Spectrophotometric determination of a substance trifusol in a veterinary suppository

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A – research concept and design; B – collection and/or assembly of data; C – data analysis and interpretation; D – writing the article; E – critical revision of the article; F – final approval of the article

The purpose of the work was to develop of a method for the quantitative determination of piperidinium 2-[5-(2-furyl)-4-phenyl-1,2,4-triazol-3-ylthio] acetate (trifuzol) as part of a dosage form – an effervescent intrauterine suppository using spectrophotometry in the ultraviolet region spectrum and its validation according to State Pharmacopoeia of Ukraine.

Materials and methods. The study was used a working standard sample of trifuzol, intrauterine effervescent veterinary suppositories, 1.0 g of trifuzol, as a solvent – purified water. Analytical equipment: Specord 200 spectrophotometer, electronic scales ABT-120-5DM, ultrasonic bath ELMASONICE 60 H, class A measuring dishes. Method of spectrophotometric analysis was used.

Results. A spectrophotometric method has been developed and validated for the quantitative determination of trifuzol in a dosage form – an intrauterine effervescent veterinary suppository, based on measuring the absorption of an aqueous solution of the drug at 278 nm.

The methodology tally the requirements of State Pharmacopoeia of Ukraine for such validation characteristics as specificity, linearity, precision, correctness, and robustness. The analysis of the predicted total uncertainty of the analysis was showed the reproducibility of the method and the possibility of its application in other laboratories.

Conclusions. A method for the quantitative determination of trifuzol in the composition of the dosage form, an effervescent intrauterine suppository, according to the requirements of State Pharmacopoeia of Ukraine, was developed and validated. It was proved that according to such validation characteristics as linearity, specificity, precision, correctness, and robustness, the technique is correct.

Key words: spectrophotometry, trifuzol, veterinary suppository, quantitative determination.

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Endometritis of different etiologies is the most common form of postnatal pathology in cows, which can take a mass character and cause significant economic damage to both the economy and the state as a whole. Against the background of the intensification of all branches of animal breeding, animal diseases involving damage to the sexual sphere prompted the search, creation, and improvement of existing medicines.

The use of the substance piperidinium 2-[5-(2-furyl)-4-phenyl-1,2,4-triazole-3-iltil] acetate (trifusol), as an active substance in the composition of the effervescent intruterine suppository, provides a sufficiently high therapeutic efficacy in the treatment, particularly of purulent postpartum cattle endometritis. Its use can improve the course of the pathological process and individual blood parameters [5,6].

The development of new accurate and sensitive methods for quantifying trifusol in new dosage forms is an immediate need at the stage of ensuring proper quality control of drugs, in terms of safe, rational and effective therapy.

For other dosage forms, an express and easy-to-use method of trifusol quantitative determination have already been proposed [1]. Any analytical technique (including the technique of quality control of a particular drug) that may be proposed for inclusion in a normative document or by means of which official tasks will be performed (for example, official opinion on the quality of the object) should be validated [2–4]. Only a specific method for a specific dosage form can be validated.

Aim

Thus, the purpose of our work was to develop a method for quantitative determination of piperidinium 2-[5-(2-furyl)-4-phenyl-1,2,4-triazole-3-iltil] acetate in the dosage form of effervescent intruterine suppository using spectrophotometry in the ultraviolet region of the spectrum and its validation, according to the State Pharmacopeia of Ukraine.

Materials and methods

Study objects, solvents and equipment. The objects of the investigation were intruterine effervescent veterinary suppositories, 1.0 g of trifusol substance. The indicated dosage form was prepared extemporally according to the prescription [5]. Purified water was used as a solvent. As working standards of trifusol were used.

Analytical equipment: spectrophotometer Specord 200, electronic scales AVT-120-5DM, ultrasonic bath ELMASONICE 60 H, A class measuring ware.

General methodology for quantification of trifusol substance. Aliquot of trifusol (0.050 g) was placed in a measuring flask of 100.00 ml and bring to the mark with purified water. Then 1.00 ml of the resulting solution was transferred to a measuring flask with a capacity of 25.00 ml, and diluted with a solvent to the mark. Optical density was measured on the background of a compensation solution (purified water) at an analytical wavelength of 278 nm.

Results

Method for quantitative determination of trifusol in a veterinary effervescent suppository for intruterine use. I suppository was placed in a glass of 50 ml, added 25 ml of distilled water and heated in an ultrasonic bath until the suppository was completely melted, then cooled and decanted in a measuring flask of 100.00 ml. This operation was repeated twice. The content of the flask was diluted with water to the mark and stirred thoroughly. 5.00 ml of the resulting solution was transferred to a measuring flask with a capacity of 25.00 ml, and diluted with a solvent to the mark. Optical density was measured on the background at wavelength 278 nm. At the same time, a 1.00 ml of 0.05 % trifusol comparison solution was determined. The calculation of the active substance content was done according to
the usual formula.

Prediction of complete methodology uncertainty. To verify that the technique would be replicated in other laboratories, it was not enough just the results of the validation in one laboratory, the level of equipment which may be much higher than allowed by the State Pharmacopoeia of Ukraine. Calculations of the forecast of complete methodology uncertainty in accordance with the requirements of the State Pharmacopoeia of Ukraine were created specifically for this purpose. The total uncertainty of the analysis technique was based not only on the real total uncertainty of sample preparation but also on the maximum uncertainty for a specific equipment type [7,8].

According to the requirements of the State Pharmacopoeia of Ukraine, uncertainty calculations were performed to the maximum permissible errors for measuring dishes, scales, and spectrophotometer as a final analytical operation in analysis (ΔFAO = 0.70 %) (Table 1).

It can be seen from Fig. 1 that the most significant uncertainty in sample preparation is introduced by operations 6 – taking a standard sample of trifusol, as well as 2, 4 and 8 – taking aliquots by pipettes 1.00 and 5.00 ml. This distribution of uncertainty in sample preparation is quite typical for the quantitative determination of drugs.

Thus, the projected total uncertainty of the analysis results (ΔAs = 1.53 %) does not exceed the critical value (Δmax As = 3.20), i.e. the technique will be reproducible and correct in other laboratories as well.

Discussion

The specificity of the technique was established by determining the effect of auxiliary substances (Table 2), which are part of the studied dosage form, on the results of trifusol quantitative determination. For this purpose, the placebo solution (As) absorption in purified water was measured in the concentrations of the corresponding prescription [5], making 3 measurements with cuvette extraction. At the same time the optical density of the comparison solution (Ast) was measured. The following was found out: Ablank = 0.0031; Ast

### Table 1. Uncertainty calculation of sample preparation for quantitative determination of trifusol content in the intrauterine suppository

| Sample preparation operation | Calculation formula parameter | Uncertainty,% |
|------------------------------|-------------------------------|---------------|
| Test solution                |                               |               |
| Bringing up to volume in a measuring flask with a capacity of 100 ml | 100 | 0.12 |
| Taking aliquots dilution with a pipette in 5 ml | 5 | 0.69 |
| Bringing up to volume in a measuring flask with a capacity of 100 ml | 100 | 0.12 |
| Taking aliquots dilution with a pipette in 1 ml | 1 | 0.74 |
| Bringing up to volume in a measuring flask with a capacity of 25 ml | 25 | 0.23 |

| Reference solution           |                               |               |
|------------------------------|-------------------------------|---------------|
| Taking a weighted amount of trifusol standard design sample | m0 | 0.2 mg / 50 mg·100 % = 0.40 |
| Bringing up to volume in a measuring flask with a capacity of 100 ml | 100 | 0.12 |
| Taking aliquots dilution with a pipette in 1 ml | 1 | 0.74 |
| Bringing up to volume in a measuring flask with a capacity of 25 ml | 25 | 0.23 |

\[ \Delta SP = \sqrt{(0.40^2 + 0.12^2 + 0.74^2 + 0.23^2 + 0.12^2 + 0.69^2 + 0.12^2 + 0.74^2 + 0.23^2 + 0.12^2 + 0.74^2 + 0.23^2} = 1.37 \%

### Table 2. Composition of excipients in intrauterine suppositories with trifusol

| Excipients            | Quantity, g |
|-----------------------|-------------|
| Sodium lauryl sulfate | 0.2         |
| Polyethylene oxide mixture 1500 i 400 – 1:9 | Up to 20 |

Fig. 1. Uncertainty distribution of sample preparation by operation for quantitative determination of trifusol in intrauterine suppository.
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The concentration was calculated using a standard formula.

The placebo contribution to the total absorption was $\delta_{\text{exc}} = 100 \times \frac{0.0031}{0.9542} = 0.32\%$ and was insignificant, because there is a ratio: $\delta_{\text{exc}} \leq 0.033 \times B = 0.32 \leq 0.33$, therefore, the method had a sufficient specificity [7,8].

Linearity for suppositories was determined within 73–127 % of the nominal trifusol concentration. For this purpose was prepared a solution of suppository in purified water according to the method of quantitative determination of trifusol in veterinary effervescent suppository for intrauterine use, later it was used to obtain nine dilutions. The absorption of the obtained solutions at 278 nm was measured and a graph of the dependence of optical density on the concentration of the studied substance in the sample was drawn up (Fig. 2, Table 3).

So, it was calculated numerical indicators testify that all requirements of the State Pharmacopoeia of Ukraine on parameters of linear dependence are fulfilled and linearity of a technique was confirmed in the chosen range of concentration [2–4].

To establish the accuracy of the technique nine parallel definitions (three weighed quantities of the studied drug form, three repetitions) were performed. The absorption of the comparison solution was measured simultaneously. The gram content of trifusol in the dosage form was calculated using a standard formula.

**Table 3. Main characteristics of linear dependence**

| Data         | Value          | Criteria (For tolerances 90.0–110.0%, number of points 9) | Conclusion |
|--------------|----------------|------------------------------------------------------------|------------|
| $b \pm (s_b)$ | 1.0122 ± (0.0073) | --                                                         | --         |
| $a \pm (s_a)$ | -1.1938 ± (0.7116) | $|a| \leq \Delta a = t(95\%, 7) \cdot S_a = 1.348$         | corresponds |
| $S_{xy}$     | 0.3859         | $\Delta a(\%) / t(95\%, 7) = 1.690$                       | corresponds |
| $r$          | 0.9998         | $\geq 0.9555$                                              | corresponds |

**Table 4. Determination of the convergence of the quantitative determination results in intrauterine suppositories with trifusol**

| $Z_\%$ | $S_{z_\%}$ | $\Delta_\%$ | $\Delta_{As_\%}$ |
|--------|------------|--------------|-------------------|
| 109.4  | 0.464      | 0.863        | 3.20              |

**Table 5. Determination of the correctness of the results of the quantitative determination of trifusol in intrauterine suppositories by the standard addition method**

| Index             | Criteria (For tolerances 90–110 %) | Value and conclusion |
|-------------------|--------------------------------------|----------------------|
| Avg. $Z_\%$       |                                      | 99.88                |
| Relative standard deviation, $S_{z_\%}$ | $\leq 1.69$                           | 0.519 Corresponds    |
| Relative confidence interval $\Delta_\% = t(95\%, 8) \cdot S_{z_\%}$ | $\leq 3.20$               | 0.965 Corresponds    |
| Systematic error $\delta_{z_{\%}} = [Z - 100]$ | --                               | 0.120               |
| Criterion of insignificance of systematic error $\delta_{z_{\%}} \leq \Delta_\%/3$ | $\leq 0.322$               | Corresponds          |
A new sensitive, economic and express spectrophotometric method for quantitative determination of piperidinium 2-[5-(2-furyl)-4-phenyl-1,2,4-triazol-3-ylthioacetate in its dosage form – veterinary intrauterine effervescent suppository with its own absorption has been developed. The proposed method was validated and its compliance with the requirements of the State Pharmacopoeia of Ukraine was proved for the main validation characteristics: linearity, convergence, correctness and robustness.

Conclusions

A new sensitive, economic and express spectrophotometric method for quantitative determination of piperidinium 2-[5-(2-furyl)-4-phenyl-1,2,4-triazol-3-ylthio] acetate in its dosage form – veterinary intrauterine effervescent suppository with its own absorption has been developed. The proposed method was validated and its compliance with the requirements of the State Pharmacopoeia of Ukraine was proved for the main validation characteristics: linearity, convergence, correctness and robustness.