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AQbD-driven UHPLC method for simultaneous quantification of budesonide, glycopyrronium bromide, and salbutamol sulphate: a unified approach for inhalation product and bioanalytical applications

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Inhalation therapies often combine budesonide, glycopyrronium bromide, and salbutamol sulphate, necessitating analytical methods capable of their simultaneous quantification. Conventional UHPLC methods typically address these active pharmaceutical ingredients (APIs) individually, leading to inefficiencies in development, validation, and regulatory alignment. Analytical Quality by Design (AQbD) has transformed pharmaceutical analytical method development through systematic risk assessment and structured optimization, but its application in the bioanalytical domain remains limited. The challenge addressed in this study is the creation of a single robust UHPLC method that meets both pharmaceutical and bioanalytical requirements within regulatory frameworks. A unified UHPLC method was developed using AQbD principles, employing Design of Experiments to identify and optimize critical method parameters. The optimized method achieved baseline separation of all three APIs on a YMC UltraHT Hydrosphere C18 (2.1 × 100 mm; 2.0 μm) column under gradient elution with methanol and 0.1% formic acid in 10 mM ammonium formate. Robustness was established through a defined Method Operable Design Region. The method was first validated under ICH Q2 (R2) for pharmaceutical applications, confirming accuracy, precision, and sensitivity. Subsequently, it was extended to the bioanalytical domain and validated under ICH M10 guidelines in simulated lung fluid, demonstrating reproducibility in complex matrices. This dual validation highlights the method's versatility and regulatory robustness, underscoring AQbD's ability to unify pharmaceutical and bioanalytical method development into a single lifecycle appropriate platform. This study demonstrates the first AQbD-driven UHPLC method validated under both ICH Q2 (R2) and ICH M10, bridging pharmaceutical and bioanalytical applications. Extending AQbD principles into bioanalysis provides a regulatory-relevant framework that enhances robustness, lifecycle flexibility, and compliance. The work establishes a unified strategy for inhalation therapies and beyond, supporting broader adoption of science- and risk-based analytical development.

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1. Introduction

Chronic obstructive pulmonary disease (COPD) and asthma are two of the most prevalent and debilitating respiratory disorders globally, affecting hundreds of millions of individuals and contributing significantly to healthcare burdens.^{1,2} COPD ranks as the fourth leading cause of death worldwide, account-

ing for approximately 3.5 million deaths in 2021,³ while asthma impacted over 262 million people and was responsible for 455 000 deaths in 2019.⁴ Although both conditions involve airflow obstruction and inflammation, they differ in pathophysiology: COPD is defined by persistent and progressive airflow limitation driven by chronic inflammation and alveolar destruction, whereas asthma is characterized by variable airflow limitation, airway hyperresponsiveness, and typically reversible bronchoconstriction triggered by allergens or environmental stimuli.⁵

To manage these complex diseases, especially in moderate to severe stages, current therapeutic strategies often rely on combination therapies that target multiple underlying mechanisms. Among the most widely used agents are short-acting β₂-

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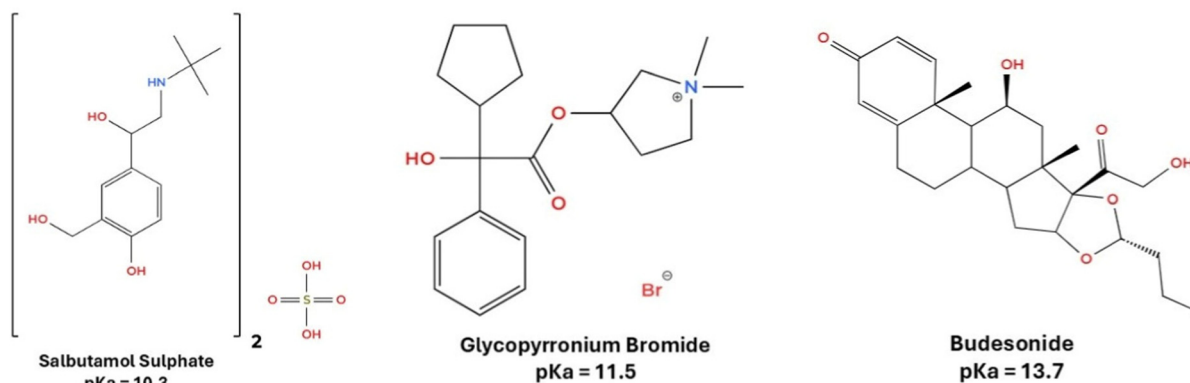


Fig. 1 Chemical structures of the selected APIs with the relative pK_a (strongest acid).^{13–15}

agonists (SABAs) like salbutamol sulphate (SBS), which provide rapid bronchodilation;⁶ long-acting muscarinic antagonists (LAMAs) such as glycopyrronium bromide (GB), which reduce cholinergic-mediated bronchoconstriction;⁷ and inhaled corticosteroids (ICS) like budesonide (BUD), which attenuate airway inflammation.⁸ These three APIs are frequently co-formulated in fixed-dose combinations (*e.g.*, ICS-LAMA, ICS-SABA), offering a synergistic approach to bronchodilation and inflammation control.^{9,10} Their combination is particularly relevant in treating asthma-COPD overlap (ACO), a phenotype associated with frequent exacerbations and poor treatment response when managed with monotherapy alone.^{11,12} The molecular structures and corresponding pK_a values of the APIs are depicted in Fig. 1.

Inhalation-based delivery of these agents ensures targeted action and reduced systemic side effects but also presents analytical challenges. Ensuring precise quantification and consistent performance of combination inhalation products requires highly sensitive, selective, and robust analytical methodologies.¹⁶ This is particularly important given the low doses involved and the need to differentiate among multiple APIs with diverse chemical properties.¹⁷ Furthermore, the inhalation products' performance is nowadays carried out according to the United States Pharmacopeia (USP) chapter <601>, where typically a cascade impactor is recommended for the testing.¹⁸ These cascade impactors (*e.g.*, Next Generation Impactor – NGI) artificially simulate the lung surface, and the powder's particle size distribution can be measured. Usually, an analytical methodology is developed and dedicated to powder recovery in the different stages of the impactor.¹⁹

To address these needs, we developed a novel ultra-high-performance liquid chromatography (UHPLC) method to quantify BUD, GB, and SBS simultaneously. The technique was designed and optimized using AQbD principles, a modern approach that integrates risk assessment, critical method attributes (CMAs) identification, and design of experiments (DoE) to build quality into the method from its inception.²⁰ The method was validated following ICH Q2 (R2) guidelines,²¹ ensuring suitability for quality control of pharmaceutical inhalation products.

Importantly, this work extends the AQbD paradigm to the bioanalytical field, an area where conventional method development practices, typically linear and empirical, still dominate. Bioanalytical methods, governed by ICH M10²² and other regulatory guidelines,²³ are essential for quantifying drug levels in biological matrices for pharmacokinetic, toxicokinetic, and bioequivalence studies.²⁴ However, these methods often suffer from matrix effects, high variability, and limited flexibility due to their empirical design.^{25,26}

Applying AQbD to bioanalytical method development represents a transformative shift. It enables systematic understanding of critical method parameters, such as extraction efficiency, matrix interference, and ion suppression. While using DoE and risk assessment tools (*e.g.*, Ishikawa diagrams, Failure Mode and Effect Analysis), AQbD facilitates the creation of robust methods^{27,28} that perform reliably across variable biological conditions and over extended study durations.

In this study, we also developed and validated a parallel bioanalytical method for quantifying the same three APIs in a lung fluid scenario. The method was optimized and validated under ICH M10,²² demonstrating regulatory compliance and suitability for clinical applications. Implementing AQbD across pharmaceutical and bioanalytical workflows underscores its versatility and the value of a unified, science-based development framework.

This work serves as a first step toward broader integration of AQbD into bioanalytical science, demonstrating its potential to improve method robustness, streamline development timelines, and elevate the reliability of pharmacokinetic assessments. By bridging pharmaceutical and bioanalytical method development through a single, scientifically justified platform, we offer a pathway for more unified, lifecycle-based analytical strategies in drug development.

2. Materials and methods

2.1. Reagents and consumables

The ammonium formate used for the mobile phase was acquired from Sigma-Aldrich GmbH (Vienna, Austria). The



Methanol (HPLC gradient) used for the mobile phases and diluent was purchased from M&B Stricker Laborfachhandel GbR (Bernried am Starnberger See, Germany). The purified water for all analyses was obtained from Triton UV purification equipment from Neptec (Elbtal, Germany). The formic acid ($\geq 98\%$) used for pH modifications was purchased from Carl Roth (Karlsruhe, Germany). Additional solvents used for forced degradation studies, such as 30% w/v hydrogen peroxide, sodium hydroxide solution, and hydrochloric acid, were acquired from Sigma-Aldrich GmbH (Vienna, Austria). The chloroform for the simulated lung fluid (SLF) preparation was acquired from Sigma-Aldrich GmbH (Vienna, Austria). The buffers used for the Phosphate Buffer Saline (PBS) preparation were purchased from VWR International GmbH (Vienna, Austria). All sample solutions were filtered before being injected into the chromatographic systems using PTFE syringe filters (0.45 μm) from YETI Merz Brothers GmbH (Haid, Austria).

2.2. Standards, samples, and excipients

The standards for quantification of Budesonide (Pharmaceutical Secondary Standard), glycopyrronium bromide (European Pharmacopeia Reference Standard), and Salbutamol Sulphate (Albuterol Sulfate, Pharmaceutical Secondary Standard) were acquired from Sigma-Aldrich GmbH (Vienna, Austria). The identical standards batches were employed for sample preparation and method validation. Dipalmitoylphosphatidylcholine (DPPC) and dipalmitoylphosphatidylglycerol (DPPG), included in the SLF preparation, were obtained from Avanti Polar Solutions, Inc. (Alabama, USA). A complete protocol for the SLF preparation is provided in the SI (chapter 1.1).

2.3. Equipment and software

A Reversed-Phase UHPLC H-Class from Waters Corp. (Milford, MA, USA) coupled with a Photo-Diode Array Detector (PDA) and a QDa single-quadrupole Mass Detector (QDa) was used to acquire chromatographic data. The Empower 3 (v.3.8.0) software from Waters Corp. (Milford, MA, USA) was used to control the HPLC and acquire method development and validation data. The screening DoE was performed using the Acquity HSS T3 (2.1 \times 100 mm; 1.8 μm) column from Waters Corp. (Milford, MA, USA), the Triart C18 (2.1 \times 100 mm; 1.9 μm), the Triart Phenyl (2.1 \times 100 mm; 1.9 μm), and the UltraHT Hydrosphere C18 (2.1 \times 100 mm; 2.0 μm) columns provided by YMC Europe GmbH (Dinslaken, Germany). The robustness DoE included a Kinetex C18 (2.1 \times 100 mm; 1.7 μm) from Phenomenex. The DoEs' conceptualization and statistical analysis were performed on Design Expert v.13 (Stat-Ease Inc., Minneapolis, MN, USA). The regression analysis was performed on OriginPro 2023b v.10.0.5.157 (OriginLab Corp., Northampton, MA, USA). The Centrivap centrifugal concentrator used for SLF preparation was obtained from LabConco Corporation (Kansas City, MO, USA). The FiveEasy™ FP20 pH meter used for mobile phase preparation was acquired from Mettler Toledo GmbH (Columbus, OH, USA).

2.4. Analytical method validation by ICH Q2 (R2)

2.4.1. Specificity. Specificity was evaluated to demonstrate the method's ability to unequivocally assess the analyte in the presence of expected components, such as impurities, degradation products, excipients, and matrix constituents.²¹ Chromatographic analyses were conducted for the mobile phase, blank (methanol/water), and matrix constituent (SLF). The resulting chromatograms were assessed for co-eluting peaks and interferences at the retention time of the analytes. Additionally, a forced degradation study was performed to determine the stability-indicating feature of the method,^{21,29} with applied conditions described in the SI (chapter 1.2).

2.4.2. Linear response, limit of detection, and limit of quantification. Linearity was assessed over a concentration range containing eight independent levels from 100 to 500 $\mu\text{g ml}^{-1}$, including 70, 100, and 130% of the target analyte concentrations. Each API was independently weighed and dissolved in a unique volumetric flask to generate a stock solution. Calibration standards were prepared by serially diluting a stock solution using the designated diluent. Each level was analyzed in triplicate. Calibration curves were constructed by plotting the peak area against the corresponding nominal concentration. Linearity was evaluated using the adjusted correlation coefficient (R^2 -adj), residuals' normality, and the y -intercept assessment for statistical significance.

Limit of detection (LOD) and quantification (LOQ) were calculated using the signal-to-noise ratio approach. The signal-to-noise ratio is determined by comparing measured signals from samples with known low analyte concentrations with those of blank injections. A signal-to-noise ratio of 3 : 1 was considered acceptable for estimating LOD; for LOQ, a ratio of at least 10 : 1 was acceptable.²¹

2.4.3. Accuracy. The accuracy included a mixture of the APIs at three concentration levels of 70, 100, and 130% of the declared content/labeled claim, which were prepared by weighing ($n = 3$) the APIs and dissolving them in the designated diluent, then further analyzed. Accuracy was reported as a mean percent recovery of each API, and the acceptance criteria were established to be $\pm 2\%$ of the nominal concentration.³⁰

2.4.4. Precision (repeatability and intermediate precision). The precision included six 100% label claim sample solutions, which were prepared independently and analyzed against a freshly prepared calibration curve to prove repeatability. Another identical set of samples ($n = 6$) was prepared by a different analyst on a different day under the same conditions to prove intermediate precision. Precision was reported as Relative Standard Deviation in percentage (RSD%), and the acceptance criterion was set to $< 2\%$.³⁰

2.4.5. Robustness. Robustness was proven with a series of multivariate experiments combined with a statistical approach, such as DoE, which should be adopted to test robustness, as suggested by the last update of the ICH Q14 guideline.³¹ Therefore, we deployed an I-optimal response surface design for quadratic effects, comprising 30 experimental runs, including replicates, to ensure adequate statistical power for robust-



ness assessment. The evaluation was conducted using a Type III ANOVA.

2.5. Analytical method validation by ICH M10

2.5.1. Selectivity, specificity, and matrix effect. Selectivity was evaluated by injecting six different lots of SLF, prepared with individual batches of DPPC and DPPG from the same supplier, and quantifying the response of interfering peaks at the same retention time of the analytes. The response attributable to such components should not exceed 20% of the analyte's responses at the Lower Limit of Quantification (LLOQ).²² Specificity was measured by injecting a pure reference standard solution containing the APIs against an artificially prepared solution at the LLOQ.²² The mass-to-charge ratios (m/z) were recorded for both injections *via* Mass Spectrometry (MS) detection to confirm the identity of compounds. The Matrix Effect was performed by injecting three independent replicates of low and high Quality Control Samples (QCs), each prepared using six different lots of SLF, the same ones used for evaluating selectivity. For each matrix source tested, the measured accuracy, expressed as Relative Error (RE%), should fall within $\pm 15\%$ of the nominal concentration, and the precision, expressed as RSD%, should not exceed 15%.²²

2.5.2. Calibration curve and range. A calibration curve was generated with a blank sample and six concentration levels from 1 to 20 ng ml⁻¹, including the LLOQ and the Upper Limit of Quantification (ULOQ). Subsequently, two more linearities were prepared over different days to guarantee reproducibility.²² The back-calculated concentrations of the calibration standards were measured, and the accuracy of each level should have been within $\pm 15\%$ of the nominal concentration, and $\pm 20\%$ at the LLOQ. At least 75% of the calibration standards for each generated curve should meet the above criteria.²² The calibration curves were freshly prepared by serially diluting a stock solution in methanol/water to a corresponding level in SLF. The solutions were stirred and afterward filtered through a PTFE 0.45 μm filter.

2.5.3. Accuracy and precision. Accuracy and precision were proven by preparing fresh QCs, storing them under fridge conditions ($\sim 4\text{ }^\circ\text{C}$) for 48 hours, and subsequently analyzing them. In our study, QCs were produced at four distinct concentration levels within the linear range: the LLOQ, three times the LLOQ (low QC), 50% of the ULOQ (mid QC), and 75% of the ULOQ (high QC). The levels were determined to be 1, 3, 10, and 15 ng ml⁻¹.

Accuracy and precision were determined by analysing the QCs within each run (within-run) and in two different runs (between-run). Within-run accuracy and precision were evaluated by analysing five replicates at each QC concentration level. Between-run accuracy and precision were assessed by examining each QC concentration level in three analytical runs over two days ($n = 15$ total). The accuracy (RE%) at each concentration level should be within $\pm 15\%$ of the nominal concentration, except at the LLOQ, where it should be within $\pm 20\%$.²² The precision (RSD%) of the concentrations determined at

each level should not exceed 15%, except at the LLOQ, which should not exceed 20%.²²

During this stage, one analytical run used for accuracy and precision was injected again on a different day to prove reinjection reproducibility, following the same acceptance criteria.²²

2.5.4. Carry-over. During validation, carry-over was assessed by injecting a blank sample following the ULOQ calibration standard. The analyte response in this blank should not exceed 20% of the response observed at the LLOQ.²²

2.5.5. Dilution integrity. Dilution integrity was proven with a diluted QC sample that was prepared in SLF at a concentration of 100 ng ml⁻¹, and the following dilutions to the ULOQ were applied in replicate 5 times. The mean accuracy (RE%) of the dilution QCs should be within $\pm 15\%$ of the nominal concentration, and the precision (RSD%) should not exceed 15%.²²

2.5.6. Stability of solutions. To prove stability, three replicates at the ULOQ concentration level were stored at room temperature, under fridge conditions ($\sim 4\text{ }^\circ\text{C}$), and freezing conditions ($\sim -20\text{ }^\circ\text{C}$). The solutions were thawed after 48 hours, analyzed, and their recovery evaluated.

3. Results and discussion

3.1. Method development by AQbD principles

The AQbD workflow consists of multiple steps that align with the most recent version of the USP <1220> chapter and ICH Q14 revisions, including an extended review of the analytical procedure development and lifecycle.^{31,32} Following this rigorous process ensures that the developed analytical methodology fits the intended scope by producing robust and reliable results.^{27,33} An overview of the steps involved in the AQbD stream is provided in Fig. 2.

The initial stage involves defining the Analytical Target Profile (ATP), which outlines the intended performance criteria that the analytical method is expected to achieve. The developed method should detect the three selected APIs with desirable chromatography standards, including adequate peak resolution, a fundamental property to ensure specificity and peak purity, without sacrificing the fast analysis setup provided by UHPLC. The solubility pattern of the APIs presents another challenge in terms of retention: SBS and GB are very hydrophilic, while BUD is strongly hydrophobic due to the steroid groups (Fig. 1). In that context, the choice of mobile phase, gradient, and column is pivotal to secure a correct elution of the analytes.³⁴ Sensitivity-wise, the method should reach interesting LOD and LOQ values, laying a solid basis for future tuning *via* MS. Inhalation performance testing often requires further dilution (*e.g.*, *via* NGI) to recover a quantity of sample to test afterward *via* HPLC, making reaching low sensitivity standards a fundamental feature for this method. Therefore, in a later stage, the technique was asked to robustly quantify the analytes in the range of a few ng ml⁻¹, especially given the bioavailability of these drugs once they reach the lungs and then the bloodstream.^{35–38} This study aimed to



Analytical Quality by Design Workflow



Fig. 2 Analytical quality by design workflow used for method development.

Table 1 Analytical target profile for the determination of BUD, GB, and SBS³⁹

ATP element	Target	Requirement Ref.
USP signal-to-noise	Maximized	30 and 40
Peak tailing	0.8–1.8	30 and 40
Peak purity	Acceptable	30 and 41
Capacity factor (<i>k'</i>)	>2	30
USP resolution	>2	30

develop a robust method with trustworthy performance across a broad range of applications, ensuring its suitability for translation into an industrial setting, where consistency and routine operation still play an essential role. The ATP with the performance characteristics and its criteria is shown in Table 1.

Regarding the detection setup, the PDA mode simultaneously captured the full UV spectrum during each injection, with validation data evaluated at a selected wavelength of 225 nm. A baseline filter was applied to minimize futile noise. Continuing with the AQbD process, the identification of critical method parameters (CMPs) focused on factors such as column selection, column temperature, flow rate, injection

volume, organic modifier, and mobile phase buffer. Each factor was optimized to ensure compatibility with potential MS-based method development. A Quality Risk Management (QRM) tool like the Ishikawa fishbone diagram was employed to assess the intrinsic relationship between the CMPs and CMAs (Fig. 3).

To complement the Ishikawa diagram, a semi-quantitative risk matrix was constructed to prioritize method parameters based on their potential influence on CMAs; it is included in the SI (section 2.1.; Table S2). Due to the inherent complexity of developing a method with broad applicability, an initial DoE was conducted to screen and evaluate key chromatographic characteristics and identify significant factors that influence method performance. Moreover, the DoE serves as a preliminary screening tool to establish a baseline for the method in terms of both detection and retention properties.⁴² A set of four different C18 columns was explored to achieve optimal retention, given the diverse separation behavior of the chosen substances, including an HSS T3 from Waters and a Triart Phenyl from YMC, which are standard options for retaining more hydrophilic APIs and can take up to a 100% aqueous mobile phase composition. Three MS-compatible buffers were examined to provide the best setup for sensitivity purposes,

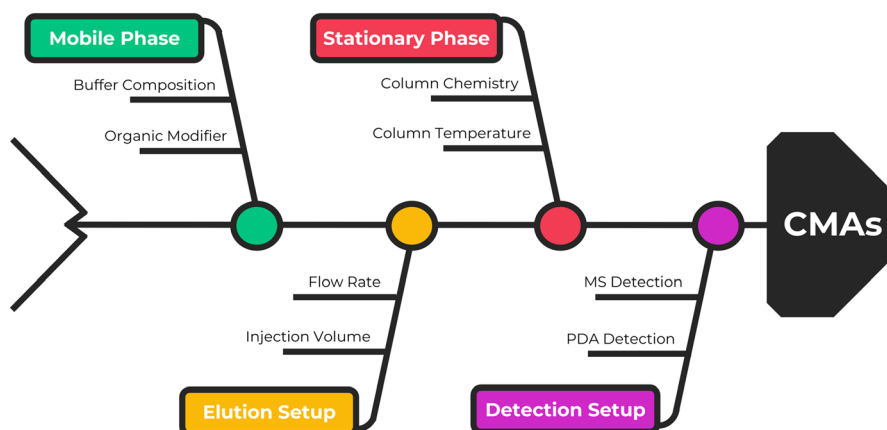


Fig. 3 Proposed Ishikawa diagram for the analytical method assessment.



Table 2 Factors and levels of the design of experiments for the analytical method development

Analytical column	Organic modifier (%)	Mobile phase buffer	Flow (mL min ⁻¹)	Injection volume (μL)	Column temperature (°C)
(A)	Acetonitrile methanol	10 mM ammonium formate	0.3–0.5	1–10	30–60
(B)		10 mM ammonium Acetate			
(C)		0.1% formic acid			
(D)					

Column letters corresponding to (A) Acquity HSS T3 (2.1 × 100 mm; 1.8 μm); (B) Triart C18 (2.1 × 100 mm; 1.9 μm); (C) Triart Phenyl (2.1 × 100 mm; 1.9 μm); (D) UltraHT Hydrosphere C18 (2.1 × 100 mm; 2.0 μm).

providing a broader scenario of retention compatibility with our APIs. The tested gradient started with a 95% aqueous buffer phase and proceeded towards the opposite side of the organic phase, to shape a scouting gradient, an ideal choice when the retention behavior of the analytes remains unknown.⁴³ The gradient speed was kept linear. Table 2 provides a complete overview of the chosen factors and levels of the development DoE.

3.2. Statistical evaluation of the development DoE

A response surface methodology (RSM) approach was employed to systematically optimize the HPLC method following Quality by Design principles. An I-optimal design for quadratic effects was implemented, comprising 50 experimental runs to ensure adequate model fitting and prediction capability. Due to the practical constraints of changing column temperature and organic modifier during the experimental sequence, these factors were designated as “hard-to-change” parameters, necessitating a split-plot design structure. The experimental design was organized into seven whole plots with two levels of organic modifier, where column temperature settings were nested within each organic modifier level. The split-plot arrangement resulted in acetonitrile and methanol being tested as separate whole plots with varying column temperature profiles. The chromatographic performance was evaluated through four critical response variables for the three APIs: USP signal-to-noise (USP *s/n*), *k'*, peak tailing, and USP resolution, currently listed in Table 1. The optimization strategy prioritized compliance with the selected ATP (Table 1) which is based on USP pharmacopeial requirements for *k'*, tailing, and resolution, while USP *s/n* served as a secondary objective for method robustness enhancement.⁴⁰

3.2.1. Desirability function approach. Multi-response optimization was performed using the desirability function methodology implemented in the Design Expert software. This approach transforms each response into a dimensionless desirability score ranging from 0 (completely undesirable) to 1 (entirely desirable), with the overall desirability calculated as the geometric mean of individual desirability. For each response, the desirability transformation was defined based on the optimization goal: maximization for USP *s/n* ratios, and target achievement for ATP-related parameters. All responses were weighted equally in the overall desirability calculation, ensuring balanced optimization across all CMA.

3.2.2. Optimization results and modeling validation. The optimization process identified optimal chromatographic conditions yielding an overall desirability of 0.689: flow rate of 0.478 mL min⁻¹, column temperature of 30 °C, injection volume of 10 μL, YMC UltraHT Hydrosphere C18 column, ammonium formate buffer, and methanol as organic modifier. The desirability profiles for individual factors are presented in Fig. 4.

Model predictions were validated through 13 independent confirmation runs using the optimized conditions, with injection volume adjusted to 5 μL to fulfil a satisfactory chromatographic performance aligned with our ATP (Table 1). Statistical evaluation compared predicted mean values against experimental results using 95% prediction intervals (PI), with results summarized in the SI (chapter 2.2.1). The optimization results derived from the development DoE, bridged with the robustness results from method validation, helped us establish the Method Operable Design Region (MODR). MODR is a multidimensional space that represents a scientific warrant that the method perform robustly and efficiently.

The I-optimal response surface design effectively identified optimal chromatographic conditions (flow rate 0.478 mL min⁻¹, column temperature 30 °C, injection volume 5 μL, YMC UltraHT Hydrosphere C18 column, ammonium formate buffer, methanol organic modifier), yielding an overall desirability of 0.689. Factor effect analysis revealed column type and mobile phase buffer composition as the most critical parameters influencing separation performance, with the YMC Ultra HT Hydrosphere C18 column providing superior chromatographic characteristics across the investigated design space.

Model validation through confirmation runs demonstrated that while not all predicted values fell within 95% prediction intervals, all experimental results remained well within the ATP frame (Table 1). This outcome highlights the practical success of optimization despite minor prediction deviations, confirming that the DoE approach delivered a specification-compliant analytical method.

3.3. Analytical method definition

As a result of the DoE evaluation, the final conditions of the analytical method were assessed (Table 3) and the same parameters were utilized for the validation according to ICH Q2 (R2) and M10 guidelines. An exemplary chromatogram obtained with this validated methodology is given in Fig. 5, where complete separation could be achieved in 9 minutes.



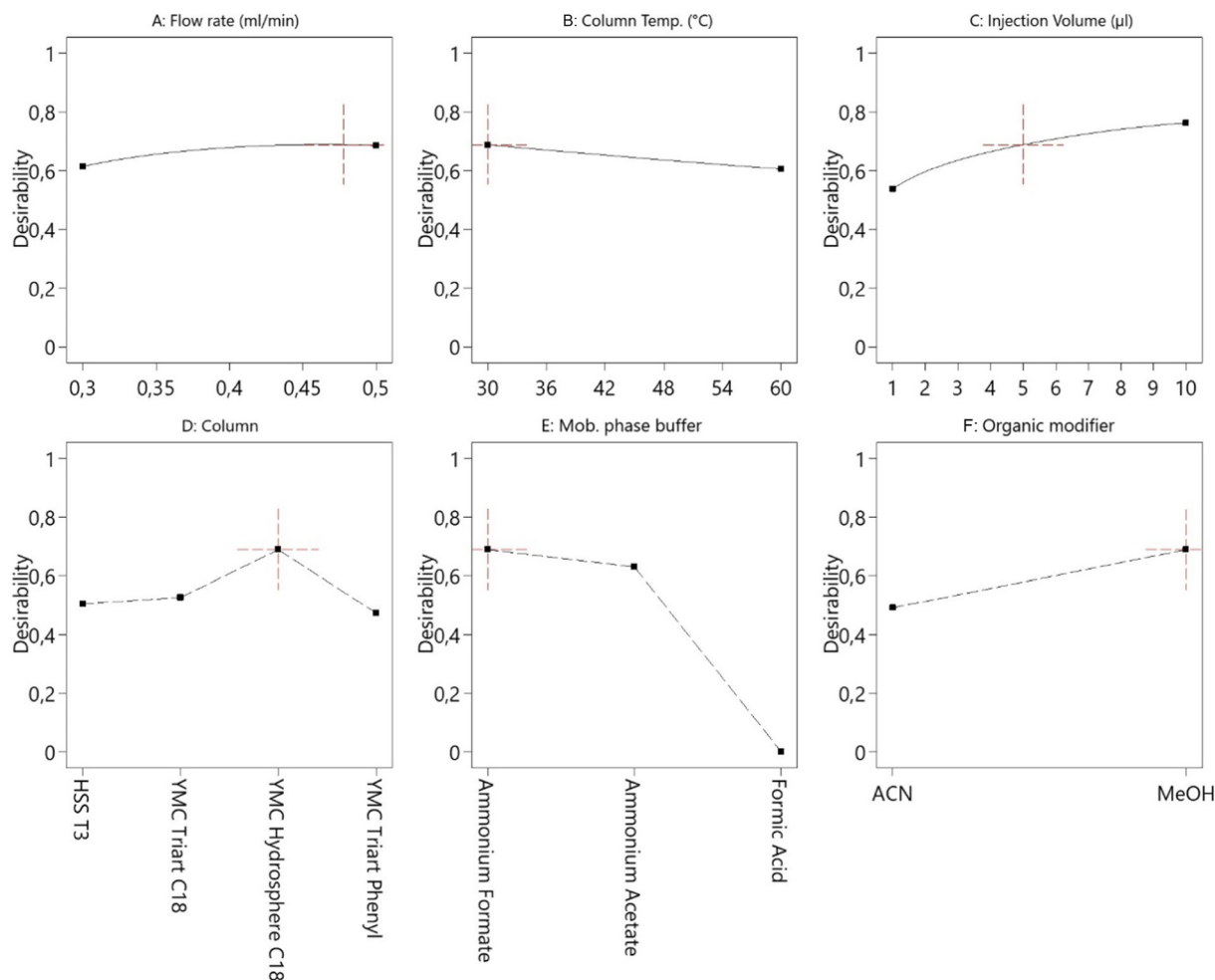


Fig. 4 Individual desirability functions show the influence of each factor on method performance: (A) flow rate; (B) column temperature; (C) injection volume; (D) column; (E) mobile phase buffer; and (F) organic modifier. The red cross indicates the optimal factor setting, and the desirability scale ranges from 0 (completely undesirable) to 1 (entirely desirable).

Table 3 Analytical method conditions used for validation

Validation conditions	
Column	YMC UltraHT hydrosphere C18
Column temperature	30 °C
Flow rate	0.478 mL min ⁻¹
Gradient	%B: 0.0 min 5%, 0.5 min 5%, 6.0 min 95%, 6.5 min 95%, 9.0 min 5%
Injection volume	5 μL
Mobile phase buffer (A)	10 mM ammonium formate
Organic modifier (B)	Methanol
Sample diluent	Methanol/water (50 : 50)
Wavelength	225 nm

3.4. Analytical method validation results ICH Q2 (R2)

3.4.1. Specificity. Specificity was proven by injecting the blank solution (equivalent to the sample diluent) and the

mobile phase into the chromatographic system. The results confirm that the analyte peaks were found pure and without external interference from other components at the elution RT across the analysis wavelength (225 nm). The chromatograms are depicted in Fig. 6.

The forced degradation studies and purity plots analysis confirmed the absence of interference under distinct stressing conditions, and the results highlighting the samples' purity are presented in the SI (chapter 2.3.1).

3.4.2. Linear response, LOD, and LOQ. Response data showed linear proportionality in the range between 100 μg mL⁻¹ and 500 μg mL⁻¹ for three independent linearities prepared for each API, providing a resulting R^2 -adj of 0.994 for combined regression data for SBS, 0.999 for GB, and 0.999 for BUD. Furthermore, the Shapiro-Wilk test on standardized residuals against predicted values from the regression model verified that they are normally distributed.⁴⁴ The p -value was assessed at 0.06 for SBS, 0.60 for GB, and 0.47 for BUD (acceptance criterion p -value > 0.05). The data is presented in the SI (chapter 2.3.2).



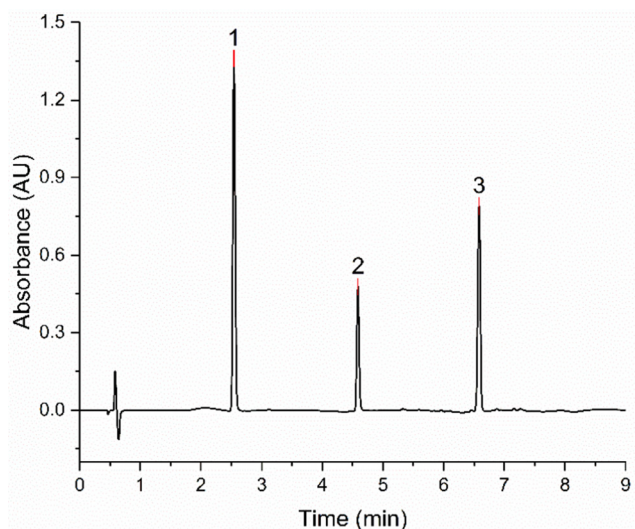


Fig. 5 Chromatogram of a $250 \mu\text{g mL}^{-1}$ injection of the three APIs mixture obtained with the final method conditions. (1) Salbutamol sulphate, (2) glycopyrronium bromide, (3) budesonide.

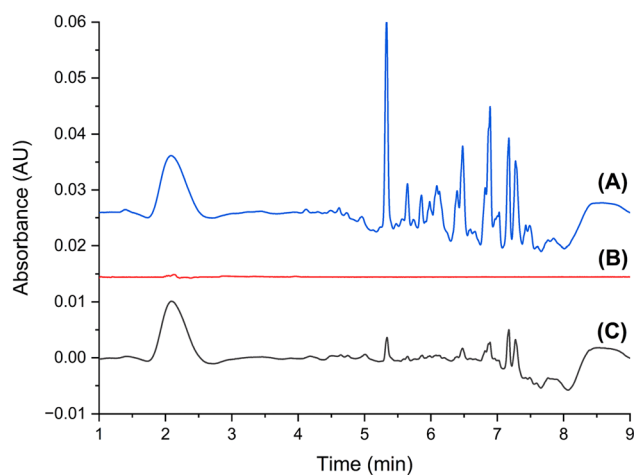


Fig. 6 Chromatograms of specificity studies. (A) Blank solution injection; (B) mobile phase injection; (C) SLF injection.

The LOD and LOQ were assessed by estimating the USP s/n ratio of prepared solutions at the lowest calibration point ($100 \mu\text{g mL}^{-1}$) against a blank injection in the interested region of the analyte. The results are summarized in the SI (chapter 2.3.2).

3.4.3. Accuracy and precision (repeatability and intermedicate precision). The accuracy was established by evaluating the mean recovery from at least nine replicate measurements across three concentration levels, 70%, 100%, and 130% of the labeled content, corresponding to 175, 250, and $325 \mu\text{g mL}^{-1}$, respectively. The method demonstrated compliant accuracy, with recoveries consistently falling within $100 \pm 2\%$ (Table 4).

The developed method exhibited high precision and reproducibility, with RSD below 2% for all measurements (Table 5).

Table 4 Accuracy results from method validation

API	Percent of target (%)	Recovery (%)	Mean (%)	RSD (%)
Salbutamol sulphate	70%	99.1	99.3	0.2
		99.3		
		99.5		
	100%	102.0	102.3	0.8
		101.7		
		103.2		
	130%	102.4	102.8	0.8
		102.2		
		103.7		
70%	101.0	99.9	1.2	
	98.6			
	100.1			
	100.6			
	100.2			
	100.2			
Glycopyrronium bromide	100%	100.6	100.4	0.4
		100.7		
		99.9		
	130%	100.6	100.4	0.3
		100.6		
		100.6		
70%	99.7	99.9	1.2	
	99.7			
	99.9			
	99.9			
	99.9			
	99.9			
Budesonide	100%	99.9	100.4	0.4
		99.7		
		99.9		
	130%	100.4	100.2	0.2
		100.2		
		100.0		

Table 5 Precision results from method validation

API	Parameter	Analyst I	Analyst II
Salbutamol sulphate	Mean	101.4	98.3
	SD	1.4	1.7
	RSD	1.4	1.8
	SEM	0.6	0.8
	n	6	6
	p -Value	1.0	
Glycopyrronium bromide	Mean	99.7	99.3
	SD	0.5	1.6
	RSD	0.5	1.6
	SEM	0.2	0.6
	n	6	6
	p -Value	0.5	
Budesonide	Mean	100.1	101.4
	SD	1.8	2.0
	RSD	1.8	2.0
	SEM	0.7	0.7
	n	6	6
	p -Value	0.3	

A Student's t -test revealed no statistically significant difference between the two analysts' results ($p > 0.05$), indicating insufficient evidence to conclude that the datasets differ.⁴⁵

3.4.4. Robustness. Following the optimization study, a comprehensive robustness evaluation was conducted to assess method stability under deliberate variations of critical parameters. An I-optimal response surface design for quadratic



effects was employed, comprising 30 experimental runs including replicates to ensure adequate statistical power for robustness assessment.

Three factors were systematically varied within realistic operational ranges: sample diluent composition (A: 40–60% methanol in sample preparation), formic acid concentration in mobile phase (B: 0.05–0.15%), and column variability (C: two different C18 stationary phases). The column factor included the optimized YMC UltraHT Hydrosphere C18 column alongside a Phenomenex Kinetex C18 column. The same response variables used in the optimization study were monitored: USP *s/n*, *k'*, peak tailing, and USP resolution for each API, with the same limits described in the ATP (Table 1) serving as acceptance criteria for the evaluation. The robustness assessment demonstrated that the optimized method maintained acceptable performance across all tested conditions, without violating the selected specifications throughout the 30 experimental runs. This outcome provides strong evidence for method robustness under the investigated parameter variations. The data is shown in the SI (chapter 2.3.3, Table S8).

Statistical modeling revealed varying degrees of complexity across the different API responses, as summarized in the SI (chapter 2.3.3, Tables S9–S19).

SBS responses presented the most challenging modeling scenarios:

- USP *s/n* ratio (adjusted $R^2 = 0.53$): required complex higher-order terms including A^2B and A^2C interactions to achieve an acceptable model fit. Despite including these quadratic interaction terms, the lack-of-fit test remained marginally significant ($p = 0.074$). The large difference between adjusted R^2 (0.53) and predicted R^2 (0.33) indicated potential issues with model adequacy. However, attempts at model reduction consistently resulted in a significant lack of fit, necessitating the retention of the complex model structure.

- Capacity factor (adjusted $R^2 = 0.62$): modeling required inclusion of the B^2 term for adequate fit, despite the main factor B (formic acid concentration) showing no significant effect ($p = 0.663$). This counterintuitive result highlights the complex response surface behavior.

- Tailing factor (adjusted $R^2 = 0.93$): achieved excellent model fit but required a complex nine-term model including multiple quadratic and interaction terms (A, C, AC, BC, A^2 , B^2 , A^2C , B^2C).

GB and BUD responses demonstrated much simpler and more interpretable patterns:

- For GB, the column was the dominant significant factor across all responses: *k'* (adjusted $R^2 = 0.97$), USP *s/n* (adjusted $R^2 = 0.49$), tailing (adjusted $R^2 = 0.14$), and resolution (adjusted $R^2 = 0.44$).

- For BUD, column type similarly dominated: USP *s/n* (adjusted $R^2 = 0.96$), *k'* (adjusted $R^2 = 0.20$), and tailing (adjusted $R^2 = 0.82$). The exception was resolution (adjusted $R^2 = 0.61$), where diluent composition (A, $p = 0.044$) and the AC interaction ($p = 0.049$) were also statistically significant alongside column type.

Technical investigation suggested that sample diluent composition (factor A) may contribute to baseline variability,

potentially affecting USP *s/n* ratios and peak tailing. This observation provides one possible explanation for the complex modeling behavior observed, particularly with SBS responses. All experimental results remained within the specification limits across the entire investigated parameter space, demonstrating maintenance of acceptable analytical performance under the tested robustness conditions.

The subsequent robustness evaluation across 30 experimental runs confirmed method stability under realistic operational variations. Despite encountering complex statistical modeling challenges, particularly for Salbutamol responses requiring higher-order polynomial terms, the fundamental robustness criterion was consistently met: all analytical results remained within USP pharmacopeial limits across the entire investigated parameter space.

The statistical complexity observed, especially the requirement for quadratic interaction terms in Salbutamol models and the counterintuitive significance of B^2 terms without corresponding main effects, likely reflects the influence of sample diluent composition on chromatographic baseline behavior. These findings underscore the multifactorial nature of chromatographic systems and the value of comprehensive experimental design in revealing such interactions.

The successful demonstration of method robustness across different C18 stationary phases (YMC Hydrosphere and Kinetex) addresses regulatory requirements for method transferability and provides confidence for implementation across multiple laboratory environments. The consistent performance of glycopyrronium and budesonide responses, contrasted with the more complex behavior of Salbutamol, suggests compound-specific susceptibility to analytical conditions that should be considered during method transfer protocols.

This study demonstrates both the power and limitations of statistical modeling in analytical method development. The varying model adequacy across robustness responses (*e.g.*, R^2 -adj ranging from 0.14 to 0.97) reflects the different sensitivity levels of individual analytical parameters to experimental variability. While ideally robustness studies would show no significant effects, the detection of significant but poorly predictable effects (*e.g.*, glycopyrronium tailing, R^2 -adj = 0.14) indicates inherent analytical variability that, while statistically detectable, remains within acceptable specification limits.

While complex response surfaces may challenge conventional modeling approaches, the ultimate measure of method suitability remains compliance with analytical specifications. The QbD approach successfully guided method development decisions despite statistical modeling complexities, emphasizing the importance of focusing on practical performance criteria alongside statistical rigor.

The comprehensive experimental design provided valuable insights into factor interactions that would be difficult to detect through traditional one-factor-at-a-time approaches, justifying the investment in systematic DoE methodology for critical analytical method development projects.



3.5. Analytical method validation results ICH M10

3.5.1. LC-MS method optimization. One of the potential benefits of applying AQbD to method development is the ability to translate an effective chromatographic setup from one detection system to another, enabling interchangeability among different technologies.^{46,47} In this study, the conditions provided by previous validation were further optimized using MS to achieve lower LODs and LOQs, supporting a more advantageous approach for bioanalytical applications. Therefore, while the retention setup remained the same, the three analyzed APIs could be well ionized in positive mode, at 240.1, 318.1, and 431.2 *m/z*, respectively. These were the most intense peaks observed in the MS scan, and thus, they were used for quantification. Satisfying sensitivity was achieved using a QDa single-quadrupole MS detector with a fixed probe temperature of 450 °C, a capillary voltage of 1.0 kV, and a cone voltage of 15 CV. The recorded ion mass spectra are displayed in the SI (chapter 2.4.1).

3.5.2. Selectivity, sensitivity, and matrix effect. The method exhibited no significant interferences at the RT of SBS (2.61 min), GB (5.18 min), and BUD (6.56 min) following six injections of individually prepared lots of SLF. In the SI (chapter 2.4.2), Fig. S10 shows typical LC-MS chromatograms obtained by spiking the analytes in blank SLF, demonstrating the absence of interference at the APIs' LLOQ compared to the blank. The response in the blanks was below 20% of the analyte response at LLOQ (Table S20). The LLOQ was determined to be 1 ng mL⁻¹ for all the APIs, with a USP *s/n* ratio > 10. The chromatograms are extracted from the MS spectra at pre-selected Signal Ion Recordings (SIRs), chosen from the abundance maximum (*e.g.*, 240.1 for SBS).

Specificity was proven by injecting a 1 µg mL⁻¹ solution prepared in working diluent (methanol/water) and recording the signal at the same SIR where the analytes were processed, as per selectivity. These injections were compared to the analytes' response at the LLOQ in SLF diluent. The chromatograms con-

firmed identical retention times, while the MS scan verified the compounds' identities (Fig. S11).

Regarding the matrix effect, no significant interference was observed when different lots of prepared SLF were used, defining a stable analysis regardless of the sample composition. The quantification showed no evidence of matrix effects, ion suppression, or ion enhancement, as the accuracy and precision fell within the target of ±15% of nominal concentration and RSD% (Table S21).

3.5.3. Calibration curve and range. The calibration curve plots for each performed linearity test demonstrate an acceptable correlation, with SBS showing an *R*²-adj of 0.999, while GB and BUD show an *R*²-adj of 0.997. The linearities were individually prepared and analyzed on different days, proving the reliability of the developed method. The plots are displayed in the SI with the analysis of the residuals (chapter 2.4.3). The back-calculated concentrations for each linearity (*n* = 3) fell within the established accuracy acceptance criteria, and more than 75% of the calibration samples respected the same criteria Table 6.

3.5.4. Accuracy and precision. Table 7 displays the accuracy and precision results for the method validation. The values complied with the acceptance criteria, providing a consistent scenario for routine applications. Moreover, these results prove the reinjection reproducibility.

3.5.5. Carry-over. The validated method did not exhibit any carry-over phenomena after injecting the ULOQ samples in the validation analytical run for the calibration range, considering that the response never exceeded the acceptance criteria.

3.5.6. Dilution integrity. The dilution integrity met the acceptance criteria, confirming that appropriate dilutions can be used when preparing calibration standards and QCs. The average accuracy of the dilution was +2.5%, expressed as RE%, and the precision was 5.89%, complying with the acceptance criteria.

3.5.7. Stability of solutions. The stability solutions were analyzed after the storage cycle, and the accuracy, expressed as

Table 6 Back-calculated concentrations from the calibration runs

API	Calibration level (ng mL ⁻¹)	Mean back-calculated (ng mL ⁻¹)	Accuracy (RE, %)	Precision (RSD, %)
Salbutamol sulphate	1.0	1.1	+10.7	3.0
	2.5	2.5	-1.6	1.2
	5.0	5.0	-0.5	1.4
	10.0	10.1	+1.4	2.5
	15.0	14.9	-1.0	0.7
	20.0	20.3	+1.3	0.9
Glycopyrronium bromide	1.0	1.0	+0.7	10.9
	2.5	2.5	-2.1	6.7
	5.0	5.4	+8.3	8.7
	10.0	10.3	+3.4	5.0
	15.0	16.1	+7.3	4.0
	20.0	21.5	+7.6	5.4
Budesonide	1.0	1.1	+8.0	11.0
	2.5	2.7	+6.5	6.6
	5.0	5.0	+0.8	4.8
	10.0	10.4	+3.6	2.4
	15.0	16.0	+6.9	0.2
	20.0	20.3	+1.6	0.8



Table 7 Accuracy and precision results from the method validation

API	Spiked concentration (ng ml ⁻¹)	Intra-day (n = 5)			Inter-day (n = 10)		
		mean (ng ml ⁻¹)	Accuracy (RE, %)	Precision (RSD, %)	Mean (ng ml ⁻¹)	Accuracy (RE, %)	Precision (RSD, %)
Salbutamol sulphate	1.0	1.0	-0.3	12.8	1.0	3.9	4.8
	3.0	3.2	+6.3	8.3	3.2	5.4	11.0
	10.0	10.4	+4.2	10.7	9.8	-1.9	2.1
	15.0	15.8	+5.3	8.7	14.7	-2.1	5.1
Glycopyrronium bromide	1.0	1.2	+8.1	7.4	1.0	-5.3	10.5
	3.0	3.3	+5.3	10.2	3.3	9.6	7.5
	10.0	11.3	+9.1	11.7	10.1	0.5	6.9
	15.0	16.9	+7.8	11.4	15.7	4.2	5.4
Budesonide	1.0	1.1	+9.9	11.5	1.1	9.1	7.3
	3.0	2.7	-12.4	6.3	3.2	4.8	7.9
	10.0	9.8	-1.8	6.0	9.8	-2.2	7.1
	15.0	15.4	+2.8	5.0	16.0	6.1	5.3

RE, relative error; RSD, relative standard deviation.

mean recovery (%), was evaluated. The results showed that the lowest recovery occurred after 48 hours at room temperature, although it did not drop below 5%. The data is presented in SI (chapter 2.4.4).

4. Conclusions

A unified UHPLC method was successfully developed and validated for the simultaneous quantification of budesonide, glycopyrronium bromide, and salbutamol sulphate, key APIs in inhalation therapy for asthma, COPD, and ACO. Using AQbD principles, the method was optimized with a strong understanding of critical parameters and validated according to ICH Q2 (R2) for pharmaceutical use. The approach was further extended to the bioanalytical field, with successful adaptation and validation in human plasma following ICH M10 guidelines. This demonstrates the feasibility and value of applying AQbD to bioanalysis, enhancing method robustness, flexibility, and regulatory confidence. By integrating AQbD into both pharmaceutical and bioanalytical workflows, this work supports more efficient, consistent, and science-based analytical development across the drug lifecycle.

Author contributions

Alessio Gaggero: methodology, data curation, investigation, visualization, formal analysis, writing – original draft preparation. Dalibor Jeremic: formal analysis, investigation, writing – original draft preparation. Anna Fedorko: investigation, formal analysis, methodology, writing – reviewing and editing. Jesús Alberto Afonso Urich: conceptualization, visualization, investigation, supervision, project administration, resources, writing – original draft preparation.

Conflicts of interest

The authors declare no conflict of interest.

Data availability

All data and materials are present in the manuscript and supplementary information (SI). Supplementary information is available. See DOI: <https://doi.org/10.1039/d5an01086a>.

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