# Comprehensive Chemical Characterization of Hydrocarbons in NIST Standard Reference Material 2779 Gulf of Mexico Crude Oil 

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## (S) Supporting Information


#### Abstract

Comprehensive chemical information is needed to understand the environmental fate and impact of hydrocarbons released during oil spills. However, chemical information remains incomplete because of the limitations of current analytical techniques and the inherent chemical complexity of crude oils. In this work, gas chromatography (GC)-amenable $\mathrm{C}_{9}-\mathrm{C}_{33}$ hydrocarbons were comprehensively characterized from the National Institute of Standards and Technology Standard Reference Material (NIST SRM) 2779 Gulf of Mexico crude oil by GC coupled to vacuum ultraviolet photoionization mass spectrometry (GC/VUV-MS), with a mass balance of $68 \pm 22 \%$. This technique overcomes one important limitation faced by  traditional GC and even comprehensive 2D gas chromatography ( $\mathrm{GC} \times \mathrm{GC}$ ): the necessity for individual compounds to be chromatographically resolved from one another in order to be characterized. VUV photoionization minimizes fragmentation of the molecular ions, facilitating the characterization of the observed hydrocarbons as a function of molecular weight (carbon number, $N_{\mathrm{C}}$ ), structure (number of double bond equivalents, $N_{\text {DBE }}$ ), and mass fraction ( $\mathrm{mg} \mathrm{kg}^{-1}$ ), which represent important metrics for understanding their fate and environmental impacts. Linear alkanes ( $8 \pm 1 \%$ ), branched alkanes ( $11 \pm 2 \%$ ), and cycloalkanes ( $37 \pm 12 \%$ ) dominated the mass with the largest contribution from cycloalkanes containing one or two rings and one or more alkyl side chains ( $27 \pm 9 \%$ ). Linearity and good agreement with previous work for a subset of $>100$ components and for the sum of compound classes provided confidence in our measurements and represents the first independent assessment of our analytical approach and calibration methodology. Another crude oil collected from the Marlin platform ( 35 km northeast of the Macondo well) was shown to be chemically identical within experimental errors to NIST SRM 2779, demonstrating that Marlin crude is an appropriate surrogate oil for researchers conducting laboratory research into impacts of the DeepWater Horizon disaster.


## INTRODUCTION

In the three months following the explosion and loss of the Deepwater Horizon (DWH) drilling platform on April 20, 2010, an estimated 5 million barrels of sweet light crude oil were released from the Macondo well located in the Mississippi Canyon lease block 252 (MC252) into the Gulf of Mexico. ${ }^{1,2}$ The oil ascended from a depth of 1.5 km to the sea surface, forming an oil slick that eventually impacted more than 1000 km of coastline. ${ }^{3}$ The released oil was a complex mixture of many thousands of different organic compounds with different molecular structures. The environmental fate from different weathering pathways, i.e., evaporation, dissolution, photooxidation, and biodegradation, are dependent on their chemical and physical properties, which are influenced by molecular structure. ${ }^{4,5}$ A considerable number of oxygenated products were formed during some of these weathering processes, many of which are believed to be from saturated precursors, although the exact species remain largely unknown. ${ }^{6,7}$ Molecular structure also substantially influences the oxidation chemistry and secondary organic aerosol (SOA) yields of
the volatile components that evaporate from the surface slicks. ${ }^{8-11}$ Therefore, comprehensive information on the chemical composition of the released hydrocarbons is essential for evaluating the fates and impacts of the emitted species in the environment. Additionally, without a better accounting of the oil composition, especially that of the persistent components of weathered oil, it is challenging to determine what oxygenated products could form and what their potential impacts on marine and coastal ecosystems might be.

Traditionally, gas chromatography (GC) coupled to flame ionization detection (GC-FID) or mass spectrometry (GC-MS) has been used to chemically characterize crude oils. ${ }^{12,13}$ These chromatograms typically exhibited a large, raised baseline often referred to as the unresolved complex mixture (UCM), ${ }^{4,14,15}$ composed of

[^0]many thousands of constitutional isomers of aliphatic hydrocarbons that are difficult or impossible to separate using conventional GC-based techniques. ${ }^{16}$ Additionally, the significant fragmentation observed upon electron impact (EI) ionization yields many smaller fragments, which prevents the determination of molecular mass. The analytical challenge of deciphering the complexity of the UCM has meant that historically the majority of chemical analysis of crude oil have focused on components that can be distinguished from the bulk of the mass in the UCM typically $n$-alkanes, polycyclic aromatic hydrocarbons (PAHs), and sterane and hopane biomarkers. These components make up $<20 \%$ of the total mass of the Macondo crude oil. ${ }^{17}$

More recently, comprehensive 2D gas chromatography (GC×GC) has been successfully used to identify and quantify many more individual compounds and classes of compounds in fresh and weathered crude oils. ${ }^{4,5,14,18}$ However, with the number of possible constitutional isomers increasing exponentially with increasing carbon number, ${ }^{19}$ it becomes increasingly difficult to resolve individual species even with high-resolution capillary GC $\times$ GC. Previously, this limitation has prevented a comprehensive understanding of the mass distribution of all compounds that are present. Determining the mass contribution of every constitutional isomer is extremely challenging and probably an unnecessary task because this information is likely to be too detailed for current models. However, being able to characterize the total mass distribution as a function of volatility and structure would provide useful constraints for environmental modeling of the released hydrocarbons.

GC with vacuum ultraviolet ionization mass spectrometry (GC/VUV-MS) overcomes the limitations of isomer resolution in GC for complex mixtures. VUV photoionization results in substantially reduced fragmentation of the molecular ion, which facilitates the classification of hydrocarbon compounds by carbon number $\left(N_{\mathrm{C}}\right)$, number of double-bond equivalents in their structure ( $N_{\text {DBE }}$ ), and degree of branching ${ }^{20}$ determined from their molecular weight and GC retention times. The advantage of this technique relative to $\mathrm{GC} \times \mathrm{GC}$ is that individual compounds do not necessarily need to be chromatographically resolved from one another to be characterized. Additionally, compound classes, e.g., alkanes and cycloalkanes, are unambiguously separated from one another as a result of molecular mass differences, and data processing times are reduced because individual peak assignments are not necessary. As a result, this technique provides a more comprehensive methodology than was previously possible, especially for species with more than 10 carbon atoms. This analytical methodology has previously been shown to provide a more comprehensive approach for the characterization of complex organic mixtures of petrochemical products derived from crude oil. ${ }^{10,20,21}$ The main objectives of this work are to use these advanced techniques and classification schemes to provide a more comprehensive characterization of the insoluble semivolatile hydrocarbons released during the DWH disaster and to provide a comprehensive assessment of the chemical composition of National Institute of Science and Technology (NIST) Standard Reference Material (SRM) 2779 Gulf of Mexico crude oil for the research community to utilize.

## EXPERIMENTAL METHODOLOGY

Description of Oil Samples. Analysis of two crude oil samples are presented in this work: NIST SRM 2779 Gulf of Mexico Crude Oil and surrogate oil (SURR). Both these crude oils are being widely used by the research community. SURR is being provided to supplement the high demand and limited supply
of Macondo well crude oil. The NIST SRM 2779 crude oil was collected on May 21, 2010 on the drillship Discoverer Enterprise from the insertion tube receiving oil directly from the Macondo well during response operations. The water was separated from the oil, and the resulting oil was homogenized before being transferred into 2 mL amber glass ampules. Certified mass fractions of 21 PAHs and reference mass fractions for an additional 22 PAHs and 31 alkylated PAHs were reported by NIST (available online at https://www-s.nist.gov/srmors/certificates/2779.pdf). Certified mass fractions were weighted means obtained via 2-5 different analytical methods at NIST and represent values with the highest level of confidence in their accuracy, in that all known or suspected sources of bias have been investigated or taken into account. Reference values for the additional 22 PAHs were weighted means obtained via 2-5 different analytical methods at NIST. The reference mass fractions of the 31 alkylated PAHs were the means of results obtained using one analytical technique from an interlaboratory intercomparison exercise of 24 laboratories, although not all returned data for each analyte. The reference mass fractions are noncertified values that are estimates of the true value. However, these values do not meet NIST's criteria for certification and were provided with associated uncertainties that may reflect only measurement precision, may not include all sources of uncertainty, or may reflect a lack of sufficient statistical agreement among multiple analytical methods. SURR, classified as a dead oil because it does not contain any dissolved gases, was collected from the Marlin Platform of the Dorado field, 35 km northeast of the Macondo well. The oil was stored in 350 gallons tanks that were filled on site. The SURR oil was made available to Gulf of Mexico Research Initiative (GoMRI)-funded researchers by BP and was obtained from BP's Knox Storage Facility in Denver, Colorado, by submitting a sample request to Architecture, Engineering, Consulting, Operations, and Maintenance (AECOM), the contracted managers of the facility.

Analytical Methods. Samples were analyzed using GC/ VUV-HRTOFMS. Oil samples were diluted ( $50: 1$ ) in chloroform (HPLC grade, Sigma-Aldrich), directly injected via a septumless inlet into a liquid nitrogen cooled inlet (CIS4, Gerstel Inc.), and cyrofocused at $-25^{\circ} \mathrm{C}$ on a quartz wool packed inlet liner. Analytes were transferred to the gas chromatograph (GC, Agilent 7890 ) by rapid heating of the CIS to $300^{\circ} \mathrm{C}$ under a helium flow. Analytes were separated on a nonpolar primary column ( $60 \mathrm{~m} \times$ $0.25 \mathrm{~mm} \times 250 \mu \mathrm{~m}$ Rxi-5Sil-MS; Restek) with a carrier gas flow rate of $2 \mathrm{~mL} \mathrm{~min}{ }^{-1}$ of helium. The GC temperature program was $40^{\circ} \mathrm{C}$ hold for $2 \mathrm{~min}, 3.5^{\circ} \mathrm{C} \mathrm{min}{ }^{-1}$ until $320^{\circ} \mathrm{C}$, and hold for 10 min . Analytes were ionized with 10.5 electron volts $(\mathrm{eV})$ vacuum ultraviolet photoionization (VUV) using a high-resolution ( $m / \Delta m \approx 4000$ ) time-of-flight mass spectrometer (HTOF, Tofwerk). The frequency of data collection on the mass spectrometer was 200 Hz , which was averaged to 0.5 Hz to improve identification and deconvolution of high-resolution mass spectral peaks. The transfer line from the GC to the TOF was maintained at $270^{\circ} \mathrm{C}$. To minimize fragmentation in VUV, the ion source was operated at $170{ }^{\circ} \mathrm{C}$. The VUV beam (tunable from $8-30 \mathrm{eV}$ but typically used at $10.5 \pm 0.2$ or $9.0 \pm 0.2 \mathrm{eV}$ and photon flux $\approx 10^{15}$ photons $\mathrm{s}^{-1}$ ) was generated by the Chemical Dynamics Beamline 9.0.2 of the Advanced Light Source (ALS) at Lawrence Berkeley National Laboratory. In our group, we have been pioneering the use of synchrotron radiation as the source of VUV photons because of the high photon flux. However, we acknowledge that this limits the accessibility of this technique beyond our research group. However, it should be noted that the first application of VUV ionization utilized a novel excimer VUV light source ${ }^{22}$ and


Figure 1. (a) Mass fraction distribution of the NIST SRM 2779 Gulf of Mexico crude oil as a function of carbon number ( $N_{\mathrm{C}}$ ) and the number of doublebond equivalents $\left(N_{\text {DBE }}\right) . N_{\text {DBE }}=0$ and 1 are separated into straight and branched. For simplicity, all branched alkanes are colored red, and all compounds with $N_{\text {DBE }} \geq 7$ are colored black. The data is provided in Table 1. The contribution of each compound class to the total mass for all observed carbon numbers $\left(\mathrm{C}_{9}-\mathrm{C}_{33}\right)$ is also shown. (b) Mass fraction distribution of the branched components of $N_{\text {DBE }}=0$ (red bars from a) separated by number of methyl groups (B1-6) observed on the carbon chain. (c) Mass fraction distribution of the polycyclic aromatic hydrocarbon (PAH) and thiophene components with $N_{\text {DBE }} \geq 7$ (black bars from a) separated by $N_{\text {DBE }}$. This data is provided in Table S1.
that alternative soft ionization technologies are also now emerging, e.g., a variable electronvolt ion source (Markes International Ltd. UK) and field-ionization mass spectrometry (JEOL USA Inc.). In particular, these recent instrumental developments will likely lead to much wider use of our analytical methodology within the research community.

Because the ionization energies of most organic compounds are between 8 and 11 eV , the minimal excess energy of 10.5 eV VUV photons (compared to EI at 70 eV ) limits fragmentation of ionized molecules, allowing for significant detection of the molecular ions. ${ }^{20}$ In this work, as in previous work, the molecular ion mass and chromatographic retention time were used to classify compounds comprising the UCM by $N_{\mathrm{C}}, N_{\text {DBE }}$, and degree of branching. ${ }^{20,21}$ High-mass-resolution data processing was employed to distinguish hydrocarbons from any GC-amenable oxygen containing molecular ions. ${ }^{11}$ All data processing and visualization of data were carried out using custom code written in Igor 6.3.6 (Wavemetrics) adapted from high-resolution aerosol mass spectrometer data analysis code. ${ }^{24}$

Calibration Methodology. The calibration methodology used in this work was based on the method described in Isaacman et al..$^{20}$ and Chan et al. ${ }^{25}$ In brief, the molecular ion signals for linear, branched, cyclic, and aromatic hydrocarbons under VUV ionization are used as the basis for quantification. The sensitivity of the molecular ion of any given compound is a function of its thermal transfer efficiency, ionization efficiency, and degree of fragmentation. Owing to ionization efficiency and extent of fragmentation, the relative molecular ion signal increases with increasing $N_{\text {DBE }}$ because of reduced fragmentation. Authentic
standards of more than 80 compounds were used for calibration; these included $n$-alkanes, branched alkanes, $n$-alkyl cyclohexanes, $n$-alkyl benzenes, hopanes, and steranes, PAHs, and alkylated PAHs. These species were selected to span both the $N_{\mathrm{C}}$ and $N_{\text {DBE }}$ ranges of the crude oil samples. Because compounds with very similar structures fragment almost identically after undergoing photoionization by VUV, they yield essentially the same massresponse calibration curves meaning that mass response is strongly a function of structure and less dependent on carbon number. This resulted in $N_{\text {DBE }}$-specific calibration curves for $N_{\text {DBE }}=0,1,4$, and 7+ because of the limited availability of authentic standards for the other $N_{\text {DBE }}$ 's. The dependence of fragmentation on $N_{\text {DBE }}$ was previously characterized using a standard mixture consisting of isomers of the same carbon number but different $N_{\text {DBE }},{ }^{\prime 25}$, and this was used to interpolate for $N_{\text {DBE }}$ 's for which very limited authentic standards are available.

The thermal transfer efficiency is not linear with carbon number because early eluting components and late eluting components show lower responses relative to components eluting between them as a result of challenges in efficiently transferring all the high- and low-volatility material. A series of perdeuterated $n$-alkanes ( $n$-octane, $n$-decane, $n$-dodecane, $n$-tetradecane, $n$-hexadecane, $n$-octadecane, $n$-eicosane, $n$-docosane, $n$-tetracosane, $n$-hexacosane, $n$-octacosane, $n$-triacontane, $n$-dotriacontane, and $n$-tetratriacontane; $\mathrm{C} / \mathrm{D} / \mathrm{N}$ isotopes) added as an internal standard to all samples were used to generate a relationship between thermal transfer and retention time. This relationship was used for all species because the nature of the transfer efficiency is related to volatility and is not chemically specific. These were
Table 1. Mass Fractions ( $\mathrm{mg} \mathrm{kg}^{-1}$ ) of Components Present in the NIST SRM 2779 Gulf of Mexico Crude Oil ${ }^{a}$

|  | $N_{\mathrm{DBE}}$ |  |  |  |  |  |  |  |  |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | 0 |  |  |  |  |  |  | 1 |  | 2 | 3 | 4 | 5 | 6 | 7+ |
| $N_{\text {C }}$ | B0 | B1 | B2 | B3 | B4 | B5 | B6 | B0 | B |  |  |  |  |  |  |
| 9 | $8600 \pm 1200$ | $13200 \pm 1830$ | $9910 \pm 1377$ |  |  |  |  | $4240 \pm 590$ | $25700 \pm 8490$ | $3070 \pm 1020$ |  | $17300 \pm 5210$ | $144 \pm 53$ |  | $21 \pm 20$ |
| 10 | $6790 \pm 944$ | $7920 \pm 1100$ | $7300 \pm 1014$ |  |  |  |  | $2820 \pm 391$ | $17600 \pm 5810$ | $7580 \pm 1740$ | $177 \pm 113$ | $7030 \pm 2320$ | $9270 \pm 305$ |  | $942 \pm 160$ |
| 11 | $6390 \pm 888$ | $4930 \pm 685$ | $5280 \pm 733$ |  |  |  |  | $2000 \pm 277$ | $13700 \pm 4520$ | $10300 \pm 3420$ | $690 \pm 292$ | $3660 \pm 1210$ | $1740 \pm 656$ |  | $2650 \pm 895$ |
| 12 | $5700 \pm 792$ | $3930 \pm 545$ | $3050 \pm 424$ |  |  |  |  | $1550 \pm 252$ | $10820 \pm 3580$ | $6810 \pm 2250$ | $1210 \pm 435$ | $2420 \pm 800$ | $1960 \pm 649$ | $119 \pm 60$ | $4730 \pm 1510$ |
| 13 | $5090 \pm 707$ | $2960 \pm 411$ | $3360 \pm 466$ |  |  |  |  | $1300 \pm 245$ | $9640 \pm 3190$ | $6070 \pm 2010$ | $1610 \pm 544$ | $1660 \pm 619$ | $2060 \pm 680$ | $493 \pm 164$ | $4720 \pm 1620$ |
| 14 | $4620 \pm 642$ | $2550 \pm 354$ | $2050 \pm 284$ | $2260 \pm 314$ |  |  |  | $1100 \pm 237$ | $8510 \pm 2810$ | $5470 \pm 1810$ | $1990 \pm 656$ | $1560 \pm 573$ | $1800 \pm 595$ | $840 \pm 273$ | $3310 \pm 1890$ |
| 15 | $4530 \pm 629$ | $2290 \pm 318$ | $1610 \pm 253$ | $1910 \pm 265$ |  |  |  | $996 \pm 231$ | $7720 \pm 2550$ | $5110 \pm 1690$ | $2150 \pm 710$ | $1600 \pm 588$ | $1490 \pm 538$ | $979 \pm 324$ | $4310 \pm 1810$ |
| 16 | $4290 \pm 596$ | $2090 \pm 291$ | $1440 \pm 249$ | $2170 \pm 302$ |  |  |  | $871 \pm 223$ | $7260 \pm 2400$ | $4640 \pm 1530$ | $2190 \pm 724$ | $1650 \pm 613$ | $1280 \pm 445$ | $1020 \pm 338$ | $3840 \pm 1600$ |
| 17 | $4010 \pm 557$ | $1730 \pm 257$ | $1030 \pm 233$ | $964 \pm 229$ |  |  |  | $753 \pm 213$ | $6690 \pm 2210$ | $4320 \pm 1430$ | $2150 \pm 710$ | $1640 \pm 608$ | $1220 \pm 422$ | $992 \pm 329$ | $3510 \pm 1450$ |
| 18 | $3490 \pm 485$ | $1510 \pm 251$ | $645 \pm 201$ | $1890 \pm 262$ |  |  |  | $658 \pm 202$ | $6350 \pm 2100$ | $4310 \pm 1430$ | $2150 \pm 709$ | $1620 \pm 601$ | $1240 \pm 428$ | $933 \pm 307$ | $3580 \pm 1560$ |
| 19 | $3380 \pm 469$ | $1370 \pm 247$ | $654 \pm 202$ | $695 \pm 206$ | $2550 \pm 354$ |  |  | $622 \pm 198$ | $5940 \pm 1960$ | $4260 \pm 1410$ | $2180 \pm 720$ | $1750 \pm 662$ | $1240 \pm 428$ | $905 \pm 297$ | $3610 \pm 1600$ |
| 20 | $3240 \pm 451$ | $1490 \pm 250$ | $712 \pm 208$ | $516 \pm 182$ | $1790 \pm 258$ |  |  | $578 \pm 191$ | $5630 \pm 1860$ | $4190 \pm 1380$ | $2120 \pm 701$ | $1640 \pm 610$ | $1250 \pm 431$ | $863 \pm 282$ | $3190 \pm 1400$ |
| 21 | $3050 \pm 424$ | $1130 \pm 238$ | $376 \pm 155$ | $307 \pm 138$ | $752 \pm 212$ |  |  | $534 \pm 185$ | $5350 \pm 1770$ | $4110 \pm 1360$ | $2070 \pm 684$ | $1770 \pm 674$ | $1240 \pm 429$ | $870 \pm 284$ | $2830 \pm 1170$ |
| 22 | $2830 \pm 394$ | $1040 \pm 234$ | $241 \pm 121$ | $205 \pm 109$ | $535 \pm 185$ |  |  | $432 \pm 167$ | $5080 \pm 1680$ | $3610 \pm 1190$ | $1960 \pm 648$ | $1720 \pm 649$ | $1160 \pm 395$ | $818 \pm 266$ | $2460 \pm 968$ |
| 23 | $2570 \pm 357$ | $887 \pm 224$ | $178 \pm 100$ |  | $633 \pm 199$ |  |  | $379 \pm 156$ | $4620 \pm 1530$ | $3260 \pm 1080$ | $1880 \pm 620$ | $1510 \pm 549$ | $1080 \pm 365$ | $795 \pm 258$ | $2140 \pm 866$ |
| 24 | $2450 \pm 340$ | $699 \pm 207$ | $77 \pm 54$ |  | $285 \pm 133$ | $467 \pm 174$ |  | $319 \pm 142$ | $4150 \pm 1370$ | $3030 \pm 1000$ | $1810 \pm 598$ | $1410 \pm 501$ | $1050 \pm 353$ | $778 \pm 252$ | $1420 \pm 620$ |
| 25 | $2220 \pm 308$ | $693 \pm 206$ | $89 \pm 62$ |  | $142 \pm 86$ | $481 \pm 176$ |  | $292 \pm 135$ | $3810 \pm 1260$ | $2810 \pm 929$ | $1770 \pm 590$ | $1350 \pm 474$ | $1020 \pm 338$ | $772 \pm 250$ | $1080 \pm 487$ |
| 26 | $2010 \pm 280$ | $585 \pm 192$ |  |  |  | $305 \pm 139$ |  | $251 \pm 124$ | $3460 \pm 1150$ | $2640 \pm 872$ | $1680 \pm 565$ | $1340 \pm 473$ | $982 \pm 325$ | $784 \pm 254$ | $869 \pm 346$ |
| 27 | $1690 \pm 255$ | $434 \pm 168$ |  |  |  | $221 \pm 114$ |  | $205 \pm 109$ | $3070 \pm 1020$ | $2370 \pm 784$ | $1660 \pm 559$ | $2080 \pm 688$ | $1130 \pm 383$ | $796 \pm 258$ | $133 \pm 65$ |
| 28 | $1430 \pm 248$ | $298 \pm 137$ |  |  |  | $326 \pm 144$ |  | $154 \pm 91$ | $2630 \pm 869$ | $2130 \pm 703$ | $1550 \pm 528$ | $2110 \pm 697$ | $1030 \pm 345$ | $780 \pm 253$ | $57 \pm 36$ |
| 29 | $1350 \pm 246$ | $372 \pm 155$ |  |  |  | $136 \pm 84$ | $252 \pm 124$ | $203 \pm 108$ | $2380 \pm 785$ | $1940 \pm 640$ | $1510 \pm 518$ | $2270 \pm 751$ | $1180 \pm 382$ | $718 \pm 232$ |  |
| 30 | $1180 \pm 240$ | $325 \pm 144$ |  |  |  | $34 \pm 30$ | $254 \pm 124$ | $43 \pm 36$ | $2260 \pm 746$ | $1860 \pm 616$ | $1450 \pm 501$ | $1690 \pm 633$ | $1500 \pm 543$ | $748 \pm 242$ |  |
| 31 | $1060 \pm 235$ | $260 \pm 126$ |  |  |  |  | $197 \pm 106$ | $69 \pm 52$ | $2160 \pm 714$ | $1770 \pm 589$ | $1250 \pm 446$ | $1260 \pm 438$ | $1160 \pm 396$ | $767 \pm 248$ |  |
| 32 | $830 \pm 219$ | $196 \pm 106$ |  |  |  |  | $136 \pm 83$ |  | $2280 \pm 753$ | $1820 \pm 601$ | $1560 \pm 532$ | $1300 \pm 454$ | $1190 \pm 409$ | $811 \pm 263$ |  |
| 33 | $660 \pm 202$ | $108 \pm 71$ |  |  |  |  | $107 \pm 71$ |  | $2490 \pm 821$ | $2310 \pm 764$ | $1670 \pm 563$ | $1170 \pm 399$ | $1190 \pm 407$ | $746 \pm 242$ |  |
|  | mponents | parated by | on | $\mathrm{N}_{\mathrm{C}}$ ) fo | $\mathrm{C}_{3}$ | es, number of | of double bo | nd equiva | ( $N_{\text {DBE }}$ ), and | r |  | ranching (B0 | = straight; B | all bran | d, B1 = | ${ }^{a}$ Components are separated by carbon number $\left(N_{\mathrm{C}}\right)$ for $\mathrm{C}_{9}-\mathrm{C}_{33}$ species, number of double bond equivalents $\left(N_{\mathrm{DBE}}\right)$, and, where possible, degree of branching ( $\mathrm{B} 0=$ straight; $\mathrm{B}=$ all branched, $\mathrm{B} 1=$ one

methyl group; $\mathrm{B} 2=$ two methyl groups, etc). The contributions of higher $N_{\mathrm{DBE}}$ components $\left(N_{\mathrm{DBE}}=7-15\right)$ to $N_{\mathrm{DBE}}=7+$ total are given in Table S 1 .
selected to span the volatility range of the analytes of interest. Because aliphatics including the perdeuterated $n$-alkanes were not ionized at 9 eV , perdeuterated polycyclic aromatic hydrocarbons (phenanthrene, pyrene, chrysene, perylene, dibenzo $[a, h]$ anthracene; $\mathrm{C} / \mathrm{D} / \mathrm{N}$ isotopes) were used to correct for thermal transfer efficiency in the 9 eV runs. The perdeuterated $n$-alkanes were also used to postcalibrate the high-resolution mass spectral data. However, high-resolution peak fitting was not conducted at 9 eV because mass calibration was not possible as a result of an insufficient number of ions spanning a wide enough mass range to generate a robust relationship between exact mass and time-offlight. This is a result of the much lower fragmentation of the PAHs relative to the $n$-alkanes. For these runs the nominal mass data was used. A comparison between the nominal and high-resolution mass data for the samples collected at 10.5 eV yielded very similar results.

Total analytical uncertainty includes contributions from transfer efficiency, structural differences in fragmentation within a $N_{\text {DBE }}$ class, and uncertainties in calibration curves. The uncertainty in calibrating response to mass, determined from calibration curves of authentic standards, was structurally and massdependent with larger uncertainties for lower $N_{\text {DBE }}$ species and smaller mass fractions (Figure S1). In general, the total analytical uncertainty was $<40 \%$ for all species at mass fractions above $1000 \mathrm{mg} \mathrm{kg}^{-1}$, increasing to $<75 \%$ at mass fractions of $100 \mathrm{mg} \mathrm{kg}^{-1}$ (Table S2). Periodic system blanks were analyzed, and these showed that background levels were negligible relative to observed levels of analytes in the samples. Estimated detection limits were $<10$ and $<1 \mathrm{mg} \mathrm{kg}^{-1}$ for aliphatic and aromatic species, respectively, and analytical precision was $<10 \%$ for all species as determined from replicate analysis of the NIST SRM 2779 crude oil.

## RESULTS

Comprehensive Mass Fraction Distributions of NIST SRM 2779 Oil Components. Figure 1a shows the comprehensive mass fraction ( $\mathrm{mg} \mathrm{kg}^{-1}$ ) distribution of the NIST SRM 2779 Gulf of Mexico crude oil for components with 9 to 33 carbon atoms $\left(N_{\mathrm{C}}=\mathrm{C}_{9}-\mathrm{C}_{33}\right)$ separated by the $N_{\text {DBE }}$. Contributions of branched and straight compounds to $N_{\text {DBE }}=$ 0 and 1 are also shown because they are clearly distinguishable in the chromatograms (Figure S2). The total contribution of each $N_{\text {DBE }}$ class to the total mass divided by the total mass injected (mass balance) is also shown in Figure 1a as a percentage for all $\mathrm{C}_{9}-\mathrm{C}_{33}$ compounds observed. For clarity, all branched compounds for $N_{\text {DBE }}=0$ and 1 are shown as a sum. The mass distributions of the branched components of $N_{\text {DBE }}=0$ separated by the number of methyl groups observed on the carbon chain are shown in Figure 1b. Again, for clarity, all components with $N_{\text {DBE }} \geq 7\left(N_{\text {DBE }}=7+\right)$ are presented as a sum, and the contribution of each $N_{\text {DBE }}$ class from 7 to 15 is shown in Figure 1c. The data used to construct Figure 1 are shown in Tables 1 and S1, along with the total analytical uncertainty.

The lower $N_{\text {DBE }}$ components dominate with contributions from $N_{\text {DBE }}=0,1,2$, and 3 making up approximately $50 \%$ of the mass of the oil (Figure 1a). These $N_{\text {DBE }}$ classes represent straight and branched acyclic alkanes and mono-, bi- and tricyclic alkanes. From the complexity of the observed parent ion chromatograms and knowledge of the $N_{\mathrm{C}}$ and $N_{\text {DBE }}$, we infer that the vast majority of the cycloalkane mass is dominated by those with one or more alkyl side chains on the ring or rings. This is supported by the GC retention time data, which are consistent with the literature for where branched alkyl cyclohexanes elute relative to
the $n$-alkyl cyclohexane isomers. ${ }^{21}$ The majority of these compounds have not previously been reported. The straight-chain alkanes and the branched alkanes both contribute approximately similar fractions of $\sim 10 \%$. There are substantially fewer alkane isomers than are statistically possible, and the observed isomer distributions are consistent with previous work on diesel fuel and crude oil that suggest branched alkanes in these fuels are primarily comprised of methylated isomers. ${ }^{5,26,27}$ On average, methyl alkanes (B1) comprised about half of the branched alkane mass with the majority of the remaining mass being comprised of the most branched isoprenoid alkanes, e.g., for $N_{\mathrm{C}}=18$ this is 3 methyl groups (B3), but for $N_{\mathrm{C}}=19$, this is 4 methyl groups (B4) (Figure S3). This is consistent with previous work on diesel fuel, ${ }^{10}$ but the observation in crude oil implies that it is not a result of the refining process and is related to the formation from the concatenation of isoprene $\left(\mathrm{C}_{5} \mathrm{H}_{8}\right)$ subunits to form more complex molecules in living organisms prior to crude oil formation. ${ }^{28}$ It is therefore highly likely that the alkylated side chains of the cyclic alkanes are also restricted to being only methyl branched isomers derived from isoprene subunits, though this cannot be confirmed from their mass spectra. However, many more isomers are observed for the cycloalkanes relative to those observed for acyclic alkanes (Figure S2) because there are a larger number of possible configurations resulting from the different positions for alkyl substitution on the ring. The most likely cycloalkane ring structures are cyclopentane and cyclohexane because of the minimal ring strain of these configurations. This is supported by the retention time data that is consistent with these structures, similar to previous work on lubricating oil, ${ }^{21}$ and not with larger cycloalkane rings that elute later. Alkene, diene, and triene moieties would also have $N_{\text {DBE }}$ of 1 , 2 , and 3 . However, unsaturated chains are relatively unstable and are thus very unusual in crude oils. As a result, we expect contributions from these species to be negligible, consistent with previous work on refined petrochemical products. ${ }^{21,29}$

Higher $N_{\text {DBE }}$ 's contribute a smaller fraction of the mass ( $N_{\text {DBE }} \geq 4, \sim 15 \%$ ) and include contributions from tetra-, penta-, and hexacyclic alkanes, benzenes, tetralins, indanes, and PAHs including the alkylated varieties. Naphthalene and alkylated naphthalenes $\left(N_{\text {DBE }}=7\right)$ and phenanthrenes and their alkylated homologues $\left(N_{\text {DBE }}=10\right)$ and $N_{\text {DBE }}=8$ were the dominant PAH components present (Figure 1c). It should be noted that at $10.5 \mathrm{eV} N_{\text {DBE }}=8$ was heavily impacted by fragment ions of aliphatic compounds (Figure S4) that prevented reliable quantitation. At 9 eV , the interference from these fragments was minimal because aliphatic compounds have ionization energies higher than $9 \mathrm{eV}^{30}$ A calibration curve at 9 eV was not available, so we estimate the mass of $N_{\text {DBE }}=8$ by converting the raw signal at 9 eV to mass at 10.5 eV through comparison to $N_{\text {DBE }}=7$ and 9 (Figure S4). The $N_{\text {DBE }}=8$ values listed in Table S1 therefore include an additional uncertainty of $10 \%$.

In this work, we do not explicitly report the aromatic and aliphatic contributions to $N_{\text {DBE }}=4,5$, and 6 because of difficulties in distinguishing them using our methodology due to the IE of the species being similar and thus unable to be differentiated in 9 versus 10.5 eV runs. (See Supporting Information text and Figures S5-S7.) However, we can rationalize that a greater proportion of the $N_{\mathrm{DBE}}=4,5$, and 6 mass at lower $N_{\mathrm{C}}(9-20)$ is likely aromatic. This is because the complexity (many coeluting peaks) observed in the parent ion chromatograms for these $N_{\text {DBE }}$ 's imply that there are a large number of structural isomers present. For example, consider compounds with $N_{\mathrm{C}}=15$ : For a monoaromatic compound, this represents a benzene ring with a nonyl side chain that could be distributed in hundreds of different
configurations, whereas for a tetracyclic compound with two cyclohexane rings and two cyclopentane rings, only one configuration is possible. This suggests a larger contribution from aromatic compounds at lower carbon numbers $\left(\leq C_{20}\right)$ because statistically there are more possible isomer structures.

Table 2 compares the mass fractions eluting between different $n$-alkane windows observed in this work with values reported by

Table 2. Comparison of the Eluting Mass Fractions (\%) from This Work and High-Temperature Stimulated Distillation of the Gas-Chromatography-Amenable Fraction of the Macondo Well Oil Reported by Reddy et al. ${ }^{17}$

| eluting fraction | Reddy et al. | this work |
| :--- | :---: | :---: |
| before $n$-nonane $\left(<\mathrm{C}_{9}\right)$ | 15 |  |
| $n$-nonane to $n$-undecane $\left(\mathrm{C}_{9}-\mathrm{C}_{11}\right)$ | 10 | $13 \pm 4$ |
| $n$-dodecane to $n$-octadecane $\left(\mathrm{C}_{12}-\mathrm{C}_{18}\right)$ | 25 | $27 \pm 8$ |
| $n$-nonadecane to $n$-triacontane $\left(\mathrm{C}_{19}-\mathrm{C}_{30}\right)$ | 25 | $24 \pm 8$ |
| after $n$-triacontane $\left(>\mathrm{C}_{30}\right)$ | 25 | $4 \pm 2$ |
| total | 100 | $68 \pm 22$ |

Reddy et al. ${ }^{17}$ for oil sampled directly from the Macondo well by a remotely operated vehicle on June 21, 2010, during response efforts to the DWH disaster. The very good agreement within each of these elution bins provides confidence in our results. Small compounds are not quantitatively captured by our inlet, and elution of large compounds is limited by temperatures within the mass spectrometer. Within the range of compounds studied in this work $\left(\mathrm{C}_{9}-\mathrm{C}_{33}\right)$, we account for $68 \pm 22 \%$ of the mass.

Comparison with Other Published Data. Figure 2 shows the comparison between this work and published mass fraction


Figure 2. Comparison of mass fractions ( $\mathrm{mg} \mathrm{kg}^{-1}$ ) for individual compounds or groups of isomers observed in this work (listed in Table S3), with reported values from (a) $\mathrm{WHOI}^{17}$ and (b) NIST. A comparison between NIST and WHOI is also shown in c. Data are color-coded by $N_{\text {DBE }}$ class, consistent with the color scheme used in Figure 1. Solid lines indicate 1:1, and dashed lines 2:1 and 1:2.
( $\mathrm{mg} \mathrm{kg}^{-1}$ ) data for selected species reported by Woods Hole Oceanographic Institution (WHOI, ${ }^{17}$ Figure 2a) and NIST (Figure 2b). Figure 2c shows the comparison between NIST and WHOI. All compounds included in these comparisons are shown in Table S3 along with their measured mass fractions and
uncertainties if available. As part of the comparison, the elution order of specific isomers was confirmed through comparison of reported Kovats retention indices (KI) for temperature-programmed GC ${ }^{31}$ ( or Lee retention indices (LI) ${ }^{32}$ converted to KI) from the literature to those observed in this work (Table S3). In general, there is very good agreement and linearity ( $r^{2}=0.84$, $n=67$ ) between this work and WHOI (Figure 2a) with all data agreeing to within a factor of 2 , except for the $\mathrm{C}_{9}$ aromatics (Figure S8), with a relative mean standard deviation (RMSD) of $18 \%$. There is also good agreement between this work and NIST ( $r^{2}=0.64, n=55$, RMSD $=32 \%$; Figure $2 b$ ), although the mass fractions observed in this work were typically higher for most compounds (on average by a factor of 2, Figure S8), though most were less than a factor of 4 with the exception of the dimethyl decalins (Figure S8). A comparison between NIST and WHOI also agreed very well ( $r^{2}=0.95, n=30, \mathrm{RMSD}=32 \%$; Figure 2c). The mass fractions reported by WHOI were also higher than those reported by NIST with a similar magnitude to that observed in this work. This intercomparison represents the first independent assessment of our calibration methodology, and the good agreement provides confidence in our analytical approach.

Table 3 shows the total mass fraction (\%) and total mass released from the Macondo well during the DWH disaster for all $N_{\text {DBE }}$ classes reported in this work for compounds with $N_{\mathrm{C}}=$ $9-33\left(\mathrm{C}_{9}-\mathrm{C}_{33}\right)$. Table 3 also shows the GC-amenable percent mass (\%) data reported by Reddy et al. ${ }^{17}$ separated by broad chemical class. There is excellent agreement for $n$-alkanes as well as for the sums of the branched alkanes, cycloalkanes, monoaromatics, and PAHs, with both data sets reporting the same total mass fractions within the reported uncertainties. However, although the sums of the branched and cycloalkanes agree well, the contribution of those compound classes differ. In this work, we observe fewer (by approximately a factor of 2) branched alkanes and almost three times more cycloalkanes than did Reddy et al. ${ }^{17}$ Some of these discrepancies might be from contributions from branched and/or cycloalkanes that were not characterizeable using their method and that were reported in a category labeled as "other" ( $18 \%$ ). As such, we suggest that the results presented here provide a more accurate representation of the total emission of branched and cyclic alkanes released during the DWH disaster (Table 3).

Intercomparison of NIST SRM 2779 and Surrogate Oil from the Marlin Platform. With many scientific studies into the impacts of the DWH disaster planned for at least the next 10 years, there is expected to be continued demand for Macondo source oil. Demand may exceed the supply of the available source oil such that surrogate crude oils will be necessary. Oil from the BP-owned and operated Marlin platform was selected as surrogate oil on the basis of its similar geography, oil family, composition, crude oil chemistry, and aquatic toxicity. Previous chemical analyses of the composition were limited to total petroleum hydrocarbons (TPH), $n$-alkanes, selected aromatics including benzene, toluene, xylenes (BTEX), polycyclic aromatic hydrocarbons (PAH), and geochemical biomarkers. This data was presented at the 32nd annual meeting of the Society of Environmental Toxicology and Chemistry (SETAC) in 2011, but to the best of our knowledge, this work has not been published in the peer-reviewed literature.

Figure 3 shows an intercomparison of the comprehensive chemical composition of the Marlin surrogate and NIST SRM 2779 (Macondo well) crude oils and clearly demonstrates that they are chemically very similar ( $r^{2}=0.86, n=236, \mathrm{RMSD}=10 \%$ ) with almost all $N_{\text {C }}$ and $N_{\text {DBE }}$ classes reported in this work being

Table 3. Comparison of the Mass Fractions for Analyte Classes from This Work with Those Reported by Reddy et al. ${ }^{17 a}$

${ }^{a}$ Also shown are the calculated total releases from the Macondo well for each analyte class reported here. ${ }^{b}$ Calculated with a net oil emission of $4.9 \times$ $10^{6}$ barrels. ${ }^{1,2}{ }^{c}$ Including the traditional sterane $\left(N_{\mathrm{DBE}}=4\right)$ and hopane biomarkers $\left(N_{\mathrm{DBE}}=5\right)$.


| $N_{\text {DBE }}=0(\mathrm{~B} 0)$ |
| :--- |
| $N_{\text {DBE }}=0(\mathrm{~B} 1)$ |
| $N_{\text {DBE }}=0(\mathrm{~B} 2)$ |
| $N_{\text {DBE }}=0(\mathrm{~B} 3)$ |
| $N_{\text {DBE }}=0(\mathrm{~B} 4)$ |
| $N_{\text {DBE }}=0(\mathrm{~B} 5)$ |
| $N_{\text {DBE }}=0(\mathrm{~B} 6)$ |
| $N_{\text {DBE }}=1($ (straight $)$ |
| $N_{\text {DBE }}=2$ |
| $N_{\text {DBE }}=3$ |
| $N_{\text {DBE }}=4$ |
| $N_{\text {DBE }}=5$ |
| $N_{\text {DBE }}=6$ |
| $N_{\text {DBE }}=7+$ |
| thiophenes |

Figure 3. Comparison of mass fractions in the NIST SRM 2779 Gulf of Mexico crude oil and surrogate oil for $N_{\text {DBE }}=0$ (left panel) colored by number of methyl groups ( $n$-alkane, B0; methyl, B1; dimethyl, B2; trimethyl, B3; tetramethyl, B4, pentamethyl, B5, and hexamethyl, B6) and $N_{\mathrm{DBE}}>0$ colored by $N_{\text {DBE }}$ (right panel). For simplicity, all components with $N_{\text {DBE }} \geq 7$ are colored black. Solid lines indicate 1:1, and dashed lines indicate 2:1 and 1:2.
within the error of unity. This is perhaps not too surprising given that BP made a concerted effort to identify a similar crude oil that would make a suitable surrogate. However, what is interesting is
how similar these two crude oils are considering that their sources are located 35 km apart which underscores the challenges of studying in the Gulf of Mexico. As a result of the similarity in these
crude oils, the comprehensive chemical composition data reported in this work (Tables 1 and S1) will be of interest to those researchers conducting research using authentic Macondo well source oil, NIST SRM 2779 oil, and/or the surrogate oil from the Marlin Platform.

Environmental Implications. In this work, the chemical composition of hydrocarbons present in Macondo well crude oil was characterized using GC/VUV-MS, with a mass balance of $68 \pm 22 \%$ for $\mathrm{C}_{9}-\mathrm{C}_{33}$ hydrocarbon compounds. The chemical classification scheme used in this work provides data in a format that can be utilized by environmental models to better understand the comprehensive chemical evolution of crude oil during weathering processes. One such example is to estimate the speciated mass fluxes of evaporating components from surface oil slicks and to predict the impact of these evaporating hydrocarbons on air pollutant concentrations, i.e., particulate matter and ozone, with implications for air quality in the vicinity of an oil spill. ${ }^{10,33}$ Analysis by GC/VUV-MS is particularly well-suited for the determination of saturated semivolatile compounds where the presence of large numbers of constitutional isomers inhibits the ability to chromatographically resolve individual peaks, which would otherwise prevent their detection. This is especially true for cycloalkanes where in addition to skeletal isomerism of the alkyl side chain or chains there are many positional isomers due to different arrangements of those alkyl side chains on the ring or rings. Recent work has identified these hydrocarbons as being important precursors to observed oxygenated hydrocarbons formed via photooxidation and biodegradation, although their identities have not been categorically determined. ${ }^{6,7}$ The application of GC/VUV-MS to crude oil samples with varying degrees of environmental weathering could provide invaluable insights into the identities of the key precursors and therefore the likely identity of the oxygenated products formed during the weathering process.

## - ASSOCIATED CONTENT

## (s) Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.est.5b03472.

Additional information as noted in the text and Figures S1-S8 and Tables S1-S3. (PDF)

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## Notes

The authors declare no competing financial interest.

## - ACKNOWLEDGMENTS

The data presented in Table 1 and Table S 1 are also available from the GRIIDC data archive, accessible via the Internet at http:// ezid.cdlib.org/id/doi:10.7266/N7K64G1H. This work was funded by the Gulf of Mexico Research Initiative (GoMRI) as part of the Gulf Integrated Spill Response (GISR) consortium under contract

SA12-09/GoMRI-006. G.I.-V.W. was supported by the National Science Foundation (NSF) Graduate Research Fellowship (NSF grant: DGE 1106400). The Advanced Light Source, the Chemical Dynamics Beamline, and K.R.W. were supported by the Director, Office of Science, Office of Basic Energy Sciences, of the U.S. Department of Energy under contract no. DE-AC02-05CH11231.

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[^0]:    Received: July 17, 2015
    Revised: October 8, 2015
    Accepted: October 13, 2015

