The study of the qualitative composition and the quantitative content of phenolic compounds in dietary supplements with lingonberry

Abstract
Aim. Today, there are a lot of dietary supplements with lingonberry at the pharmaceutical market of Ukraine; therefore, the analysis and quality control of these products are relevant. In this connection, the aim of the research was to study the qualitative composition and determine the quantitative content of phenolic compounds in dietary supplements with lingonberry.

Materials and methods. Such dietary supplements as “Extract of lingonberry” (MEDAGROPROM), “Lingonberry” (Danikafarm), “Lingonberry nano” (LSS SYSTEM) were chosen for the study. The qualitative analysis was performed by thin layer chromatography (TLC), spectrophotometry was used for the quantitative determination.

Results and discussion. Hydroquinone derivatives, flavonoids and hydroxycinnamic acids were found in the dietary supplements analyzed. The total content of phenolic compounds was 8.70, 0.26, 0.30 %, flavonoids – 6.37, 0.15, 0.12 %, hydroxycinnamic acids – 0.94, 0.06, 0.13 %, and hydroquinone derivatives – 1.01, 0.04, 0.03 % in such dietary supplements as “Extract of lingonberry” (MEDAGROPROM), “Lingonberry” (Danikafarm), “Lingonberry nano” (LSS SYSTEM), respectively.

Conclusions. The qualitative and quantitative analysis of the dietary supplements with lingonberry analyzed has been performed. “Extract of lingonberry” (MEDAGROPROM) dietary supplement meets the requirements of the State Pharmacopoeia of Ukraine 2.0, whereas “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) do not. Based on the results of the study it can be concluded that the problem of compliance of dietary supplements is relevant today and requires the introduction of regulatory documentation for the detection and determination of biologically active substances in dietary supplements.

Keywords: lingonberry; spectrophotometry; analysis; phenolic compounds; dietary supplements
Introduction

Lingonberry (Vaccinium vitis idaea) is small shrubs belonging to the genus Vaccinium, of Ericaceae family. Its berries mature from July to September, while the ripeness time is highly affected by the site conditions, precisely altitude, and the habitat type. Usually, higher altitudes generate later plant ripening when compared with lower elevations [1].

The main biologically active substances of lingonberry leaves and berries are phenolic compounds (arbutin, hydroquinone, gallo- and ellagotannins), flavonols (luteolin, kempferol and quercetin), hydroxycinnamic acids (chlorogenic, cumaric and ferulic acids), coumarins, organic acids [2–4].

The chemical composition of leaves and berries are approximately the same, but berries contain higher amount of phenolic compounds and organic acids; moreover, unlike leaves, berries accumulate anthocyanidin derivatives [5].

Owing to the presence of various phenolic compounds, lingonberry leaves and berries possess the antibacterial [6], anti-inflammatory [7], and antioxidative activities [8]. Extracts of lingonberry leaves have shown to possess multiple beneficial properties e.g., diuretic, analgesic and urinary antiseptic ones. They also display the anti-cough, phlegm removing, anti-inflammatory, neuroprotective, and antioxidant activity [9–11].

Nowadays at the pharmaceutical market of Ukraine one can find a number of dietary supplements for treating urological diseases. One of reasons is that 10% of the Ukrainian population has signs of chronic diseases of the genitourinary system. Another reason is a high consumer interest in purchasing dietary supplements. According to the Law of Ukraine [12] dietary supplements are not subjected to the qualitative and quantitative analysis, as a result, the quality of these products are casted doubt.

Thus, the aim of study was to determine the qualitative composition and the quantitative content of phenolic compounds in dietary supplements with lingonberry.

Materials and methods

Rutin, arbutin, hydroquinone and chlorogenic acid were of analytical grade and purchased in Sigma-Aldrich.

Three dietary supplements with lingonberry of different Ukrainian manufactures were chosen for the analysis.

“Extract of lingonberry” contains the extract of lingonberry fruits according to the information on the label; the dosage form is drops (30 mL), the manufacturer is “MEDAGROPROM”, Dnipro.

“Lingonberry” contains leaves and shoots of lingonberry according to the information on the label; the dosage form is tablets (100 pcs), the manufacturer is “Danikafarm”, Kharkiv.

“Lingonberry nano” contains leaves and fruits of lingonberry, intercellular and intracellular liquid of lingonberry leaves and fruits according to the information on the label; the dosage form is tablets (100 pcs), the manufacturer is “LSS SYSTEM”, Kharkiv.

All solvents and reagents were of analytical grade. “Sorbfil-PTSH-AF-A-UV” plates were used for the TLC analysis.

The quantitative analysis of biologically active compounds was performed on an UV-1000 UV-spectrophotometer (China) with a matched 1 cm quartz cell.
Preparation of standard solutions for the TLC analysis
0.010 g (accurate weight) of each arbutin, rutin, chlorogenic acid and hydroquinone was transferred into four 50 mL measuring flasks, dissolved in 96% ethanol and diluted to the volume with the same solvent.

Preparation of the standard solution of rutin for the spectrophotometric assay
0.050 g (accurate weight) of rutin was transferred into a 100.0 mL measuring flask, dissolved in 96% ethanol and diluted to the volume with the same solvent. 1.0 mL of the solution prepared was taken, transferred into a 25.0 mL measuring flask and diluted to the volume with 96% ethanol.

Preparation of the standard solution of arbutin for the spectrophotometric assay
0.015 g (accurate weight) of arbutin was transferred into a 100.0 mL measuring flask, dissolved in distilled water and diluted to the volume with the same solvent.

Preparation of the solution of berlin blue
10.0 g of iron(III) chloride and 0.5 g of potassium hexacyanoferrate were dissolved in 100.0 mL of distilled water.

TLC analysis assay
The powder of 7 crushed tablets of each dietary supplement was completely dissolved in 96% ethanol, filtered into a 50.0 mL measuring flask, dissolved in 96% ethanol and diluted to the volume with the same solvent. 5.0 mL of “Extract of lingonberry” (MEDAGROPROM) was taken, placed into a 25.0 mL measuring flask, and diluted to the volume with 96% ethanol.

To identify arbutin, the developing system consisting of ethyl acetate/methanol/water (100:13.5:10) was used; for flavonoids and hydroxycinnamic acids the developing system was ethyl acetate/methanol/water (100:13.5:10). To identify rutin, the developing system was ethyl acetate/methanol/water (100:10:40). To identify chlorogenic acid and hydroquinone was transferred into a 50.0 mL measuring flask, dissolved in 96% ethanol and filtered into a 100.0 mL measuring flask. 5.0 mL of “Extract of lingonberry” (MEDAGROPROM) was dissolved in a 25.0 mL measuring flask and diluted to the volume with 96% ethanol. An aliquot of the solutions prepared was mixed with 1.0 mL of 1 M Folin-Ciocalteu reagent, the mixture was diluted to the volume of 25.0 mL with 20% NaOH solution. The optical density of the solutions was measured at 760 nm in 30 min after preparation. The calibration curve was plotted using gallic acid, the calibration equation was \( y = 0.1055x + 0.1745 \) \( (r^2 = 0.9951) \). The total amount of phenolic compounds in “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) dietary supplements in 1 tablet was calculated by the following equation and expressed with reference to gallic acid:

\[
X(\%) = \frac{C_s \cdot K_{\text{dil}} \cdot m_{\text{aver tab}} \cdot 100 \cdot 100}{m_s \cdot (100 - W)},
\]

where \( C_s \) – is the concentration of gallic acid according to the calibration curve, \( C \times 10^{-6} \), g mL\(^{-1}\); \( m_s \) – is the sample weight, g; \( m_{\text{aver tab}} \) – is the average mass of a tablet, g; \( K_{\text{dil}} \) – is the coefficient of dilution; \( W \) – is the percentage of moisture, %.

The total content of phenolic compounds in “Extract of lingonberry” (MEDAGROPROM) dietary supplement in the total volume of drops was calculated by the equation and expressed with reference to gallic acid:

\[
X(\%) = \frac{C_s \cdot K_{\text{dil}} \cdot V_{\text{drops}} \cdot 100}{V_{\text{al}}},
\]

where \( C_s \) – is the concentration of gallic acid according to the calibration curve; \( C \times 10^{-6} \), g mL\(^{-1}\); \( V_{\text{al}} \) – is the volume of an aliquot, mL; \( V_{\text{drops}} \) – is the total volume of drops, mL; \( K_{\text{dil}} \) – is the coefficient of dilution.

Assay for the total content of flavonoids
3.2 g (accurate weight) of “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) was dissolved in 96% ethanol and filtered in a 50.0 mL measuring flask. 5.0 mL of “Extract of lingonberry” (MEDAGROPROM) was added to a 25.0 mL measuring flask and diluted to the volume with 96% ethanol.
ethanol (Solution A). An aliquot of Solution A was mixed with 1.0 mL of 2% AlCl₃ solution in methanol and diluted to the volume of 25.0 mL with 5% solution of glacial acetic acid in methanol. The solution prepared was allowed to stand for 30 min, and its optical density was measured at 417 nm. An aliquot of Solution A diluted to the volume of 25.0 mL with 5% solution of glacial acetic in methanol was used as a compensation liquid [14]. The total amount of flavonoids in “Lingonberry” (Danika-farm) and “Lingonberry nano” (LSS SYSTEM) dietary supplements in 1 tablet was calculated by the equation and expressed with reference to rutin:

\[
X(\%) = \frac{A \cdot K_{\text{dil}} \cdot m_{\text{aver tab}} \cdot 100 \cdot 100}{A_s \cdot m_s \cdot (100 - W)},
\]

where \(A\) – is the absorbance of the test solution; \(A_s\) – is the absorbance of the standard solution of rutin; \(m_s\) – is the sample weight, g; \(m_{\text{aver tab}}\) – is the average mass of a tablet, g; \(K_{\text{dil}}\) – is the coefficient of dilution; \(W\) – is the percentage of moisture, %.

The total content of flavonoids in “Extract of lingonberry” (MEDAGROPROM) dietary supplement in the total volume of drops was calculated by the equation and expressed with reference to rutin:

\[
X(\%) = \frac{A \cdot V_{\text{drops}} \cdot 100}{A_s \cdot V_{s}},
\]

where \(A\) – is the absorbance of the test solution; \(A_s\) – is the absorbance of the standard solution of rutin; \(V_{s}\) – is the volume of an aliquot, mL; \(V_{\text{drops}}\) – is the total volume of drops, mL; \(K_{\text{dil}}\) – is the coefficient of dilution.

**Assay for the total content of hydrocin-

namic acids**

3.2 g (accurate weight) of “Lingonberry” (Danika-farm) and “Lingonberry nano” (LSS SYSTEM) was dissolved in 96% ethanol and filtrated in a 50.0 mL measuring flask. 5.0 mL of “Extract of lingonberry” (MEDAGROPROM) was dissolved in a 25.0 mL measuring flask and diluted to the volume with 96% ethanol. An aliquot of the solutions prepared was mixed with 2.0 mL of 0.5 M HCl, 2.0 mL of 10% NaNO₂, 2.0 mL of 10% Na₂MoO₄, 2.0 mL of 8.5% NaOH and diluted to the volume of 25.0 mL with distilled water. The optical density of the solutions was measured at wavelength of 525 nm in 30 min after preparation. The compensation liquid was a mixture of 1.0 mL of the extract solution, 2.0 mL of 0.5 M HCl, 2.0 mL of 8.5% NaOH diluted with distilled water to the volume of 25.0 mL [14]. The total amount of hydroxycinnamic acids in “Lingonberry” (Danika-farm) and “Lingonberry nano” (LSS SYSTEM) dietary supplements in 1 tablet was calculated by the following equation and expressed with reference to chlorogenic acid:

\[
X(\%) = \frac{A \cdot K_{\text{dil}} \cdot m_{\text{aver tab}} \cdot 100}{188 \cdot m_s \cdot (100 - W)},
\]

where \(A\) – is the absorbance of the test solution; \(188\) – is the specific adsorption coefficient of chlorogenic acid; \(m_s\) – is the sample weight, g; \(m_{\text{aver tab}}\) – is the average mass of a tablet, g; \(K_{\text{dil}}\) – is the coefficient of dilution; \(W\) – is the percentage of moisture, %.

The total content of hydroxycinnamic acids in “Extract of lingonberry” (MEDAGROPROM) dietary supplement in the total volume of drops was calculated by the equation and expressed with reference to chlorogenic acid:

\[
X(\%) = \frac{A \cdot K_{\text{dil}} \cdot V_{\text{drops}}}{188 \cdot V_{s}},
\]

where \(A\) – is the absorbance of the test solution; \(188\) – is the specific adsorption coefficient of chlorogenic acid; \(V_{s}\) – is the volume of an aliquot, mL; \(V_{\text{drops}}\) – is the total volume of drops, mL; \(K_{\text{dil}}\) – is the coefficient of dilution.

**Assay for the total content of hydroqui-

none derivatives**

3.2 g (accurate weight) of “Lingonberry” (Danika-farm) and “Lingonberry nano” (LSS SYSTEM) was dissolved in 96% ethanol and filtrated in a 50.0 mL measuring flask. 5.0 mL of “Extract of lingonberry” (MEDAGROPROM) was dissolved in a 25.0 mL measuring flask and diluted to the volume with 96% ethanol. 20.0 mL of the aliquot of the solution obtained was evaporated to dryness on a water bath at 80°C. The residue was dissolved in 20.0 mL of water, filtered through a paper filter, transferred into a separating funnel, than mixed with 1 mL of 2% 4-amino-2,3-dimethyl-1-phenylpyrazolin-5-one solution, 0.5 mL of 3.3% ammonia solution, 1 mL of 8% potassium ferri-

nycide and extracted twice with 20.0 mL of chloro-

form for 10 min. The absorbance was measured
at wavelength of 455 nm in 30 min after extraction. The compensation liquid was chloroform [14]. The total content of hydroquinone derivatives in “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) dietary supplements in 1 tablet was calculated by the equation and expressed with reference to arbutin:

\[ X(\%) = \frac{A \cdot K_{\text{dil}} \cdot m_{\text{aver tab}} \cdot 100 \cdot 100}{A_{\text{st}} \cdot m_{\text{a}} \cdot (100 - W)} , \]

where \( A \) – is the absorbance of the test solution; \( A_{\text{st}} \) – is the absorbance of the standard solution of arbutin; 
\( m_{\text{a}} \) – is the sample weight, g; 
\( m_{\text{aver tab}} \) – is the average mass of a tablet, g; 
\( K_{\text{dil}} \) – is the coefficient of dilution; 
\( W \) – is the percentage of moisture, %.

The total content of hydroquinone derivatives in “Extract of lingonberry” (MEDAGROPROM) dietary supplement in the total volume of drops was calculated by the equation and expressed with reference to arbutin:

\[ X(\%) = \frac{A \cdot K_{\text{dil}} \cdot V_{\text{drops}} \cdot 100}{A_{\text{st}} \cdot V_{\text{al}}} , \]

where \( A \) – is the absorbance of the test solution; 
\( A_{\text{st}} \) – is the absorbance of the standard solution of arbutin; 
\( V_{\text{al}} \) – is the volume of an aliquot, mL; 
\( V_{\text{drops}} \) – is the total volume of drops, mL; 
\( K_{\text{dil}} \) – is the coefficient of dilution.

## Results and discussion

The TLC method was applied for the qualitative analysis of dietary supplements. Hydroquinone derivatives were detected in the ethyl acetate/methanol/water (100:13.5:10) developing system. Substances were detected at wavelengths of 254 and 325 nm. The chromatogram showed the dominant bands with the value of \( R_f = 0.40 \) (arbutin), \( R_f = 0.80 \) (hydroquinone). The chromatogram was then sprayed by 10% solution of Berlin blue; blue bands with the same \( R_f \) values were also detected.

For identification of flavonoids and hydroxycinnamic acids in the dietary supplements analyzed the ethyl acetate/glacial acetic acid/formic acid/water (100:11:11:26) developing system was used. The TLC plate was sprayed with 10% solution of NaOH; as the result, yellow bands with values of \( R_f = 0.35 \) (rutin), \( R_f = 0.40 \) (chlorogenic acid) and \( R_f = 0.55 \) (hyperoside) appeared.

The total content of phenolic compounds was determined by the Folin-Ciocalteu method. As shown in Table, the highest amount of phenolic compounds was found in “Extract of lingonberry” (MEDAGROPROM) (8.70%); at the same time, only minor amounts of phenolic compounds were found in “Lingonberry nano” (LSS SYSTEM) (0.30%) and “Lingonberry” (Danikafarm) (0.26%).

“Extract of lingonberry” (MEDAGROPROM) was much richer in flavonoids (6.37%) than “Lingonberry” (Danikafarm) (0.15%) and “Lingonberry nano” (LSS SYSTEM) (0.12%).

The highest content of hydroxycinnamic acids was found in “Extract of lingonberry” (MEDAGROPROM) (0.94%), followed by “Lingonberry nano” (LSS SYSTEM) (0.13%) and “Lingonberry” (Danikafarm) (0.06%).

The total content of hydroquinone derivatives was dominant in “Extract of lingonberry” (MEDAGROPROM) (1.01%), while “Lingonberry” (Danikafarm) (0.03%) and “Lingonberry nano” (LSS SYSTEM) (0.04%) were quite poor in hydroquinones.

All dietary supplements analyzed contained hydroquinone derivatives, flavonoids and hydroxycinnamic acids. However, the content of biological active substances is in minor amounts in “Lingonberry” (Danikafarm) and “Lingonberry nano”

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Total phenolic compounds</th>
<th>Total flavonoids</th>
<th>Total hydroxycinnamic acids</th>
<th>Total hydroquinone derivatives</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>% (a)</td>
<td>mg/L</td>
<td>%</td>
<td>mg/L</td>
</tr>
<tr>
<td>“Lingonberry” (Danikafarm)</td>
<td>0.26</td>
<td>0.78 mg</td>
<td>0.15</td>
<td>0.59 mg</td>
</tr>
<tr>
<td>“Lingonberry nano” (LSS SYSTEM)</td>
<td>0.30</td>
<td>1.10 mg</td>
<td>0.12</td>
<td>0.44 mg</td>
</tr>
<tr>
<td>“Extract of lingonberry” (MEDAGROPROM)</td>
<td>8.70</td>
<td>87 mg mL⁻¹</td>
<td>6.37</td>
<td>63.7 mg mL⁻¹</td>
</tr>
</tbody>
</table>

Notes: [a] the percentage of the biologically active compounds in 1 tablet (“Lingonberry” and “Lingonberry nano”) and in the total volume of drops (“Extract of lingonberry”); [b] the content of the biologically active compounds in 1 tablet (“Lingonberry” and “Lingonberry nano”) and in the total volume of drops (“Extract of lingonberry”).
(LSS SYSTEM) dietary supplements, and this fact casts doubt on the quality of these dietary supplements.

According to the section of “Dietary supplements” of the State of Pharmacopoeia of Ukraine (2.0) [14] the minimum content of each vitamin and/or mineral substance (nutrients) in the recommended daily amount of dietary supplements should be at least 15% of the recommended daily intake. The regulation document of daily intake of nutrients “Norms of physiological needs of the population of Ukraine in basic nutrients and energy” [14] declares that daily intake of flavonoids is 250 mg. In this regard the recommended daily amount of flavonoids taken with a dietary supplement is 37.5 mg. According to the instructions for use of dietary supplements, 10 drops of “Extract of lingonberry” (MEDAGROPROM) are taken in 200 mL of water 3 times a day, 2 tablets of “Lingonberry” (Danikafarm) are taken 3 times a day, and 2 tablets of “Lingonberry nano” (LSS SYSTEM) are taken 3 times a day. Thus, a daily amount of flavonoids in “Extract of lingonberry” (MEDAGROPROM) is 110.49 mg, in “Lingonberry” (Danikafarm) – 3.52 mg, and in “Lingonberry nano” (LSS SYSTEM) – 2.61 mg. The results obtained show that “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) dietary supplements do not meet the requirements of the State of Pharmacopoeia of Ukraine 2.0 in relation to the recommended daily amount of nutrients of dietary supplements, therefore, these dietary supplements are of poor quality and cannot be recommended for application.

**Conclusions**

“Extract of lingonberry” (MEDAGROPROM) dietary supplement meets the requirements of the State of Pharmacopoeia of Ukraine 2.0, whereas “Lingonberry” (Danikafarm) and “Lingonberry nano” (LSS SYSTEM) do not comply with the requirements. Based on the results of the study it can be concluded that the problem of compliance of dietary supplements is relevant today and requires the introduction of regulatory documentation for the detection and determination of biologically active substances in dietary supplements.

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