Integrating ion mobility spectrometry into mass spectrometry-based exposome measurements: what can it add and how far can it go?

Measuring the exposome remains a challenge due to the range and number of anthropogenic molecules that are encountered in our daily lives, as well as the complex systemic responses to these exposures. One option for improving the coverage, dynamic range and throughput of measurements is to incorporate ion mobility spectrometry (IMS) into current MS-based analytical methods. The implementation of IMS in exposomics studies will lead to more frequent observations of previously undetected chemicals and metabolites. LC-IMS-MS will provide increased overall measurement dynamic range, resulting in detections of lower abundance molecules. Alternatively, the throughput of IMS-MS alone will provide the opportunity to analyze many thousands of longitudinal samples over lifetimes of exposure, capturing evidence of transitory accumulations of chemicals or metabolites. The volume of data corresponding to these new chemical observations will almost certainly outpace the generation of reference data to enable their confident identification. In this perspective, we briefly review the state-of-the-art in measuring the exposome, and discuss the potential use for IMS-MS and the physico-chemical property of collisional cross section in both exposure assessment and molecular identification.

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The majority of human disease (>90%) is attributable to a combination of individual genetic factors and nongenetic ‘environmental factors’ [1–3]. The former can be readily assessed using rapid genome sequencing technologies, whereas the latter can be assessed or estimated using a variety of analytical methods [3,4]. To represent these non-genetic environmental factors, the concept of the ‘exposome’ was defined by Christopher Wild as the sum of all exposures over a lifetime to chemical, social and biological agents that influence human health [1]. The exposome was proposed as the natural complement to the genome, with the recognition that the phenome (sum of individual traits) was a product of both the genome and an exposome – the environment within which genes are expressed.

The exposome embraces the totality of both external and internal exposure. External exposure is mainly to chemicals in the air, water, soil, diet or consumer products, and internal exposures correlate to molecular changes within body compartments. Furthermore, these internal exposures can be to products of normal metabolism or in response to external exposures, including chemical, infectious or social [5]. External exposure measurements are often more convenient and cost effective, particularly when the goal is to identify sources of exposure. Internal exposures are often preferred when the objective is to identify exposure–disease interactions.

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relationships because they better represent concentrations at or near the biological site of action. The extensive set of sample matrices – from blood, breath, hair and nails to soil and water – as well as the range of complexity or ‘space’ comprising xenobiotic chemicals and endogenous molecules, is a challenge for analytical chemists attempting to measure the exposome. As the exposome is the ‘totality’ of exposure, each measurement will provide a snapshot (or piece of the puzzle) that can be used to build the bigger picture.

The use of ‘omics’ technologies to characterize the exposome [2,6–9], originally proposed by Wild [1], considers two distinct approaches: direct measurement of chemical exposures by measuring parent compounds and their transformation products in environmental samples or parent compounds and their metabolites in biofluids and tissues using similar analytical approaches as implemented in metabolomics, and inference of an exposure based on biological signatures (i.e., the ‘phenome’) obtained by one or more complementary approaches, such as transcriptomics or proteomics. A significant difference among these omics approaches is that genomics, transcriptomics and proteomics focus on a narrow chemistry – polymers of nucleotides and amino acids – whereas direct measurement of chemical exposure and of the metabolome requires dedicated or, better, complimentary approaches suitable for a very diverse chemical space [10–15] (Figure 1). Although the average molecular composition of a peptide [16] is not significantly different from that of a metabolite [17], the peptide chemistry is constrained to the typical peptide bonding between a fixed set of amino acids. In contrast, the metabolite chemistry is constrained only by the organism (for endogenous metabolites) and the boundaries of chemistry itself, while some xenobiotic compounds of anthropogenic origin even push the boundaries of chemistry.

In the search for causes of human disease, the genomics revolution brought unprecedented ability to obtain genetic information across individuals and populations, and a deeper appreciation for the importance of exposure as causative agent, yet tools for measuring the exposome were far less developed. For example, while advancements are continually being made in methods for broadly and accurately identifying metabolites in metabolomics studies [18–31], these developments are still relatively immature compared with the ease, robustness and throughput with which the proteomics community confidently identifies peptide sequences [32–34]. The expansion of more conventional analytical approaches to handle higher sample throughput, and to measure thousands of chemicals, will allow comprehensive characterization of chemical exposures across larger populations. Combined with emerging untargeted analytical methods (e.g., in metabolomics), there is the expectation that previously unknown or unexpected exposures will be identified, improving the search for environmental drivers of disease and providing exposure data to guide chemical selection for toxicity testing. Applied to both human (e.g., blood and urine) and environmental samples (e.g., water, air, dust and soil), and linked through geographical information systems, these new analytical approaches will also advance the search for sources of exposure, even when the identity of some chemicals involved remain unknown.

In this perspective paper, we will briefly review the state-of-the-art in measuring the small, organic molecular components of the exposome and discuss the potential use for ion mobility spectrometry-mass spectrometry (IMS-MS) and the physicochemical property of collisional cross section (CCS) in both exposure assessment and molecular identification in metabolomics studies.

**State-of-the-art in measuring the exposome**

Current exposure science leverages many disciplines to assess level(s) of environmental exposure and risk thereof, including epidemiology, toxicology and analytical chemistry, with the latter being used for environmental and biological monitoring [5]. In this section, state-of-the-art approaches used in environmental and biomonitoring are illustrated. Specifically, methods used to identify molecular signatures of exposure, whether evidence of parent compounds and their metabolites (i.e., direct measurement) or of pathogens, or the host’s complex biological response to these molecules and organisms (i.e., indirect measurement) will be discussed. We further limit our discussion to direct measurements of chemical exposure and indirect inference of an exposure based on biological signatures contained in the metabolome. Macromolecules and metals will not be covered.

Regardless of whether direct or indirect measures of exposure are performed, the technical approaches used largely fall under one of two categories: targeted or untargeted. The next two sections will provide an overview of these two approaches, citing specific examples and covering benefits and caveats of each.

**Targeted analysis**

Methods for targeted analysis in assessment of the exposome focus on a single analyte, a class of chemically similar analytes or a set of analytes whose chemistries are sufficiently similar to allow their measurement in a single analysis, all of which aim to address specific questions of exposure. The analytical protocols for such methods, including sample preparation,
have been optimized to provide accurate identification and quantification with high sensitivity (i.e., true positive rate) and specificity (i.e., true negative rate) in select matrices [35]. Select examples of targeted analysis include a multipanel LC-MS assay to assess exposure to pesticides, veterinary drugs and parabens [36], GC-MS analysis of flame retardants after flow-through air sampling [37] and the measurement of estrogens in wastewater samples using GC-MS [38]. The assays used for targeted analysis can be as simple as a glucose colorimetric [39] kit to determine the degree of hyperglycemia [40] associated with the response to streptozotocin [41], or certain drugs [42], to as complex as chromatography coupled with MS to quantify oxygenated polycyclic aromatic hydrocarbons [43]. Regardless of the degree of complexity of the assay, targeted analysis is rooted in the fundamentals of basic analytical chemistry: an analyte is selected for quantification in a given matrix; experiments are performed to optimize the detection of the analyte by the chosen measurement platform, based on analyses of authentic chemical standards; if necessary, experiments are performed to optimize the extraction of the chosen analyte from a given matrix, again using authentic chemical standards and with determination of extraction efficiency; finally, an approach for appropriate quantitative data analysis is established, usually based on calibration curves constructed through analyses of authentic chemical standards in the matrix of interest and, ideally using stable isotope-labeled internal standards. The benefits of targeted analysis include the accurate, quantification of the analyte(s) of interest, based on the optimization of all aspects of the analysis method; low limits of quantification (e.g., sufficient to quantify pM levels of chemicals in blood [44]), again due to the optimization of the method parameters; and usually absolute confidence in the identity of the analyte. Caveats of targeted approaches include the narrow snapshot of chemistry measured; the level of effort required to fully optimize and validate the analytical pipeline [43,45], and the relatively low analysis throughput, depending on the specifics of the method.

Untargeted analysis

In contrast to targeted analysis, untargeted analysis does not focus on a specific analyte but instead seeks to comprehensively measure all analytes in a sample. Untargeted analysis of small molecules is increasingly popular in metabolomics and environmental studies [46–57]. For general metabolomics studies, NMR spectroscopy [58] and MS [59] have been the primary analytical techniques employed. NMR is
inherently quantitative and offers the ability to elucidate molecular structures, but suffers from low measurement sensitivity and throughput [60,61]. The utility of NMR in exposome studies is limited to measuring relatively high-abundance xenobiotics and to assessing biological response to exposure. On the other hand, MS analyses when coupled with chromatography are highly sensitive, but not absolutely quantitative unless stable-isotope-labeled internal standards, matrix-matched external standards or the standard addition method are used. However, the high dynamic range and low of MS approaches tend to outweigh the typical limitation to semiquantitative analyses (if desired, full quantitation is possible using any of the approaches mentioned above) and offer the potential to measure both xenobiotic chemicals and the biological responses to exposure. Chromatography (e.g., LC; GC) coupled with MS is thus the most widely used analytical platform for exposome studies [46–52]. A typical workflow for untargeted analysis using chromatography coupled with MS involves SPE or liquid–liquid extraction of the sample, drying of the extracts and reconstitution in a suitable buffer or solvent, followed by analysis. If GC-MS is used, then chemical derivatization of extracted molecules may be required to increase their volatility and enhance their separation. For GC-MS, electron ionization (EI) is typically used to impart the required charge to the analyte for detection by the mass spectrometer, although chemical ionization can also be used. Molecular fragmentation occurs simultaneously with EI, and commercial and open access reference libraries of spectra can be used to identify detected molecules [18,22,62], often in conjunction with other information such as retention time and/or retention indices [22]. For LC-MS, ESI is most commonly used to introduce analyte ions to the instrument, and data are typically collected by repeatedly scanning over a wide mass range (e.g., 100–1000 m/z) to detect as many molecules as possible. In most LC-MS experiments, intact molecular ions are fragmented using collision induced dissociation to produce MS/MS spectra, which can be used to identify the structures of detected molecules in conjunction with libraries of reference spectra [20,28,63] or tentatively identify candidates using in silico fragmentation approaches combined with candidate look-up [26,64–66]. The benefit of untargeted analysis is that it is not biased by a priori assumptions and so offers the best opportunity to discover novel markers of exposure or to characterize the response to exposure by measuring a broad swath of chemical space. Caveats, however include possible artifacts in the data due to the lack of optimization of sample preparation procedures, difficulty detecting very low-abundance analytes in the presence of high-abundance analytes (particularly when using LC-MS due to ESI suppression) and an incomplete representation of chemical space in spectral reference libraries, leading to, in some cases, limited confidence in the identification of detected species where reference standards are not available.

Molecular identification confidence

As described above, the identification confidence of molecules measured in targeted analyses is generally very high, since methods are optimized for the chemistry of interest and the use of authentic standards ensures that the correct molecule is identified each time. Therefore, the discussion in this section will focus on the confidence in identification of metabolites and chemicals detected in untargeted, MS-based analyses.

The use of untargeted acquisition techniques brings a whole set of identification challenges with it as, unlike targeted acquisition, no preliminary hypothesis as to the number and identity of analytes of interest has been generated. It is possible (and increasingly common) to perform target analysis on data acquired with untargeted methods, and this is the best starting point for any untargeted investigation. This requires the availability of in-house reference standards (measured with the same technique) and, for quantitative results, internal standards and appropriate calibration curves [54,67]. Confident identification requires the match of two orthogonal pieces of experimental data between standard and sample, such as matching exact mass of the molecular ion and fragments as well as retention time. Typically, targets are ‘screened’ in data from untargeted analysis by searching for the exact mass in peak lists or via extracted ion chromatograms, hence the term ‘target screening’ [54].

In cases where investigations wish to pursue a substance of interest for which the reference standard is not available in house, a so-called ‘suspect screening’ can be performed. The exact masses of the ions relating to the substance(s) of interest can be used to search the data from untargeted analyses either via peak lists or extracted ion chromatograms, as is performed for target screening. Unlike target screening, however, additional steps need to be undertaken to prove (or disprove) the identity of the suspect, since a standard is not available. This is discussed further below. The lists used for suspect screening can vary widely and depend on the study question. One could consider screening the Human Metabolome Database (HMDB) [68] or other metabolite-specific databases as suspect screening for metabolomics. In environmental studies, one could use a list of registered pharmaceuticals [69] or pesticides [70], surfactants [53] or even all substances registered under the Registration, Evaluation,
probable identification with all evidence indicating unknown compounds – molecular formula is. In addition, many of these molecules trans putative annotation based on physicochemical properties and spectral matching (MSI level 2, Eawag Level 2a).

• Tentative identification/putative compound class – tentative identification using predictive techniques, multiple structures are possible, or insufficient evidence to eliminate other structures; substance class only is clear (MSI level 3, Eawag Level 3).

• Unknown compounds – molecular formula is unequivocal (Eawag Level 4) or exact mass only (Eawag Level 5; both MSI Level 4). These can be traced in samples and correspond to ‘detected features’ in the analyses, but the identity remains uncertain.

To date, the evidence for coupled chromatographic-MS systems has been based on the mass spectral information (exact mass and fragments for MS/MS, EI-MS spectrum for GC-MS) plus retention time information (either as retention time or retention indices) as the orthogonal information. However, coupled chromatographic systems alone are not ideally suited to high-throughput studies due to the time required per analysis. Later in this article, the potential for IMS and CCS to provide orthogonal identification evidence together with (or in place of) the retention time in coupled chromatographic systems is discussed.

Introducing IMS as a new tool for the exposomics toolbox

Comprehensive characterization of exposure is extremely challenging since individuals are generally subject to thousands of structurally and physicochemically diverse chemical agents per day (e.g., pharmaceuticals, pesticides, personal care products, industrial chemicals), with various systemic responses to these. Measuring these chemicals in clinical samples is very difficult since they typically occur from picomolar to millimolar concentrations in complex matrices. In the following sections, we will focus the discussion on the benefits of DTIMS and how it can be used to address the challenges of exposomics.
Figure 2. Anthropogenic chemicals and endogenous metabolite concentrations in blood. A wide dynamic range of concentrations of chemicals and metabolites are observed in human blood, making the comprehensive measurement of these in an exposomics approach extremely challenging. Reproduced with permission from [44].

DTIMS increases the dynamic range of existing LC-MS methods

In DTIMS platforms, ions travel through a drift tube under the influence of a weak attractive electric field while colliding with a stationary buffer gas (Figure 3A). In this way, ions with small CCSs spend less time inside the drift tube than other equally charged ions with larger CCS values [83,84], as described in the following equation:

\[
CCS = \frac{3}{16Nk_B} \sqrt{\frac{e^2 z^2}{\mu k_B T}}
\]

where \( e \) is the elementary charge, \( z \) is the charge on the ion, \( N \) the gas density number (mol/m\(^3\)), \( K \) is the mobility of the ion, \( \mu \) is the reduced mass of the collision gas and ion, \( k_B \) is the Boltzmann constant and \( T \) is absolute temperature. While the \( m/z \) of an ion strongly influences the time it spends in the drift tube, the CCS value directly reflects the mobility of each ion and correlates with the shape of the molecule based on its ion-neutral interaction potential [85]. As the \( m/z \) ratios of ions can be measured following the DTIMS separation using a mass spectrometer, this allows CCSs to be directly determined for each ion via the fundamental zero-field (i.e., Mason Schamp equation [86]). In general, this orthogonality between the separation mechanisms of IMS and MS is key to the utilization of this combination of techniques, particularly for the separation of isobaric compounds.

The ability to resolve isomers that are difficult to distinguish using LC-MS alone is an inherent strength of DTIMS, particularly in small molecule analysis since many metabolites and other chemicals have the same molecular formula but play very different roles in biological systems. Indeed, DTIMS has been used to separate some important classes of isomers [87–91]. Stephen et al. used library CCS values to confirm the identities of the isomeric tramadol and desvenlafaxine in wastewater samples [92], which have the same molecular formula and hence the same
Figure 3. Ion mobility spectrometry and its incorporation into LC-MS workflows. (A) In ion mobility spectrometry (IMS), ions are separated based on their size and shape. In the example shown, the bile acids β-muricholic, urocholic and α-muricholic acids separate in order of fastest to slowest based on relative gas-phase ion shape. (B) The millisecond acquisitions of IMS are easily coupled between the minute and microsecond timescales of LC separations and TOF detection, since the time scale of each allows rapid sampling of the preceding separation. (C) Integration of IMS into LC-MS workflows provides an additional dimension of separation. Incorporating IMS provides greater coverage of the sample for the same LC separation length, or allows for decreasing the LC separation length while maintaining the same coverage. The example shows a human plasma total lipid extract analyzed by LC-IMS-MS. To indicate the increased separation capability obtained with the combination of LC and IMS, IMS-MS spectra were summed across three 10-s regions of the LC separation, creating three dimensional plots of m/z, drift time and intensity. Figure 3C adapted with permission from [125] © Elsevier (2008).
DTIMS separations are ultra-high throughput

While using chromatographic separations prior to the IMS stage will continue to play a role in exposome studies, DTIMS-MS alone lends itself extremely well to coping with the high-throughput demands of measuring large population cohorts. Interfacing DTIMS with high-resolution MS, such as TOF (99) permits simultaneous separation of sample constituents and acquisition of structural information and high-accuracy MS data, and when coupled with quadrupole-TOF MS, also allows for acquisition of tandem mass spectra in targeted, data-dependent or data-independent modes. IMS separations can also be coupled with trapping MS instruments such as Orbitrap and Fourier Transform Ion Cyclotron Resonance. However, in these cases the millisecond IMS separations would be faster than the second timescale sampling of the mass spectrometers, and so MS data acquisition manipulations would be required. Thus, TOF-based instruments that sample on the microsecond timescale are more practical to retain the IMS measurement throughput. In DTIMS-TOF MS analysis, a single separation typically occurs in 10–100 ms and the TOF MS pulser samples each peak approximately every 100 μs, so DTIMS separations can be reconstructed with sufficient points-per-peak to allow for reliable discrimination of features in 3D space (drift time, m/z, abundance). DTIMS transients are normally summed for at least 0.5 s to acquire enough signal for reproducible IMS peaks and repeatable determination of peak apices, which is essential for accurate determination of CCS.

DTIMS separations do not require derivatization and can be combined with different ionization sources for analysis of distinct compound types such as polar metabolites (ESI in positive or negative ionization modes) and as well as compounds for which other ionization modes are required (e.g., APCI in positive or negative modes for analysis of polyaromatic hydrocarbons), allowing for multiple analyses to obtain high coverage of the exposome. Without chromatographic or other preseparation prior to ESI, ionization suppression due to the presence of highly abundant chemicals has previously limited the utility of DTIMS-MS for analysis of complex mixtures. However, a front-end automated SPE system was recently coupled directly to DTIMS-MS and different cartridge chemistries were utilized to provide a means to reduce ion suppression in analysis of complex mixtures. This automated SPE-IMS-MS system provided a 10-s sample-to-sample duty cycle and a theoretical maximum throughput of >8000 injections per day (100,101). This is 2–3 orders of magnitude higher throughput than conventional GC-MS or LC-MS methods and makes the study of large patient cohorts practical. When necessary, DTIMS-MS can also be coupled with GC or LC to address prohibitively complex samples and reduce front-end analysis times (98).

Incorporating CCS into molecular identification workflows

The implementation of IMS in exposomics studies will lead to increasingly more frequent observations of previously undetected chemicals and metabolites. LC-IMS-MS will provide increased overall measurement dynamic range, resulting in detections of lower abundance molecules, and the throughput of IMS-MS alone will provide the opportunity to analyze many thousands of longitudinal samples over lifetimes of exposure, capturing evidence of transitory accumulations of chemicals or metabolites. The volume of data corresponding to these new chemical observations will almost certainly outpace the development of reference data to enable their confident identification. Thus, the question remains as to how these molecules can then be identified in exposomics studies.

We here explore the possibility of using computationally predicted CCS to assist in the identifications of newly observed anthropogenic chemicals and metabolites, analogous to previous and ongoing efforts to predict retention times or tandem mass spectra during analysis of small molecules (29,30,102,103). In this
approach, CCSs are computationally predicted from the chemical structures of the molecules themselves, as captured using molecular modeling and quantum chemical calculations. In addition, unlike chromatographic retention times, the CCS parameter recorded under low drift field conditions is influenced by far fewer physicochemical parameters, meaning that excellent correspondence between measured values and library entries can be expected across different instruments.

To broadly predict CCS values, an automated computational pipeline is required as depicted in Figure 4. First, International Chemical Identifiers (InChIs; [104]) are converted into 2D structures, and then analyzed using the Marvin pK plugin (Marvin 15.9.14, 2015, ChemAxon) to predict protonation/deprotonation states and adduct sites [105]. Initial geometry optimizations are performed using the Merck molecular force field [106] implemented in Avogadro (v.1.1.1) [107] and final geometry optimizations are performed using a density functional theory implemented in NWChem [108] at the B3LYP/6–31g* level [109–111]. Finally, the theoretical CCS values are calculated based on the geometry-optimized structures using the N$_2$-optimized trajectory method implemented in the MOBCAL software [112–114]. In this way, the generation of accurate predicted CCS values can facilitate the broad identification of detected molecules in combination with accurate mass and MS/MS spectra, when available, and possibly with just accurate mass. The power of this approach is that metabolites can be annotated rapidly since only a chemical structure (e.g., in InChI format) obtained from either one of the many databases (e.g., HMDB, CHEBI, ChemSpider, PubChem) or drawn by hand in chemical structure software is required for predicting theoretical CCS values to be matched with experimental numbers, together with other metrics, such as accurate mass and/or MS/MS spectra.

By utilizing a theoretical pipeline such as defined above, the feasibility of broadly predicting CCS for small molecules was determined by calculating theoretical CCSs for 5068 metabolites from the HMDB, in their protonated, deprotonated and sodiated forms based on pK$_a$ analysis, yielding predicted CCSs for a total of 11,046 unique ionized structures. To improve throughput for predicting so many structures, CCS was calculated using the IMPACT software [105,115], parameterized for helium as the drift gas. On average, it required approximately 0.5 s to calculate the CCS of a molecule on a Windows desktop (with 16 GB RAM, Intel® Xeon® 1.6 GHz CPU); while the InChI to 2D/3D structure conversions, the pK$_a$ calculations and the construction of the ionized structures required about 3.3, 2.9 and 26.3 s per molecule, respectively, for 1465 molecules. The DFT calculations required approximately 1 h per molecule to run on a single computer node of a 3.4 petaflop Linux cluster that has 1440 compute nodes and 16 cores per node. From these results, the chemical search space for possible molecules with a given accurate monoisotopic mass could be reduced by a minimum of 79% using CCS with a standard deviation (SD) of 2% (compared with a 0% reduction if using only m/z to find a match) within the HMDB library. For example, 76 different metabolites had the molecular formula C$_{35}$H$_{30}$O$_6$ (Figure 5B) plotted versus their corresponding accurate masses. As the searched databases grow in size (i.e., by including ChemSpider or others), and as novel undocumented molecules are considered, this problem will only increase in magnitude (Figure 5C).

In terms of the required agreement between predicted and measured CCS values, the combination of instrumental measurement precision and accuracy of CCS prediction is critical to the success of the described workflow. As discussed above, extremely good instrumental precision (<0.5%) is now readily possible on individual instruments, and interlaboratory exercises indicate that DTIMS-MS instrumentation can provide reproducible consensus CCS values with an RSD of less than 1% [93,94] for a wide range of biologically relevant molecules, which is a marked improvement over raw chromatographic retention time matching across different laboratories. This latter point is particularly critical to the success of the proposed approach, as a sound understanding of the uncertainty of CCS measurements must be ascertained in order to improve the identification potential. In an initial evaluation, experimental and theoretical CCS values of selected ionized structures (protonated, deprotonated and sodiated) of 11 metabolites were compared, resulting in an average error of approximately 2% (Figure 5D) and indicating the potential of predicted CCS as a metric for supporting chemical identification in exposomic analysis.

The approach of using IMS to compare experimental predicted CCS values is not new and dates back to the early 1990s [116]. More recently, the community has been moving in the direction of routine implementation of IMS and CCS in their methods for chemical analysis on a larger scale. Paglia et al. integrated TWIMS into an LC-MS method and established a retention time, CCS and accurate mass database of 125 common metabolites; CCSs were...
Figure 4. Workflow for the identification of small molecules based on experimental and theoretical collision cross sections and accurate mass. Untargeted SPE-ion mobility spectrometry-MS measurements (green box) provide experimental collision cross section (CCS) and accurate mass m/z values for sample molecules, as well as their isotopic signatures. Independently, theoretical CCS calculations (blue box) augment the contents of a CCS and accurate mass reference library. Separately, molecules that cannot be identified by matching to the CCS and accurate mass reference library or to MS/MS libraries are analyzed based on their masses to generate candidate molecular formulas, which are then converted into 3D structures for CCS prediction (purple box).

Conclusion
In summary, comprehensive characterization of exposure is challenging due to the thousands of structurally and physicochemically diverse chemicals one encounters per day, and which may be present at any given time in some amount in body compartments. Complicating this, when employing metabolomics approaches, is the presence of the endogenous metabolome and its localized or systemic response to exposure. Thus, the potential chemical space that can be measured is vast, and untargeted data acquisition approaches are best suited to comprehensively measure the exposome.

Advancing measurement technologies, including those based on or incorporating DTIMS, are enabling increasingly higher coverage of sample molecular compositions. The volume of data corresponding to these new chemical observations will require new informatics methodologies to confidently assign as many identifications as possible. To address this challenge, we propose that accurate, computationally predicted CCS values can facilitate the broad identification of detected molecules in combination with accurate mass and MS/MS spectra, when available, and possibly with just accurate mass...
Figure 5. Performance of the molecular modeling and quantum chemical calculation pipeline. (A) A plot of calculated collision cross section (CCS) versus m/z for the protonated forms of the molecules shown in (B), illustrating the degree of orthogonality of the two metrics; (B) metabolite structures from the Human Metabolome Database corresponding to the formula C₆H₁₃NO₂; (C) a plot of CCS versus m/z for 200 metabolites, out of the >11 k solved structures obtained from the Human Metabolome Database. Different colors correspond to CCSs for different metabolites with identical molecular formulae. The data for C₆H₁₃NO₂ from (B) are outlined in the dashed box, and two anthropogenic chemicals with the formula C₁₅H₂₄O, 9-bisabolatrien-11-ol and α-3-copaen-8-ol are shown to have very different CCS; and (D) the theoretical CCS values obtained for selected +H, -H and +Na forms of 11 small molecules show good agreement with experimental CCS values derived from an SPE-ion mobility spectrometry (IMS)-MS platform, with average error of approximately 2%. All CCS values were predicted using He as the buffer gas.
The pentasaccharides cellopentaose (blue, collision cross section [CCS] 250 Å²) and mannopentaose (green, CCS 256 Å²) were measured as their sodiated forms using (A) an Agilent 6560 ion mobility spectrometry-quadrupole TOF and (B) a 31-m Structures for Lossless Ion Manipulations (SLIM)-TOF arrangement. The additional features (i.e., peak shoulders) can be putatively assigned to either different sodium cation binding locations or separation of α and β anomers. Estimating that roughly six peaks can fit in the SLIM-TOF separation space, the resolution of the measurement is at least less than half the difference between two of the sugars. The cellopentaose/maltopentaose and maltopentaose/mannopentaose pairs each differ by approximately 1% in CCS, and the cellopentaose/ mannopentaose pair differs by approximately 2% in CCS. The additional features resolved in the SLIM separation show a resolution of at least less than 0.5% difference in CCS.

**Future perspective**

Unsurprisingly, some analytical and data processing challenges remain for full-scale implementation of IMS in standard exposomics workflows, including dealing with the annotation of adducts and dimers separated by DTIMS in untargeted analysis, which need to be reconciled with the principal monoisotopic ion for correct annotation. Similarly, the nature of the DTIMS separation necessitates an ion gating or trap-and-release mechanism that entails a loss of duty cycle and (in the latter case) the possibility of losses of ions or interactions between different ions in the trapping environment. As such, careful analytical method development must be pursued in practice to validate the robustness and minimize the false annotation potential for exposome studies.

**The next generation of IMS instrumentation: structures for lossless ion manipulations**

A current limitation of DTIMS technology is its limited resolution for a given fixed length drift tube, and since resolution is proportional to the length of the device for a given weak electric field [118], making the devices longer than several meters is not practical. Recently, ultra-high resolution IMS devices were constructed with Structures for Lossless Ion Manipulations (SLIM), utilizing traveling waves. In their present form, SLIM are ion conduits formed by confining electric fields using a pair of parallel printed circuit boards. Radiofrequency potentials prevent ion losses to the circuit boards and direct current potentials prevent lateral ion losses. SLIM has separated isomers from multiple molecular classes, including peptides, natural products, lipids and oligosaccharides that have not been fully resolved with currently available IMS systems. In initial evaluations, SLIM has separated compounds with differences in CCS < 0.5% (Figure 6). Recently, a 13-m length, serpentine path SLIM device was shown to have about fivefold higher resolution separations than present commercially available DTIMS or TWIMS platforms [119]. SLIM research at Pacific Northwest National Laboratory currently focuses on multipass separations where it should be possible to obtain resolutions at least fivefold higher than the current SLIM, with a reasonable number of passes (<50), and a future vision includes km-long path length devices. Though the separation resolution is increased, the sensitivity of these SLIM measurements is maintained. In addition, when coupled with TOF MS, SLIM separations are still carried out on a rapid (~1s) time scale, providing the opportunity for ultra-high throughput studies.

Traveling wave separations are the result of the abilities of ions to keep up with a direct current wave pushing them throughout the drift cell while undergoing collisions with the drift gas. Ions with larger CCS will be passed over by traveling waves more frequently than ions with smaller CCS. Due to this mechanism, the resulting separation is not linearly correlated to CCS, and the Mason Schamp equation does not hold. Thus, one caveat of TWIMS-based SLIM devices is that they do not allow for direct measurement of CCS. TWIMS does allow for calibration of CCS against standards that have been measured by DTIMS; however, calibrated ions should be from the same biomolecule class, should overlap in m/z space and should have the same charge states as the target ions, otherwise large errors in CCS (e.g., up to >5%) can result [82,120–123]. However, since accurate CCSs are measured from the DTIMS workflow, the identified compounds can be used as internal calibrants for SLIM measurements. Importantly, SLIM should allow CCS measurements to be made for many more mixture components and with greater precision due to the far greater resolution achieved [124], and potentially allowing CCS with much greater accuracy as better standards are developed. In this manner, small molecules that are not distinguishable by present DTIMS should be much better separated in SLIM-based IMS-MS and assigned with more accurate CCS values in a combined DTIMS/SLIM-MS pipeline, allowing more effective identification. Although the initial cost of devices and data management will be expensive, the time cost for measurement would be much lower than the traditional targeted/suspect screening using LC-MS/MS or GC-MS/MS analysis.
Executive summary

Background
- Human disease is a combination of individual genetic factors and nongenetic environmental factors.
- The ‘exposome’ complements the genome and is the sum of all nongenetic exposures over a lifetime to chemical, social and biological agents.
- Omics technologies can assist in characterizing the exposome by directly measuring chemical exposures and by inferring exposure based on biological signatures obtained by one or more complementary approaches.
- As new analytical methodologies for measuring small molecules advance, there will be a need for computational approaches to rapidly and comprehensively identify chemicals and metabolites.

State-of-the-art in measuring the exposome
- The technical approaches used to measure the exposome largely fall under one of two categories: targeted or untargeted.
- Targeted analyses focus on a limited number of analytes, have high quantitative accuracy and low limits of quantification, but only measures a narrow snapshot of the sample molecular composition.
- Untargeted analyses do not focus on a specific analyte but instead seek to comprehensively measure all analytes in a sample and offer the best opportunity to discover novel markers of exposure. Caveats include possible artifacts in the data due to the lack of optimization of sample preparation procedures, difficulty detecting very low-abundance analytes in the presence of high-abundance analytes and an incomplete representation of chemical space in spectral reference libraries.
- NMR spectroscopy and LC-MS coupled with MS are typically employed in untargeted analysis of chemicals and metabolites.
- Communicating the confidence in chemical identification can be a challenge and several systems exist for small molecules, including the Metabolomics Standards Initiative, an LC-high resolution MS/MS specific set from Eawag and many more. The essences of these are:
  - Confirmed identification with two orthogonal matching properties to an authentic reference standard measured in-house (MSI Level 1, Eawag Level 1).
  - Probable identification with all evidence indicating only one structure is possible, but authentic standard is not available for confirmation (Eawag Level 2a/b).
  - Probative annotation based on physicochemical properties and spectral matching (MSI Level 2, Eawag Level 2a).
  - Tentative identification/Probative compound class – tentative identification using predictive techniques, multiple structures are possible or insufficient evidence to eliminate other structures; substance class only is clear (MSI Level 3, Eawag Level 3).
  - Unknown compounds – molecular formula is unequivocal (Eawag Level 4) or exact mass only (Eawag Level 5; both MSI Level 4). These can be traced in samples and correspond to ‘detected features’ in the analyses, but the identity remains unclear.

Introducing ion mobility spectrometry as a new tool for the exposomics toolbox
- Drift tube ion mobility spectrometry (DTIMS) shows great promise in small molecule measurements because it is able to directly determine molecular structural information.
- The ability to resolve isomers that are difficult to distinguish using LC-MS alone is an inherent strength of DTIMS, particularly in small molecule analysis.
- Because DTIMS instruments depend only on drift cell pressure, temperature and length, molecular collisional cross section (CCS) measurements are extremely reproducible. Furthermore, measurements from different instruments in different laboratories have also been compared and their values normally agree within <2% error, with recent DTIMS instruments yielding values with reproducibility precision of <1%.
- The high reproducibility and speed of DTIMS allows it to be easily nested between LC and MS to provide additional separation power and dynamic range of detection in measuring the exposome.
- When coupled with TOF time-of-flight MS, DTIMS-MS analysis can be ultra-high throughput, with a single ion mobility spectrometry (IMS) separation typically occurring in 10–100 ms.
- A novel, automated SPE sample introduction system coupled with DTIMS-MS provides a 10-s sample-to-sample duty cycle and a theoretical maximum throughput of >8000 injections per day.
- Accurate, computationally predicted CCS values can facilitate the broad identification of detected molecules in combination with accurate mass and MS/MS spectra, when available, and possibly with just accurate mass.

Future perspective
- Analytical and data processing challenges remain for full-scale implementation of IMS in standard exposomics workflows.
- Ultra-high resolution IMS devices constructed with Structures for Lossless Ion Manipulations are pushing the separation resolution possibilities of IMS.
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