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1	Analytical Chemistry Solutions to Hazard Evaluation of
2	Petroleum Refining Products
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14	
15	Abbreviations:
16	API, American Petroleum Institute; ASTM, American Society for Testing and Methods; CAS
17	Chemical Abstract Service; CCS, collision cross section; DBE, double bond equivalents; ECHA,
18	European Chemical Agency; EINECS, European Inventory of Existing Commercial Chemical
19	Substances; US EPA, United States Environmental Protection Agency; FID, Flame ionization
20	detection; FT-ICR MS, Fourier transform ion cyclotron resonance mass spectrometry; FWHM, full
21	width at half maximum; GC, gas chromatography; GC×GC, two-dimensional gas chromatography;
22	GC-MS, gas chromatography coupled to mass spectrometry; HPLC, high-performance liquid
23	chromatography; IMMS, ion mobility mass spectrometry; IR, infrared; KMD, Kendrick mass
24	defect; LC, liquid chromatography; MS, mass spectrometry; NMR, nuclear magnetic resonance;
25	REACH, Registration, Evaluation, Authorisation and Restriction of Chemicals; TIMS, trapped ion
26	mobility spectrometry; TOF MS, time-of-flight mass spectrometry; UV, ultraviolet; UVCBs,
27	substances of unknown, variable composition, complex reaction products, or biological materials.

28 Abstract

Products of petroleum refining are substances that are both complex and variable. These 29 substances are produced and distributed in high volumes; therefore, they are heavily scrutinized in 30 terms of their potential hazards and risks. Because of inherent compositional complexity and 31 variability, unique challenges exist in terms of their registration and evaluation. Continued 32 dialogue between the industry and the decision-makers has revolved around the most appropriate 33 approach to fill data gaps and ensure safe use of these substances. One of the challenging topics 34 has been the extent of chemical compositional characterization of products of petroleum refining 35 36 that may be necessary for substance identification and hazard evaluation. There are several novel analytical methods that can be used for comprehensive characterization of petroleum substances 37 and identification of most abundant constituents. However, translation of the advances in 38 analytical chemistry to regulatory decision-making has not been as evident. Therefore, this 39 review's goal is to bridge the divide between the science of chemical characterization of petroleum 40 41 and the needs and expectations of the decision-makers. Collectively, mutual appreciation of the regulatory guidance and the realities of what information these new methods can deliver should 42 facilitate the path forward in ensuring safety of the products of petroleum refining. 43

44 Introduction

Crude oils are naturally occurring and highly complex substances which vary considerably 45 in molecular composition according to their origins; they comprise a myriad of constituents, 46 primarily hydrocarbons, but also other organic and inorganic molecules (Smith et al., 1959). Close 47 48 to one hundred billion barrels of crude oils are annually extracted, distributed, and processed into a wide variety of refined petroleum products (Kaiser, 2017; Salvito et al., 2020). The chemical 49 composition of petroleum refining products therefore depends on both the type (*i.e.*, origins) of 50 crude oil from which it was derived, and the refining process (*i.e.*, fractional distillation and/or 51 52 cracking followed by additional processing through solvent extraction, hydro-desulfurization, or 53 hydrogenation) used to meet performance characteristics of the end-products (McKee et al., 2015). Products of petroleum refining are high production volume substances and thus are heavily 54 scrutinized in terms of their potential human and environmental health hazards and risks. Because 55 56 of inherent compositional complexity and variability, petroleum substances are prototypical representatives of a diverse class known as substances of unknown, variable composition, complex 57 reaction products, or biological materials (UVCBs); these substances present unique challenges to 58 59 regulatory agencies, especially in terms of characterization of their chemical composition (Clark et al., 2013; ECHA, 2017c; Lai et al., 2022). It is worth noting that the different fields within the 60 61 academic, industrial, and regulatory science communities can use differing terminology. Analytical researchers often refer to petroleum refining products as "hydrocarbon mixtures" which are, in turn, 62 part of the broader "complex mixtures" family of samples. The term "mixture" is avoided by the 63 industry and decision-makers as they reason that most substances in commerce that are made from 64 65 oil are products of refining, rather than mixing, and thus to differentiate from the mixtures found in the environment, such substances are called petroleum UVCBs. 66

Studies of molecular composition of crude oils and petroleum refining products have a long 67 history spanning over 80 years (Figure 1). The analytical characterization of petroleum substances 68 69 historically tracked the physico-chemical properties that pertained to the functionality of the product, such as flash points and vapor pressure. With the advent of spectroscopy and mass 70 spectrometry techniques, there came the possibility to gradually gain more detailed understanding 71 72 of composition; however, the granularity of information on the constituents in registered petroleum products is still lacking. Recent improvements have been made in the resolution of mass 73 spectrometers, ionization methods in order to access a wider range of components, separation 74

methods to offer structural insights, and developments in data visualization through standardized 75 diagrams (Palacio Lozano et al., 2020). As a result of these advances, new opportunities emerged 76 to provide comprehensive characterization of these complex substances and satisfy regulatory needs 77 on the composition, quantity of potentially hazardous constituents, and the extent of variability 78 79 among manufacturing batches of these products. The range and types of mass spectrometry techniques that can be used for the analysis of petroleum-related samples is quite extensive; 80 collectively, the methods for study of petroleum are now being referred to as "petroleomics," a sub-81 field of analytical chemistry aiming to identify the totality of constituents of crude oil and petroleum 82 83 refining products using high resolution mass spectrometry methods (Hsu et al., 2011; Marshall and 84 Rodgers, 2004; Palacio Lozano et al., 2020).

Despite major advances in the ever-improving analytical resolution of individual molecules 85 and their classes in oils and complex petroleum UVCBs (Wise et al., 2022), there has been relatively 86 little use of the data from these new methods and instruments in regulatory submissions, with the 87 88 exception of GC×GC-FID-derived data (Redman et al., 2014; Ventura et al., 2011), or even their mention in the reviews or original research publications (Figure 2A). The naming conventions and 89 90 approaches to identification of complex petroleum UVCBs remain rather imprecise (Rasmussen et al., 1999); only general compositional characteristics are used to define broad manufacturing 91 92 categories (Salvito et al., 2020). While such information is generally sufficient for naming and identification of petroleum UVCBs (ECHA, 2017a), it is often not sufficient for evaluation of human 93 94 health and environmental hazards, a prerequisite to registration and authorization for their use 95 (Salvito et al., 2020).

96 The regulatory science and analytical chemistry fields run in parallel and both are highly specialized, requiring significant expertise. Consequently, intricacies of the legislative mandates 97 governing regulatory decision-making are often unfamiliar to the researchers who develop and 98 refine advanced methods for petroleomics. Similarly, decision-makers may not be aware of the latest 99 100 opportunities that analytical chemistry has to offer. Collectively, there is a considerable gap in the translation of knowledge from petroleomics-focused analytical laboratories to applied decision-101 making. This review aims to first summarize the regulatory guidance for characterization of the 102 chemical composition of petroleum UVCBs and then demonstrate how existing petroleomics 103 techniques could be applied to address these needs. Recent additions to the European Union (EU) 104 Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) regulatory guidance 105

106 for demonstrating the composition of complex UVCB (ECHA, 2022), together with recent 107 advancements in petroleomics applications that are poised to address these needs (Palacio Lozano 108 et al., 2020), create a unique opportunity to bridge the divide. Here, we highlight the opportunities 109 that are already within reach for using modern analytical and data analysis/visualization techniques 110 for impactful decisions on petroleum UVCBs. We reason that it is imperative for decision-makers 111 to be aware of the possibilities and limitations of current analytical approaches so that regulations 112 can be sufficiently strict, yet realistic in terms of their attainment using best available science.

113

114 What Information do the Decision-Makers Seek on Petroleum UVCBs?

115 Most impactful guidance to the industry on petroleum UVCBs has been produced in the United States and the European Union. These include the United States Environmental Protection 116 Agency (US EPA) High Production Volume (HPV) Challenge Program, a voluntary industry-117 government information sharing effort that was launched in 1998 (Petroleum HPV Testing Group, 118 119 2017), and the REACH legislation and associated guidance documents in the European Union (ECHA, 2017c) which is now being adopted in other countries around the globe. Because REACH 120 121 is the most recent and stringent legislative regime, it has become a *de facto* global driver for the regulatory scrutiny of both new and existing chemicals, including petroleum UVCBs. Therefore, 122 123 most decision-making contexts discussed herein pertain to EU REACH regulation and its implementation by the European Chemical Agency (ECHA) through guidance documents. 124

125 Simply put, REACH-based guidance for UVCBs states that data provided by the registrants shall enable (i) identification of the substances that are submitted for registration, and (ii) evaluation 126 127 of the potential hazards to human health and the environment (ECHA, 2017c). In both instances, 128 information on chemical composition of a substance is required; however, in a slightly different context (Table 1). For the former, data requirements are typically less stringent because complex 129 substances may be registered based on the manufacturing process, intended use, and/or physical-130 131 chemical properties. For the latter, the molecular identification (both elemental composition and structure) data requirements are far greater because the individual constituents that may be present 132 in a "representative" sample, their amounts, and presence of known or suspected hazardous 133 substances must be reasonable ascertained to enable grouping and read-across among petroleum 134 UVCB substances. Based on these two initial components of REACH, an EU regulation that came 135 into force in June of 2007 (European Council, 2007), decisions on authorization of use or 136

restriction(s) are made. An additional challenge for petroleum-derived and other UVCBs is that their 137 composition is inherently variable from batch to batch of the nominally the "same" product; this is 138 also true for product-to-product variability within broader categories (FuelsEurope, 2015). This 139 variability may be due to the source crude oil used in manufacturing; petroleomics-related research 140 141 has revealed significant differences in compositions of petroleum according to its origins, slight variations in manufacturing processes among refineries, and naturally occurring degradation and 142 weathering processes. Because safety testing is typically conducted with a "representative" sample 143 of a UVCB product or category, the confidence with which these data can be extrapolated to other 144 samples of the same product or to other products that are "similar," especially when sample 145 composition data may be commercially sensitive, depends on how much and what type of data is 146 available to ascertain such similarity. 147

For the purpose of substance identification, the regulatory frameworks in the United States 148 149 and the European Union have historically (Figure 1) named, grouped and categorized petroleum 150 UVCBs based on the manufacturing processes used in oil refining, as well as physical-chemical properties and other broad chemical fingerprinting data (Dimitrov et al., 2015; ECHA, 2008; 151 Rasmussen et al., 1999). Manufacturing process-centric naming conventions for these complex 152 substances were originally developed by the American Petroleum Institute (API) and the US EPA 153 154 for the purpose of creating an inventory of petroleum products under the Toxic Substances Control Act inventory (API, 1983; EPA, 1995; U.S. EPA, 1978). Both Chemical Abstract Service (CAS) 155 156 and European Inventory of Existing Commercial Chemical Substances (EINECS) identifications have been assigned to a large number of petroleum UVCBs, even though the descriptions of each 157 158 of these substances under either one of these "unique identifiers" are rather imprecise and far from being unique (Rasmussen et al., 1999). The broad substance categories were somewhat refined 159 under the US EPA's HPV Challenge Program through addition of more detailed information on 160 physical-chemical properties, as well as some human and environmental hazards data (Petroleum 161 162 HPV Testing Group, 2017).

Far greater information requirements, with respect to both chemical composition and potential hazards, were imposed by REACH (European Council, 2007). Petroleum UVCBs are high production volume substances that were subject to the earliest deadline for registration and the most stringent requirements for hazard evaluation. From 2007 to 2010, about 8,000 registrations were submitted in the EU for petroleum substances that were produced or imported at >100 tons/year

(CONCAWE, 2022). Subsequently, the number of registrations of petroleum products was reduced 168 to 191 substances through consolidation of redundant submissions and further grouping of 169 substances deemed to be "similar" based on a variety of considerations. To put the scale of the 170 challenge in context, tens of thousands or even hundreds of thousands of unique molecular formulae 171 can be observed in a single fraction of a petroleum sample using ultrahigh resolution mass 172 spectrometry, indicating that potentially millions of different structures are present in individual 173 crude oils when allowing for isomers (Palacio Lozano et al., 2019b; Palacio Lozano et al., 2020). 174 While the registration submissions for petroleum UVCBs were completed more than 10 years ago 175 176 and are regularly updated (CONCAWE, 2021), discussions between trade associations and regulatory agencies are ongoing to determine the most sensible ways to improve the dossier quality 177 and ensure the information is compliant with the REACH regulation, primarily by generating more 178 testing information and reinforcing read-across and category approaches. Still, industry's attempts 179 to waive animal testing requirements through read-across have been rejected by ECHA because of 180 181 considerable data gaps in hazard assessment and compositional characterization (ECHA, 2020a; ECHA, 2020b; ECHA, 2021). 182

183 For registration under REACH (Annex VI, Section 2), the data should be sufficient to enable substance identification (Table 1). For petroleum substances, ECHA guidance is that the following 184 185 data should be provided: (1) accepted nomenclature; (2) appropriate identifiers such as source/feedstock, refining history, boiling and carbon number ranges, physio-chemical 186 187 characteristics, chromatographic or spectral information, flash point, and viscosity; and (3) compositional information including identification and concentration of the individual constituents 188 189 present at >10% and that are known to be hazardous, persistent and/or bioaccumulative, identification of any additives, and generic description of unknown constituents (CONCAWE, 190 2012; ECHA, 2017a). Additionally, Articles 7(2) and 33 of REACH have defined a concentration 191 threshold of 0.1% w/w for constituents classified as "substance of very high concern" (ECHA, 192 193 2017b). Based on these guidance documents, the registrants (companies or trade associations) have traditionally relied on a wide range of analytical techniques (Figure 2A) to furnish the information 194 on petroleum substance identification (Clark et al., 2013; CONCAWE, 2012; CONCAWE, 2014; 195 196 CONCAWE, 2020).

197 For the <u>evaluation</u> step, REACH specifies (Annex XI, Section 1.5) that substances may be 198 grouped based on "*structural similarity between substances which results in a likelihood that the*

substances have similar physicochemical, toxicological and ecotoxicological properties so that the 199 substances may be considered as a group or category." Next, prediction of possible hazards of data-200 poor substances are made through the application of "read-across" from an analogous substance that 201 has been tested in a requisite assay and is deemed "similar." For this step, REACH regulation states 202 203 that "it is required that the relevant properties of a substance within the group may be predicted from data for reference substance(s) within the group (read-across approach)." The registrant shall 204 establish a read-across hypothesis which explains what structural similarities or differences exist 205 between the source and target substance(s) to which read-across is applied and why a prediction for 206 207 a toxicological or ecotoxicological property can be made with confidence.

208 Recently, REACH regulation (Annex XI, Section 1.5) has been amended to state that for the application in grouping, "structural similarity for UVCB substances shall be established on the basis 209 of similarities in the structures of the constituents, together with the concentration of these 210 constituents and variability in the concentration of these constituents" (European Commission, 211 212 2021). Once such information becomes available, and in cases where structural differences are present between the source and target substances, the read-across hypothesis should explain why 213 214 the differences in the chemical structures within a group will not influence the toxicological or ecotoxicological properties or may do so in a regular pattern. 215

216 Even though the regulatory language above may seem rather straightforward, in practice the bar on establishing "structural similarity" is very elusive in the case of petroleum UVCBs. It is 217 218 widely acknowledged that the chemical complexity of petroleum substances far exceeds the capabilities of any one method, even the highest resolution mass spectrometers; therefore, a 219 220 combination of techniques and approaches is often employed. However, despite the considerable scientific advances achieved in the past decade (Figure 1) in both molecular separation 221 222 (encompassing chromatography, spectrometry and ionization) and detection (various modalities of mass spectrometry, flame ionization detection, and spectroscopy), many challenges with precision 223 224 and confidence in comprehensive molecular characterization of petroleum samples persist.

For example, a recent study using Fourier transform ion cyclotron resonance mass spectrometry (FT-ICR MS) revealed nearly a quarter of a million molecular formulae in a fraction from one petroleum sample (Palacio Lozano et al., 2019b). This study is an example of the reality that millions of structures (as opposed to formulae, as structure also determines toxicity) are present in petroleum UVCBs, an intractable challenge both for the chemical analysts to resolve and identify,

as well as for the decision-makers to evaluate. Indeed, even the REACH regulation itself (Annex 230 XI, Section 1.5) acknowledges that "if it can be demonstrated that the identification of all individual 231 constituents is not technically possible or impractical, the structural similarity may be demonstrated 232 by other means, to enable a quantitative and qualitative comparison of the actual composition 233 234 between substances" (European Commission, 2021). Clarification regarding the definition of "other means" was recently released May 2022 (ECHA, 2022), where specific justification for lack of 235 knowledge about constituents, lack of published methods to identify constituents, and explanation 236 about technical hindrance to resolution and identification of constituents comprising >20% of the 237 238 substance is necessary to apply such means for characterization. Where "other" means are justified, 239 quantitative comparison of constituents in common between source and target substances should be included, as well as a qualitative comparison of structures that vary between the substances. 240 Fingerprinting, for example, can be used if the following are addressed: information on >95% of all 241 constituents, information on the constituents of high concern, and high analytical resolution to 242 243 enable accurate alignment, quantitation, variability, and structural data (beyond molecular formulas) of constituents between substances (ECHA, 2022). Therefore, the subsequent sections of this review 244 245 are framed around three overarching critical needs/questions that REACH regulation challenges the registrants of petroleum UVCBs to address to gain regulatory acceptance of the grouping and read-246 247 across hypotheses that are addressable by means of standardized analytical chemistry methods:

• Critical Need 1: Providing detailed information on the *structure* of the constituents;

• Critical Need 2: Providing information on the *concentration* of the individual constituents; and

• Critical Need 3: Demonstrating *compositional similarity* of complex petroleum UVCBs through

other means when the identification of all individual constituents is not technically possible orimpractical.

253

254 Conventional Methods for Characterization of Petroleum Substance Identity and

255 Composition

There are many methods for characterization of physical-chemical properties and chemical composition of oil, that have also been applied to petroleum UVCBs (**Figure 2A**). Multiple techniques are used due to the broad range of substances with widely different composition, volatility, and polarity (CONCAWE, 2014; Stout and Wang, 2007; Wang et al., 2011). Most publications concerning characterization of petroleum UVCBs have explored approaches to define

elemental composition, physical properties, and gross structural information using nuclear magnetic 261 resonance (NMR) or infrared spectra of these substances. The elemental analyses for 262 characterization of petroleum UVCBs evaluate the concentrations of major elements ranging from 263 carbon and heteroatoms to metals (CONCAWE, 2019). There are specific methods for assessing 264 265 physical properties, which typically relate to the quality of the product and are thus more regularly measured during manufacturing and more available. For example, measurements of specific gravity 266 using American Society for Testing and Methods (ASTM) methods such as ASTM D287 Standard 267 Method for API Gravity of Crude Petroleum and Petroleum Products are in wide use (Giles, 2016). 268 269 Other physical-chemical information, such as boiling and carbon number ranges, is typically deduced through physical (e.g., ASTM methods D86, D1160 and D2892), or simulated 270 (CONCAWE, 2019; CONCAWE, 2020) distillation methods. 271

Spectroscopic techniques have been widely employed to obtain broad compositional 272 273 information for regulatory characterization and identification of UVCB substances but their utility 274 for the analysis of petroleum UVCBs has been questioned (CONCAWE, 2020). NMR methods (IP392, ASTM D5292) measure the percent of carbon or hydrogen atoms in an aromatic ring 275 (CONCAWE, 2020). Infra-red spectroscopy measures the presence of functional groups to define 276 the degree of saturation in the constituents (CONCAWE, 2012). Ultra-violet spectroscopic analysis 277 278 quantifies compounds by detecting unsaturated bonds such as those in olefins and aromatics, as well as ketonic and heteroatom groups, but is limited in resolution for other constituents. 279

280 More detailed compositional information, which gives greater insight into the chemical classes and carbon ranges of the substance, is obtained using chromatographic techniques that 281 282 enable separation of constituent groups in complex petroleum UVCBs, these include gas- and liquidbased approaches. Gas chromatography (GC)-based analyses predominate; gas-based analysis of 283 hydrocarbons were first published in the 1960s leading to the development of a standardized method 284 (ASTM 2887-84) for determination of Boiling Range Distribution of Petroleum Fractions by Gas 285 286 Chromatography (Giles, 2016). GC is a powerful tool used for the separation and semi-quantitative assessment of non-polar constituents such as hydrocarbons and polycyclic aromatic hydrocarbons 287 (PAH) (CONCAWE, 2012). It offers better separation than liquid chromatography (LC) but is 288 limited by the boiling point of compounds (affecting the accessible mass range), which is a greater 289 hindrance for the characterization of the heavy petroleum that is increasingly relied upon, and 290 sometimes compounds must be derivatized to ensure GC compatibility. LC is dependent on the 291

polarities of the constituents present, predominantly used to characterize less volatile polar 292 compounds. High-performance liquid chromatography (HPLC; e.g., ASTMD6379 and IP391 293 methods) is used to quantify mono-, di- and tri-aromatic hydrocarbons (CONCAWE, 2012; 294 CONCAWE, 2019). Meanwhile, thin layer or liquid column chromatography (ASTM D2007) 295 296 separation generates information on basic chemical properties (CONCAWE, 2012; CONCAWE, 2019). Gas chromatography coupled to mass spectrometry detection (GC-MS) is widely used in 297 forensic fingerprinting (US EPA 8270 and 8051B) and to characterize the composition of petroleum 298 UVCBs (US EPA, 1996; US EPA, 2014). Flame ionization detection (FID) has been coupled with 299 300 both LC and GC for the detection and quantification of hydrocarbons (CONCAWE, 2012).

301 However, conventional standardized methods detailed above are insufficient to establish truly comprehensive compositional characterization of UVCBs as needed by REACH to accept 302 grouping and read-across hypotheses from registrants. Recent decisions by ECHA on testing 303 proposals provided several reasons as to why that is the case (ECHA, 2020a; ECHA, 2020b; ECHA, 304 305 2021). In these decisions, ECHA noted that (i) physical-chemical characterization of whole complex substances does not demonstrate similarity of chemical constituents of these substances; (ii) 306 elemental and other traditional analysis methods do not provide information on the identity and 307 concentration of individual chemical constituents, but rather provide physical-chemical 308 309 characterization of the substance as a whole; and (iii) standard methods used in the submissions provided insufficient information to estimate the variability of constituents both within and among 310 311 substances and groups. ECHA deems these criteria necessary to establish the applicability domain of a given category to confirm membership of the source substance(s) and enable subsequent read-312 313 across to the target substance(s) (ECHA, 2020a; ECHA, 2020b; ECHA, 2021).

314 Collectively, despite the use of a battery of analytical assays, expending large sample quantities on some of these analyses, using specialized sample preparation techniques, and incurring 315 considerable costs to acquire these data, the registrants did not establish substance characterization 316 that would be acceptable by ECHA. Indeed, the industry itself acknowledges that spectroscopic 317 techniques are limited to bulk characterization and "most substances [i.e., petroleum UVCBs] 318 cannot be effectively differentiated from each other by UV, IR, ¹H-NMR or ¹³C-NMR 319 spectroscopies" (CONCAWE, 2020). Further, even though chromatography-based methods provide 320 considerable amount of information for characterization of nonpolar and relatively volatile 321 compounds (e.g., ASTM D2134, D6729, D6730, among others (CONCAWE, 2012), as well as 322

aliphatic and aromatic fractions (Reddy and Quinn, 1999; Wang and Fingas, 2003), their limited
resolution leaves much of the complex substance uncharacterized (Wang et al., 2011; Weng et al.,
2015). When assessing the needs for analysis of complex substances, it is important to consider that
different analytical approaches are complementary and bring their own advantages and
disadvantages. With the limitations of many of the more routine methods, there has been increasing
use of ultrahigh resolution mass spectrometry and other advanced methods.

More recently, two-dimensional GC (GC×GC) technique has been applied to petroleum substances because it allows for an even greater separation of the multitude of constituents. Specifically, the coupling of GC×GC with FID allows qualitative constituent information at the level of carbon number and chemical class (ASTM International, 2011); this has been useful for simplifying the composition of petroleum substances by binning molecules by "hydrocarbon block" (Redman et al., 2012). Coupling GC×GC with mass spectrometry provides more structural information on the specific constituents (Jennerwein et al., 2014; Mao et al., 2009).

336

337 High and Ultrahigh Resolution Mass Spectrometry Techniques

338 Advancements in mass spectrometry over the past 80 years (Figure 1) have spawned the application of high-resolution approaches for the study of petroleum substances at a molecular level 339 (Palacio Lozano et al., 2019a; Wise et al., 2022). Resolving power is one of the key performance 340 metrics of any mass spectrometry and is typically defined as $\frac{m}{\Delta m}$, where m is the m/z of the ion of 341 interest and Δm is the width of the peak at half its height, using the full width at half maximum 342 343 (FWHM) definition. In essence, the higher the resolving power, the more peaks can be observed for complex samples, as this reduces overlap of peaks (Phillips et al., 2022). "High resolution" has 344 345 typically been accepted to mean a resolving power of >10,000 (Xian et al., 2012), but it has become well-established over the past two decades that "ultrahigh resolution," which often refers to a 346 347 resolving power of >100,000, is essential for characterization of the most complex samples such as 348 petroleum.

The second performance metric that should always be considered is that of mass accuracy. An instrument that offers high mass accuracy indicates that it typically provides data with low mass errors. When an elemental composition (molecular formula) has been assigned to an observed peak, the observed m/z and the theoretical m/z for the assignment can be used to calculate the mass error, measured in parts per million (ppm), using the following equation:

$\frac{m_{observed} - m_{theoretical}}{m_{theoretical}} \times 1,000,000$

355 Note that a negative mass error indicates the peak appears at a lower m/z than the theoretical value and a positive mass error indicates the peak appears at a higher m/z than the theoretical value; for 356 357 each peak, the closer the mass error is to zero, the greater the confidence in the given assignment of the molecular formula. It is also important to note that, while researchers desire resolving power to 358 359 be as high as possible, they strive to keep mass errors as low as possible. Thus, the highest resolving 360 powers afford researchers the ability to observe more components within complex samples, while the highest mass accuracies (lowest mass errors) afford greater confidence in the assignments of 361 elemental composition and structures, together establishing detailed compositional "profiles," 362 "fingerprints," or "signatures" for complex samples. Within the field, there has been discussion 363 364 about the most appropriate terminology to use, drawing parallels with how fingerprints do not normally change but that signatures do, and how this understanding may be related to compositions 365 366 of complex substances changing when subject to anthropogenic or environmental processes.

The comprehensive characterization of molecular composition of petroleum through high 367 368 resolution mass spectrometry is an active area of investigation and includes several approaches to precise detection, naming, and structural characterization of the individual constituents (Hsu et al., 369 370 2011; Marshall and Rodgers, 2004; Niyonsaba et al., 2019; Palacio Lozano et al., 2019a; Palacio 371 Lozano et al., 2019b; Palacio Lozano et al., 2020; Roman-Hubers et al., 2022; Xian et al., 2012). 372 Modern time-of-flight (TOF) mass spectrometers, which are widespread and considered high resolution, offer resolving powers typically in the range of 10,000-60,000. By contrast, FT-ICR MS 373 374 is the highest performance variety of mass spectrometer and is considered to offer researchers ultrahigh resolution, at one or two orders of magnitude higher performance. FT-ICR MS is based 375 376 upon ions orbiting inside a cell, which is in turn housed within the bore of a superconducting magnet, and the technique offers ultrahigh resolving power ($\sim 10^6$ FWHM) and mass accuracy (sub-ppm). 377 Orbitrap mass spectrometers are a newer variety of mass spectrometer based upon a Kingdon trap 378 design, rather than using magnetic fields; these instruments typically offer a resolving power of $\sim 10^5$ 379 FWHM. FT-ICR MS and Orbitrap MS offer differing degrees of ultrahigh resolution, with FT-ICR 380 381 MS offering the highest performance; however, the advantage of TOF MS is in rapid acquisition time which allows for coupling with additional separation techniques such as two-dimensional gas 382 chromatography and ion mobility spectrometry (Palacio Lozano et al., 2019a). These three 383 384 analytical techniques (Figure 2B) are actively used in petroleomics analyses because they offer

somewhat different approaches to determining molecular formulae present in complex substances
(Palacio Lozano et al., 2019b; Rodgers and McKenna, 2011), but have not yet been used for detailed

- 387 petroleum substance characterization for registration or evaluation purposes.
- 388

389 Ultrahigh Resolution MS Data Processing and Visualizations

Petroleum substances contain highly homologous series of hydrocarbon molecules; thus, 390 complex substance analysis can be facilitated by exploiting the patterns of various chemical groups. 391 The method of Kendrick mass defect (KMD) analysis facilitates sorting molecules into homologous 392 393 series (Kendrick, 1963). The composition of complex petroleum substances can be visualized using 394 the KMD approach because most molecules belong to homologous series comprised of (CH₂) alkyl groups and other functional groups (Figure 3A), this method has been widely used in petroleomics 395 (Hughey et al., 2001; Marshall and Rodgers, 2004; Marshall and Rodgers, 2008; Palacio Lozano et 396 al., 2020). Due to the high resolution and mass accuracy, the molecular composition assigned to the 397 398 ions that fall in or out of the homologous series can be used to predict their elemental content (HC[#], O_#, N_#, O_#, S_#), carbon number, and double bond equivalents (DBE = C_# - H_#/2 + N_#/2 + 1). A 399 number of other visualizations have been proposed to express the molecular composition of various 400 401 substances based on the rings and double bonds in the carbon framework (i.e., DBE) of the 402 constituents that can be plotted against their carbon number (Figure 3B). Van Krevelen diagrams (Figure 3C) are used to display the degree of aromaticity and oxidation of constituents by plotting 403 404 the H/C versus O/C ratio of the organic compounds in the complex substance (Kim et al., 2003; Van Krevelen, 1950; Van Krevelen, 1984). Relative abundance of various classes of compounds in the 405 406 samples is typically plotted as a chemical class bar chart (Figure 3D), or as two-dimensional "hydrocarbon blocks" (Figure 3E). 407

Molecular-level analyses using ultrahigh resolution MS typically generate voluminous 408 datasets even after processing by the software provided by instrument manufacturers. A number of 409 410 third-party software packages have been developed recently to facilitate data analysis and visualizations. Peak-by-Peak fee-based software (Spectroswiss, 2019) is tailored for processing FT-411 MS raw data on transients and mass spectra and generates output files for follow-up statistical 412 413 analyses. PetroOrg (Riches et al., 2015) and Composer (Sierra Analytics, 2022), launched in 2014 and 2008, respectively, are two stand-alone fee-based packages that allow processing and 414 visualization of the molecular composition of petroleum substances data acquired through high 415

resolution MS. UltraMassExplorer is an open source web-based package that uses R Studio 416 (Leefmann et al., 2019). It applies static formula libraries for molecular formula assignment of the 417 molecular formulas based on neutral masses coupled to PubChem searches for putative structural 418 assignment. Data visualization is enabled by van Krevelen, KMD and DBE plots. DropMS is 419 420 another web-based tool that facilitates high resolution MS data processing and molecular assignments with its corresponding DBE, error, signal intensity, as well as a number of 421 visualizations (Rosa et al., 2020). KairosMS utilizes an R Shiny interface to process complex data 422 sets produced through hyphenated MS experiments that, when combined with software for formula 423 424 assignments, can be used for visualization, comparison, and statistical analyses for both direct infusion and hyphenated data set, using a wide variety of approaches (Gavard et al., 2020). 425 Interactive van Krevelen (Kew et al., 2017) and Open van Krevelen (Brockman et al., 2018) are 426 two packages that offer interactive diagrams for molecular-level exploration of the data from high 427 428 resolution MS.

429

430 The Regulatory Needs for Characterizing Chemical Composition of Petroleum UVCBs

431 General considerations and examples of the application of high resolution MS for the 432 analysis of petroleum substances have been reviewed elsewhere (Hsu and Shi, 2013; Niyonsaba et 433 al., 2019; Palacio Lozano et al., 2019a; Palacio Lozano et al., 2019b; Palacio Lozano et al., 2020; Rodgers and McKenna, 2011; Xian et al., 2012). However, these previous reviews did not 434 435 specifically place the application of these techniques in the context of the regulatory needs for registration and evaluation of petroleum UVCBs. Comprehensive characterization, or at least more 436 437 detailed information on some specific types of constituents, of complex substances based on high resolving power and mass accuracy of FT-ICR MS, Orbitrap MS and TOF MS affords the 438 opportunity to attain valuable information on both molecular composition and relative abundance 439 of the constituents. Such data can also be used to quantitatively evaluate both variability within a 440 substance or group of substances, and similarity between substances. Then, it should be possible to 441 442 conduct read-across and complete hazard characterization. To provide specific examples of how each technique can be used to address specific regulatory needs, we performed a systematic 443 444 literature search (with end date of December 2021) focused on the application of each technique to petroleum substances (Figure 4). We found that even though there are many dozens to hundreds of 445 publications on each technique in general, and on their use to analyze petroleum samples in 446

particular, few studies presented data or drew conclusions in a manner relevant to address each
regulatory need. The Web of Science search terms and results of the literature search are included
in Supplemental Table 1.

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451 *Critical Need 1: Providing detailed information on the structure of the constituents.*

High mass measurement accuracy and resolving power of modern mass spectrometry analyzers are key steps to address this regulatory need for petroleum UVCBs. Regulatory agencies have repeatedly stressed that "*broad*" compositional information is insufficient to justify groupings and support read-across hypotheses (ECHA, 2014; ECHA, 2020a). While many publications tout ever increasing resolution as an advance in the science of analytical chemistry, we highlight recent examples of studies that focused on comprehensive structural information of the individual constituents in petroleum samples.

459 With the advances in the performance of mass spectrometers and development of new 460 methodologies, significant strides have been made with respect to characterization of mixtures and petroleum samples. For the analysis of whole crude oils and refined products, FT-ICR MS could 461 462 resolve hundreds of molecular formulae in the late 1990s (Rodgers et al., 1998) and several thousand 463 in the early 2000s (Hughey et al., 2002; Qian et al., 2001). By 2019, newer FT-ICR MS 464 instrumentation and experiment design led to the identification of ~245,000 molecular formulae in a non-distillable fraction of the maltenes from a heavy petroleum sample, using a resolving power 465 466 of more than 3,000,000 across the entire m/z range (Palacio Lozano et al., 2019b). Thus, in less than two decades, there was a 60-fold increase in resolving power and 80-fold increase in the number of 467 468 molecular formulae being assigned within a single sample.

469 While ultrahigh resolution mass spectrometry has offered unprecedented levels of insight into highly complex samples, such instrumentation typically affords molecular formulae but does 470 not lead to definitive identification of structures. Experimental parameters, such as polarity and 471 472 ionization behavior, can yield information about probable functionalities of the components, but true structural identification would require additional data. This could be obtained by fragmentation 473 patterns associated with individual peaks (and hence molecular formulae), acquired during "tandem 474 mass spectrometry" (also known as "MS/MS") experiments. As complex samples may comprise 475 tens of thousands or even hundreds of thousands of molecular formulae, performing hundreds of 476

thousands of tandem mass spectrometry experiments to target all of the peaks is not viable, due tothe time required and workload due to data analysis.

Combination with an orthogonal separation method, such as a form of chromatography or 479 ion mobility, is an alternative approach for accessing information about functional groups and 480 481 isomeric contributions. Optimization of these experiments (such as choice of columns, temperature program, etc.) is similarly non-trivial, however, and resolution of components in complex 482 substances can be challenging; the concept of "unresolved complex mixtures", also referred to as an 483 "UCM hump" on the chromatograms, is well-known (Gough and Rowland, 1990). Furthermore, 484 485 while chromatography is a well-known method for distinguishing isomers, this is typically combined with a need for authentic standards (e.g., to determine expected retention times on the 486 chromatography column) and databases, although certainly not all available compounds are found 487 within databases. Where there may be dozens of isomers per molecular formula, this would in 488 practice mean a requirement for millions of authentic standards to be run in order to address the 489 490 most complex petroleum samples that have been characterized by ultrahigh mass spectrometry. This is, again, impractical due to the amount of time required. It is also worth drawing attention to a 491 subtle distinction between molecular formulae (e.g., C₁₈H₃₆O₂) and the concept of "peaks;" a peak 492 may represent a single molecular formula in a direct infusion experiment by mass spectrometry (and 493 494 data is represented by m/z and intensity only), but in combination with mass spectrometry with chromatography, there will be an additional dimension to the data (m/z and intensity, but now also 495 496 time) and multiple peaks per molecular formula may occur due to the presence of isomers. Combining mass spectrometry with chromatography can yield more peaks by counting of isomers, 497 498 but not necessarily more molecular formulae.

499 The isobaric information afforded by ultrahigh resolution direct injection FT-ICR MS has played a significant role in comprehensive understanding of petroleum substance composition; 500 however, the presence of a large number of isomeric hydrocarbons in petroleum substances 501 502 challenges the utility of ultrahigh resolution methods that do not provide other means for separation 503 of isomeric species. Some studies addressed a growing need to define the isomeric composition of petroleum substances by coupling FT-ICR MS with separation techniques. Gas chromatography 504 505 prior to FT-ICR MS was used for separation of isomers in petroleum by resolving individual 506 constituents (Barrow et al., 2014; Palacio Lozano et al., 2022). Information of the structural composition and the isomeric diversity can be also attained by coupling with trapped wave ion 507

mobility spectrometry (TWIMS), a variety of ion mobility spectrometry, to FT-ICR MS (Maillard
et al., 2021). Multidimensional separation with ultrahigh resolution allows for characterization of
individual constituents with accurate mass measurements and structural features. Additionally, high
mass accuracy and ultrahigh resolution measurements have elucidated structural information of the
individual constituents through post-instrumental data analysis (Cho et al., 2011; Hu et al., 2018).

A number of studies have coupled chromatography to Orbitrap MS for analysis of complex 513 environmental samples (MacLennan et al., 2018; Pereira et al., 2013; Sorensen et al., 2019; Yang et 514 al., 2019). Some studies applied this technique for the analysis of oils or fractions thereof. A 515 516 combination of GC separation with different ionization methods for Orbitrap MS was used to study volatile components of a petroleum refining product, gas condensate. The authors showed that 517 separation of different isomeric compounds could be achieved using this hyphenated method thus 518 aiding in deeper characterization of a complex substance (Kondyli and Schrader, 2019). More 519 520 recently, a reverse-phase liquid chromatography method was applied to the analysis of petroleum 521 refining-derived UVCBs and compared to direct injection Orbitrap and FT-ICR MS (Xia et al., 2021). The authors showed that not only could they obtain elemental formulas for a large number 522 523 of hydrocarbon and heteroatom species, but chromatography-informed retention patterns could also be used to distinguish among isomeric species, hence increasing confidence in structural 524 525 identification of the individual constituents in petroleum substances.

TOF MS detection, despite its lower mass resolution as compared to FT-ICR and Orbitrap 526 527 MS, affords an advantage of rapid data acquisition; this technique has been explored extensively by coupling with chromatography or ion mobility spectrometry separations (Palacio Lozano et al., 528 529 2019a). The most common type of front-end chromatography in petroleum substance analysis is GC×GC. The additional separation afforded by this technique offers substantial improvement for 530 531 resolving isomeric constituents and compounds that would otherwise coelute (Ball and Aluwihare, 2014; Luna et al., 2014a; Ngo et al., 2012; Rowland et al., 2011). For example, this technique was 532 533 used to identify a large number of isomeric molecules, both aliphatic and aromatic, that could not be resolved by a typical GC-MS technique in a complex milieu of hydrocarbons containing a wide 534 (C₁₂-C₃₆) range of carbon numbers (Alam et al., 2016). Similarly, the ability GC×GC-TOF MS to 535 resolve isomers of classical and sulfur-containing naphthenic acids in oil-contaminated 536 environmental samples enabled contamination-source fingerprinting (Bowman et al., 2019) and 537 identification of potentially hazardous substances (Bowman et al., 2020). TOF MS has also been 538

coupled with ultrahigh pressure liquid chromatography to isolate isomeric species in petroleum
derivatives (Lv et al., 2013; Mahmoud and Dabek-Zlotorzynska, 2018).

The compositional characterization of petroleum samples has typically relied upon either the 541 use of chromatography or, more recently, the use of ultrahigh resolution mass spectrometry. The 542 543 separation on the basis of retention time using gas chromatography can be sufficient for monitoring targeted components (e.g., steranes for differentiation of sample origins) and revealing the most 544 significant contributions. As samples become more complex, however, particularly with the 545 increasing use of heavy petroleum sources, there is the increasing probability of coeluting 546 547 components, as mentioned above, where compounds have the same retention time and therefore 548 cannot be distinguished by one-dimensional chromatography alone. The use of a second column of a different type to result in a two-dimensional approach, GC×GC, can significantly improve the 549 separation and therefore the number of compounds observed. 550

551 Due to the temperature ranges of the ovens, gas chromatography-based methods have limited 552 retention times that are accessible, as these are, in turn, linked to the boiling points of the individual 553 compounds. Direct infusion methods coupled with ultrahigh resolution mass spectrometry (*i.e.*, 554 where samples are injected directly into an ion source, with no preceding chromatography) do not 555 have the limitations associated with boiling point, and ultrahigh resolution approaches with direct 556 injection delivery have become increasingly significant, especially for heavy petroleum samples.

There has also been limited coupling of chromatography with ultrahigh resolution mass 557 558 spectrometry, which affords researchers the ability to separate compounds on the basis of two different dimensions: retention time and m/z. In this way, co-eluting compounds (*i.e.*, the same 559 560 retention time associated with the GC column) can be separated due to the additional use of the m/zdimension (Barrow et al., 2014). The data sets acquired can be large (e.g., ~25-50 GB) and therefore 561 562 present data processing and data analysis challenges. The combination of orthogonal approaches, yielding so-called "hyphenated" techniques, represents means by which to access the advantages of 563 564 the individual methods. Ultrahigh resolution mass spectrometry affords the ability to resolve a greater number of peaks and assign many thousands of unique molecular formulae with confidence, 565 while chromatography or ion mobility affords the ability to separate isomers (same molecular 566 formulae and therefore same m/z, but differing structures due to different arrangements of the 567 atoms). While GC and low resolution GC-MS methods have seen widespread usage for many years, 568

it can be expected that multidimensional GC and the coupling of orthogonal methods with ultrahighresolution mass spectrometry will both become increasingly used.

Obtaining structural isomeric information is also possible using ion mobility mass 571 spectrometry (IMMS), a post-ionization separation technique, often coupled to TOF MS and used 572 573 for the analysis of petroleum samples (Santos et al., 2015). IMMS provides structural information that is complementary to the observed m/z of a compound by characterizing the spatial conformation 574 of individual constituents via their drift time through an inert gas and the subsequent derivation of 575 collision cross section (CCS) values (Dodds and Baker, 2019). CCS values can also be compared 576 577 with those determined through computational means to determine structures for observed 578 compounds. Currently, this is laborious for complex samples and so not viable for samples with, for example, tens of thousands of molecular formulae and their associated isomers. The combination of 579 ion mobility, mass spectrometry, and computational chemistry does, however, hold potential for 580 providing greater structural insights during characterization of samples. IMMS allows for separation 581 582 of isomeric compounds by their structural composition (Hoskins et al., 2011; Lalli et al., 2015; Lalli et al., 2017; Mahmoud and Dabek-Zlotorzynska, 2018). The high mass accuracy and resolution of 583 584 TOF MS, coupled with structural characterization using ion mobility, has allowed comprehensive 585 elucidation of the composition of complex petroleum substances and byproducts (Lalli et al., 2015; 586 Lalli et al., 2017). Confident molecular formula assignment to the IMMS-derived features in a gasoline standard and a crude oil sample was demonstrated through the use of KMD analyses based 587 588 on CH₂ and H functional units (Roman-Hubers et al., 2021).

While instrumentation has increased in performance, the data analysis methods have often struggled to keep pace. It is evident that highly complex samples are being characterized better than ever before. The field is acutely aware that handling the increasingly complex data is already challenging, but that also there is a need to obtain greater structural insights, going beyond molecular formulae alone and/or ensuring all of the many thousands of peaks are associated with a definitive structure, not only those that are mostly easily targeted. While the progress with complex substance analysis has been remarkable in recent years, there remain mountains to climb.

596

597 *Critical Need 2: Providing information on the <u>concentration</u> of the individual constituents.*

598 Quantification of individual constituents in complex and multi-constituent substances is a 599 required step for hazard evaluation to ensure no underestimation of the potential human health and

environmental hazards (CONCAWE, 2012; ECHA, 2017a). Thus, the ability of high resolution 600 mass spectrometers to determine the thousands of individual constituents is not sufficient without 601 determination of their abundance. Traditional quantitative approaches relying on mass spectrometry 602 detection require the use of various extraction and detection standards, preferably isotopically-603 604 labelled ones (Urban, 2016). However, because of the complexity of petroleum substances, the use of standards for absolute quantitation of numerous, rather than a small number of targeted 605 constituents, would potentially mean the need for millions of authentic standards, which is both 606 impractical and impossible. Instead of quantifying absolute concentrations of multiple individual 607 608 constituents, traditional analysis techniques such as GC-MS and GC×GC-FID derive relative (e.g., 609 fraction of total) amounts for groups of compounds based on a limited number of standards for key classes of constituents. For example, quantitation of individual n-alkanes, selected isoprenoids, 610 polycyclic aromatic and alkyl polycyclic aromatic hydrocarbons, and biomarker compounds is 611 possible by GC-MS (Wang et al., 1994); however, the "UCM hump" of high molecular weight 612 613 hydrocarbons limits the utility of this technique for hazard evaluation. GC×GC-FID technique is also commonly used for petroleum analyses to derive "hydrocarbon blocks" (ASTM International, 614 2011). This technique is more amenable for hazard evaluation as it can separate polycyclic 615 compounds with known or suspected hazardous properties (Bierkens and Geerts, 2014); however, 616 617 this technique is not considered sufficiently informative by some decision-makers, especially for the substances that contain C30 or greater hydrocarbons (ECHA, 2020a). Therefore, there is a great 618 619 need to determine the ability of high resolution MS techniques to provide quantitation of the individual constituents in petroleum substances. 620

621 There are significant challenges with respect to quantifying the abundance of constituents in petroleum UVCBs. The signal observed for a given compound will be influenced by a number of 622 experimental parameters. One of these is, of course, concentration. Other factors include solubility 623 in the chosen solvents, pH of the sample solution, the suitability of ionization method chosen, the 624 625 polarity of the ion source (e.g., when using electrospray ionization, acidic species will be observed in negative-ion mode while basic species will be observed in positive-ion mode), and instrument 626 tuning, amongst other variables. As one example, alkanes can represent as much as half or more of 627 628 the composition of some petroleum samples and yet if performing analysis using electrospray ionization, a widespread ionization method which is suitable for observation of polar and ionic 629 species, then the alkanes would essentially go unobserved. 630

The matter of quantification is also closely linked to structure, as previously discussed. 631 Different isomers of a molecular formula may have differing functional groups which, in turn, may 632 influence the ionization response of the compounds. For instance, an organic molecule containing 633 two oxygen atoms may incorporate two hydroxyl groups or a carboxylic acid, where the carboxylic 634 acid would ionize much more readily than the former, and therefore give a much stronger signal. 635 For this reason, quantification typically involves the coupling of chromatography to mass 636 spectrometry, in order that isomers are separated the signals associated with each isomer can be 637 measured and considered separately. Such experiments are labor intensive, however. Authentic 638 639 standards must be used for each isomer of each molecular formula, as mentioned previously, but the 640 standards must also be prepared as a series of sample solutions spanning a range of concentrations to measure the signal intensity as a function of concentration, leading to calibration curves. The 641 needs for authentic standards and for calculation of the calibration curves leads to increases in 642 researcher workload akin to orders of magnitude. "Untargeted" analyses could necessitate the need 643 644 for potentially millions of experiments for quantification purposes, depending on the objectives. It is much more common that "targeted" analyses are instead used, where a short list of compounds of 645 646 concern are searched for, such as benzene, toluene, ethylbenzene, and xylene compounds. Many advanced mass spectrometry approaches which are commonly used are accepted to be semi-647 648 quantitative, balancing sample complexity, experimental design, data processing, and time.

Most studies that use FT-ICR MS for the analysis of petroleum samples report tens to 649 650 hundreds of thousands of detectable constituents; however, this technique is semi-quantitative where abundance of each molecule would depend on multiple factors (e.g., solubility, concentration, 651 652 ionization response, tuning, etc.). FT-ICR MS studies typically report relative abundances of various hydrocarbon or heteroatom classes rather than that of individual constituents (Bae et al., 2010; Chen 653 et al., 2012; Jennerwein et al., 2014; Kim et al., 2015; Oldenburg et al., 2014; Walters et al., 2015). 654 Some publications focused on detection and quantitation of specific constituents, such as organic 655 656 sulfur compounds (Lu et al., 2013), or metalloporphyrin complexes (Cho et al., 2014); however, these constituents have uncertain relevance for the purpose of hazard evaluation. The ultrahigh 657 resolution Orbitrap MS technique has not been used for the characterization of the individual 658 659 constituents; instead, the abundances of the detected ions with inferred elemental composition are used for semi-quantitative evaluation of various fractions and broad chemical classes (Castiblanco 660 et al., 2020; Liu et al., 2020; Rodrigues Covas et al., 2020; Vanini et al., 2020). 661

Similarly, quantification of the individual constituents using TOF MS is challenging and 662 most studies focused on quantifying abundances of broad classes of compounds (Scarlett et al., 663 2008). A combination of GC×GC-TOF MS and GC×GC-FID was used to conduct qualitative and 664 quantitative analysis of polycyclic aromatic hydrocarbons for several petroleum UVCBs; however, 665 666 quantitation was performed only for the constituents for which chemical standards were used (Ristic et al., 2018). Even though absolute quantitation of the individual constituents in complex petroleum 667 samples may be unattainable, the combination of confident molecular formula assignments aided 668 by structural information provided by IMMS and data on relative abundance does enable 669 670 quantitative evaluation of the most abundant constituents (Roman-Hubers et al., 2021). For example, a study of variability in chemical composition of petroleum UVCBs used the average relative 671 abundance of each constituent in a product across production cycles to determine what molecules 672 may be present at relatively high (e.g., REACH threshold of concern at 0.1% (ECHA, 2017b)) 673 674 amounts and whether those molecules vary significantly between production cycles (Roman-Hubers 675 et al., 2022). This study is an example of how high resolution MS can be used to not only characterize individual constituents, but also to determine their abundance for consideration as 676 potential substances of concern in hazard evaluation. Indeed, the ability to quantify, even in relative 677 terms, the abundance of the identifiable constituents that comprise most of the petroleum sample is 678 679 of utmost interest under the REACH framework (Table 1). The most recent advice from ECHA states that "all constituents present in a concentration at or above 1% must be identified" when 680 681 grouping or read-across is proposed (ECHA, 2022). The bar is even greater for specific constituents that may possess hazardous properties, i.e., "0.1% for constituents that are classified as 682 683 carcinogenic or mutagenic and 0.3% for substances that are toxic to reproduction or development." Collectively, "identified constituents above the thresholds given above must account for a minimum 684 of 80% of the mass of a UVCB substance" (ECHA, 2022). 685

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687 Critical Need 3: Demonstrating compositional similarity of complex petroleum UVCBs through 688 other means when the identification of all individual constituents is not technically possible or 689 impractical.

Even though challenges remain in the confident identification of structures for individual
 constituents in complex substances, the multi-dimensional data from high resolution MS analyses
 (FT-ICR, Orbitrap and TOF MS) is highly valuable for evaluating broad similarities among complex

petroleum UVCB substances and for identifying the degree of variability within a class of substances or between production batches of the same substance (**Figure 4**). Indeed, a number of studies have used these data to perform statistical analyses and visualize the relationships between samples in large datasets to demonstrate that even broad molecular compositional data can be effective as a means for screening, evaluating similarity and determining what constituents may be variable to prioritize selected samples and constituents that may warrant further targeted quantitative analyses.

The variation in composition of hydrocarbons and heteroatoms resolved using FT-ICR MS 700 701 has been explored for the analysis of crude oils (Hosseini et al., 2021; Rocha et al., 2018; Silva et 702 al., 2020) and refining products (Benassi et al., 2013; Mennito and Qian, 2013; Oldenburg et al., 2014; Oldenburg et al., 2017; Orrego- Ruíz, 2018; Silva et al., 2020; Walters et al., 2015; Wang et 703 al., 2020), including studies that used such data for categorization of new products (Abib et al., 704 705 2020; Hourani et al., 2013; Liu et al., 2014). Through detailed chemical profiles, compositional 706 characterization of different environmental samples allowed detection of the asphaltenes (Neumann et al., 2021; Ruger et al., 2015), fulvic acids (Stenson et al., 2003), oil sands process-affected water 707 708 (Barrow et al., 2016), bitumen (Lacroix-Andrivet et al., 2021), biochar-derived organic matter (Li 709 et al., 2022a), as well as soil and sedimentary organic matter (Zhong et al., 2011). Studies of oil 710 weathering (Wozniak et al., 2019) and transformation in the environment (Jaggi et al., 2019; Li et al., 2022b; Wozniak et al., 2019) enabled to not only determine the trends in compositional changes, 711 712 but also to group samples, allow for forensic identification of the related samples, predict the potential environmental impact of oil spills, and designing mitigation strategies. 713

714 While there will be limitations to the lowest m/z that can be detected by FT-ICR MS, based upon the magnetic field strength and highest frequency that can be detected (e.g., $\sim m/z$ 37 for a 715 modern 12 T instrument), with appropriate instrument tuning, both Orbitrap and FT-ICR mass 716 spectrometers can offer comprehensive insights into the lower molecular weight compounds of the 717 718 chemical profile of complex substances, not afforded by other high resolution MS instruments (Chen et al., 2018; Cheng and Hous, 2021; Cho et al., 2017; Headley et al., 2011). Through the application 719 of Orbitrap MS the variations of the mass spectrum profiles can be traced at a molecular level with 720 721 low mass error to determine the overall chemical composition (Castiblanco et al., 2020; Dong et al., 2019; Porto et al., 2019; Silva et al., 2019; Vanini et al., 2020). Liquid chromatography separation 722 techniques coupled to Orbitrap MS have provided qualitative and quantitative monitoring of the 723

organic species through the chemical profile complex substances. (Folkerts et al., 2019; Miles et al.,
2020; Sorensen et al., 2019; Xia et al., 2021)

When designing future strategies for screening petroleum-related compounds, it is important 726 to consider what is being measured by different approaches. For example, with electrospray 727 728 ionization, acidic species would typically deprotonate and be observed using negative-ion mode, while basic species would protonate and be observed using positive-ion mode; acidic and basic 729 species would not be observed in the same experiment, therefore. While this might initially be 730 considered a disadvantage because of the need for two experiments instead of one, on the other hand 731 732 this can be used to the researcher's advantage when needing to differentiate by functionality. An 733 organic molecule with a heteroatom content of only one nitrogen atom (e.g., C_cH_hN₁) could be pyrrolic (weakly acidic) if observed in negative-ion mode, but would more likely be pyridinic (basic) 734 if observed in positive-ion mode. Where electrospray ionization is suitable for polar and ionic 735 compounds, it does not readily ionize non-polar species and so, for these, an alternative ionization 736 737 method, such as atmospheric pressure photoionization, would be used. To cover acidic, basic, and non-polar species in an untargeted manner, three experiments may be used, but the number of 738 739 methods employed could be reduced if instead adopting a targeted approach. Through an awareness 740 of the advantages and disadvantages of the different ionization methods, combined with clear 741 objectives of the screening process (which could become of a targeted nature), it is possible to tailor a system which balances required information and workload. 742

743 Multidimensional GC separation coupled with high resolution detection in TOF MS has been widely employed to map out the composition of complex substances. The high-throughput 744 745 acquisition has been readily employed to screen for saturate and aromatic hydrocarbons to characterize the broad compositional profile (Haitao et al., 2013; Hao et al., 2017; Kulkarni and 746 Thies, 2012; Luo et al., 2016; Qian et al., 2004; Rui et al., 2012). Nevertheless, to achieve a better 747 compositional information and distribution patterns of complex volatile constituents, TOF MS has 748 749 been coupled with gas chromatography for qualitative and quantitative assessment (Haitao et al., 2013; Hao et al., 2017). When coupled with an HPLC, non-volatile components can be readily 750 separated based on isomeric and isobaric information for comprehensive high resolution 751 752 characterization (Cao et al., 2020). The drawbacks observed from the above addressed separation 753 techniques prompted the coupling of capillary electrophoresis with TOF MS for qualitative assessment of high molecular weight constituents in a heavy gas oil (Nolte et al., 2013). Compared 754

to other TOF MS hyphenations, GC×GC offers multidimensional high resolution characterization 755 to map out the composition of complex substances (Alam et al., 2018; Frenzel et al., 2010; Gabetti 756 et al., 2021; Luna et al., 2014b; Muller et al., 2020; O'Reilly et al., 2019; Qian and Wang, 2019; 757 Ristic et al., 2018; Zhu et al., 2020). The plots generated from GC×GC-TOF MS analysis plotting 758 constituents in a two-dimensional space (1st retention time versus 2nd retention time) help elucidate 759 the composition of complex substances and facilitates the molecular classification of compounds 760 (Damasceno et al., 2014). Notably, physical characteristics (*i.e.*, density) can be directly correlated 761 based on the detailed chemical composition defined through GC×GC-TOF MS (Vozka et al., 2019). 762

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Petroleum UVCB composition: A regulatory challenge, but what are the solutions?

As summarized above, novel high resolution mass spectrometers offer comprehensive 765 characterization of petroleum substances to qualitatively assess and measure the broad chemical 766 767 composition of a substance and its individual constituents. The information available through high 768 resolution characterization of the chemical composition of UVCBs can address the shortcomings with regards to the prediction of toxicological properties when practicing read-across assessment. 769 770 Comprehensive characterization can provide sufficient information on the molecular composition 771 and their relative abundance within a substance to define the commonality between substances and 772 their similarity at a molecular level. With this information, it is then possible to determine constituents which may be used to infer human health hazard properties of the whole substance. 773

774 Still, complex UVCBs, especially those produced by refining of oil, remain to be an evolving competency in regulatory science. In the recent past, both decision-makers and industry relied on 775 776 rather imprecise substance categorization and read-across hypotheses to predict toxicological properties (Clark et al., 2013; McKee et al., 2015; Salvito et al., 2020). Recognizing the lack of 777 clarity in the original regulations and guidelines (ECHA, 2017c), and persistent issues identified in 778 regulatory submissions of petroleum UVCBs (ECHA, 2020a; ECHA, 2020b; ECHA, 2021), 779 780 additional advice was recently provided with respect to the information on chemical composition for UVCB substances (ECHA, 2022). Indeed, there appears to be a major gap in what information 781 is perceived as sufficient in terms of chemical characterization of petroleum UVCBs by either 782 783 decision-makers or industry. At the same time, the science of analytical chemistry delivered a number of major advances in terms of novel techniques and visualizations to aid in deep 784 characterization of complex petroleum UVCBs. Scores of research studies have been published, 785

reviews written, and lectures delivered. A "meeting in the middle" is required between those
working in regulatory and analytical disciplines, so that the translation of petroleomics science into
the practice of decision making on petroleum UVCBs can begin to be realized.

One barrier to such a translation could be the difference of opinions on what "science" is 789 790 "evidence" and whether the scientific data are "sufficient" to satisfy the regulatory requirements for making decisions (National Research Council, 2009). What is sufficient for a peer-reviewed 791 792 scholarly publication in a specialized scientific journal may not be sufficient for a regulatory decision. For example, in the EU, it is generally noted that "available scientific and technical data" 793 794 shall be taken into account "in preparing its policy on the environment" (Allio et al., 2006). It was noted that this legislative mandate does not call for the available data to be of "best available" 795 quality, even though some of the agencies are required to "provide the EU institutions and the 796 Member States with the best possible scientific opinions" (Allio et al., 2006). In the US, the concept 797 798 of "best available science" has been more defined as a statutory requirement for risk evaluation of 799 chemical substances (US EPA, 2017). Specifically, the statute defines best available science as "science that is reliable and unbiased. Use of best available science involves the use of supporting 800 801 studies conducted in accordance with sound and objective science practices, including, when 802 available, peer reviewed science and supporting studies and data collected by accepted methods or 803 best available methods (if the reliability of the method and the nature of the decision justifies use of the data). Additionally, EPA will consider as applicable: (1) The extent to which the scientific 804 805 information, technical procedures, measures, methods, protocols, methodologies, or models 806 employed to generate the information are reasonable for and consistent with the intended use of the 807 information; (2) The extent to which the information is relevant for the Administrator's use in 808 making a decision about a chemical substance or mixture; (3) The degree of clarity and completeness with which the data, assumptions, methods, quality assurance, and analyses employed 809 to generate the information are documented; (4) The extent to which the variability and uncertainty 810 in the information, or in the procedures, measures, methods, protocols, methodologies, or models, 811 are evaluated and characterized; and (5) The extent of independent verification or peer review of 812 the information or of the procedures, measures, methods, protocols, methodologies or models." In 813 principle, all of these can apply to characterization of the chemical composition of petroleum 814 UVCBs when data are submitted to the authorities for a decision. 815

Related to the concept of best available science is the concept of "reasonably available 816 information" which means information that the agency "possesses or can reasonably generate, 817 obtain, and synthesize for use in risk evaluations, considering the deadlines [...] for completing 818 such evaluation" (US EPA, 2017). To relate the latter concept to the data on chemical composition 819 820 of petroleum UVCBs, one may argue that a range of technically valid analytical methods applicable to the challenge of characterizing their composition and variability are already available and the data 821 could be generated in a reasonable period of time. There is a balance between cost, availability, 822 time, and the level of information required; the most advanced techniques which ultimately yield 823 824 the fullest possible understanding may not be suitable for routine screening, whilst more common 825 techniques may afford incomplete information or increased risk of misinterpretation. In this regard, novel analytical advances, such as those described herein, have greatly increased our ability to 826 access valuable information on the composition of UVCBs at a molecular level. At the same time, 827 the advances have revealed in much greater detail the compositional complexity of petroleum and 828 829 areas where development is still required in order to fully access the range of components and, most significantly, establish structures. Still, strong arguments have been made about the impractical and 830 831 potentially unnecessary regulatory requirements to deconvolute every possible constituent in a UVCB, even if at some arbitrary abundance cut-off level(s) (Table 1), and that hazard assessment 832 833 may be sufficiently informed by using bulk compositional data and hydrocarbon blocks (Redman et al., 2012; Salvito et al., 2020). Clearly, the fields of analytical chemistry and regulatory science have 834 835 not always worked in tandem.

To bridge this chasm, all sides will need to meet in the middle to establish baseline 836 837 requirements with an understanding of what is analytically possible (Figure 4), including considerations of instrumentation availability, cost and time required for the analysis and data 838 processing. It has been well-documented that different analytical methods, when applied to 839 petroleum substances, have different windows of applicability with respect to the types of molecules 840 they may ionize and/or detect, and may suffer from being semi-quantitative (Aeppli, 2022; 841 Fernandez-Lima et al., 2009; McKenna et al., 2013; Prince and Walters, 2022; Rodgers and 842 McKenna, 2011). Indeed, Figure 4 illustrates the point that novel high resolution analytical 843 844 methods (FT-ICR, Orbitrap-MS and TOF-MS) have been already applied by a number of authors to the research questions that are directly relevant to the specific regulatory needs in petroleum 845 UVCB space. However, the number of such publications remains very small as compared to studies 846

of the traditional analytical approaches (Figure 2), and seldom do the authors acknowledge the 847 potential for their work to be applied in the regulatory context(s). It is clear that the analytical 848 chemists could better appreciate the regulatory issues and strive to provide solutions that not only 849 advance the science but also address specific challenges. For example, the focus on the critical needs 850 851 identified herein will serve well both the researchers and the ultimate end-users if targeted collaborations and regulatory-informed case studies to build confidence in their performance and 852 utility. In addition, cross-validation collaborative trials are needed to achieve standardization of the 853 new techniques and analysis methods to build confidence among stakeholders, a relevant example 854 855 comes from the evolution of the analytical methods used for oil spill response (Faksness et al., 2002).

Concomitantly, the government bodies tasked with enforcing REACH and other relevant 856 regulations, while providing additional useful guidance about the details of chemical 857 characterization of petroleum UVCBs (ECHA, 2022), should have realistic expectations as to what 858 859 is achievable using even the most advanced analytical methods. The most comprehensive analytical 860 information may or may not be actionable in terms of hazard evaluation in absence of anchoring to the toxicological data. In this regard, existing data on a handful of known hazardous components in 861 petroleum UVCBs, be it "priority PAHs" (ATSDR, 2005) or other constituents with existing human 862 hazard evaluations, are widely regarded as insufficient for assessment of the whole substance(s). 863 864 Thus, additional data from in vitro and other "bioactivity" data streams may be needed to determine the toxicologically relevant compositional features and variability within and among substances and 865 866 categories (Grimm et al., 2016; House et al., 2022; House et al., 2021).

Concomitantly, the size and complexity of data sets pertaining to sample composition, which 867 868 are increasingly accompanied with orthogonal high-dimensional information such as (eco)toxicological data, present challenges with respect to data handling and subsequent 869 interpretation. These comprehensive high-/multi-dimensional datasets will typically be subject to 870 dimensionality reduction and used to support grouping and/or classification of the individual 871 872 petroleum UVCBs. A recent study demonstrated that the choices of the data analysis and visualization methods can not only potentially aid in the communication of "sufficient similarity" 873 among complex substances, but also yield different outcomes in terms of grouping and classification 874 (Onel et al., 2019). The proverbial "black box" of data processing and analysis may create "a variety 875 of methodological and scientific concerns which mean that it is impossible to independently assess 876 the methods and results" (ECHA, 2020a) in terms of reliably predicting the properties of the 877

substances that are being evaluated. Hence, it is important to involve not only the analytical chemists when examining the data, but those in other scientific disciplines, such as statistics, mathematics, and computer science, and ensure that the data processing and analysis methods are transparent, and that the decision-makers are sufficiently familiar with the bioinformatics aspects of the information presented to them. Such involvement can expedite assessments of which components are particularly relevant for regulatory needs and may warrant greater focus and analytical precision.

Overall, while statutory and advisory language in the government agencies-produced 884 documents may seem clear, the analytical science may not be available to fully identify and quantify 885 886 all, or even most, components within complex petroleum UVCBs. It is being acknowledged that "it 887 is not required to provide detailed structural information on all constituents of a UVCB substance, but there must be sufficient characterization of constituents so as to demonstrate structural 888 similarity, and consecutively provide a basis for predicting the properties of the substance in read-889 across" (ECHA, 2020a). Even though additional clarifications have been recently made with respect 890 891 to what "sufficient" may mean in the context of petroleum UVCBs (ECHA, 2022), ultimately, the flexibility is needed to consider new science and determine if it is "best available" and also fit for a 892 893 specific decision-context purpose so that the statutes and regulations are ultimately enforceable. 894 While adherence to the most common and established methods is understandable, the industry needs 895 to be more open to the application of the modern analytical chemistry methods to the analysis of the samples and inclusion of such data into regulatory submissions. The traditional approaches that are 896 897 used for broad characterization of petroleum UVCBs still have an important role to play with respect to substance identification, greater consideration of nontargeted approaches (and how these may 898 899 also shed new light on emerging hazards), followed by targeted approaches in some cases, is needed. It is also important, however, to recognize that the measurement of the composition of a complex 900 substance is very much influenced by the methods and techniques used; comparisons of data from 901 different laboratories must demonstrate appreciation of such factors if the comparisons are to be 902 903 meaningful. Therefore, detailed characterization of the technical performance of novel analytical 904 methods is required, in addition to the generation of data for a wide range of samples and applications. 905

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909 Declaration of Competing Interest

- 910 The authors declare that they have no known competing financial interests or personal 911 relationships that could have appeared to influence the work reported in this paper.
- 912

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918 Table 1. Summary of the chemical characterization needs for UVCB that are registered and919 evaluated under REACH in the European Union.

	Substance Identification for <u>Registration</u> (REACH Annex VI, Section 2)	Hazard Characterization for <u>Evaluation</u> (REACH Annex XI, Section 1.5)		
Infor- mation Needed		Read-Across Assessment Framework: Considerations on multi-constituent substances and UVCBs (ECHA, 2017c)		Advice on Using Read-Across for UVCB substances
		Category Approach	Analogue Approach	(ECHA, 2022)
Overall Substance Identification	 Substance name [from a trade association and/or nomenclature system] Substance identifiers [source, manufacturing process, carbon and boiling range, phys-chem properties, etc.] 	 Substance(s) to be grouped in a category Compositions to be included Manufacturing process description 	 Source substance(s) and target substance Compositions to be included Manufacturing process description 	 Similarity may be established based on: (i) presence of <i>identical</i> constituents OR (ii) variation in concentration and variability in constituents Constituents present at >1% must be identified; lower thresholds if constituents of concern are present (>0.1% for carcinogenic/mutagenic, >0.3% for repro/developmental) >80% of constituents in the substance must be identified
Composition Characterization	 Constituents present >10% Constituents <10% that may be impacting hazard classification 	Category "domain" needs to be defined (constituent- specific concentration determination not specified)	 Source substance(s) Target substance (constituent- specific concentration determination not specified) 	 Must be characterized up to 100% When full characterization is impractical/impossible need to provide (i) justification AND (ii) demonstration of similarity "by other means"
Variability	Not Required	 Structural similarity for category based on worst-case scenario Determine if quantitative differences or patterns in predicted properties may be reflected in structural similarity 	 Structural Similarity between source & target based on worst- case scenario Determine if quantitative differences or patterns in predicted properties may be reflected in structural similarity 	 Structural similarity explained based on quantitative and qualitative comparison of composition When full characterization is impractical/impossible need to provide (i) comparison of constituents AND (ii) demonstration of similarity "by other means" (e.g. analytical information for >95% constituents, constituents of high concern, high resolution for confidence fingerprinting) Analysis of at least 5 independent (i.e., production batches) samples analyzed from ALL registrants of a substance

922	References
923 924 925 926	Abib, G. A. P., et al., 2020. Assessing raw materials as potential adsorbents to remove acidic compounds from Brazilian crude oils by ESI (-) FT-ICR MS. An Acad Bras Cienc. 92, e20200214.
927	Aeppli, C., 2022. Recent advance in understanding photooxidation of hydrocarbons after oil spills.
928	Curr Opin Chem Eng. 36, 100763.
929 930	Alam, M. S., et al., 2016. Using Variable Ionization Energy Time-of-Flight Mass Spectrometry with Comprehensive GCxGC To Identify Isomeric Species. Anal Chem. 88, 4211-20.
931	Alam, M. S., et al., 2018. Mapping and quantifying isomer sets of hydrocarbons
932	(≥ C ₁₂) in diesel exhaust, lubricating oil and diesel fuel samples
933	using GC × GC-ToF-MS. Atmos Meas Tech. 11, 3047-3058.
934 935	Allio, L., et al., 2006. Enhancing the role of science in the decision-making of the European Union. Regul Toxicol Pharmacol. 44, 4-13.
936	API, Petroleum Process Stream Terms Included in the Chemical Substances Inventory Under the
937	Toxic Substances Control Act (TSCA). In: Control, H. S. R. c. T. F. o. T. S., (Ed.).
938	American Petroleum Institute, 1983.
939	ASTM International, UOP Method 990-11: Organic Analysis of Distillate by Comprehensive
940	Two-Dimensional Gas Chromatography with Flame Ionization Detection. ASTM
941	International, West Conshohocken, PA, 2011.
942 943	ATSDR, Toxicology profile for polyaromatic hydrocarbons. CRC Press Boca Raton City, FL, 2005.
944	 Bae, E., et al., 2010. Identification of about 30 000 Chemical Components in Shale Oils by
945	Electrospray Ionization (ESI) and Atmospheric Pressure Photoionization (APPI) Coupled
946	with 15 T Fourier Transform Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS)
947	and a Comparison to Conventional Oil. Energ Fuel. 24, 2563-2569.
948	Ball, G. I., Aluwihare, L. I., 2014. CuO-oxidized dissolved organic matter (DOM) investigated
949	with comprehensive two dimensional gas chromatography-time of flight-mass
950	spectrometry (GC × GC-TOF-MS). Org Chem. 75, 87-98.
951 952	Barrow, M. P., et al., 2014. An added dimension: GC atmospheric pressure chemical ionization FTICR MS and the Athabasca oil sands. Anal Chem. 86, 8281-8.
953	Barrow, M. P., et al., 2016. Effects of Extraction pH on the Fourier Transform Ion Cyclotron
954	Resonance Mass Spectrometry Profiles of Athabasca Oil Sands Process Water. Energy &
955	Fuels. 30, 3615-3621.
956 957	Benassi, M., et al., 2013. Petroleum crude oil analysis using low-temperature plasma mass spectrometry. Rapid Commun Mass Spectrom. 27, 825-34.

958 959	Bierkens, J., Geerts, L., 2014. Environmental hazard and risk characterisation of petroleum substances: a guided "walking tour" of petroleum hydrocarbons. Environ Int. 66, 182-93.
960 961 962	Bowman, D. T., et al., 2019. Profiling of individual naphthenic acids at a composite tailings reclamation fen by comprehensive two-dimensional gas chromatography-mass spectrometry. Sci Total Environ. 649, 1522-1531.
963	Bowman, D. T., et al., 2020. Isomer-specific monitoring of naphthenic acids at an oil sands pit
964	lake by comprehensive two-dimensional gas chromatography-mass spectrometry. Sci Total
965	Environ. 746, 140985.
966 967	Brockman, S. A., et al., 2018. Van Krevelen diagram visualization of high resolution-mass spectrometry metabolomics data with OpenVanKrevelen. Metabolomics. 14, 48.
968 969 970 971	Cao, X. H., et al., 2020. Sequential thermal dissolution of two low-rank coals and characterization of their structures by high-performance liquid chromatography/time-of-flight mass spectrometry and gas chromatography/mass spectrometry. Rapid Commun in Mass Spectrom. 34, e8887.
972 973	Castiblanco, J. E. B., et al., 2020. Molecular behavior assessment on initial stages of oil spill in terrestrial environments. Environ Sci Pollut Res. 28, 13595-13604.
974	Chen, X., et al., 2018. Separation and Molecular Characterization of Ketones in a Low-
975	Temperature Coal Tar. Energy Fuels. 32, 4662-4670.
976	Chen, X. B., et al., 2012. Characterization and Comparison of Nitrogen Compounds in
977	Hydrotreated and Untreated Shale Oil by Electrospray Ionization (ESI) Fourier Transform
978	Ion Cyclotron Resonance Mass Spectrometry (FT-ICR MS). Energy & Fuels. 26, 1707-
979	1714.
980	Cheng, X., Hous, D., 2021. Characterization of Severely Biodegraded Crude Oils Using Negative-
981	Ion ESI Orbitrap MS, GC-NCD and GC-SCD: Insights into Heteroatomic Compounds
982	Biodegradation. Energies. 14.
983	Cho, Y., et al., 2017. Extension of the Analytical Window for Characterizing Aromatic
984	Compounds in Oils Using a Comprehensive Suite of High-Resolution Mass Spectrometry
985	Techniques and Double Bond Equivalence versus Carbon Number Plot. Energy & Fuels.
986	31, 7874-7883.
987	Cho, Y., et al., 2011. Planar limit-assisted structural interpretation of
988	saturates/aromatics/resins/asphaltenes fractionated crude oil compounds observed by
989	Fourier transform ion cyclotron resonance mass spectrometry. Anal Chem. 83, 6068-73.
990	Cho, Y., et al., 2014. Evaluation of Laser Desorption Ionization Coupled to Fourier Transform Ion
991	Cyclotron Resonance Mass Spectrometry To Study Metalloporphyrin Complexes. Energy
992	& Fuels. 28, 6699-6706.
993 994	Clark, C. R., et al., 2013. A GHS-consistent approach to health hazard classification of petroleum substances, a class of UVCB substances. Regul Toxicol Pharmacol. 67, 409-20.

995 996	CONCAWE, REACH – Analytical characterisation of petroleum UVCB substances. Brussels, Belgium, 2012.
997 998	CONCAWE, Guidance on Reporting Analytical Information for Petroleum Substances in REACH Registration Dossiers V2. Brussels, 2014.
999 1000	CONCAWE, Concawe Substance Identification Group Analytical Program Report (Abridged Version). Brussels, Belgium, 2019.
1001 1002	CONCAWE, Guidance to Registrants on Methods for Characterisation of Petroleum UVCB Substances for REACH Registration Purposes. Brussels, Belgium, 2020.
1003	CONCAWE, Petroleum Substances and REACH. 2021.
1004	CONCAWE, REACH Background. 2022.
1005 1006	Damasceno, F. C., et al., 2014. Characterization of naphthenic acids using mass spectroscopy and chromatographic techniques: study of technical mixtures. Anal Methods. 6, 807-816.
1007 1008	Dimitrov, S. D., et al., 2015. UVCB substances: methodology for structural description and application to fate and hazard assessment. Environ Toxicol Chem. 34, 2450-62.
1009 1010 1011	Dodds, J. N., Baker, E. S., 2019. Ion Mobility Spectrometry: Fundamental Concepts, Instrumentation, Applications, and the Road Ahead. J Am Soc Mass Spectrom. 30, 2185- 2195.
1012 1013 1014	Dong, X., et al., 2019. Evaluation of elemental composition obtained by using mass spectrometer and elemental analyzer: A case study on model compound mixtures and a coal-derived liquid. Fuel. 245, 392-397.
1015 1016	ECHA, Guidance on information requirements and chemical safety assessment. Chapter R.6: QSARS and grouping of chemicals. 2008.
1017 1018 1019	ECHA, Decision on a testing proposal set out in a registration pursuant to Article 40(3) of regulation (EC) No 1907/2006 for Asphalt, CAS No 8052-42-4 (EC No 232-490-9). European Chemical Agency, Helsinki, Finland, 2014.
1020 1021	ECHA, Guidance for identification and naming of substances under REACH and CLP. Vol. 2.1. European Chemical Agency, Helsinki, Finland, 2017a.
1022 1023	ECHA, Guidance on requirements for substances in articles. Vol. Version 4.0. European Chemicals Agency, Helsinki, Finland, 2017b.
1024 1025	ECHA, Read-Across Assessment Framework (RAAF) - considerations on multi-constituent substances and UVCBs. European Chemical Agency, Helsinki, Finland, 2017c.
1026 1027	ECHA, Testing Proposal Decision on Substance EC 295-332-8 "Extracts (petroleum), deasphalted vacuum residue solvent". European Chemicals Agency, Helsinki, Finland, 2020a.

- ECHA, Testing Proposals Decision on Substance EC 265-110-5 "Extracts (petroleum), residual
 oil solvent". 2020b.
- ECHA, Testing Proposal Decision on Substance EC 265-182-8 "Gas oils (petroleum),
 hydrodesulfurized". European Chemicals Agency, Helsinki, Finland, 2021.
- ECHA, Advice on using read-across for UVCB substances obligations arising from Commission
 Regulation 2021/979, amending REACH annexes. European Chemicals Agency, Helsinki,
 Finland, 2022.
- EPA, U. S., Toxic Substances Control Act Inventory Representation for Chemical Substances of
 Unknown or Variable Composition, Complex Reaction Products and Biological Materials:
 UVCB Substances. 1995.
- European Commission, COMMISSION REGULATION (EU) 2021/979 of 17 June 2021
 amending Annexes VII to XI to Regulation (EC) No 1907/2006 of the European
 Parliament and of the Council concerning the Registration, Evaluation, Authorisation and
 Restriction of Chemicals (REACH). Official Journal of the European Union, Brussels,
 Belgium, 2021.
- European Council, Corrigendum to Regulation (EC) No 1907/2006 of the European Parliament
 and of the Council of 18 December 2006 concerning the Registration, Evaluation,
 Authorisation and Restriction of Chemicals (REACH), establishing a European Chemicals
 Agency, amending Directive 1999/45/EC and repealing Council Regulation (EEC) No
 793/93 and Commission Regulation (EC) No 1488/94 as well as Council Directive
 76/769/EEC and Commission Directives 91/155/EEC, 93/67/EEC, 93/105/EC and
 2000/21/EC. Vol. L 136/3. European Commission, Brussels, Belgium, 2007.
- Faksness, L. G., et al., 2002. Round Robin Study—Oil Spill Identification. Environ Forensics. 3,
 279-291.
- Fernandez-Lima, F. A., et al., 2009. Petroleum crude oil characterization by IMS-MS and FTICR
 MS. Anal Chem. 81, 9941-7.
- Folkerts, E. J., et al., 2019. Toxicity in aquatic model species exposed to a temporal series of three
 different flowback and produced water samples collected from a horizontal hydraulically
 fractured well. Ecotoxicol Environ Saf. 180, 600-609.
- Frenzel, M., et al., 2010. Complications with remediation strategies involving the biodegradation
 and detoxification of recalcitrant contaminant aromatic hydrocarbons. Sci Total Environ.
 408, 4093-4101.
- FuelsEurope, FuelsEurope position paper on REACH and the Refining industry. Brussels,
 Belgium, 2015.

Gabetti, E., et al., 2021. Chemical fingerprinting strategies based on comprehensive two dimensional gas chromatography combined with gas chromatography-olfactometry to

capture the unique signature of Piemonte peppermint essential oil (Mentha x piperita var 1064 Italo-Mitcham). J Chromatogr A. 1645, 462101. 1065 Gavard, R., et al., 2020. KairosMS: A New Solution for the Processing of Hyphenated Ultrahigh 1066 Resolution Mass Spectrometry Data. Anal Chem. 92, 3775-3786. 1067 Giles, H. N., 2016. Crude Oil Analysis: History and Development of Test Methods From 1854 to 1068 2016. Mater Perform Charact. 5, 1-169. 1069 Gough, M. A., Rowland, S. J., 1990. Characterization of unresolved complex mixtures of 1070 hydrocarbons in petroleum. Nature. 344, 648-650. 1071 Grimm, F. A., et al., 2016. A chemical-biological similarity-based grouping of complex 1072 substances as a prototype approach for evaluating chemical alternatives. Green Chem. 18, 1073 4407-4419. 1074 Haitao, S., et al., 2013. Hydrocarbon composition of different VGO feedstocks and the correlation 1075 with FCC product distribution. China Petrol Proc Petrochem Techn. 15. 1076 Hao, J., et al., 2017. Thermal cracking behaviors and products distribution of oil sand bitumen by 1077 TG-FTIR and Py-GC/TOF-MS. Energy Convers Manag. 151, 227-239. 1078 Headley, J. V., et al., 2011. Preliminary fingerprinting of Athabasca oil sands polar organics in 1079 environmental samples using electrospray ionization Fourier transform ion cyclotron 1080 resonance mass spectrometry. Rapid Commun Mass Spectrom. 25, 1899-909. 1081 1082 Hoskins, J. N., et al., 2011. Architectural Differentiation of Linear and Cyclic Polymeric Isomers by Ion Mobility Spectrometry-Mass Spectrometry. Macromolecules. 44, 6915-6918. 1083 Hosseini, S. H., et al., 2021. Characterization of crude oils derived from carbonate and siliciclastic 1084 source rocks using FTICR-MS. Org Geochem. 159, 104286. 1085 Hourani, N., et al., 2013. Atmospheric pressure chemical ionization Fourier transform ion 1086 cyclotron resonance mass spectrometry for complex thiophenic mixture analysis. Rapid 1087 Commun Mass Spectrom. 27, 2432-8. 1088 House, J. S., et al., 2022. Grouping of UVCB substances with dose-response transcriptomics data 1089 1090 from human cell-based assays. ALTEX. 39, 388-404. 1091 House, J. S., et al., 2021. Grouping of UVCB substances with new approach methodologies (NAMs) data. ALTEX. 38, 123-137. 1092 Hsu, C. S., et al., 2011. Petroleomics: advanced molecular probe for petroleum heavy ends. J Mass 1093 1094 Spectrom. 46, 337-43. Hsu, C. S., Shi, Q., 2013. Prospects for petroleum mass spectrometry and chromatography. Sci 1095 China Chem. 56, 833-839. 1096

- Hu, M., et al., 2018. Collision cross section (CCS) measurement by ion cyclotron resonance mass
 spectrometry with short-time Fourier transform. Rapid Commun Mass Spectrom. 32, 751 761.
- Hughey, C. A., et al., 2001. Kendrick mass defect spectrum: a compact visual analysis for
 ultrahigh-resolution broadband mass spectra. Anal Chem. 73, 4676-81.
- Hughey, C. A., et al., 2002. Resolution of 11,000 compositionally distinct components in a single
 electrospray ionization Fourier transform ion cyclotron resonance mass spectrum of crude
 oil. Anal Chem. 74, 4145-9.
- Jaggi, A., et al., 2019. Composition of the dissolved organic matter produced during in situ
 burning of spilled oil. Org Geochem. 138, 103926.
- Jennerwein, M. K., et al., 2014. Complete Group-Type Quantification of Petroleum Middle
 Distillates Based on Comprehensive Two-Dimensional Gas Chromatography Time-of Flight Mass Spectrometry (GC×GC-TOFMS) and Visual Basic Scripting. Energy Fuels.
 28, 5670-5681.
- 1111 Kaiser, M. J., 2017. A review of refinery complexity applications. Pet Sci. 14, 167-194.
- 1112 Kendrick, E., 1963. A Mass Scale Based on CH2 = 14.0000 for High Resolution Mass
 1113 Spectrometry of Organic Compounds. Anal. Chem. 35, 2146-2154.
- 1114 Kew, W., et al., 2017. Interactive van Krevelen diagrams Advanced visualisation of mass
 1115 spectrometry data of complex mixtures. Rapid Commun Mass Spectrom. 31, 658-662.
- Kim, D., et al., 2015. Combination of ring type HPLC separation, ultrahigh-resolution mass
 spectrometry, and high field NMR for comprehensive characterization of crude oil
 compositions. Fuel. 157, 48-55.
- 1119 Kim, S., et al., 2003. Graphical method for analysis of ultrahigh-resolution broadband mass
 1120 spectra of natural organic matter, the van Krevelen diagram. Anal Chem. 75, 5336-44.
- Kondyli, A., Schrader, W., 2019. High-resolution GC/MS studies of a light crude oil fraction. J
 Mass Spectrom. 54, 47-54.
- Kulkarni, S. U., Thies, M. C., 2012. Quantitative analysis of polydisperse systems via solvent-free matrix-assisted laser desorption/ionization time-of-flight mass spectrometry. Rapid
 Commun Mass Spectrom. 26, 392-398.
- Lacroix-Andrivet, O., et al., 2021. Molecular Characterization of Aged Bitumen with Selective and Nonselective Ionization Methods by Fourier Transform Ion Cyclotron Resonance Mass Spectrometry. 2. Statistical Approach on Multiple-Origin Samples. Energy & Fuels.
 35, 16442-16451.
- Lai, A., et al., 2022. The Next Frontier of Environmental Unknowns: Substances of Unknown or
 Variable Composition, Complex Reaction Products, or Biological Materials (UVCBs).
 Environ Sci Technol. 56, 7448-7466.

1133 1134	Lalli, P. M., et al., 2015. Isomeric Separation and Structural Characterization of Acids in Petroleum by Ion Mobility Mass Spectrometry. Energy Fuels. 29, 3626-3633.
1135 1136 1137	Lalli, P. M., et al., 2017. Functional Isomers in Petroleum Emulsion Interfacial Material Revealed by Ion Mobility Mass Spectrometry and Collision-Induced Dissociation. Energy & Fuels. 31, 311-318.
1138 1139	Leefmann, T., et al., 2019. UltraMassExplorer: a browser-based application for the evaluation of high-resolution mass spectrometric data. Rapid Commun Mass Spectrom. 33, 193-202.
1140 1141	Li, S., et al., 2022a. Molecular characteristics of biochar-derived organic matter sub-fractions extracted by ultrasonication. Sci Total Environ. 806, 150190.
1142 1143	Li, Y., et al., 2022b. Comprehensive chemical characterization of dissolved organic matter in typical point-source refinery wastewaters. Chemosphere. 286, 131617.
1144 1145	Liu, D., et al., 2014. Direct hydro-liquefaction of sawdust in petroleum ether and comprehensive bio-oil products analysis. Bioresour Technol. 155, 152-60.
1146 1147	Liu, Y., et al., 2020. The acid and neutral nitrogen compounds characterized by negative ESI Orbitrap MS in a heavy oil before and after oxidation. Fuel. 277.
1148 1149 1150	Lu, H., et al., 2013. Geochemical Explication of Sulfur Organics Characterized by Fourier Transform Ion Cyclotron Resonance Mass Spectrometry on Sulfur-Rich Heavy Oils in Jinxian Sag, Bohai Bay Basin, Northern China. Energy Fuels. 27, 5861-5866.
1151 1152 1153	Luna, N., et al., 2014a. Identification And Characterization Of Sulfur Compounds In Straight-Run Diesel Using Comprehensive Two-Dimensional Gas Chromatography Coupled To Time- Of-Flight Mass Spectrometry. China Pet Process Petrochem. 16, 10-18.
1154 1155 1156	Luna, N., et al., 2014b. Identification and Characterization of Sulfur Compounds in Straight-Run Diesel Using Comprehensive Two-Dimensional GC Coupled with TOF MS. China Petrol Proc Petrochem Techn. 16, 10-18.
1157 1158	Luo, T., et al., 2016. A novel characterization of furfural-extract oil from vacuum gas oil and its application in solvent extraction process. Fuel Process Technol. 152, 356-366.
1159 1160 1161	Lv, H., et al., 2013. Application of UPLC-Quadrupole-TOF-MS Coupled with Recycling Preparative HPLC in Isolation and Preparation of Coumarin Isomers with Similar Polarity from Peucedanum praeruptorum. Chromatographia. 76, 141-148.
1162 1163 1164	MacLennan, M. S., et al., 2018. Characterization of Athabasca lean oil sands and mixed surficial materials: Comparison of capillary electrophoresis/low-resolution mass spectrometry and high-resolution mass spectrometry. Rapid Commun Mass Spectrom. 32, 695-702.
1165 1166 1167 1168	Mahmoud, M. Y., Dabek-Zlotorzynska, E., 2018. Investigation of isomeric structures in a commercial mixture of naphthenic acids using ultrahigh pressure liquid chromatography coupled to hybrid traveling wave ion mobility-time of flight mass spectrometry. J. Chromatogr. A. 1572, 90-99.

- Maillard, J. F., et al., 2021. Structural analysis of petroporphyrins from asphaltene by trapped ion
 mobility coupled with Fourier transform ion cyclotron resonance mass spectrometry.
 Analyst. 146, 4161-4171.
- Mao, D., et al., 2009. Combining HPLC-GCXGC, GCXGC/ToF-MS, and selected ecotoxicity
 assays for detailed monitoring of petroleum hydrocarbon degradation in soil and leaching
 water. Environ Sci Technol. 43, 7651-7.
- Marshall, A. G., Rodgers, R. P., 2004. Petroleomics: the next grand challenge for chemical
 analysis. Acc Chem Res. 37, 53-9.
- Marshall, A. G., Rodgers, R. P., 2008. Petroleomics: chemistry of the underworld. Proc Natl Acad
 Sci U S A. 105, 18090-5.
- McKee, R. H., et al., 2015. Characterization of the toxicological hazards of hydrocarbon solvents.
 Crit Rev Toxicol. 45, 273-365.

McKenna, A. M., et al., 2013. Expansion of the analytical window for oil spill characterization by
 ultrahigh resolution mass spectrometry: beyond gas chromatography. Environ Sci Technol.
 47, 7530-9.

- Mennito, A. S., Qian, K., 2013. Characterization of Heavy Petroleum Saturates by Laser
 Desorption Silver Cationization and Fourier Transform Ion Cyclotron Resonance Mass
 Spectrometry. Energy & Fuels. 27, 7348-7353.
- Miles, S. M., et al., 2020. Oil sands process affected water sourced Trichoderma harzianum
 demonstrates capacity for mycoremediation of naphthenic acid fraction compounds.
 Chemosphere. 258, 127281.
- Muller, H., et al., 2020. Innate Sulfur Compounds as an Internal Standard for Determining
 Vacuum Gas Oil Compositions by APPI FT-ICR MS. Energy Fuels. 34, 8260-8273.
- 1192 National Research Council, 2009. Science and Decisions: Advancing Risk Assessment. National
 1193 Academies Press, Washington, DC.
- Neumann, A., et al., 2021. Investigation of Island/Single-Core- and Archipelago/Multicore Enriched Asphaltenes and Their Solubility Fractions by Thermal Analysis Coupled with
 High-Resolution Fourier Transform Ion Cyclotron Resonance Mass Spectrometry. Energy
 & Fuels. 35, 3808-3824.
- Ngo, H. I., et al., 2012. Improved synthesis and characterization of saturated branched-chain fatty
 acid isomers. Eur J Lipid Sci Technol. 114, 213-221.
- Niyonsaba, E., et al., 2019. Recent Advances in Petroleum Analysis by Mass Spectrometry. Anal
 Chem. 91, 156-177.
- Nolte, T., et al., 2013. Desulfurized Fuels from Athabasca Bitumen and Their Polycyclic Aromatic
 Sulfur Heterocycles. Analysis Based on Capillary Electrophoresis Coupled with TOF MS.
 Energy Fuels. 27, 97-107.

O'Reilly, K. T., et al., 2019. Oxygen-Containing Compounds Identified in Groundwater from Fuel 1205 Release Sites Using GCxGC-TOF-MS. Ground Water Monit R. 39, 32-40. 1206 Oldenburg, T. B. P., et al., 2014. The impact of thermal maturity level on the composition of crude 1207 oils, assessed using ultra-high resolution mass spectrometry. Org Geochem. 75, 151-168. 1208 1209 Oldenburg, T. B. P., et al., 2017. The controls on the composition of biodegraded oils in the deep subsurface – Part 4. Destruction and production of high molecular weight non-hydrocarbon 1210 species and destruction of aromatic hydrocarbons during progressive in-reservoir 1211 biodegradation. Org Geochem. 114, 57-80. 1212 Onel, M., et al., 2019. Grouping of complex substances using analytical chemistry data: A 1213 framework for quantitative evaluation and visualization. PLoS One. 14, e0223517. 1214 Orrego- Ruíz, J. A., 2018. Finding a relationship between the composition and the emulsifying 1215 character of asphaltenes through FTICR-MS. Cienc Tecn Fut. 8, 45-52. 1216 Palacio Lozano, D. C., et al., 2019a. Chapter 32 | Mass Spectrometry in the Petroleum Industry," 1217 in Fuels and Lubricants Handbook: Technology, Properties, Performance, and Testing. 1218 MNL37-2ND-EB Fuels and Lubricants Handbook: Technology, Properties, Performance, 1219 and Testing. 2. 1220 Palacio Lozano, D. C., et al., 2019b. Pushing the analytical limits: new insights into complex 1221 mixtures using mass spectra segments of constant ultrahigh resolving power. Chem Sci. 1222 10, 6966-6978. 1223 Palacio Lozano, D. C., et al., 2022. Revealing the Reactivity of Individual Chemical Entities in 1224 Complex Mixtures: the Chemistry Behind Bio-Oil Upgrading. Anal Chem. 94, 7536-7544. 1225 1226 Palacio Lozano, D. C., et al., 2020. Petroleomics: Tools, Challenges, and Developments. Annu Rev Anal Chem (Palo Alto Calif). 13, 405-430. 1227 Pereira, A. S., et al., 2013. Characterization of oil sands process-affected waters by liquid 1228 chromatography orbitrap mass spectrometry. Environ Sci Technol. 47, 5504-13. 1229 Petroleum HPV Testing Group, HPV Challenge Overview. API, 2017. 1230 1231 Phillips, A. L., et al., 2022. A Framework for Utilizing High-Resolution Mass Spectrometry and Nontargeted Analysis in Rapid Response and Emergency Situations. Environ Toxicol 1232 Chem. 41, 1117-1130. 1233 Porto, C. F. C., et al., 2019. Characterization of organosulfur compounds in asphalt cement 1234 samples by ESI(+)FT-ICR MS and 13C NMR spectroscopy. Fuel. 256, 115923. 1235 Prince, R. C., Walters, C. C., 2022. Modern analytical techniques are improving our ability to 1236

- Qian, K., et al., 2004. The Coupling of Supercritical Fluid Chromatography and Field Ionization
 Time-of-Flight High-Resolution Mass Spectrometry for Rapid and Quantitative Analysis
 of Petroleum Middle Distillates. Eur J Mass Spectrom. 10, 187-196.
- Qian, K., et al., 2001. Reading chemical fine print: Resolution and identification of 3000 nitrogen containing aromatic compounds from a single electrospray ionization Fourier transform
 ion cyclotron resonance mass spectrum of heavy petroleum crude oil. Energy & Fuels. 15,
 492-498.
- Qian, K., Wang, F. C., 2019. Compositional Analysis of Heavy Petroleum Distillates by
 Comprehensive Two-dimensional Gas Chromatography, Field Ionization and High resolution Mass Spectrometry. J Am Soc Mass Spectrom. 30, 2785-2794.
- Rasmussen, K., et al., 1999. Compilation of EINECS: Descriptions and definitions used for UVCB
 substances: Complex reaction products, plant products, (post-reacted) naturally occurring
 substances, micro-organisms, petroleum products, soaps and detergents, and metallic
 compounds. Toxicol Environ Chem. 69, 403-416.
- Reddy, C. M., Quinn, J. G., 1999. GC-MS analysis of total petroleum hydrocarbons and
 polycyclic aromatic hydrocarbons in seawater samples after the North Cape oil spill. Mar
 Pollut Bull. 38, 126-135.
- Redman, A. D., et al., 2014. PETRORISK: a risk assessment framework for petroleum substances.
 Integr Environ Assess Manag. 10, 437-48.
- Redman, A. D., et al., 2012. PETROTOX: an aquatic toxicity model for petroleum substances.
 Environ Toxicol Chem. 31, 2498-506.
- Riches, E., et al., Ion Mobility & PetroOrg Software: Novel Techniques for Petroleomics
 Investigations In: Waters, (Ed.), 2015.
- Ristic, N. D., et al., 2018. Compositional Characterization of Pyrolysis Fuel Oil from Naphtha and
 Vacuum Gas Oil. Energy Fuels. 32, 1276-1286.
- Rocha, Y. D. S., et al., 2018. Geochemical characterization of lacustrine and marine oils from off shore Brazilian sedimentary basins using negative-ion electrospray Fourier transform ion
 cyclotron resonance mass spectrometry (ESI FTICR-MS). Org Geochem. 124, 29-45.
- 1266 Rodgers, R. P., McKenna, A. M., 2011. Petroleum analysis. Anal Chem. 83, 4665-87.
- Rodgers, R. P., et al., 1998. Resolution, elemental composition, and simultaneous monitoring by
 Fourier transform ion cyclotron resonance mass spectrometry of organosulfur species
 before and after diesel fuel processing. Anal Chem. 70, 4743-4750.

Rodrigues Covas, T., et al., 2020. Fractionation of polar compounds from crude oils by hetero medium pressure liquid chromatography (H-MPLC) and molecular characterization by ultrahigh-resolution mass spectrometry. Fuel. 267, 117289.

Roman-Hubers, A. T., et al., 2021. Data Processing Workflow to Identify Structurally Related 1273 Compounds in Petroleum Substances Using Ion Mobility Spectrometry-Mass 1274 Spectrometry. Energy Fuels. 35, 10529-10539. 1275 Roman-Hubers, A. T., et al., 2022. Characterization of Compositional Variability in Petroleum 1276 1277 Substances. Fuel. 317, 123547. Rosa, T. R., et al., 2020. DropMS: Petroleomics Data Treatment Based in Web Server for High-1278 Resolution Mass Spectrometry. J Am Soc Mass Spectrom. 31, 1483-1490. 1279 Rowland, S. J., et al., 2011. Identification of individual acids in a commercial sample of 1280 naphthenic acids from petroleum by two-dimensional comprehensive gas 1281 chromatography/mass spectrometry. Rapid Commun Mass Spectrom. 25, 1741-1751. 1282 Ruger, C. P., et al., 2015. Hyphenation of Thermal Analysis to Ultrahigh-Resolution Mass 1283 Spectrometry (Fourier Transform Ion Cyclotron Resonance Mass Spectrometry) Using 1284 1285 Atmospheric Pressure Chemical Ionization For Studying Composition and Thermal Degradation of Complex Materials. Anal Chem. 87, 6493-9. 1286 Rui, D., et al., 2012. Molecular Characterization of Hydrotreated Atmospheric Residue Derived 1287 from Arabian Heavy Crude by GC FI/FD TOF MS and APPI FT-ICR MS. China Pet 1288 Process Petrochem. 14, 80-88. 1289 1290 Salvito, D., et al., 2020. Improving the Environmental Risk Assessment of Substances of Unknown or Variable Composition, Complex Reaction Products, or Biological Materials. 1291 Environ Toxicol Chem. 39, 2097-2108. 1292 Santos, J. M., et al., 2015. Petroleomics by ion mobility mass spectrometry: resolution and 1293 characterization of contaminants and additives in crude oils and petrofuels. Anal Methods. 1294 7,4450-4463. 1295 Scarlett, A., et al., 2008. Chronic sublethal effects associated with branched alkylbenzenes 1296 1297 bioaccumulated by mussels. Environ Toxicol Chem. 27, 561-567. Sierra Analytics, Composer: State of the art visualization software for petrolemics. 2022. 1298 1299 Silva, R. C., et al., 2020. Mechanistic insights into sulfur rich oil formation, relevant to geological carbon storage routes. A study using (+) APPI FTICR-MS analysis. Org Geochem. 147. 1300 Silva, R. V. S., et al., 2019. Comprehensive study of the liquid products from slow pyrolysis of 1301 1302 crambe seeds: Bio-oil and organic compounds of the aqueous phase. Biomass Bioenerg. 123, 78-88. 1303 Smith, H. M., et al., 1959. Keys to the mystery of crude oil. Proc. Amer. Petrole. Inst. 39, 433-1304 1305 465. Sorensen, L., et al., 2019. Establishing a link between composition and toxicity of offshore 1306 produced waters using comprehensive analysis techniques - A way forward for discharge 1307 monitoring? Sci Total Environ. 694, 133682. 1308

- Spectroswiss, FTMS Data Processing Tools PeakByPeak FTMS Data Analysis. In: Spectroswiss,
 (Ed.), 2019.
- Stenson, A. C., et al., 2003. Exact masses and chemical formulas of individual Suwannee River
 fulvic acids from ultrahigh resolution electrospray ionization Fourier transform ion
 cyclotron resonance mass spectra. Anal Chem. 75, 1275-84.
- Stout, S. A., Wang, Z., 2007. Chemical fingerprinting of spilled or discharged petroleum —
 methods and factors affecting petroleum fingerprints in the environment. in: Wang, Z.,
 Stout, S. A., (Eds.), Oil Spill Environmental Forensics: Fingerprinting And Source
 Identification. Academic Press, Cambridge, MA, pp. 1-53.
- U.S. EPA, Toxic Substance Control Act (TSCA) PL 94-469 Candidate List of Chemical
 Substances Addendum 1 Generic Terms Covering Petroleum Refinery Process Streams.
 US Environmental Protection Agency,, Washignton, D.C., 1978.
- Urban, P. L., 2016. Quantitative mass spectrometry: an overview. Philos Trans A Math Phys Eng
 Sci. 374, 20150382.
- US EPA, Method 8051B: Non-halogenated organics using GC-FID. US Environmental Protection
 Agency, Washington, DC, 1996.
- US EPA, Method 8270E (SW-846): Semivolatile Organic Compounds by Gas Chromatography/
 Mass Spectrometry (GC/MS). US Environmental Protection Agency, Washington, DC,
 2014.
- US EPA, Procedures for Chemical Risk Evaluation Under the Amended Toxic Substances Control
 Act. 82 FR 33726, Vol. 40 CFR 702. US Environmental Protection Agency, Washington,
 DC, 2017.
- 1331 Van Krevelen, D. W., 1950. Graphical-statistical method for the study of structure and reaction
 1332 processes of coal. Fuel. 29, 269-284.
- 1333 Van Krevelen, D. W., 1984. Organic geochemistry old and new. Org Geochem. 6, 1-10.
- Vanini, G., et al., 2020. Characterization of nonvolatile polar compounds from Brazilian oils by
 electrospray ionization with FT-ICR MS and Orbitrap-MS. Fuel. 282, 118790.
- 1336 Ventura, G. T., et al., 2011. Analysis of petroleum compositional similarity using multiway
 principal components analysis (MPCA) with comprehensive two-dimensional gas
 chromatographic data. J Chromatogr A. 1218, 2584-92.
- 1339 Vozka, P., et al., 2019. Jet fuel density via GC x GC-FID. Fuel. 235, 1052-1060.
- Walters, C. C., et al., 2015. Petroleum alteration by thermochemical sulfate reduction A
 comprehensive molecular study of aromatic hydrocarbons and polar compounds. Geochim
 Cosmochim Acta. 153, 37-71.

- Wang, X., et al., 2020. Characterization of wastewater effluent organic matter with different solid
 phase extraction sorbents. Chemosphere. 257, 127235.
- Wang, Z., et al., 1994. Fractionation of a Light Crude Oil and Identification and Quantitation of
 Aliphatic, Aromatic, and Biomarker Compounds by GC-FID and GC-MS, Part I. J
 Chromatogr Sci. 32, 361-366.
- Wang, Z., et al., 2011. Forensic fingerprinting and source identification of the 2009 Sarnia
 (Ontario) oil spill. J Environ Monit. 13, 3004-17.
- Wang, Z. D., Fingas, M., 2003. Fate and identification of spilled oils and petroleum products in
 the environment by GC-MS and GC-FID. Energy Sources. 25, 491-508.
- Weng, N., et al., 2015. Insight into unresolved complex mixtures of aromatic hydrocarbons in
 heavy oil via two-dimensional gas chromatography coupled with time-of-flight mass
 spectrometry analysis. J Chromatogr A. 1398, 94-107.
- Wise, S. A., et al., 2022. Advances in Chemical Analysis of Oil Spills Since the Deepwater
 Horizon Disaster. Crit Rev Anal Chem. 1-60.
- Wozniak, A. S., et al., 2019. Rapid Degradation of Oil in Mesocosm Simulations of Marine Oil
 Snow Events. Environ Sci Technol. 53, 3441-3450.
- Xia, Y., et al., 2021. Characterization of nitrogen-containing compounds in petroleum fractions by
 online reversed-phase liquid chromatography-electrospray ionization Orbitrap mass
 spectrometry. Fuel. 284, 119035.
- 1362 Xian, F., et al., 2012. High resolution mass spectrometry. Anal Chem. 84, 708-19.
- Yang, L., et al., 2019. Gas chromatography-Orbitrap mass spectrometry screening of organic
 chemicals in fly ash samples from industrial sources and implications for understanding
 the formation mechanisms of unintentional persistent organic pollutants. Sci Total
 Environ. 664, 107-115.
- Zhong, J. Y., et al., 2011. Combining advanced NMR techniques with ultrahigh resolution mass
 spectrometry: A new strategy for molecular scale characterization of macromolecular
 components of soil and sedimentary organic matter. Org Geochem. 42, 903-916.
- 1370 Zhu, G. Y., et al., 2020. Discovery and Molecular Characterization of Organic Caged Compounds
 1371 and Polysulfanes in Zhongba81 Crude Oil, Sichuan Basin, China. Energy Fuels. 34, 6811 1372 6821.
- 1373

Figure 1. A timeline of major developments in the fields of analytical chemistry and data analysis
of petroleum UVCB, and the concomitant evolution of the regulatory frameworks for registration
and hazard classification of these substances See abbreviations in the text.



OAnalytical Methods OData Processing/Visualization OLegistaion/Regulations

Figure 2. Literature review of the major analytical methods and their application for the analysis of petroleum substances. (**A**) A dendrogram of the major searches. The numbers indicate the quantity of publications for each search. See Supplemental Table 1 for information on the exact search terms and hyperlinks to the publications. (**B**) Cumulative histograms indicating the number of publications across time through December 2021. Colors correspond to the methods indicated in the inset.



Figure 3. Example visualizations commonly used to represent hydrocarbon composition of 1389 petroleum substances analyzed using high resolution MS techniques. (A) A Kendrick mass defect 1390 (KMD) plot demonstrating repetitive patterns of CH₂-containing molecules in a petroleum sample. 1391 (B) A DBE vs carbon number plot indicating the relative proportions of molecules varying by their 1392 degree of aromaticity and carbon number in a sample. (C) van Krevelen plot display the degree of 1393 oxidation by plotting the H/C versus O/C ratio in a sample. (D) A stacked bar plot showing relative 1394 proportions of constituents from various chemical groups. (E) A plot of relative amounts of various 1395 hydrocarbon blocks in a sample. 1396



Figure 4. The use of various novel MS methods to address specific regulatory needs identified in 1398 this review. (A) Scholarly publications that were identified as relevant to each regulatory 1399 need/question (identified by colors). A total number of publications identified by a literature search 1400 is listed in the first circle (see Supplemental Table 2 for details). Upon examination of each study's 1401 content, a number of publications were deemed not relevant (red circles); the remaining studies are 1402 shown in green circles. (B) The number of relevant publications as a function of the high-/ultra-high 1403 MS technique. Top left, a stacked bar graph indicating the number of publications as they pertain to 1404 each regulatory need/question in (A). Remaining stacked bar plots show the number of studies that 1405 used various separation (HPLC, GC, GC×GC or IMMS) or direct injection with each MS technique. 1406 1407

