IDENTIFICATION OF THE IMPURITIES OF BUDESONIDE USING SMALL PARTICLE LIQUID CHROMATOGRAPHY AND Q-TOF MASS SPECTROMETRY

THE SCIENCE OF WHAT'S POSSIBLE.™

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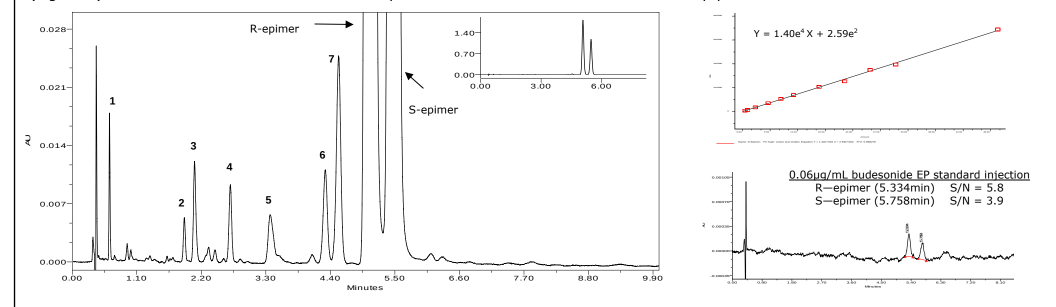
INTRODUCTION

The demands on the analytical laboratory to qualitatively and quantitatively determine active pharmaceutical ingredients and degradants continues to increase. The FDA regulations require companies to develop methods for their analysis and characterization of the APIs, as well as the impurities/degradants that could arise from the synthesis process, raw material provider, and/or storage conditions. The utilization of UV or PDA data alone for these analyses is often inadequate. Complications arise in many situations where compounds have (a) poor to no UV absorbance whether from lack of chromophores or (b) low level impurity concentrations exhibit poor UV spectral quality and peak purity becomes more difficult to identify when co-elution occurs. Mass spectral analysis becomes more essential as FDA regulations for impurity content reporting continues to decrease from 0.1% a few years ago to 0.05% today with expectations to decrease as instrumentation detection limits and techniques continue to become more sensitive. The importance of exact mass and MS fragmentation products to determine the structure of degradants/ impurities provides a higher understanding to the relationship of the impurity/degradant origin. We will demonstrate the utility of a UPLC-PDA-MS system and show the significant benefits in resolution, speed, and sensitivity using the ACQUITY UPLC[™] System and how this configuration will impact the identification of pharmaceuticals and their related substances.

To best illustrate this concept, we will analyze the pharmaceutical drug substance; budesonide. Budesonide is a glucocorticosteriod used for the treatment of asthma via various matrices and inhalation mechanisms. The official European Pharmacopeial assay was used as a guidance for the redevelopment of the budesonide assay and related substances for use with UPLC-MS. The new method will be used to determine various required qualitative system performance (eg; resolution, S/N, theoretical plates, tailing, symmetry factors). We will also demonstrate the quantification benefits (eg; limits of detection, limits of quantification) of this configuration. The impurity profiles of multiple batches from three manufacturers of pharmaceutical grade budesonide will be assessed and tested for EP related impurity compliance. Exact mass MS data will be collected to determine similarities/differences between the impurity profiles of the vendor batches. This increased performance makes UPLC[™]/PDA/MS the ideal tool for purity profiling.

UPLC-MS METHOD COMPATIBILITY Resulting Method

There are three published methods for the separation of budesonide and the related impurities.⁽¹⁻³⁾ Two of these methods are not "MS compatible" due to the use of phosphate buffers.^(2,3) The European Pharmacopoeia requires the following system suitability specifications based on a 500µg/mL budesonide test solution and reference solutions: (a) The resolution between the R-epimer and S-epimer is not less than 1.5 (b) run time 1.5x the retention of S-epimer, (c) the symmetry factor for the R-epimer peak is less than 1.5, (d) the theoretical plates calculated for the R-epimer peak is at least 4000, (e) and after six injections of the 500µg/mL reference solution the %RSD of the sum of the peak areas of the two epimers is at most 1.0%.⁽²⁾ In converting the method to UPLC we evaluated four currently available hybrid UPLC column chemistries, various pH, buffer concentrations, and temperatures. There was no significant change in separation when the pH was varied from 2.5 - 4.0, neither for pH 9.0 10mM ammonium bicarbonate nor for buffer concentrations from 10mM to 25mM. The resulting method showed the ACQUITY BEH C₁₈ 2.1 x 100mm column provided the best separation of the impurities from the budesonide API at 40^oC (Figure 1). 20mM ammonium formate buffer to pH 3.2 with formic acid was chosen to keep pH consistant with the EP method.



<u>Figure 1</u>: A 5µL full loop injection of 500µg/mL budesonide EP grade (Spectrum Quality Products) standard solution. UPLC method conditions: ACQUITY BEH C_{18} 2.1 x 100mm; 1.7µm, 68% 20mM ammonium formate buffer to pH 3.2: 32% acetonitrile; wavelength at 240nm; temperature at 40°C; flow rate at 0.6mL/min; 11,500psi. The additional peaks 1 thru 7 are identified as impurity peaks above 0.05% area.

Suitability Results

The %RSD of the sum of the areas of both epimers was 0.3% (n=6 injections) for the $500\mu g/mL$ budesonide EP grade standard solution The results in the table below are for the budesonide EP $500\mu g/mL$ standard solution. The requirements of the minimum European Pharmacopoeia specifications are met.

| Name | Retention Time | Resolution | Symmetry Factor | Signal/ Noise | EP Plates |
|-----------|-------------------|------------|--------------------|------------------|--------------|
| R— epimer | 5.073 | N/A | 1.05 | 10262 | 17011 |
| S— epimer | 5.476 | 2.46 | 1.02 | 6646 | 17390 |

IMPURITY PROFILE COMPARISONS

EP Related Substances Test

| European Pharmacopoeia Related Substances Test Specification | <u>Spectrum</u> EP grade 98% - 102% | <u>Sigma</u> >99% purity | Molekula 100.2% | <u>Spectrum</u> Research grade (no spec) |
|---|---|-----------------------------|--------------------|--|
| Individual Impurities (x < 2.5μg/mL Σof epimers areas) | Fail | Fail | Pass | Fail |
| Total Impurities (x < 7.5μg/mL Σof epimers areas) | Fail | Fail | Pass | Fail |
| R—epimer/S—epimer Ratio (S—epimer is 40.0% to 51% Σof epimers areas) | 59.24%/ 40.76% | 50.49%/ 49.51 | 51.38%/ 48.62 | 58.66%/ 41.34 |
| Purity | 98.24% | 97.99% | 99.52% | 98.07% |

Materials:

<u>Budesonide:</u> Spectrum quality products (New Brunswick, NJ); lot numbers: UI0628 (EP Rx grade); 98.0% - 102.0% and lot # RB2362 (research grade). Sigma Chemical Co. (St. Louis, MO); lot 81K1654; 99%. Molekula; batch# 52459 (Dorset, UK).

EXPERIMENTAL

<u>Reagents:</u> Acetonitrile Optima; Fisher Scientific (Fairlawn, NJ); Lot#050580. Ammonium formate 97%; Sigma-Aldrich (St. Louis, MO); batch # 04507AC. Formic acid 98%-100%; Reidel deHaën.

UPLC Conditions

| Instrument: ACQU | ITY UPLC |
|-------------------|---|
| Column: | ACQUITY UPLC [™] BEH C ₁₈ |
| Dimensions: | 100 x 2.1mm, 1.7µm |
| Mobile Phase: | 68% 20mM Ammonium formate |
| | (pH 3.6)/32% acetonitrile |
| Flow Rate: | 0.60 mL/min |
| Temperature: | 40 ⁰ C |
| Injection Volume: | 5 µL; full loop injection mode |
| Weak Wash: | 68:32 600µL (water: acetonitrile) |
| Strong Wash: | none |
| Detection: | ACQUITY PDA @ 240 nm with |
| | High sensitivity flow cell |
| Software: | Empower™ 2 CDS |
| | |

MS Conditions

Instrument: Waters Q-Tof Premier XE Software: Masslynx 4.1

Tune Page Parameters:

Source: ES+ Capillary (V): 3.2 Sample Cone (V): 35 for reference 20 for analyte Extraction Cone (V): 4.5 Desolvation Temp (⁰C): 350.0 Source Temp (⁰C): 150.0 Cone Flow (L/Hr): 0.0 Desolvation Flow (L/Hr): 800.0

Tof Settings

Acquisition Range: 100 - 800Da Scan Time: 0.20s Interscan delay: 0.05s Lock mass: 100fmol leucine/enkephalin @ ~20µL/min

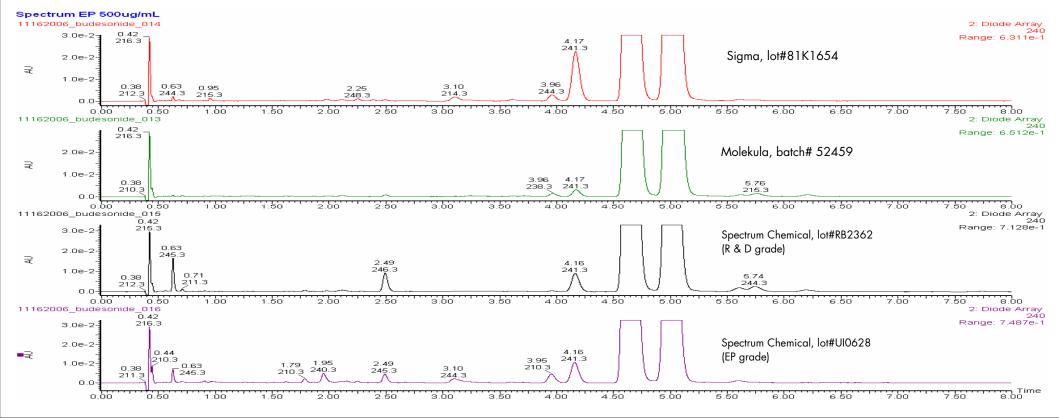
CONCLUSIONS

An Ultra Performance LC-MS method for budesonide and the related impurities was developed. European Pharmacopoeia specifications were met using the UPLC method. The resolution between the budesonide epimers was 2.5, EP plate count was 17,000, symmetry factor for the R-epimer was 1.05, and the area %RSD for six replicate injections of the 500µg/mL test solution was 0.3%. These results far exceeds the EP specifications and any other published HPLC methodology. A linear regression was constructed from the calibration curve to determine the LOD and LOQ of the method. It was determined that the LOD was 0.06µg/mL and the LOQ was 0.19µg/mL. The developed method was used for impurity profile comparisons of four different budesonide batches. Peak purity was used to determine method completion while using MS for peak identification. The method proved to be very specific in determining the differences and similarities between each of the suppliers batches. Exact mass was performed to yield a mass accuracy below 3ppm for each of the impurity peaks. An unknown mass of 447amu was determined to be a hydration products of one of the impurities D/impurity E and the respective epimer. The system configuration was shown to be sensitive and accurate for the determination and identification of impurities related to pharmaceutical drug substances.

The EP related substances test as described in the Budesonide EP monograph was performed on four different batch lots of budesonide which were purchased from three different vendors. Test solutions ($500\mu g/mL$) were prepared for each batch lot. Each test solution was diluted to yield two reference solutions each with concentrations of $2.5\mu g/mL$ and $7.5\mu g/mL$ with each representing 0.5% and 1.5% of the $500\mu g/mL$ mL solution, respectively.

Impurity Profile Comparisons

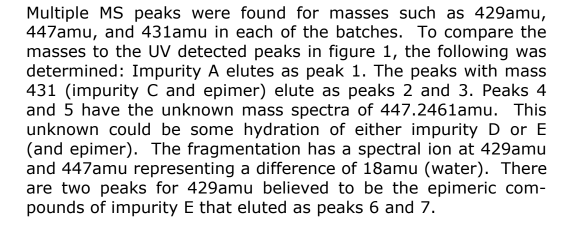
The impurity profiles for each lot was compared using the 500µg/mL test solution. There were varied impurities detected and at different levels for each sample when prepared fresh. It was observed after exposure to light and time, the profiles demonstrated more similarities.

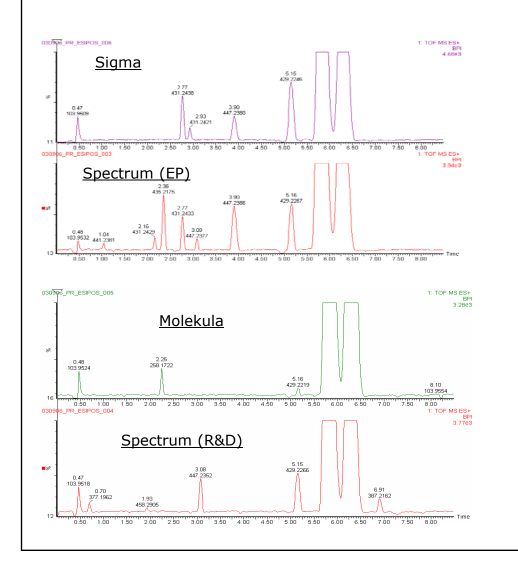


EXACT MASS

<u>MS Utility</u>

Using LC/MS in purity profiling experiments aids in peak tracking during method development and facilitates a high level of confidence with known analyte identification when exact mass is employed. ACS requires < 5ppm mass accuracy for patent submission and publication. The mass accuracy of the known peaks in each of the budesonide batches are less than 3ppm (below) . By coupling exact mass with tools like elemental composition, it is possible to predict molecular formulas for the unknown analytes such as the 447.2380amu.





| <u>Compound</u> | <u>[M+H]</u> | <u>Spectrum</u> | <u>Molekula</u> | <u>Sigma</u> | <u>Spectrum</u> |
|------------------|--------------|---|-----------------|--|-----------------|
| | | (EP) | | | (R&D) |
| Known Impurity | | | | | |
| Impurity A | 377.1964 | Image: A set of the set of the | | ~ | ✓ |
| Impurity B and F | 403.2120 | | | | |
| Impurity C | 431.2433 | ~ | < | Image: A second s | |
| Impurity D and E | 429.2277 | > | < | ~ | √ |
| Impurity G | 433.2512 | | ~ | | |
| Unknown Impurity | | | | _ | |
| 1 | 447.2461 | > | | _ | ✓ |
| 2 | 258.1722 | | | | |
| 3 | 435.2175 | | | | |

<u>Table 3 :</u> Each manufacturers' budesonide substance was analyzed by MS. The resulting table represents known and unknown impurities found in 500μ g/mL test solutions.

References

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