Identification and Quantification of Designer Drugs in Urine by LC-MS/MS

Using SCIEX QTRAP® LC-MS/MS Systems and Technology

Recently, trends have been seen within the drugs of abuse arenas to suggest that attempts are being made to bypass controlled substance laws, with novel compounds appearing on the market that are similar in structure to current drugs of abuse. These “designer drugs” or “legal highs” have caused concern due to their unknown quantity in terms of potency, side effects, health consequences and potential for abuse. As the number of new designer drugs is constantly rising, methods which can be easily expanded and have a non-selective sample preparation are needed.

The method presented here uses a simple dilute and shoot sample preparation, using a QTRAP® system to quantify and identify a number of these compounds in urine, specifically ketamine, norketamine, dehydronorketamine, 3-methoxyeticyclidine (3-MeO-PCE), 3-methoxyphencyclidine (3-MeO-PCP) and methoxetamine using a Scheduled MRM™ Algorithm.

To increase the reliability of identification by acquiring automated triggered MS/MS spectra and spectral library comparison, a second method was established utilizing QTRAP® technology. For this approach an information dependent acquisition (IDA) method was set up with one MRM transition per compound for detection and on-the-fly acquisition of EPI scans to obtain highly sensitive MS/MS spectra of the analytes for identification.

Key Features of SCIEX QTRAP® LC-MS/MS Systems for the identification and quantification of designer drugs.

- The QTRAP system offers high sensitivity data acquisition strategies for both quantitative and qualitative experiments
- System sensitivity allows small sample volumes and simplified sample processing procedures
- High-throughput quantitative analysis is achieved using MRM (Multiple Reaction Monitoring) and the Scheduled MRM™ Algorithm
- MRM triggered EPI (Enhanced Product Ion) scans generate high sensitivity MS/MS spectra to enhance confidence of identification of analytes, even at low concentrations.

Experimental

Sample Preparation: A simple dilution process has been chosen as sample preparation to enable a fast procedure without applying any selective extraction procedure:

- 20 μL of urine sample was added to 50 μL internal standard (D5-PCP in acetonitrile)
- Mixture was vortex mixed and centrifuged 10 minutes 15000 rpm (21500 x g)
- Supernatant diluted with 930 μL Mobile Phase A
- Solution vortex mixed and transferred to autosampler vial for injection

HPLC Conditions: LC separation was achieved on an Agilent 1290 HPLC system applying an 8 minute gradient on a Phenomenex Kinetex C18 100Å, 150 x 3 mm with KrudKatcher ULTRA HPLC In-Line Filter, 0.5 μm. Mobile phase A was water with formic acid and ammonium formate and mobile phase B was acetonitrile. The injection volume was 10 μL.
Mass Spectrometry Conditions: A SCIEX QTRAP® LC-MS/MS system equipped with a Turbo VTM Ion Source was used. Source and gas parameters were optimized for best performance, and individual compound parameters were optimized appropriately on infusion. At least 2 MRM transitions per compound were optimized.

The following Scheduled MRM™ Algorithm parameters were used: Target scan time: 0.3 sec, MRM detection window: 40 sec.

For the IDA method a target scan time of 0.1 sec was used. The EPI spectra for identification were acquired using dynamic fill time and collision energy spread to obtain detailed spectral information.

Generation of Reference Spectra: Reference EPI spectra were generated by analyzing the pure compounds applying a collision energy spread approach. This results in spectra which contain MS/MS fragments of at three different collision energies. The spectra acquired were combined into a library which was utilized for identification and confirmation.

Results and Discussion

Using this method, it was possible to analyze the compounds ketamine, norketamine, dehydronorketamine, 3-methoxyeticyclidine, 3-methoxyphencyclidine and methoxetamine with an injection - injection time of 8 minutes (Figure 2). The method showed to be very specific as no interfering signals could be observed in blank urine samples.

Quantitative performance has been demonstrated and shows accuracies within 15 % of nominal at the LOQ and %CV of 15 %, also at the LOQ and within 10 % for the higher concentrations.

An LOQ of 1 ng/mL after 1:50 dilution of the urine samples was estimated and additionally confirmed by signal to noise calculations (see Figure 3). An excellent linearity was obtained for all analytes in the applied concentration range of 1 to 200 ng/mL with an applied weighting factor of 1/x² (see Figure 4).

Figure 3. Chromatogram of all analytes in spiked urine at 1 ng/mL after 1:50 dilution. The signal to noise ratio was calculated by dividing the average background signal intensity from the peak by 3 times the standard deviation of the noise region.

Figure 4. Calibration curves of all tested designer drugs in five different urine samples.

All analytes were infused post column using a tee connector during injection of solvent A and a blank urine sample, which was diluted 1:50 and 1:10 respectively. Figure 5 clearly shows the benefit of a high dilution factor for urine analysis as this causes a significant decrease of ion suppression. At a dilution factor of 50 no ion suppression could be observed compared to the solvent injection and the injection of 1:10 diluted urine at relevant retention times.
Conclusions

An LC-MS/MS method for the analysis of 6 designer drugs was achieved utilizing a designer drug library for high confidence identification either by at least two MRM transitions per compound or performing an information dependent acquisition to generate MS/MS spectra using the linear ion trap of the QTRAP® system. The obtained MS/MS spectra can be searched against spectral libraries using the Analyst® Reporter Software for unambiguous identification and confirmation.

The use of the scheduled MRM algorithm allows inclusion of further analytes and internal standards without sacrificing cycle time and therefor data points across the chromatographic peaks.

The very simple, fast and non-selective dilution step shows to be a reliable and robust sample preparation and ensures that no analytes are lost which can be the case if a selective extraction procedure is performed instead.