

ORIGINAL ARTICLES

Development and validation of a HPLC method for quantification of degradation impurities of salbutamol sulfate with following long-term stability studies in multicomponent cough syrup

Vývoj a validace metody HPLC pro kvantifikaci nečistot degradace salbutamol-sulfátu s následujícími dlouhodobými stabilitními testy ve vícesložkovém sirupu proti kašli

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Summary

Medicines containing both a herbal extract and a synthetic substance are in high demand due to their beneficial effects and synergism. The novel combination of salbutamol sulfate and *Hedera helix* extracts seems to be prospective in terms of pharmacological activity. But for quality assurance, impurities of the synthetic component have to be determined and quantified. Plant extracts consist of various phytochemical components, therefore, it is more complicated to develop a selective analytical method due to the sample matrix. To prove the safety and efficacy of the dosage form, a new HPLC method for analysis of salbutamol sulfate impurities was developed and validated. The method was used to estimate the safety of the novel syrup by performing long-term stability studies for 24 months. Obtained

results indicated the absence in both significant reducing of the main components content and increasing of related substances level. Also, force degradation was carried out to prognosticate the possibility of impurities producing.

Key words: long-term stability study • salbutamol sulfate • impurities • multicomponent dosage form • cough syrup • HPLC

Souhrn

Léky obsahující jak bylinný extrakt, tak syntetickou látku jsou velmi žádané kvůli jejich pozitivním účinkům a synergismu. Z hlediska farmakologické aktivity se nová kombinace salbutamol-sulfátu a extraktů *Hedera helix* jeví jako perspektivní. Pro zajištění kvality je však nutné určit a kvantifikovat nečistoty syntetické složky. Rostlinné extrakty se skládají z různých fytochemických složek, proto je složitější vyvinout selektivní analytickou metodu. Z důvodu prokázání bezpečnosti a účinnosti lékové formy byla vyvinuta a validována nová metoda HPLC pro analýzu nečistot salbutamol-sulfátu. Tato metoda byla použita k odhadu bezpečnosti nového sirupu provedením dlouhodobých stabilitních testů po dobu 24 měsíců. Získané výsledky naznačily absenci jak ve významném snížení obsahu hlavních složek, tak ve zvýšení hladiny látek příbuzných. Pro předpověď možnosti tvorby nečistot byla rovněž provedena tzv. síla degradace.

Klíčová slova: dlouhodobé stabilitní testy • salbutamol-sulfát • nečistoty • vícesložková léková forma • sirup proti kašli • HPLC

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Introduction

Respiratory diseases occur in people regardless of age and social status, and coughing remains the most com-

mon complaint of patients. An original syrup has been developed based on *Hedera helix* extract with the addition of salbutamol sulfate¹⁾.

Medicines with *H. helix* leaf extract have advantages in the treatment of acute bronchitis accompanied by cough and sputum due to triterpene saponins, which show a mucolytic, mucokinetic, and antispasmodic action^{2, 3)}. It is a well-known fact that for better expectoration of sputum it is necessary to stimulate the enlargement of the bronchi. In this case, in addition to multicomponent medicine bronchodilator substances, such as salbutamol sulfate, can be quite useful in the treatment of bronchopulmonary diseases. Previously, the quality of the novel syrup was confirmed by analysing of the *H. helix* main component¹⁾.

It is worth mentioning that different components, intermediates, and reagents are used during the synthesis of active pharmaceutical ingredients (API). Thus, identification, quantification, and control of impurities in the synthetic substance and its dosage forms play a critical role in providing patients with safe medicines^{4, 5)}. The method for analysis of salbutamol sulfate is reported in EP⁶⁾. Some other literature sources propose the HPLC for quantification of related substances^{7–10)} as well as enantiomeric separation¹¹⁾ of salbutamol sulfate.

Also, the LC-MS¹²⁾ and GC-MS¹³⁾ might be used. However, none of the above-mentioned methods describe the analysis of salbutamol sulfate impurities in combination with a plant extract which can be challenging due to the content of different phytochemicals. Moreover, no publication has established stability in studying storage dosage forms with salbutamol sulfate for long terms. This study aimed to estimate the safety of the novel combination and to determine the main impurities profile for salbutamol sulfate.

Experimental part

Instrument

Liquid chromatography separation was performed using a Shimadzu Nexera X2 LC-30AD HPLC system (Shimadzu, Japan) composed of a quaternary pump, an on-line degasser, a column temperature controller, a SIL-30AC autosampler (Shimadzu, Japan); a CTO-20AC thermostat (Shimadzu, Japan) as well as a SPD-M20A diode array detector (DAD). Other instruments such as an Ultrasonic Cleaner Set for ultra-sonication using (Wise Clean WUC-A06H, Witeg Labortechnik GmbH, Germany), a Libra UniBloc AUW120D (Shimadzu Analytical Scale, Japan); a pH-meter – a Knick Electronic Battery-Operated pH Meter 911 PH (Portamess, Germany), and class A analytical volumetric flasks that meet the requirements of the SPH_U were used in the investigation.

Chemicals and reagents

HPLC grade methanol and acetonitrile, analytical grade hydrogen peroxide, hydrochloric acid, sodium hydroxide, acetic acid, and triethylamine (TEA) were from Merck (Darmstadt, Germany). Salts of ammonium acetate, sodium dihydrogen phosphate, and trifluoroacetic acid (TFA) were purchased from Sigma-Aldrich GmbH

(Steinheim, Germany). Salbutamol sulfate EPCRS, impurity-G EPCRS, impurity-D EPCRS, impurity-F EPCRS were purchased from EP commission. Purified deionized water was produced with a Millipore (Burlington, MA., USA) water purification system.

Syrup compositions

All series of syrups were prepared by Vishpha (Zhytomyr, Ukraine) as follows (amount of substance in dosage form are specified in percents): extract *H. helix* leaf (0.45%) and salbutamol sulfate (0.048%) as active pharmaceutical ingredients and sorbitol (35%), citric acid (0.045%), potassium sorbate (0.2%), natural gum (0.22%), and purified water as excipients.

Chromatographic conditions

Separation of compounds was performed with an ACE C18 column (150*4.6 mm, particle size 5 µm). The binary solvent system was used: 0.05% TEA solution adjusted to 5.5 pH with acetic acid as the mobile phase A and a mixture of methanol: acetonitrile 50 : 50 (v/v) as the mobile phase B with the flow rate of 1.0 ml/min. The following linear gradient elution was used: 95% A/5% B – 0 min, 95% A/5% B – 5 min, 10% A/90% B – 30 min, 10% A/90% B – 32 min, 95% A/5% B – 34 min. The detection of components was done at 277 nm. The column oven temperature was set at 35 °C.

The preparation of the solution for system suitability

The final concentrations of components were as follows: for salbutamol sulfate 400 µg/ml, potassium sorbate 2 mg/ml, 1.8 µg/ml for impurities, D, F, G in water.

Test solution

15.0 ml of syrup were placed into a 20.0 ml volumetric flask and made up to mark with water. All solutions were filtered through a 0.45 µm membrane filter.

Stability studies

Syrup stability was tested according to ICH guidelines, at the temperature 25 ± 2 °C and humidity 60 ± 5% maintained by a thermostat. Three different batches of syrup were stored at the mentioned conditions in amber glass bottles. The impurity profile was measured directly following preparation and after 3, 6, 9, 12, 15, 18, 21, 24 months.

Degradation studies

The concentration of salbutamol sulfate for all stress studies was the same as in test solution. Alkaline hydrolysis was performed with adding of 2 mL of 0.1 M NaOH at ambient temperature for 3 hours. The effect of acidic decomposition was tested with 2 mL of 0.1 M HCl at the ambient temperature for 3 hours. Oxidative study was performed with 2 mL of 3 % hydrogen peroxide at the ambient temperature for 3 hours. Also, the temperature degradation was used, the sample was heated at 100 °C for 3 hours. Samples were withdrawn at appropriate time and analysed with the proposed HPLC method.

Results

Method development

The possible impurities of salbutamol sulfate are similar to the main component¹⁴. To achieve a suitable

resolution among all components in the novel composition, different conditions were tested. In particular, the influence of different stationary phases sorbents (C8 and C18) as well as their particle size (3 μm and 5 μm) were studied. The C18 stationary phase with particle size 5 μm

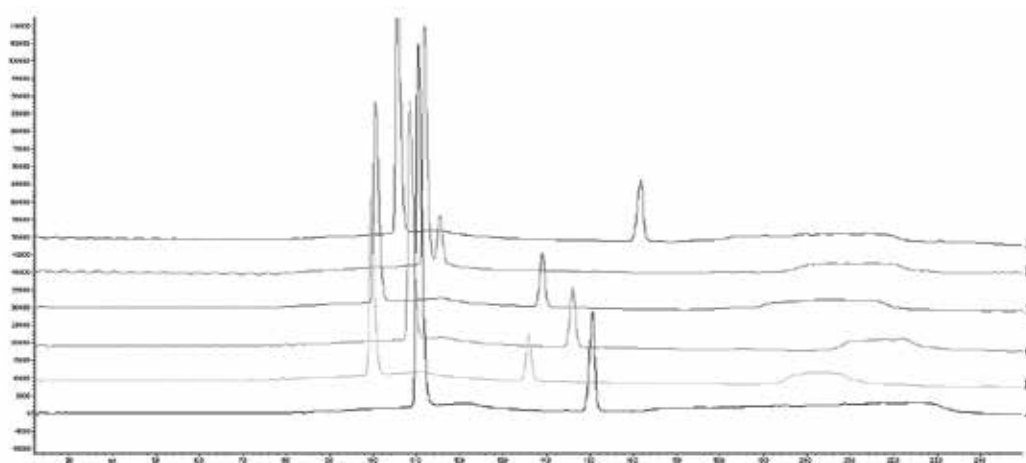


Fig. 1. Chromatograms of robustness studies (A – 0.9 mL/min, B – 1.1 mL/min, C – 30 °C, D – 40 °C, E – 6.0 pH, F – 5.0 pH)

Table 1. Effects of changed parameters in robustness study

Changes	Parameters	Salbutamol sulfate	Impurity D	Potassium sorbate	Impurity G	Impurity F
0.9 mL/min	Retention time	11.03 min	14.43 min	15.05 min	15.59 min	19.49 min
	Resolution		13.694	2.596	2.2	17.603
	Number of Theoretical Plate (NTP)	26018	66838	55667	73383	133671
	Tailing Factor	1.595	0.849	0.881	1.055	1.257
1.1 mL/min	Retention time	9.95 min	13.16 min	13.58 min	14.42 min	18.49 min
	Resolution		14.552	1.948	3.559	20.556
	NTP	27507	68813	51471	63125	196818
	Tailing Factor	1.493	1.272	0.98	1.560	1.301
30 °C	Retention time	10.85 min	14.33 min	14.60 min	15.26 min	19.43 min
	Resolution		17.437	1.372	2.674	18.386
	NTP	30670	143687	54499	62339	138268
	Tailing Factor	1.486	0.727	0.958	1.109	1.332
40 °C	Retention time	10.05 min	13.18 min	13.89 min	14.68 min	18.53 min
	Resolution		13.488	3.177	3.976	25.42
	NTP	26344	59661	54474	143619	236692
	Tailing Factor	1.493	1.437	0.963	0.966	1.418
pH 6.0	Retention time	11.18 min		11.54 min		
	Resolution			1.578		
	NTP	42542		39994		
	Tailing Factor	1.351		1.140		
pH 5.0	Retention time	10.56 min	13.69 min	13.97 min	16.16 min	17.44 min
	Resolution		11.307	0.903	7.721	2.162
	NTP	35157	27997	35222	57158	5879
	Tailing Factor	1.491		0.907		

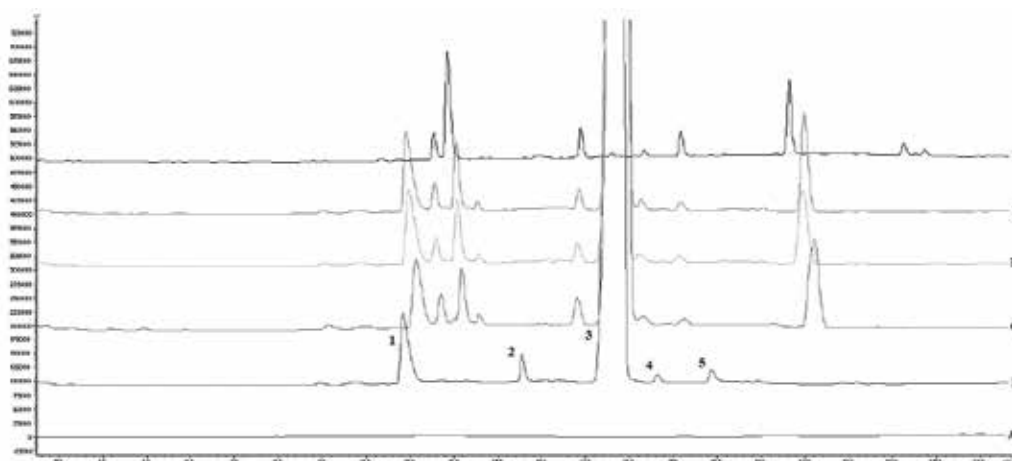


Fig. 2. Chromatograms of analysed samples of solution for system suitability performing (A – blank solution, B – solution for system suitability, C – 011 batch syrup, D – 021 batch syrup, E – 031 batch syrup, F – placebo containing ivy leaf extract; 1 – salbutamol sulfate, 2 – impurity D, 3 – potassium sorbate, 4 – impurity G, 5 – impurity F)

Table 2. Results of stability testing of salbutamol sulfate impurities

Storage terms	Series of syrup	Impurity D (%)	Impurity G (%)	Impurity F (%)	Sum of impurities (%)
0 month	011	N/D*	N/D*	0.039	0.052
	021	N/D*	N/D*	0.043	0.061
	031	N/D*	N/D*	0.041	0.055
3 months	011	N/D*	N/D*	0.04	0.057
	021	N/D*	N/D*	0.042	0.063
	031	N/D*	N/D*	0.041	0.057
6 months	011	N/D*	N/D*	0.04	0.058
	021	N/D*	N/D*	0.042	0.065
	031	N/D*	N/D*	0.042	0.060
9 months	011	N/D*	N/D*	0.04	0.060
	021	N/D*	N/D*	0.043	0.066
	031	N/D*	N/D*	0.042	0.062
12 months	011	N/D*	N/D*	0.04	0.067
	021	N/D*	N/D*	0.043	0.067
	031	N/D*	N/D*	0.041	0.065
15 months	011	N/D*	N/D*	0.04	0.067
	021	N/D*	N/D*	0.044	0.069
	031	N/D*	N/D*	0.041	0.067
18 months	011	N/D*	N/D*	0.04	0.069
	021	N/D*	N/D*	0.043	0.069
	031	N/D*	N/D*	0.041	0.070
21 months	011	N/D*	N/D*	0.04	0.070
	021	N/D*	N/D*	0.043	0.072
	031	N/D*	N/D*	0.041	0.073
24 months	011	N/D*	N/D*	0.04	0.073
	021	N/D*	N/D*	0.043	0.075
	031	N/D*	N/D*	0.041	0.075

*not detected

was recognized as the most suitable for the separation of components in the studied combination.

Moreover, the influence of the mobile phase buffer salt and additives were investigated. Two different salts in various concentrations were tested: ammonium acetate (5 mM–25 mM) and sodium dihydrogen phosphate (5 mM–50 mM) with different pH values 3.0–6.0. However, the resolution of some impurities with other components of combination remained unsatisfactory. The salts were replaced with additives such as triethylamine (TEA) and trifluoroacetic acid (TFA). Water solution with TEA adjusted to pH 5.5 showed the most suitable separation among components.

Methanol and acetonitrile were chosen as components of the mobile phase B. Pure methanol gave a poor baseline with a baseline drift and peak shapes were not acceptable. Hence acetonitrile was used to correct it. Ration 50/50 (v/v) was the most suitable.

Method validation

A new HPLC method for analysis of salbutamol impurities has been developed and validated according to

State Pharmacopeia of Ukraine¹⁵⁾ and ICH¹⁶⁾. There are many developed methods for the determination of salbutamol sulfate impurities. However, their applying for analysis of the cough syrup was not so successful, since a poor separation between the syrup matrix and salbutamol impurities has been observed.

System suitability test

The observed RSD values at a 1 % level of analyte concentration were within suitable values ($\leq 2\%$). The resolution, number of the theoretical plate, and tailing factor for all main components were determined. All values were within suitable limits.

Specificity

Specificity studied has shown that no peak either from placebo of the syrup matrix or from forced degradation studies was coeluted at the retention time of salbutamol sulfate and its impurities. The identification of components was done by comparing the retention times of peaks from the test solution with the peaks of the standard solution.

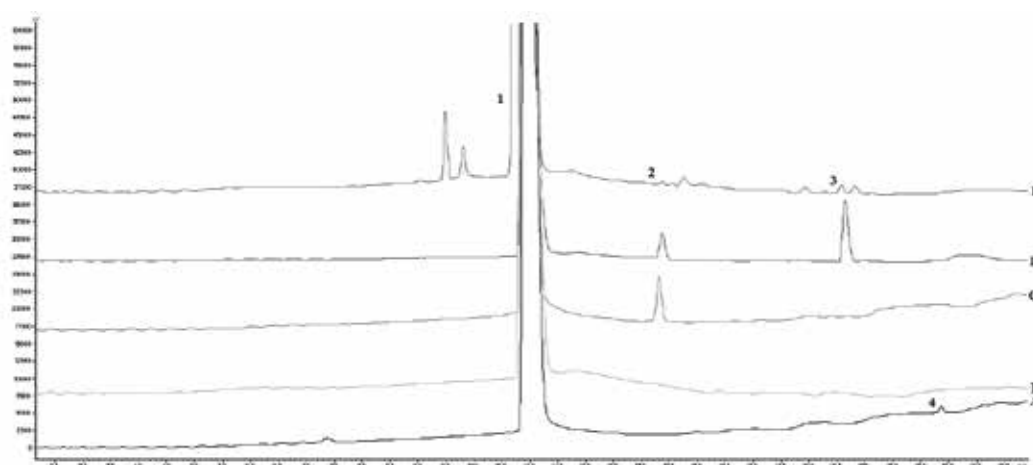


Fig. 3. The chromatogram of degradation studies (A – test, B – acidic, C – temperature, D – alkaline, E – peroxide; 1 – salbutamol sulfate, 2 – impurity D, 3 – impurity G, 4 – impurity F)

Table 3. Results of performed degradation studies

	0.1 M NaOH	0.1 M HCl	3% H ₂ O ₂	Temperature (100 °C)
Impurity 1 (14.47 min)	–	0.18%	–	–
Impurity 2 (9.69 min)	–	–	1.91%	–
Impurity 3 (10.44 min)	–	–	1.49%	–
Impurity 4 (13.59 min)	–	–	0.26%	–
Impurity 5 (13.72 min)	–	–	0.27%	–
Impurity 6 (15.99 min)	–	–	0.86%	–
Impurity 7 (16.82 min)	–	–	0.95%	–
Impurity D (13.43 min)	1.59 %	–	0.24%	2.66%
Impurity G (16.55 min)	3.15 %	–	0.78%	–
Impurity F (18.35 min)	–	–	–	–
Sum of impurities	4.74 %	0.18%	6.79%	2.66%

Limit of detection

Investigation of the sensitivity of the method has been carried out by analysis of a prepared solution with a concentration of salbutamol sulfate (36 ng/mL) which was 0.01% from its amount in the test solution. Obtained ratio of signal/noise for the main peak was greater than 3/1 hence the method was approved as sensitive.

Robustness

Examination of robustness has been performed with a changing temperature oven (30–40 °C), flow rate (0.9 ml/min to 1.1 ml/min), and pH of mobile phase A (5.0–6.0) (Table 1, Fig. 1). Obtained results showed that the main effect on the system was exerted by pH of the Mobile phase, thus at high values of pH, no separation between peaks of salbutamol sulfate and excipients was observed.

Stability studies

Long-term stability studies were performed using the proposed method. The results obtained from stability studying are shown in Table 2, Fig. 2.

According to the requirements for impurities level, there are such limits in an amount: for both impurity D and G not greater than 0.5% (from the amount of salbutamol sulfate in the dosage form), for impurity F not greater than 0.3 %, unspecified impurity not greater than 0.1%, and the sum of impurities not greater than 1.5 %.

The obtained results indicate that syrup contained only impurity F and some unspecified impurities, moreover their concentrations change insignificantly. Other impurities were not produced under storage conditions. Thus, the novel syrup was found stable for 24 months of storage, since the amount of each impurity was lesser than the allowed level.

Forced degradation study

Forced degradation was performed on salbutamol sulfate substance to prove the specificity of the method and to determine the impurities profile. Four different degradations were attempted, such as alkaline, acidic, temperature, and peroxidic. The stability of composition was carried out by preparing the sample solution as mentioned above and injected at regular parameters. From the forced degradation study it was observed and confirmed that no other formulation components and potential degradation product and impurities had interfered in the determination. The results of degradation are shown in Table 3, Fig. 3.

As can be seen, salbutamol sulfate was not stable to the effect of temperature, alkaline hydrolysis, and oxidation. Peroxide influence had the greatest effect on producing different unspecified impurities. On the other hand, the highest level of impurity D and impurity G was after the alkaline effect on the solution.

Conclusions

A new HPLC method for the determination of salbutamol sulfate impurities was developed and validated. The

developed method was used in long-term stability studies of a novel cough syrup. Obtained results showed that salbutamol sulfate is stable in syrup during all terms of storage (24 months), however, it degrades under temperature, alkaline, and oxidation actions. The main observed related substances of salbutamol sulfate were impurity D, impurity G, and impurity F. On the other hand, only impurity F was present in the syrup, while others were detected only after degradation.

Conflict of interest: none.

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