THE STUDY OF DEPENDENCE OF THE PRENYL FLAVONOID CONTENT IN THE HOP DRY EXTRACT ON TEMPERATURE PARAMETERS OF ITS OBTAINING

O.O.Dobrovolnyi
SIC “Borshchahivskiy Chemical Pharmaceutical Plant”, PJSC, Kyiv

Key words: hop cones; extractives; prenyl flavonoids; estrogenic activity; evaporation, drying

The use of plants containing phytoestrogens, with its historical length may be compared with the history of medicine in general. One of the most interesting sources of compounds with the estrogen-like action is a hop cones. The results of the pharmacological studies of hop extracts indicate the relationship between estrogenic activity of the objects and the fraction of prenyl flavonoids: xanthohumol (X), desmethylxanthohumol (DMX), isoxanthohumol (IX), 8-prenylisoflavone (8-PN) and 6-prenylisoflavone (6-PN) in the hop extract has been researched. The objects of the research were the samples of the hop liquid extract after extraction of the raw material and the dry extract after evaporation and drying. The extraction of hop cones has been carried out by the two-step filtration method extraction with the pretreatment of the raw material with steam. At the first step the hop raw material was extracted with n-hexane with the subsequent extraction of the extraction cake with water-alcohol solvent at the second step. The aqueous-ethanol extract was evaporated and dried in vacuum to prevent possible destruction of prenyl flavonoids. The quantitative content of xanthohumol, isoxanthohumol, 8-prenylisoflavone and 6-prenylisoflavone has been determined by HPLC in the samples of liquid and dry extracts. It has been found that sparing conditions of evaporation and drying used has led to decreasing the quantitative content of X, IX, 8-PN and 6-PN in the dry extract compared with the content of these compounds in the water-alcohol liquid extract in the range of 14, 16, 8, 14%, respectively.

Experimental Part

The starting materials were hop cones of “Xantha” cultivar (the product of the Polissya Institute of Agriculture at the NAASU) harvested in 2012, n-hexane, ethanol and purified water. For cyclization of chalcones X and DMX to the corresponding flavanones IX, 8-PN and 6-PN the hop cones were pretreated by water steam for 1 hour. After thermal pretreatment the remaining moisture was removed from the raw material, and hop cones were extracted in two steps. All extraction steps were performed by the method of filtration extraction at the room temperature. At the first step the hop cones treated was extracted with n-hexane to the drug extract ratio (DER) of 1:12 with the subsequent removal of the residual extractant from the extraction cake [2]. At the second step the hop semi-product (the extraction cake) was extracted with 70% (vol.) ethanol to DER of 1:10. Removal of the solvent from the liquid extract and subsequent drying were conducted in a Büchi R134 rotary evaporator at the water-bath temperature 60-63°C and at the vacuum operating range of 80-38 mBar. Before drying of the liquid extract the dry residue content and the quantitative content of X, IX, 8-PN and 6-PN were determined and the yield of extractives in it was calculated. The quantitative content of X, IX, 8-PN and 6-PN was determined in the hop dry extract obtained.

The dry residue content in the samples was determined by the method according to the State Pharmacopoeia of Ukraine [1].

The assay of X, IX, 8-PN and 6-PN in the hop dry extracts obtained was carried out by the liquid chromato-
graphy method. In the assay the approach of determination of the given compounds using secondary standards was applied [4].

Chromatographic procedure was carried out in an Ultimate 3000 chromatograph with a UV detector at the temperature of 50°C. As a mobile phase A 0.25% (v/v) formic acid R was used, and 0.25% (v/v) formic acid R in acetonitrile R was used as a mobile phase B. The chromatographic conditions were the following: the flow rate – 1 mL/min; the column – Purospher STAR RP 18-e (5 μm) 250*4.0 mm; the injection volume – 20 μL; detection – at 370 nm (X) and at 290 nm (IX, 8-PN, 6-PN). The gradient programme was 0 min, 80:20 (A:B); 3 min, 80:20; 3→25 min, 60:40; 25→37 min, 60:40; 37→55 min, 40:60; 55→56 min, 10:90. Determination of X was carried out using quercetin, and determination of IX, 8-PN, 6-PN was performed using naringenin as secondary standards.

Identification of peaks was carried out using the same conditions in an Ultimate 3000 chromatograph with MS / MS detector and literature data [3].

The yield of the extractives from the raw material extracted was calculated as:

$$ D = \frac{\omega \times V}{m} \%,$$

where: V – is the volume of the liquid extract, ml; ω – is a dry residue in the liquid extract, %; m – is the mass of the raw material used for extraction, g.

**Results and Discussion**

The hop prenyl flavonoids X, DMX, IX, 8-PN and 6-PN are relatively unstable compounds and under the certain thermodynamic conditions are capable to transform. In particular, chalcones X and DMX can be subjected to thermal isomerization, and it is confirmed by the conversion of X to IX during wort boiling in brewing [5, 6, 9].

Obtaining of a dry extract is always associated with the use of high temperatures at the stages of evaporation and drying. The nature of the thermal effects on the quantitative content of X, IX, 8-PN and 6-PN was evaluated by the content of these compounds in the water-alcohol extract before its evaporation and in the dry extract obtained. In order to prevent possible destruction of prenyl flavonoids the evaporation and drying of hop cones extracts were carried out in vacuum under the sparing conditions appropriate for subsequent manufacturing-scale production. The data obtained are presented in Table.

The experimental data show decrease of the quantitative content of X, IX, 8-PN and 6-PN in the dry extract compared with the content of these compounds in the water-alcohol liquid extract by 14.04%, 16.26%, 8.06% and 13.94%, respectively. It indicates the negative effect of evaporation and drying of the liquid extract on

| Criteria | Liquid extract | Dry extract |
|----------|----------------|-------------|
| Xanthohumol (X), % | 1.688 | 1.451 |
| Isoxanthohumol (IX), % | 0.412 | 0.345 |
| 8-prenyl naringenin (8-PN), % | 0.062 | 0.057 |
| 6-prenyl naringenin (6-PN), % | 0.208 | 0.179 |
| Yield of extractives, % | 24.55 | |

Note: the quantitative content of (X, IX, 8-PN, 6-PN) are calculated with reference to dried substance.

Fig. 1. The chromatogram of the hop water-alcoholic liquid extract at 370 nm.
quantitative characteristics of the dry extract even with the sparing conditions applied with reduced boiling temperature in the vacuum environment. The chromatographic profiles of the samples are characterized by a similar set of major peaks and differ in their intensity decrease due to the aforementioned decrease in the quantitative content of X, IX, 8-PN and 6-PN in the dry extract obtained in comparison with the water-alcohol liquid extract. Therefore, the temperature increase and/or decrease of the vacuum depth, i.e. parameters that can increase the boiling point in the process of evaporation and drying of the extract may lead to a greater reduction of the quantitative content of X, IX, 8-PN and 6-PN in the hop dry extract. The chromatograms of the test samples are presented in Fig. 1, 2, 3, 4.

![Fig. 2. The chromatogram of the water-alcoholic liquid extract at 290 nm.](image)

![Fig. 3. The chromatogram of hop dry extract at 370 nm.](image)
CONCLUSIONS
1. For the first time the influence of temperature parameters at the stages of evaporation and drying of the hop cones liquid extract on the quantitative content of xanthohumol, isoxanthohumol, 8-prenylnaringenin and 6-prenylnaringenin in the dry extract has been investigated.

2. It has been found that obtaining of the hop cones dry extract as a finished product provides the loss of the quantitative content of these prenyl flavonoids in the technological process.

3. The research results will be considered in the further development of the production process of the hop dry extract as the API.

REFERENCES
1. Державна фармакопея України / Державне підприємство «Науково-експертний фармакопейний центр». – 1-е вид. – Х: Державне підприємство «Науково-експертний фармакопейний центр», 2004. – Доп. 1. – 520 с.
2. Шматенко О.П., Добровольний О.О., Савицький В.Л., Страшний В.В. Вивчення умов екстрагування речовин ліпофільної природи з суспільд хмелю // Проблеми військової охорони здоров’я. Зб. наук. праць Української військово-медичної академії. – К.: Українська військово-медична академія, 2013. – Вип. 40. – С. 282-289.
3. Ceslova L., Holcapek M., Fidler M. et al. // J. of Chrom. A. – 2009. – Vol. 1216. – P. 7249-7257.
4. Dhooghe L., Naessens T., Heyerick A. et al. // Talanta. – 2010. – Vol. 83, Iss. 2. – P. 448-456.
5. Karabin M., Jelinek L., Kinčl T. et al. // J. Inst. Brew. – 2013. – Vol. 119. – P. 98-102.
6. Krofta K. // Kvasny Prum. – 2010. – Vol. 56, Iss. 1. – P. 2-9.
7. Milligan S., Kalita J., Pocock V. et al. // Reproduction. – 2002. – Vol. 123, Iss. 2. – P. 235-242.
8. Milligan S.R., Kalita J.C., Pocock V. et al. // J. Clin. Endocrinol. Metab. – 2000. – Vol. 85, Iss. 12. – P. 4912-4915.
9. Possemiers S., Heyerick A., Robbens V. et al. // J. Agric. Food Chem. – 2005. – Vol. 53. – P. 6281-6288.
10. Schaefer O., Hümpel M., Fritzemeier K.H. et al. // J. Steroid. Biochem. and Mol. Biol. – 2003. – Vol. 84, Iss. 2-3. – P. 359-360.

Fig. 4. The chromatogram of the hop dry extract at 290 nm.
ДОСЛІДЖЕННЯ ЗАЛЕЖНОСТІ ВМІСТУ ПРЕНИЛОВИХ ФЛАВОНОІДІВ У СУХОМУ ЕКСТРАКТІ ХМЕЛЮ ВІД ТЕМПЕРАТУРНИХ ПАРАМЕТРІВ ПРОЦЕСУ ЙОГО ОДЕРЖАННЯ

О.О.Добровольний

Ключові слова: супліддя хмелю; екстрактивні речовини; пренилові флавоноїди; естрогенна актівність; упарювання; сушка

У процесі розробки технології одержання сухого екстракту суплідь хмелю в якості АФІ з естрогеноподібною дією досліджували ключові фактори, здатні впливати на якість готового продукту. Беручи до уваги те, що пренилові флавоноїди хмелю є відносно нестійкими сполуками та при певних термодинамічних умовах здатні до перетворень або деструкції, було досліджено вплив температурних параметрів упарювання та сушки рідкої витяжки хмелю на кількісний вміст ксантохумолу (Х), ізоксантохумолу (ІХ), 8-пренилнарингеніну (8- PN) та 6-пренилнарингеніну (6- PN) в сухому екстракті. Об’єктами дослідження були зразки водно-спиртової витяжки, одержаної після екстрагування сировини та сухого екстракту після упарювання та сушки. Екстрагування супліддя хмелю здійснювалося методом фільтраційної екстракції в два ступені з попередньою термічною обробкою вихідної сировини водяною парою. На першому ступені сировину екстрагували н-гексаном, після чого на другому ступені проводили екстрагування шроту водно-спиртовим розчинником. З метою мінімізації можливої деструкції пренилових флавоноїдів одержану водно-спиртову витяжку упарювали та висушували у вакуумі. Кількісний вміст Х, ИХ, 8-PN та 6-PN в досліджуваних зразках визначали методом ВЕРХ. Встановлено, що застосовані щадні умови упарювання та сушки привели до зниження кількісного вмісту Х, ИХ, 8-PN та 6-PN в сухому екстракті порівняно з їх вмістом у водно-спиртовій витяжці в межах 14, 16, 8, 14% відповідно.

ИССЛЕДОВАНИЕ ЗАВИСИМОСТИ СОДЕРЖАНИЯ ПРЕНИЛОВЫХ ФЛАВОНОИДОВ В СУХОМ ЭКСТРАКТЕ ХМЕЛЯ ОТ ТЕМПЕРАТУРНЫХ ПАРАМЕТРОВ ПРОЦЕССА ЕГО ПОЛУЧЕНИЯ

А.А.Добровольный

Ключевые слова: соцветия хмеля; экстрактивные вещества; прениловые флавоноиды; эстрогенная активность; упаривание; сушка

В процессе разработки технологии получения сухого экстракта соцветий хмеля в качестве АФИ с эстрогеноподобным действием исследовали ключевые факторы, способные влиять на качество готового продукта. Учитывая то, что прениловые флавоноиды хмеля являются относительно нестабильными соединениями и при определенных термодинамических условиях склонны к превращениям или деструкции, было исследовано влияние температурных параметров упаривания и сушки жидкого извлечения хмеля на количественное содержание ксантохумола (Х), изоксантохумола (ІХ), 8-пренилнарингенина (8-PN) и 6-пренилнарингенина (6-PN) в сухом экстракте. Объектами исследования были образцы водно-спиртового извлечения, полученные после экстрагирования сырья и сухого экстракта после упаривания и сушки. Экстракцию соцветий хмеля проводили методом фильтрационной экстракции в две ступени с предварительной обработкой сырья водяным паром. На первой ступени сырье экстрагировали н-гексаном, после чего на второй ступени проводили экстракцию шрота водно-спиртовым растворителем. С целью минимизации возможной деструкции прениловых флавоноидов полученное водно-спиртовое извлечение упаривали и сушили в вакууме. Количественное содержание Х, ИХ, 8-PN и 6-PN в исследуемых образцах определяли методом ВЭЖХ. Установлено, что использованные щадящие условия упаривания и сушки привели к снижению количественного содержания Х, ИХ, 8-PN и 6-PN в сухом экстракте в сравнении с содержанием в водно-спиртовом извлечении в пределах 14, 16, 8, 14% соответственно.