epithelial carcinoma Hela cells, human lung carcinoma A549 cells and human colon carcinoma LS174 cells were obtained from American Type Culture Collection (Manassas, VA, USA). All cancer cell lines were cultured as monolayers in the RPMI 1640 nutrient medium with 10 % Fetal Bovine Serum (inactivated at 56 °C), 3 mM of L-glutamine, and antibiotics, at 37 °C in a humidified air atmosphere with 5 % CO₂. *In vitro* assay for cytotoxic activity of the investigated sample was performed when the cells reached 70 – 80 % confluence. A stock solution of the extract was dissolved in the corresponding medium to the required working concentrations. Neoplastic Hela, A549 and LS174 cells (5000 cells line per well) were seeded into 96-well microtiter plates, and 24 h
later, after cell adherence, five different double-diluted concentrations of the extract were added to the wells. The final concentrations of the extract were 12.5, 25, 50, 100, and 200 μg/mL, except for the control wells where only the nutrient medium was added. The cultures were incubated for the next 72 h.

The cancer cell-survival effect was determined 72 h after adding the extract, using the MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide) assay [14]. To each well was added 20 μL of MTT solution (5 mg/mL PBS) and they were further incubated in humidified air containing 5% CO₂ at 37 °C for 4 hours. Subsequently, 100 μL of 10 g/L SDS was added to solubilise the MTT formazan crystals converted by mitochondrial dehydrogenases in viable cells. The absorbance proportional to the number of viable cells was measured at 570 nm using a microplate reader (Multiskan EX, Thermo Scientific, Finland). Each experiment was performed in triplicate and independently repeated at least four times.

Cis-dichlorodiammineplatinum (cis-DDP) was used as a positive control.

Statistical analysis

All data are presented as the mean ± standard deviation (mean ± SD) of three parallel measurements. Statistical analyses were performed using Microsoft Excel and SPSS software (version 18) package. Student’s t-test was used to determine statistically significant differences which were considered significant at p < 0.05.

RESULTS

Secondary metabolites in the lichen P. acetabulum were identified using HPLC. The chromatograms for standards (salazinic, norstictic, protocetraric, evenic acid and atranorin) and lichen acetone extract eluted by HPLC are represented in Figure 1 and Figure 2.

Identification of these compounds was achieved by comparison of their retention times (tᵣ) and UV spectra (200 - 400 nm) from HPLC-UV with the standard substances previously isolated from lichens in our laboratory. The dominant peak in the chromatogram (tᵣ = 2.70 ± 0.10 min) originates from depsidone compound, norstictic acid (bryopogonic acid, 1,3-Dihydro-1,4,10-trihydroxy-5,8-dimethyl-3,7-dioxo-7H-isobenzofuro(4,5-b)(1,4)benzodioxepin-11-carboxaldehyde). The UV spectrum of norstictic acid has 3 absorption maxima (212, 239 and 320 nm). Besides norstictic acid, the tested extract of P. acetabulum contains salazinic acid (tᵣ = 1.56 ± 0.20 min), protocetraric acid (tᵣ = 3.24 ± 0.20 min), evenic acid (tᵣ = 5.08 ± 0.10 min) and atranorin (tᵣ = 14.88 ± 0.10 min) in different amounts.

Protocetraric, evenic acid and atranorin have very small peaks and present satellite substances in the chromatogram. The UV spectra of salazinic (212, 239 and 310 nm) and protocetraric (212, 242 and 320 nm) are very similar to those of norstictic acid. Norstictic,
salazinic and protocetraric acid are categorized into β-orcinoldepsidones. Absorbance maxima at 213, 270 and 305 nm are characteristic for evernic acid and at 212, 278 and 312 nm for atranorin. Salazinic and protocetraric acid belong to depsidones while atranorin and evernic acid belong to depsides. The retention times and UV absorbance maxima of the standards are shown in Table 1.

**Table 1:** Retention times of the lichen compounds and their absorbance maxima (nm)

| Compound       | Retention time (h±SD) (min) | Absorbance maxima (nm) | UV spectrum |
|----------------|-----------------------------|-------------------------|-------------|
| Salazinic acid | 1.56±0.20                   | 212, 238, 310           |             |
| Norstictic acid| 2.70±0.10                   | 212, 239, 320           |             |
| Protocetraric acid | 3.24±0.20              | 212, 242, 320           |             |
| Evernic acid   | 5.08±0.10                   | 213, 270, 305           |             |
| Atranorin      | 14.88±0.10                  | 212, 278, 312           |             |

*Values are mean ± SD (n = 3); m = minor absorbance maximum*

The structures of the detected compounds are shown in Figure 3. The antioxidant activity (scavenging DPPH radicals and reducing power) of the acetone extract is presented in Table 2.

The IC\textsubscript{50} value of the lichen extract was 151.01 μg/mL for DPPH radicals. As shown in Table 2, reducing power was concentration dependent. The values of absorbance for reducing power varied from 0.035 to 0.127. In various antioxidant activities, there was a statistically significant difference between the extract and the control (p < 0.05).

The amounts of total phenolics and flavonoids in the extract were 35.39 μg PE/mg and 12.74 μg RE/mg, respectively.

The antimicrobial activity of the lichen extract against the test microorganisms is shown in Table 3.

The MIC for the acetone extract of *P. acetabulum* fluctuated in a range of 1.25 – 20 mg/mL for bacteria and 5 – 20 mg/mL for fungi. The extract did not show inhibitory activity against *E. coli* and *A. flavus*, which have been shown to be the most resistant bacteria, and fungi. The antimicrobial activity was compared with the streptomycin and ketoconazole which were more active than the tested lichen. In a negative control, DMSO had no inhibitory effect on the tested organisms.

The data obtained for the anticancer effect of *P. acetabulum* extract are shown in Table 4.

**Table 2:** DPPH radical scavenging and reducing power of the acetone extracts of *Pleurosticta acetabulum*

| Antioxidant test        | DPPH radical scavenging IC\textsubscript{50} (μg/mL) | Reducing power Absorbance (700 nm) |
|-------------------------|-----------------------------------------------------|-----------------------------------|
|                         | 1000 μg/mL                                        | 500 μg/mL                        | 250 μg/mL | 125 μg/mL |
| *Pleurostia acetabulum* | 151.01±1.91                                       | 0.127±0.011                      | 0.055±0.005 | 0.046±0.005 | 0.035±0.001 |
| Ascorbic acid           | 6.42±0.18                                         | 2.113±0.032                      | 1.654±0.021 | 0.0957±0.008 | 0.0478±0.004 |

*Values are expressed as mean ± SD (n = 3)*
Table 3: Minimum inhibitory concentration (MIC) of the acetone extracts of *Pleurosticta acetabulum*

| Microorganism               | *Pleurosticta acetabulum* | Streptomycin | Ketoconazole |
|----------------------------|----------------------------|--------------|--------------|
| *Staphylococcus aureus*    | 20                         | 0.031        | /            |
| *Bacillus subtilis*        | 5                          | 0.016        | /            |
| *Bacillus cereus*          | 1.25                       | 0.016        | /            |
| *Escherichia coli*         | /                          | 0.062        | /            |
| *Proteus mirabilis*        | 10                         | 0.062        | /            |
| *Mucor mucedo*             | 10                         | /            | 0.156        |
| *Trichoderma viride*       | 10                         | /            | 0.078        |
| *Cladosporium cladosporiodes* | 5              | /            | 0.039        |
| *Fusarium oxysporum*       | 5                          | /            | 0.078        |
| *Alternaria alternata*     | 10                         | /            | 0.078        |
| *Aspergillus flavus*       | /                          | /            | 0.312        |
| *Aspergillus niger*        | 20                         | /            | 0.078        |
| *Candida albicans*         | 5                          | /            | 0.039        |
| *Penicillium expansum*     | 20                         | /            | 0.156        |
| *Penicillium chrysogenum*  | 10                         | /            | 0.078        |

Values given as mg/mL; Antibiotics: S – streptomycin, K – ketoconazole. Slash (/)-No activity

Table 4: Growth inhibitory activity of acetone extracts of *Pleurosticta acetabulum* on Hela, A549 and LS174 cell lines

| Sample                        | IC₅₀ (μg/mL) |
|-------------------------------|-------------|
|                               | Hela        | A549        | LS174        |
| *Pleurosticta acetabulum*     | 26.95±1.54  | 24.09±0.36  | 45.94±1.28   |
| cis-DDP                       | 0.83±0.19   | 3.56±0.23   | 2.58±0.16    |

IC₅₀ values are expressed as mean ± SD determined from the results of MTT assay in three independent experiments.

The IC₅₀ values of the lichen extract against Hela, A549 and LS174 cell lines were 26.95, 24.09 and 45.94 μg/mL, respectively.

**DISCUSSION**

The identification of secondary metabolites in the acetone extract of *P. acetabulum* and its antioxidant, antimicrobial and anticancer potentials were presented in this study.

By analyzing the composition of lichen *P. acetabulum* the presence of five secondary metabolites has been confirmed. Protocetraric and evernic acid were identified from the lichen *P. acetabulum* for the first time during this study research.

Norstictic acid as major lichen substance in the tested extract is a widespread secondary metabolite produced by lichen-forming fungi [15]. The identified metabolites could be used in the taxonomic classification of lichen species and as sources of commercial products.

In this study, the lichen extract showed relatively powerful levels of antioxidant activity. Some metabolites of lichens, including depsides, depsidones and dibenzofurans, contain phenolic groups considered to have an important role in antioxidative efficiency. The lichen used for the investigation contains secondary metabolites that have been shown to exhibit powerful antioxidant activity. Free radical scavenging and antioxidant activities of atranorin were evaluated using various *in vitro* assays for scavenging activity against hydroxyl radicals, hydrogen peroxide, superoxide radicals and nitric oxide.

Kosanic *et al* and Melo *et al* found that atranorin exerts differential effects towards reactive species production, enhancing hydrogen peroxide and nitric oxide production and acting as a superoxide scavenger, and then the activity towards hydroxyl radical production scavenging was observed [16,17].

Also, total reactive antioxidant potential and total antioxidant reactivity analysis indicate that atranorin acts as a general antioxidant, although it appeared to enhance peroxyl radical-induced lipoperoxidation *in vitro* [16]. Similarly, strong antioxidant activities were found for salazinic, protocetraric, evernic and norstictic acids [4,5,17].

In the literature sources that we examined, no data were found about the antimicrobial activity of *P. acetabulum* extracts. However, a relatively strong antimicrobial effect against numerous bacteria and fungi was found in the extract of the lichen *Parmelia acetabulum* where secondary metabolites were identified. Manojlović *et al* [5] reported about the antimicrobial activity of...
salazinic and protocetraric acids. Also, strong antimicrobial activity was found for atranorin [18,19].

The strong antimicrobial activities of evernic and norstictic acids have previously been reported [4,17]. They observed that they both exerted antibacterial activity against Gram-positive and Gram-negative bacteria, but Gram-negative bacteria were more resilient. This resistance is likely due to the fact that Gram-negative bacteria have a wall associated with an outer complex membrane, which slows down the passage of hydrophobic compounds. Lacking outer membrane, Gram-positive bacteria are more susceptible to antibiotic agents [17].

Compared to bacteria, fungi are more resistant due to the more complex structure of the cell wall [3]. Similar to this research, the acetone extract of the lichen P. acetabulum and its major compound norstictic acid were previously tested for anti-proliferative activity towards HT29 cells [20]. The results showed that acetone extracts of P. acetabulum had the strongest anticaner activity with an IC50 value of 6 µg/mL after 48h treatment. Also, other constituents of P. acetabulum (evernic, salazinic, protocetraric acids and atranorin) have been shown as promising anticancer agents [5,17,21].

CONCLUSION

The foliose lichen, Pleurosticta acetabulum, is a source of versatile bioactive compounds, which provide tremendous opportunities for the production of new antioxidant, antimicrobial and anticancer agents. Lichen is also a potentially suitable source of raw material for food and pharmaceutical industries due to their activities.

However, further studies to elucidate its mechanism of actions are necessary to establish the potential biological properties of lichen extract and its compounds.

DECLARATIONS

Acknowledgement

This work was partly financially supported by the Ministry of Science, Technology, and Development of the Republic of Serbia and was carried out within the Framework of the Projects (nos. 173032, 175011 and 172015).

Conflict of interest

No conflict of interest is associated with this work.

Contribution of authors

The authors declare that this work was done by the authors named in this article and all liabilities pertaining to claims relating to the content of this article will be borne by them.

REFERENCES

1. Morsy N. Phytochemical analysis of biologically active constituents of medicinal plants. Main Group Chem 2014; 13(1): 7-21.
2. Wasser SP. Medicinal Mushroom Science: Current Perspectives, Advances, Evidences and Challenges. Biomed J 2014; 37(6): 345-356.
3. Ristić S, Ranković B, Kosanić M, Stanojković T, Stamenković S, Vasiljević P, Manojlović I, Manojlović N. Phytochemical study and antioxidant, antimicrobial and anticancer activities of Melanella subaurifera and Melanella fuliginosa lichens. J Food Sci Tech Mys 2016; 53(6): 2804-2816.
4. Ranković B, Kosanić M, Stanojković T, Vasiljević P, Manojlović N. Biological Activities of Toninia candida and Usnea barbata Together with Their Norstictic Acid and Usnic Acid Constituents. Int J Mol Sci 2012; 13(11): 14707–14722.
5. Manojlović N, Ranković B, Kosanić M, Vasiljević P, Stanojković T. Chemical composition of three Parmelia lichens and antioxidant, antimicrobial and cytotoxic activities of some their major metabolites. Phytoled 2012; 19(13): 1166–1172.
6. Wirth V. Die Flechten Baden-Württembergs, Verbreitungsatlas, 1&2; Eugen Ulmer GmbH&Co: Stuttgart, Germany, 1995.
7. Dobson FS. Lichens. An illustrated guide to the British and Irish species, sixth ed. Richmond Publishing Co. London, 2011.
8. Dorman HJ, Bachmayer O, Kosar M, Hiltunen R. Antioxidant properties of aqueous extracts from selected Lamiaceae species grown in Turkey. J Agric Food Chem 2004; 52(4): 762-770.
9. Oyaizu M. Studies on products of browning reaction prepared from glucoseamine. Jpn J Nutr 1986; 44(6): 307-314.
10. Slinkard K, Singleton VL. Total phenolic analyses: automation and comparison with manual method. Am J Enol Viticult 1977; 28: 49-55.
11. Meda A, Lamiien CE, Romito M, Millogo J, Nacoulma OG. Determination of the total phenolic, flavonoid and proline contents in burkina fasan honey, as well as their radical scavenging activity. Food Chem 2005; 91(3): 571-577.
12. NCCLS (National Committee for Clinical Laboratory Standards). Reference method for broth dilution antifungal susceptibility testing of conidium-forming filamentous fungi: Proposed standard M38-p. Wayne, PA, USA, 1998.
13. Sarker SD, Nahar L, Kumarasamy Y. Microtitre plate-based antibacterial assay incorporating resazurin as an indicator of cell growth, and its application in the in vitro antibacterial screening of phytochemicals. Methods 2007; 42(4): 321-324.
14. Mosmann T. Rapid colorimetric assay for cellular growth and survival: application to proliferation and cytotoxicity assays. J Immunol Methods 1983; 65(1-2): 55-63.
15. Huneck S, Yoshimura I. Identification of lichen substances. Berlin Heidelberg: Springer-Verlag, 1996.
16. Melo MG, dos Santos JP, Serafini MR, Caregnato FF, Pasquali MA, Rabelo TK, da Rocha RF, Quintans Jr, Araújo AA, da Silva FA, et al. Redox properties and cytoprotective actions of atranorin, a lichen secondary metabolite. Toxicol In Vitro 2011; 25(2): 462–468.
17. Kosanić M, Manojlović N, Janković S, Stanojković T, Ranković B. Evernia prunastri and Pseudoevernia furfuraceae lichens and their major metabolites as antioxidant, antimicrobial and anticancer agents. Food Chem Toxicol 2013; 53:112-118.
18. Yilmaz Y, Turk AO, Tay T, Kivanc M. The antimicrobial activity of extracts of the lichen Cladonia foliacea and its (-)-usnic acid, atranorin and fumarprotocetraric acid constituents. Z Naturforsch C 2004; 59(3-4): 249–254.
19. Thadhani VM, Choudhary IM, Khan S, Karunaratne V. Antimicrobial and toxicological activities of some depsides and depsidones. Journal of the National Science Foundation Sri Lanka 2012; 40(1): 43-48.
20. Millot M, Delebassée S, Liagre B, Vignaud L, Sol V, Mambu L. Screening of lichen extracts on HT-29 human colon-cancer cells. Planta Med 2014; 80(16) - P1N5.
21. Bačkorova M, Jendzelovsky R, Kello M, Backor M, Mikes J, Fedoročko P. Lichen secondary metabolites are responsible for induction of apoptosis in HT-29 and A2780 human cancer cell lines. Toxicol In Vitro 2012; 26(3): 462-468.