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A ‘Periodic Table’ of mass spectrometry instrumentation and acronyms†

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Aim

The purpose of this manuscript is to describe the structure of typical MS instrumentation (sample introduction, ionisation source, mass analyser, detector, data acquisition/processing) whereby each section will be dealt with in turn, and summarised in tabulated format. With the many different types of instrumentation there is a clear need for understanding not just the relevant acronyms of the technologies involved but when and, importantly, why they are used in combination. The AMC decided in preparing such a guide that the form of a recognisable ‘Periodic Table’ (Fig. 1) is ideal for presenting the terminology to (non-expert) users of MS instrumentation. In particular it should facilitate decoding future sets of acronyms as well as those already in use. Most of the terms in the table are cross-referenced to the IUPAC document but it is not feasible to include over 500 terms in the table format. Hence, the acronym entries have been limited to those most widely used, and categories such as the atmospheric and ambient ionisation methods are ‘illustrative’ rather than comprehensive.

Introduction

Mass spectrometry is the study of ions that are counted according to their mass-to-charge ratio (m/z). The field of MS and the resulting technology has expanded rapidly in the last 30 years to accommodate a range of applications for the analysis of both organic and inorganic compounds. As a result of technological developments, the number of acronyms, abbreviations and terms encountered in the literature has also increased. Mass spectrometers are available in a range of shapes and sizes, and can perform quite different tasks depending on how they are operated. However, they typically consist of the following components:

1. Sample introduction
2. Ionisation source
3. Mass analyser
4. Detector
(5) Data acquisition/processing

Samples that are analysed by MS can be in solid, liquid or gas form providing the sample can be ionised for mass analysis. The sample introduction method and ionisation sources available depend on the particular requirement. There are also a range of mass analysers available; each mass analyser will have characteristics that will lend itself to gaining information-rich data (mass spectra), for qualitative analysis or for measuring the relative abundance of a compound for quantification. Therefore, there is a broad and varied landscape of available techniques, for the analysis of both organic and inorganic materials, that may be combined to assemble a mass spectrometer designed to meet a specific purpose.

Sample introduction

The sample introduction to a mass spectrometer is chosen according to the nature of the analyte and the complexity of the test material. Analysis of simple test materials, such as ‘pure’ substances, may use straightforward sample introduction such as direct infusion of liquids or headspace sampling of a gas. More complex samples typically require a preliminary separation, either offline or online, by using techniques such as chromatography or electrophoresis to overcome the limitations of the other components of the mass spectrometer. This flexible functionality of the sample introduction enables the advantages of MS, specifically selectivity and sensitivity, to be enhanced and provide a platform capable of trace level detection and quantitation of analytes within complex mixtures.

Ionisation source

Ionisation of target components of the sample can be carried out under a range of conditions, from atmospheric pressure to near vacuum. The ionisation source is chosen to suit the analysis of certain sample chemistries and is used in conjunction with compatible sample introduction methods. Ionisation techniques include chemical modification, thermal desorption, particle bombardment, and laser ablation, with vaporisation (if necessary) into the gas phase. Typical ionisation processes include the loss or addition of an electron, or a charged reagent species and are regarded as high energy (hard ionisation) or low energy (soft ionisation) processes causing significant or minimal fragmentation of the precursor species respectively. Appropriate selection may provide an information-rich data set of chemical structure (hard) or intact elemental/molecular information (soft).

Mass analyser (ion separation)

Mass analysis involves the separation of sample ions according to their m/z. There are a number of methods available to separate ions: spatially (by application of electric or magnetic fields), by measuring an ion’s time-of-flight (ToF), or by monitoring the frequency of ion motion. (This process of separation is carried out under low pressure (typically vacuum) and is quite different to the technique known as ion mobility, which can separate species according to shape and size and can be carried out under atmospheric conditions.) The methods of mass separation may also be classified according to how the analyser operates for the ions to be detected; typically, ions are ‘scanned’, as in a quadrupole analyser, or ‘pulsed’ as in a time-of-flight analyser. These operational considerations are suited to particular applications and are chosen for achieving specific information and combined with compatible ionisation sources and methods of detection.

Detectors

Abbreviations associated with mass spectrometers typically concern sample introduction, ionisation and mass analysis but this should not lessen the importance of ion detection methods. Ion detection will typically involve either the collection of ions (e.g., the Faraday cup), the bombardment of a charged surface to generate secondary electrons (e.g., the electron multiplier), or the monitoring of the frequency of ion movement between electrodes (e.g., image current detection) after ions have exited the mass analyser. Detectors are chosen to suit particular mass analysers and according to whether the accuracy of the ion count or the sensitivity of detection is of primary importance for the analysis.

Data acquisition/processing

Signals generated from the detection of ions require processing before viewing as an output (e.g., a mass spectrum) and this can be accomplished using different methods that involve digitising an electrical signal. Typically, this is achieved by monitoring the current from the detector. That can provide a voltage or ion counting signal (time-to-digital converter, TDC, or analogue-to-digital converter, ADC) or a Fourier transform of a frequency signal derived from an ion’s motion, and converting this into a digital signal that is further processed. Through the operational software of the mass spectrometer the user is then able to observe the mass spectrum, and interrogate and process the data for the desired application.

Operational modes

Mass spectra may be obtained using different acquisition modes covering a broad selection of m/z, such as a ‘full scan’, or specified m/z in the form of single or selected ion monitoring (SIM). Spectra may also be recorded for fragment ions of the precursor species with a range of approaches used to elicit breaking of the chemical structure. These fragmentation methods may be undertaken using suitable ionisation sources or mass analysers (i.e. a ‘hard’ electron ionisation source, or ion trap, respectively), or by using a collision cell with the mass analyser, such as those used in tandem mass spectrometers (e.g. MS/MS). Again, the combination of instrumentation components is chosen to achieve certain types of data necessary for specific applications.
Fig. 1 Periodic table of mass spectrometry (MS) terms.
Using the ‘Periodic Table’

(Note: the Table in the printed version of the Brief is for illustration only with no live links. The ESI† version contains links to access references for further information on each acronym.)

Mass spectrometry instruments and techniques are commonly named by using a string of acronyms each of which refers to one of the five major components described above. The string is usually assembled from left to right, beginning with sample introduction and finishing with the operational or data acquisition mode. The table reflects this in a series of columns moving across from left to right. Each column lists the more widely used current options for that component as a vertical series of boxes for which the key entry is the acronym of that option. Additional boxes for ion mobility separation are included in a separate row from the main table as variations of this technique may be used for sample introduction or mass analysis. Within each box is the description of the acronym and a number of abbreviations providing useful information about the component, as indicated in the legend to the table shown also in Fig. 2 below. The entry at top right in each box comprises the corresponding IUPAC number (if available) and a reference for further information. This reference can be accessed directly with electronic versions of the paper by clicking the acronym. An example of using the table to decipher technique acronyms is given at the top of the table shown also in Fig. 3 below.

Concluding remarks

The expanded range of instrumentation types available enables MS users to carry out a more targeted approach to analysis. This allows the analyst to tailor the method development to achieve the information necessary for the test material and the desired outcome. With the wide choice of instrumentation there is also great flexibility by virtue of certain modules (i.e. the sample introduction and ionisation source) often being interchangeable. This is evident in recent instrument developments offering alternative methods of ionisation within a universal housing. However, with these developments there is perfec a need to understand the relevant acronyms and when and why the technologies are used in combination. This Periodic Table of MS terms may offer some assistance in understanding these combinations.

For techniques not covered by IUPAC/terminology guides within the reference list seminal references have been cited where possible.

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This report was prepared for the Analytical Methods Committee with contributions from members of the AMC Instrumental Analysis Sub-committee and approved by the AMC on 03/07/17.

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