

Determination of Moxifloxacin in Human Plasma and Feces by LC-MS/MS

Martijn Hilhorst¹, Violaine Augustin² and Robbert Edens¹

QPS Netherlands B.V¹ and Da Volterra²

INTRODUCTION

Da Volterra devised a novel colontargeted adsorbent, DAV132, to be associated with antibiotics to prevent their impact on the microbiota, resulting in emergence and dissemination of resistant bacteria, antibiotic-associated diarrhea, and Clostridioides difficile infection. Da Volterra previously showed DAV132 ability to deliver a powerful adsorbent in the late ileum of human volunteers. Moxifloxacin is an antibiotic used to treat a number of bacterial infections. To support a clinical study of DAV132, a plasma method is developed and validated. As the amount of antibiotics in feces (unbound to feces particles) is a critical parameter in this study, a method in feces to determine the free fraction was developed and validated according to FDA and EMA guidelines.



MATERIALS AND METHODS

Plasma Sample preparation: protein precipitation was applied using 50.0 μ L sample and 400 μ L ACN:FA (100:1 v/v). After centrifuging the extracts were evaporated to dryness and reconstituted in 200 μ L Milli-Q:ACN:FA (85 : 15 : 1 (v/v/v)).

Feces sample preparation: liquid liquid extraction was applied using $100 \ \mu L$ feces filtrate and 3 mL TBME. The organic layer is evaporated to dryness and reconstituted.

Column: Phenomenex Gemini 5 µM C18 110 Å , 50 x 2 mm @ 600 µl/min. the gradient is 85% Mobile phaseThe start of A, after half a minute it decreases to 25% Mobile phase A followed by a flush step. ESI in applied positive mode was 402.3 monitoring: 1 384.4 (Moxifloxacin), 407.4 / 389.3 (Moxifloxacin-d5) using a Sciex API4000 mass spectrometer.

DIFFICULTIES WITH FECES ANALYSIS



CUSTOM-BUILT RESEARCH

VALIDATION RESULTS

Bioanalytical method validations where performed according to FDA and EMA guidelines.

Assay range: 10.0 - 5000 ng/mL for both plasma and feces extract (40.0 to 20,000 ng/gram feces).

plasma

Level	LLOQ 10.0 ng/mL	LOC 30.0 ng/mL	MQC 500 ng/mL	HQC 4000 ng/mL
Precision (%CV)	5.3%	3.8%	1.7%	1.2%
Accuracy (NRE)	-2.1%	+9.7%	-8.0%	+3.6%

Feces

Level	LLOQ 10.0 ng/mL	LOC 30.0 ng/mL	MQC 500 ng/mL	HQC 4000 ng/mL
Precision (%CV)	3.1%	1.9%	1.8%	1.6%
Accuracy (%RE)	+2.3%	+0.1%	-4.4%	+1.2%

All other validation items were also within criteria

CHROMATOGRAMS AT THE LLOQ LEVEL



CONCLUSION

The methods for the determination of moxifloxacin in human plasma and free moxifloxacin in human feces where successfully validated. The methods were applied for a clinical study (data not shown).

In this clinical study the free moxifloxacin fecal concentration as well as other clinical endpoints will be used to evaluate the clinical efficacy of DAV132 to protect the intestinal microbiota from the antibiotic-induced dysbiosis and prevent clinical manifestation of this dysbiosis in patients.

*) correspondence: martijn.hilhorst@qps.com

© 2019 QPS. Confidential. All rights reserved. QPS

