Development of a novel kinetic spectrophotometric method for the quantitative determination of Cefuroxime using Caro’s acid

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Recommendation methods for the quantitative determination of Cefuroxime substance using Caro’s acid is of great interest.

**Aim.** To study the reaction of Cefuroxime with Caro’s acid and develop a new method for the quantitative determination of Cefuroxime substance using the product of S-oxidation and perhydrolysis.

**Materials and methods.** To determine Cefuroxime substance the triple potassium salt of Caro’s acid was used as an oxidizing agent. The procedure was developed as a kinetic spectrophotometric method.

**Results and discussion.** The initial rate method can be easily applied to determine Cefuroxime substance. The data obtained showed good results in accuracy, precision, LOQ and LOD. The RSD for the substance was 1.53-2.35 %, δ = 0.55-1.40 %. The direct linear dependence was observed in a wide range of concentrations 1-7 µg mL⁻¹.

**Conclusions.** The possibility of Cefuroxime analytical determination by the biologically active part of the molecule, reproducible results and accuracy are the advantages of the method proposed. The obtained data are in good agreement with the standard pharmacopoeial HPLC method.

**Key words:** Cefuroxime; kinetic spectrophotometric method; quantitative determination; Caro’s acid

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Розробка нової кінетико-спектрофотометричної методики кількісного визначення цефуроксиму за допомогою кислоти Каро

Розробка нових простих і економічно вигідних методик кількісного визначення цефуроксиму субстанції становить великий інтерес.

**Мета** даної роботи полягає в вивченні реакції цефуроксиму з кислотою Каро та у розробці нової методики кількісного визначення цефуроксиму за продуктом S-окиснення та пергідролізу.

**Матеріали та методи.** Потрійна калієва сіль кислоти Каро була використана як окисник для визначення субстанції цефуроксиму. Методика розроблена у кінетико-спектрофотометричному варіанті.

**Результати та їх обговорення.** Метод початкової швидкості може бути застосований для визначення цефуроксиму субстанції. Отримані добрі результати точності, відтворюваності, LOQ та LOD. РSD для субстанції становить 1.53-2.35 %, δ = 0.55-1.40 %. Пряма лінійна залежність спостерігається у широкому діапазоні концентрацій 1-7 мкг /мл.

**Висновки.** Можливість аналітичного визначення цефуроксиму за біологічно активною частиною молекули, відтворюваність і точність є перевагами запропонованої методики. Отримані дані добре узгоджуються зі стандартною фармакопеєю ВФХ методикою.

**Ключові слова:** цефуроксим; кінетико-спектрофотометричний метод; кількісне визначення; кислота Каро

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Розробка нової кінетико-спектрофотометричної методики количественного определения цефуроксима с помощью кислоты Каро

Разработка новых простых и экономически выгодных методик количественного определения цефуроксима субстанции представляет большой интерес.

**Цель** данной работы заключается в изучении реакции цефуроксима с кислотой Каро и в разработке новой методики количественного определения цефуроксима субстанции, основанной на продукте S-окисления и пергидролиза.

**Материалы и методы.** Тройная калевая соль кислоты Каро была использована в качестве окислителя для определения субстанции цефуроксима. Методика была разработана в кинетико-спектрофотометрическом варианте.

**Результаты и их обсуждение.** Метод начальной скорости может быть применен для определения цефуроксима субстанции. Получены хорошие результаты точности, воспроизводимости, LOQ и LOD. RSD для субстанции составляет 1.53-2.35 %, δ = 0.55-1.40 %. Прямая линейная зависимость наблюдается в широком диапазоне концентраций 1-7 мкг /мл.

**Выводы.** Возможность аналитического определения цефуроксима по биологически активной части молекулы, воспроизводимость и точность являются преимуществами предлагаемой методики. Полученные данные хорошо согласуются со стандартной фармакопейной ВЭЖХ методикой.

**Ключевые слова:** цефуроксим; кинетико-спектрофотометрический метод; количественное определение; кислота Каро
The quantitative determination of drugs becomes more and more important. The control of the quality and quantity is one of the obligatory steps of drug production. Therefore, development of new simple and economically profitable methods is of a great interest. The methods for the quantitative determination of Cefuroxime are different.

Classical iodometric method is provided for the assay of most of the pharmacopeial β-lactams, for which iodometric titration is particularly suitable [1, 2]. The iodometric method is insensitive as all antibiotics, the initial substances for their production, as well as their degradation products, are oxidized by iodine. It is therefore not suitable for the purity control of antibiotics of this group [3]. Usually titration is a rapid and easy procedure, but unlike this, iodometric titration of cephalosporins is a long-lasting procedure (approximately 40 min) [4]. The intensive literature review has found the application of methods of chromatography [5, 6], spectrophotometry [7-9], voltammetry [10], oxidimetry [11], spectrofluorimetry [12], chemiluminescence [13].

The aim of this work is to study the reaction of Cefuroxime with Caro’s acid and develop a kinetic spectrophotometric method for the quantitative determination of Cefuroxime based on the product of S-oxidation and perhydrolysis. The possibilities of determination and peculiarities of the reaction of S-oxidation and perhydrolysis of Cefuroxime using potassium caroate as an analytical reagent were studied. The optimal conditions were determined, and the method was statistically checked.

Materials and methods

All materials were of an analytical reagent grade, and the solutions were prepared with double-distilled water.

The triple potassium salt of Caro’s acid was used as an oxidizing agent, 2KHSO₄ • KHSO₄ • K₂SO₄ (the commercial name – Oxon® manufactured by DuPont). The active substance is potassium hydrogenperoxomonosulfate, KHSO₄. The choice of the reagent was determined by its rather high oxidative activity, Eₒ = 1.84 V, availability, and satisfactory solubility in water.

Preparation of standard solutions

Preparation of 0.02 mol L⁻¹ solution of potassium caroate. Dissolve 0.615 g of 2KHSO₄ • KHSO₄ • K₂SO₄ in a 100 mL volumetric flask in double-distilled water. Check the concentration of potassium caroate by iodometric titration.

Prepare sodium hydroxide 0.51 M by dissolving 2 g of NaOH in 100 mL of double-distilled water.

The study of S-oxidation reaction kinetics. Transfer 10.0 mL of 0.02 mol L⁻¹ potassium hydrogenperoxo-

The peroxiacidic method proposed is based on the reaction of Cefuroxime oxidation by the excess of KHSO₄ with the quantitative formation of the corresponding S-oxide in the acidic medium. The KHSO₄ excess was determined by the method of iodometric titration.

The method proposed is based on the quantitative determination of Cefuroxime substance by the product of two conjugate reactions of peroxyacidic oxidation and perhydrolysis in the basic medium using Caro’s acid as an analytical reagent (KHSO₄). The reaction mechanism has been proposed on the basis of the literature review and the experimental study as shown in Fig. 1.

The redox interaction between Cefuroxime and potassium peroxomonosulfate in the acidic medium (pH = 3.5) was stoichiometrical and fast: 1 mol of Cefuroxime per 1 mol of KHSO₄ (the observation time – 1 min) was determined by the iodometric method.

In the basic medium Cefuroxime S-oxide undergoes hydrolytic cleavage. Fig. 2 shows the electronic spectra of Cefuroxime and the reaction product. The appearance of a new band with absorption λₘₐₓ = 290 nm demonstrates its formation in the reaction of alkaline hydrolysis of Cefuroxime S-oxide in the presence of potassium peroxomonosulfate (the perhydrolysis reaction).

The appearance of a new band gave the possibility of development of a new method for the quantitative determination of Cefuroxime. To solve this problem the optimal conditions should be determined.

The absorbance changed substantially with the increase of the concentration of Caro’s acid. The maximum tangent of inclination was obtained when 2.0 mL of 0.02 M of Caro’s acid was used. The maximum slope was obtained when 2 mL of 1.00 M NaOH was used. Over this volume no change in absorbance could be detected. Therefore, 2 mL of 1.00 M of NaOH was used as an optimum value.

The effect of the Cefuroxime concentration

Pipette 1 mL, 3 mL, 4 mL, 5 mL, 6 mL, 7 mL of 1 - 10⁻¹ mol L⁻¹ solution of Cefuroxime into 100 mL volumetric flasks containing 2 mL of 0.02 mol L⁻¹ KHSO₄ solution and 2 mL of 1.00 mol L⁻¹ NaOH solution. Mix the content of the mixture of each flask well and record the increase in absorbance at 290 nm as a function of time for 15 min against the reagent blank (Fig. 3). It shows the dependence of absorption of Cefuroxime alkaline solutions on time at 290 nm and the acid concentration. They have linear dependence for the first 10-15 min. The corresponding plot is given in Fig. 3.
As it is seen from the plot, the tangent of inclination depends on the Cefuroxime concentration. The kinetic curves show that the slope increases proportionally with the increase of the Cefuroxime concentration. Thus, the initial rate method can be successfully applied for determination of the content of Cefuroxime substance in the conjugated reaction between Cefuroxime and Caro’s acid in the alkaline medium using the kinetic spectrophotometric method.

The validity of the method proposed was studied by performing recovery studies. Precision and accuracy were studied by analyzing five replicates of the sample solutions at three concentration levels. The relative standard deviations calculated were all below 2.2 %, indicating the excellent precision of the method proposed.

The procedure is as follows: place 0.42 g (accurate weight) of Cefuroxime in a 100 mL volumetric flask, dilute to the volume with double-distilled water. Further dilute 1 mL of this solution to the volume of 100 mL with double-distilled water at 20 °C. Transfer 10.00 mL of the solution obtained into a 100 mL volumetric flask containing 2 mL of 0.02 mol L⁻¹ KHSO₅ solution and 2 mL of 1.00 mol L⁻¹ NaOH solution. Shake the content, and finally, dilute to the volume with double-distilled water. After addition of NaOH solution switch a stopwatch on. Transfer the solution obtained to a 1 cm cell to measure the absorbance at the wavelength of 290 nm for first the 15 min every 2 min against water.

The kinetic dependence curve of absorbance A on time (min) was obtained. The calculation was performed...
The results of calculations of Cefuroxime accuracy and precision are given in Table.

| Added, mol L⁻¹ ⋅ 10⁻³ | Found, mol L⁻¹ ⋅ 10⁻³ | Mean, mol L⁻¹ ⋅ 10⁻³ | Recovery, % | RSD, % | δ, % |
|------------------------|------------------------|------------------------|-------------|--------|------|
| 3.00                   | 2.98                   | 3.03                   | 101.00      | 2.35   | 1.00 |
| 5.00                   | 4.96                   | 5.07                   | 101.40      | 2.05   | 1.40 |
| 7.00                   | 6.97                   | 7.04                   | 100.55      | 1.53   | 0.55 |

The LOD and LOQ were calculated based on the standard deviation of response and the slope of the calibration curve. It is expressed as LOD = 3 × Sδ/b, LOQ = 10 × Sδ/b where Sδ is the standard deviation of response, b is the slope of the calibration curve.

LOD = 0.38 μg/mL, LOQ = 1.2 μg/mL.

CONCLUSIONS
1. The possibility of the Cefuroxime analytical determination by the biologically active part of the molecule (alicyclic sulfur and β-lactam ring), reproducible results and accuracy are the advantages of the method proposed.
2. The initial rate method can be easily applied to determine Cefuroxime substance, providing advantages during the experiment.
3. The method developed has a good specificity and allows determining the content of the Cefuroxime main component avoiding impurities.
4. The results of accuracy and precision are in good agreement with the results obtained in the reference method. For the pure substance RSD = 1.53-2.35 %, δ = 0.55-1.40 %. LOD = 0.38 μg/mL, LOQ = 1.2 μg/mL.

Conflict of Interests: authors have no conflict of interests to declare.

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