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Nota de aplicación

Systematic Toxicological Analysis (STA)
Using the ACQUITY QDa Mass Detector Part 1: An introduction to using mass
detection in forensic toxicology screening

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For forensic toxicology use only.

This is an Application Brief and does not contain a detailed Experimental section.

Abstract

This application brief demonstrates to evaluate the Waters STA methodology on the ACQUITY QDa Mass Detector to facilitate screening of toxicologically relevant compounds in various biological matrices.

Benefits

In the first of a two-part series on STA using the ACQUITY QDa Mass Detector, we review the strengths and weaknesses of traditional screening methods and describe a simplified screening technique for the detection of toxicologically-relevant analytes in biological matrices.

Introduction

General unknown screening or systematic toxicological analysis (STA) is an essential element of the workflow applied within forensic toxicology laboratories. The main purpose of this initial screening technique is to identify putative positive samples, i.e., those containing relevant drug substances, while simultaneously eliminating negative specimens from any subsequent analytical interrogation. Typically, urine and plasma/serum are the specimens of choice.

Traditional techniques have included immunoassay (IA), gas chromatography in conjunction with mass spectrometry (GC-MS), and liquid chromatography (LC) with ultraviolet (UV) or photodiode array (PDA). Although commonly used, each of these techniques have some significant limitations, which are summarized in Table 1.

Technique		Strengths	Weaknesses
Immunoassay		Quick and simple as no chromatographic separation required	Not a fully comprehensive screen; multiple immunoassays required for each specimen - can be costly Class-specific test only (e.g., reports 'Benzodiazepine Positive' but does not indicate which specific Benzodiazepine drug is present)
			Not available for all drugs/drug classes (e.g., NPS) Cross-reactivity for NPS with common immunoassays is unclear or variable
			Some immunoassays are subject to false positive or false negative results
Gas Chromatography	Mass Spectrometry	Chromatographic separation – good for biological samples and provides reference retention time (RT) for use in identification process Availability and reproducibility of	Generic chromatographic conditions need to be applied for a comprehensive screen – may not be optimal conditions for all analytes
			To achieve appropriate sensitivity chemical derivatization is commonly required. Different derivatization protocols may have to be applied to
		mass spectral libraries based on electron impact (EI) ionization is beneficial. Identification based on mass to charge (m/z) of the precursor and fragment ions	each specimen Thermolabile analytes such as benzodiazepines are not amenable to the high temperatures applied in GC
		Retrospective data interrogation is possible	Poor sensitivity for polar drugs, metabolites and non-volatile substances
		To possible	Degradation products and remaining derivatization compounds can complicate interpretation of spectral data
Liquid Chromatography	Mass Spectrometry	Direct analysis – chemical derivatization is not required	Generic chromatographic conditions need to be applied for a comprehensive screen – may not provide optimal conditions for all analytes
		LC amenable to widest range of analytes	
		Chromatographic separation - good for complex biological samples and provides a reference RT for use in identification	
		Availability of mass spectral libraries is beneficial and provides high specificity	
		Electrospray is a 'soft' ionization technique and yields an abundance of structural information – identification is based on matching <i>m/z</i> of precursor and fragment ions	
Liquid Chromatography	UV or PDA detection	Direct analysis - chemical derivatization is not required	Single spectra used for identification – poor sensitivity and specificity especially in complex mixture and matrices
		LC amenable to widest range of analytes	
		Chromatographic separation – good for complex biological samples and provides a reference RT for use in identification	
		Availability of UV or PDA spectral libraries is beneficial	

Table 1. A comparison of the strengths and weaknesses of various screening techniques.

In brief, immunoassays commonly generate class-specific screening results, thus multiple kits are required to provide information for the main drug classes. In addition, assays are not available for all drugs and drug classes, therefore this approach provides a very restricted screen. False positive and false negative identifications are also a significant, and well-documented, disadvantage of immunoassays.¹

For many years, gas chromatography coupled with mass spectrometry was considered the "gold-standard" comprehensive screen and the mainstay of many toxicology laboratories. The availability of large commercial libraries based on electron impact (EI) ionization containing data for xenobiotics is certainly beneficial to the user; EI is considered to be a hard ionization technique and some analytes may be poorly detected owing to excessive fragmentation. In order to improve sensitivity, chemical derivatization is often used, which is both time-consuming and can bring additional problems. For these reasons, over the last two decades, liquid chromatographic techniques have been steadily replacing GC.

In contrast to GC, LC permits direct analysis of the drug substances without the need to chemically alter the molecule and is suitable for a wider range of drugs. Techniques based on the combination of LC and MS offer a simpler, yet more sensitive approach. Mass spectrometry detection also offers superior selectivity over PDA or UV detection and provides analyte specific detection rather than the class-specific approach of immunoassay. These advantages combine to make LC-MS the ideal tool for screening in the forensic toxicology market.

Waters first introduced the combination of LC with MS detection for STA in 2007.² Over the intervening years, this methodology evolved to reflect the advancements in technology available from Waters and users can now search against a spectral library containing 1200 compounds.³⁻⁵

Waters introduced the ACQUITY QDa Mass Detector in 2013. This instrument has already been shown to be applicable to both pharmaceutical and illicit drug identification.⁶

In this technology brief, we describe the application of the same approach to biological samples for the purposes of comprehensive toxicological screening.

Results and Discussion

Screening Method

LC-MS using electrospray ionization (ESI) is ideally suited to polar, non-volatile, and thermally unstable compounds, and therefore provides a powerful means of identifying many toxicologically-relevant

compounds rapidly without the need for sample derivatization, unlike GC-MS. Electrospray is a soft ionization technique that mainly leads to protonated molecules in positive mode and to deprotonated molecules in negative mode. In order to get more specific structural information, it is possible to induce fragmentation of these molecular ions in the source region of the QDa. This can be achieved by increasing the voltage applied to the sampling cone. Molecular ions then collide with neutral molecules in the source region and fragment into characteristic ions. This is referred to as in-source collision-induced dissociation (CID). Using in-source CID it is possible to generate mass spectra of different fragmentation patterns according to the value of cone voltage applied in the source (see Figure 1). Through this process, reproducible LC-MS mass spectra can be used to produce a library of mass spectra.

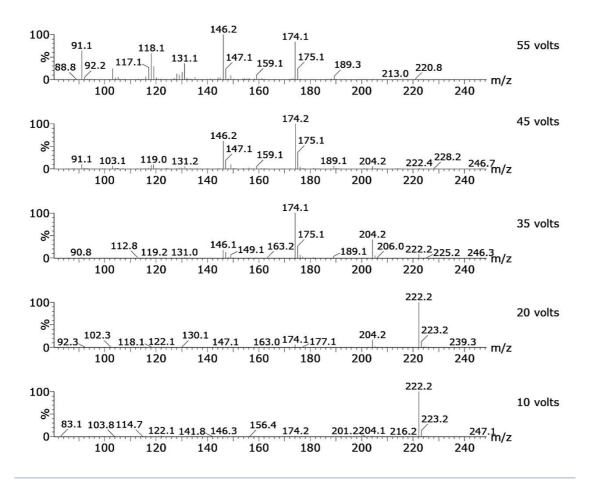


Figure 1. Spectra showing the in-source CID fragmentation pattern obtained for the Novel Psychoactive Substance (NPS) ethylone (bk-MDEA) at 200 ng/mL when different voltages are applied to the cone.

The ACQUITY QDa Mass Detector from Waters requires no specialist training or expertise to set up and all

acquisition and processing methods used here are available free of charge from Waters Marketplace. It is the only mass detector that integrates with, and even fits on top of, your instrument stack. It uses less bench space and less energy than a traditional mass spectrometer. Cleaning and routine maintenance are minimal, thereby maximizing your uptime. It can be integrated with other Waters software platforms and detection techniques, such as UV or PDA.

ChromaLynx is an application manager within MassLynx that allows for deconvolution of the mass spectral data and automatically processes and compares the data to the preconfigured library (containing 1200 compounds), which provides an identification through spectral library matching. The confidence with which a substance is identified is presented as an average match factor, which has a maximum value of 1000. All compounds are required to elute within 0.35 min of the retention time in the supplied library. Customers can create their own libraries from reference material or from authentic material data. Further, it is also possible to re-process previously acquired data, with updated libraries, without the need to prepare fresh samples.

Results

One hundred toxicologically-relevant analytes were spiked into both control human urine and human plasma at 200 ng/mL and prepared using a simple Oasis PRiME HLB µElution method and analyzed using the Waters toxicology STA screening method on a Waters ACQUITY QDa Mass Detector.⁷

In this study, a positive result was achieved by having an average match factor of >700; an example of the results obtained using ChromaLynx of a mixture of compounds spiked into human urine at 200 ng/mL is shown in Figure 2.

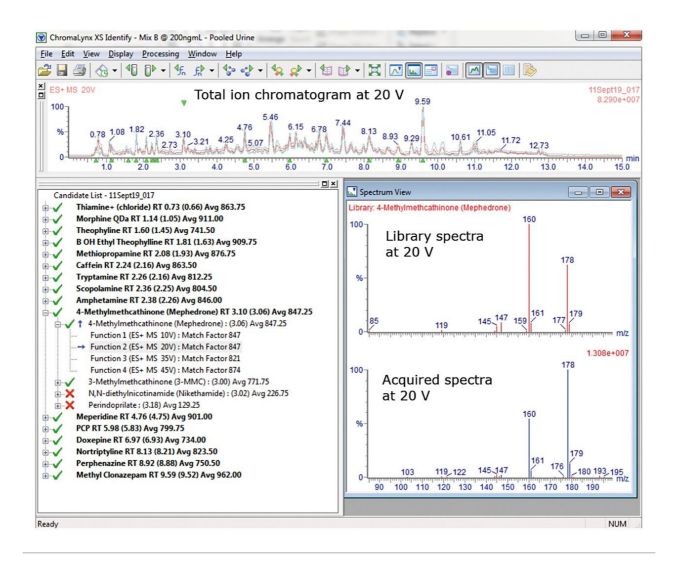


Figure 2. ChromaLynx results browser showing the analysis of a mixture of compounds spiked into human urine at 200 ng/mL, highlighting the identification of 4-methylmethcathinone (mephedrone).

Conclusion

In this technology brief a combination of liquid chromatography and ACQUITY QDa Mass Detection has been shown to provide a quick, simple, and effective way to screen for toxicologically-relevant compounds in various biological matrices. This technique offers benefits of cost savings, speed, and specificity over other established screening techniques such as immunoassay, GC-MS, and LC-UV. The same system can also be used to provide quantitative data as described by Mistry et al.⁸

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