Responses to Referee #1:

We thank Referee #1 for their comments and suggestions: all of which were helpful in improving the manuscript.

We detail our response to each item below.

The fuels themselves appear very representative; whether the emissions from the FLAME burns are representative of real-world emissions is a difficult question to answer and probably beyond the scope of this work. However, since the authors are experts in this area of research – some discussion on the validity of lab-to-field representativeness is warranted for the emissions factors reported here.

This is an important point that we were greatly concerned with in a companion paper, but did not address clearly enough in this paper. We now address this in three additional ways.

(1) We mention early on in the introduction that the representativeness of the FLAME-4 fire emissions was explored in great detail by examining agreement with field measurements in an already published companion paper (Stockwell et al. 2014). In that paper we show that much of the FLAME-4 data is representative as is and - for fuel types warranted - we suggest simple, specific adjustment procedures to make the laboratory data more representative of field data.

Changes in the text are as follows:

P22167, L15-20: Old text: “In this work, we perform, to our knowledge, the first application of PTR-TOF-MS technology to laboratory biomass burning smoke to characterize emissions from a variety of authentic globally significant fuels. We report on several new or rarely measured gases and present a large set of useful emission ratios (ERs) and emission factors (EFs) for major fuel types that can inform/update current atmospheric models.”

New text: “In a companion paper the FLAME-4 emissions were compared extensively to field measurements of fire emissions and they were shown to be representative of “real-world” biomass burning either as is or after straightforward adjustment procedures detailed therein (Stockwell et al., 2014). In this work, we describe, to our knowledge, the first application of PTR-TOF-MS technology to laboratory biomass burning smoke to characterize emissions from a variety of authentic globally significant fuels. We report on several new or rarely measured gases and present a large set of useful emission ratios (ERs) and emission factors (EFs) that can represent major fuel types and inform/update current atmospheric models.”

(2) We now better clarify that the main compilation of new recommended EF is in Table S3. This important table appears in the supplement simply because of its large size. We also better clarify that any adjustment procedures required based on the comparisons in the companion paper have been applied to the data in Table S3. One exception is for cooking fires. We did not have enough TOF data from analyzing just three fires to get a useful EF vs MCE relationship so we instead computed adjusted EF for any NMOC “X” as the average X/CH4 ratio from FLAME-4 times the literature average EF(CH4) for cooking fires from Akagi et al., (2011).
Changes in text are as follows:

P22173 L5-7, Old text: “Finally, the EFs reported in Supplement Table S3 were adjusted according to procedures established in Stockwell et al. (2014) to improve laboratory representation of real-world biomass burning emissions.”

New text: “Finally, the EFs reported in Supplement Table S3 were adjusted (when needed) according to procedures established in Stockwell et al. (2014) to improve laboratory representation of real-world biomass burning emissions. This table contains the EF we recommend other workers use and it appears in the Supplement only because of its large size.”

P22173, L17: changed “S1 and S2” to “S1-S3”

To further clarify the relative importance of Tables S3 and Table 2 we made the following change:

P22176, L14-20: Existing text: “To facilitate discussion we grouped many of the assigned (or tentatively assigned) mass peak features into categories including: aromatic hydrocarbons; phenolic compounds; furan and its derivatives; nitrogen-containing compounds; sulfur-containing compounds; and miscellaneous compounds at increasing m/z.”

New text: “To facilitate discussion we grouped many of the assigned (or tentatively assigned) mass peak features into categories including: aromatic hydrocarbons; phenolic compounds; furans; nitrogen-containing compounds; and sulfur-containing compounds. These categories do not account for the majority of the emitted NMOC mass, but account for most of the rarely-measured species reported in this work. We then also discuss miscellaneous compounds at increasing m/z.”

P22190, L1, changed “and should aid” to “and (especially the recommended values in Table S3) should aid”

(3) We found two recent additional field studies of biomass burning to compare to (Geron and Hays, 2013; Kudo et al., 2014). They both support the “realism” of our FLAME-4 EF in Table S3.

Changes in text are as follows:

We include the following sentence in the text immediately following the above mentioned sentence at P22173 L7: “In addition to the comparisons considered in Stockwell et al. (2014), we find that our EFs in Table S3 are consistent (for the limited number of overlap species) with additional, recent field studies including Kudo et al. (2014) for Chinese crop residue fires and Geron and Hays (2013) for NC peat fires.”

These additional references were added to the paper:


The authors report a conservative bound of +/- 50% measurement uncertainty about emissions by compound. This uncertainty is not surprising given the inherent limitations of PTR-MS and the fact that their calibration standards were mostly hydrocarbon based (with only a few heteroatom molecules beyond C-H-O). Perhaps the biggest weakness of this work is the choice of calibration gases. The authors report emissions factors for many previously unstudied compounds (especially oxygenates and a few S- and N-containing compounds), yet, they did not generate calibration curves for these compounds. Standards for S- and N-compounds are likely difficult to obtain (especially those that are semi-volatile), however, the development of a phenol- and furan-containing calibration mixture would have strengthened this work considerably.

P22170 lines 16-19. This sentence is awkward. “In cases where a compound contains a non-oxygen heteroatom (such as methanethiol), the mass dependent calibration factor was determined using the relationship established using the oxygenated species.” More importantly, it is unclear whether this approach is valid. What is the uncertainty associated with assuming that such heteroatomic compounds will follow the oxygenated calibration curve? The deviation of dimethylsulphide from the ‘oxygenates’calibration line (Figure 3) indicates that this assumption may not be valid.

We agree that additional calibration gases would be useful. This would be especially true for stand-alone use of the PTR-TOF-MS and we do note that we used the OP-FTIR furan data in this work. In our case, the instrument was rented for only three months and calibration gases were ordered ahead of time to meet the needs of several different experiments. We chose a variety of gases that were already known to be emitted by fires (including hydrocarbons and C-H-O heteroatoms), but operationally could not add calibration gases during the experiment. It is no longer possible to run more calibration gases, but for any group that owns or rents a TOF for smoke characterization in the future we recommend including more calibration species and measuring more fragmentation patterns. We now mention these issues in the conclusions where we summarized the lessons learned from our first attempt.

P22189 L2 before sentence beginning with: “Despite these practical…” add: “We were limited to our pre-chosen calibration mixture based primarily on gases previously observed in smoke. For future experiments we suggest adding more standards to generate more accurate calibration factors, specifically including major species such as furan and phenol and more compounds with S and N heteroatoms. In addition, measuring the fragmentation, if any, of more of the species identified in this work would be of great value.”

On the subject of our treatment of heteroatoms, such as in the case of DMS and Acetonitrile, we agree with the reviewers that perhaps this is not the best treatment of these species. One would expect that approximation of sensitivities would most accurately be calculated via grouping families of compounds (e.g. alcohols, carbonyls,
alkenes) rather than general subgroups of oxygenates and hydrocarbons. Unfortunately due to the limitations in this study and the lack of additional calibration standards as mentioned above we are forced to make some assumptions as in the case of compounds containing heteroatoms. Here we only have two calibrated examples both of which show a wide range in sensitivities. While sulfur and nitrogen containing compounds should be treated separately in a more complete analysis, we have chosen to consider these species collectively in order to increase sample size.

When considered as a group these compounds more closely follow the trend for the oxygenated family. This does result in an increased error on the calculated calibration factors which has been approximated using the scatter on the fits derived from the oxygenated + heteroatom relationship. Unfortunately without any additional information we cannot more accurately approximate the error though we do admit the cited error is likely a lower limit on the actual errors associated with this method of determination. This is a valid criticism of this work and the approximation of calibrations factors is a problem that is and will continue to be an issue with the analysis of TOF data moving forward, however, a detailed treatment is beyond the scope of this paper.

P22170, L16: Existing text: “In cases where a compound contains a non-oxygen heteroatom (such as methanethiol), the mass dependent calibration factor was determined using the relationship established using the oxygenated species.”

New text: “Sulfur and nitrogen-containing compounds were considered collectively and together they more closely followed the trend of the oxygenated species. Thus, in cases where a compound contains a non-oxygen heteroatom (such as methanethiol), the mass dependent calibration factor was determined using the relationship established using the oxygenated species.”

Then in addition to recognize the higher uncertainty for these species we added this change:

P22171, L6: We append after “prescribed” “, but with larger errors possible for compounds with N and S heteroatoms.”

Table 2 (body text). Are the numbers in parentheses standard deviations or some other measure of variability? This comment applies to nearly every Table in the manuscript; footnotes should be added to each table to explain accordingly.

Thank you for bringing this to our attention. Table 2 should include the following footnote “Note: "nm" indicates not measured; blank indicates species remained below the detection limits; values in parenthesis indicate one standard deviation”

The authors use methanol concentrations, as measured by OP FTIR, as an internal standard to account for variations associated with PTR-MS instrument. This is an innovative approach and one that will likely reduce measurement variability. Inclusion of more intercomparison data between the PTR and FTIR instruments for other shared compounds (perhaps as supplementary material) would strengthen this manuscript as these
data would shed light on PTR measurement reliability. Showing the methanol comparison plot (1:1) would be helpful, too.

Developing detailed inter-comparisons for more species for numerous fires is a large body of work that is the focus of an in-progress companion paper describing both contributions and inter-comparisons for the PTR-TOF-MS, OP-FTIR, WAS, and 2D-GC data from FLAME-4. The in-progress study comparing and identifying the strengths of four techniques promises to be more valuable than comparing two techniques. Thus, we prefer not to lengthen the current already-long paper, which is focused on expanding the amount of data concerning NMOCs emitted by BB.

The point of using methanol as the internal standard for calculating fire-integrated ERs is that it actually cancels errors even if the PTR-MS and OP-FTIR disagree on occasion for methanol due to reagent ion depletion, timing, etc. We have already included the slope and $R^2$ value for this comparison in the manuscript. The lack of overall bias between the two instruments for methanol that we observed is nice, but already a departure from the main point. As noted above, adding more comparisons or the 1:1 methanol comparison plot would lengthen the manuscript and shift focus when a separate, comprehensive paper detailing measurement issues is already in progress. Thus, while the Referees' concerns are warranted we address them elsewhere and did not add figures or comparisons at this time.

It is also important to note that depending on how we define the intercomparison (e.g. fire-integrated or real-time, etc.) the results vary a bit - and this takes time/space to describe, thus further supporting our decision to devote a separate paper to the topic of the intercomparison. As an example we show (in this response only) an FTIR/PTR methanol comparison with a slightly different filter applied than in the example quoted in the text. The slope and $R^2$ are then different, but still good. We trust the forthcoming companion paper comparing four techniques will provide the best way to gain insights into the bulk of the FLAME-4 NMOC fuel-specific emission factor results.

The good agreement for several species measured by both proton-transfer-reaction mass spectrometry and OP-FTIR spectroscopy has been demonstrated in numerous past manuscripts focused on laboratory biomass burning emissions...
Old sentence: “The agreement between these two measurements is within the uncertainties of both instruments.”

New sentence: “This result is consistent with the good agreement for several species measured by both PTR-MS and OP-FTIR observed in numerous past studies of laboratory biomass burning emissions (Christian et al., 2004; Karl et al., 2007; Veres et al. 2010; Warneke et al., 2011).”

The compound-specific emissions factors that are included as supplemental tables is a great contribution of this work; these data will likely be used by many researchers. Is the use of 3+ significant figures justified, given the uncertainty in these EF data?

We simply chose to mass-produce the data with enough places after the decimal point to account for the large range in EF/ER. The co-tabulated standard deviations should make the actual number of significant figures clear. The important table (S3) uses 3 significant figures and researchers can use less if they prefer. It seems unnecessary to go through such large tables and manually edit them because there is so much variation on how many 0s are in front of the numbers. In addition, the rounding off the CO₂ emission factors to less than four digits could be warranted, but is never done in the literature. Reporting significant figures only introduces “round-off error” into any subsequent calculations with the data.

Figure 3 is difficult to comprehend based on the information presented in the title and legend. Only a single reference to this figure appears in the text body, but that reference is not explanatory. Many questions follow. There are three categories for ‘oxygenates’, ‘phenolics’, and ‘furans’ for each fuel type. Why? What do the parentheses mean, number of independent tests? I think this plot is meant to compare FTIR to PTR measurements, but if so, I don’t know which instrument is which since there are four bars per fuel.

Thank you for pointing out that this figure needs a more detailed explanation. The data in these figures are based on the synthesis of the FTIR and PTR measurements and do not compare the two. Instead we lumped the emissions into the following categories: non-methane hydrocarbons, oxygenates containing only 1 oxygen, oxygenates containing 2 oxygen atoms, and oxygenates containing 3 oxygen atoms. To give an even greater level of detail we then indicated within each category what emissions were directly from phenolic compounds or furan and substituted furans.

To clarify these changes in the text are as follows:

Also change the figure description to add further clarity, P22210:
“Figure 3. The emission factors (g kg\(^{-1}\)) of total observed hydrocarbons and total observed species oxygenated to different degrees averaged for each fire type based on a synthesis of PTR-TOF-MS and OP-FTIR data. The patterned sections indicate the contribution to each of the above categories by selected functionalities discussed in the text (aromatic hydrocarbons, phenolics, furans). The parenthetical expressions indicate how many oxygen atoms are present.”

For the emissions factors tables, can the authors indicate which tests were true replicates? I suspect such information exists elsewhere but it would be very helpful to know how repeatable the measurements were for repeat tests (i.e., EF’s determined from the same set of experimental conditions).

If by true replicates the author means burning “the exact same fire again and seeing how the emissions changed” then we have no true replicates. This is partly because on the scale of the fires we burned (several cubic meters of fuel) no exact duplicates could be constructed, but also because we instead aimed to vary the fires (moisture, geometry, etc.) in hope of capturing the natural EF dependence on MCE. We did perform replications of the smog chamber perturbations that will be described elsewhere.

The authors might try breaking Figure 4a and 5a into two panels each. One panel would contain EF’s for those fuels whose sum exceeds > 1 g/kg and one for those fuels whose sum is less than 1 g/kg (the latter scaled appropriately). The bars for fuels with low EF’s (i.e., less than 1 g/kg) are practically unreadable in these figures.

We thank the reviewer for bringing this to our attention and we have scaled Figures 4a, 5a, and 6a accordingly.

Responses to Referee # 2:

We thank the Anonymous Referee #2 for their comments and suggestions. All of the comments have been considered and were helpful in improving the manuscript. We address the specific comments below.

Abstract Page 22165, Line 3 – There should be no hyphen between crop and residue

Page 22165, Line 5 – Suggest changing standards to standard

Page 22615, Line 10 – The abbreviation FTIR has not been defined

Page 22166, Lines 23-24 – There should be hyphens in proton transfer reactions

2.3. Proton-transfer-reaction time-of-flight mass spectrometer Page 22171, Line 8 – The abbreviation VOC has not been defined

2.4. OP-FTIR Page 22171, Line 10 – The chemical formula used has not been defined
Page 22185, Line 17 – There should be no hyphen between crop and residue

P22185 L17 We have changed “Crop-residue” to “Crop residue”

4.7 Cookstoves Page 22187, Line 18 – Envirotif is misspelled

P22187 L18 We have respelled “Envriofit” to “Envirofit”

References Page 22912, Line 6 – Believe it should be St. Clair, J.M.

P22912 L6 We have fixed our mistake concerning the location of the St. in the following reference


Page 22912, line 27 – There is an extra d at the end of European

P22912 L27 We have corrected misspelling by changing “Europeand” to “European”

Page 22195, Line 8 – There should be an and before Hjorth


Page 22199, Line 29 – There should be an and before Fanelli


General Comment: -The order of the references in the citations seems to vary through the paper of being in alphabetical or chronological order. Either way is actually fine, but it should be consistent throughout. I have tried to point out the ones I noticed below in my specific comments.

We thank the referee for noticing the alphabetical and chronological variations in our reference citations. We have addressed all discrepancies as indicated below.

P22166 L7-9 Re-ordered to read “(Reid et al., 1998; Trentmann et al., 2005; Alvarado and Prinn, 2009; Yokelson et al., 2009; Vakkari et al., 2014)”

P22174 L10 Re-ordered to read “(Akagi et al., 2011; Yokelson et al., 2013, etc.)”

P22174 L19 Re-ordered to read “(Liu et al., 2012; Pittman Jr. et al., 2012; Li et al., 2013; more citations in Table S4)”

P22174 L25 Re-ordered to read “(Lobert, 1991; Ge et al., 2011; etc.)”

P22177 L11-12 Re-ordered to read “(Phousongphouang and Arey, 2002; Ziemann and Atkinson, 2012).”

P22177 L21 Re-ordered to read “(Andreae and Merlet, 2001; Henze et al., 2008)”
Tables Table 2 - Line 2 of the caption - Suggest adding the word type after fuel
The table caption now reads “Table 2a. Emission ratios to benzene, phenol, and furan for aromatic hydrocarbons, phenolic compounds, and substituted furans in lumped fuel-type categories.

Figures Figure 1 - Suggest labeling each plot as a, b, or c since they are referred to this way in the text
Thanks for the suggestion and we have labeled the three sections of the figure a, b, and c.

Figure 3 - Suggest indicating in the caption that the number of oxygen atoms for the three various oxygenated classes are indicated in parenthesis in the legend
This has been addressed both in the text and in the figure caption as initially prompted by Anonymous Referee 1.

Figure 5 - Line 3 of the caption - Suggest changing considered the fuel average to considered in the fuel average
We have changed the caption to read “Figure 5. (a) The distribution in average fuel EF for several phenolic compounds, where compound specific contributions are indicated by color. The EFs for compounds additionally analyzed a single time for select fires are included but are not a true average. (b) The linear correlation of select phenolic compounds with phenol during an organic hay burn (Fire 119).”

Voluntary changes:

P22180 L5-6 Re-ordered to read “(Atkinson et al., 1992; Olariu et al., 2002; Harrison et al., 2005; Lauraguais et al., 2014)”
P22180 L12-13 Re-ordered to read “(Kitanovski et al., 2012; Desyaterik et al., 2013; Mohr et al., 2013; Zhang et al., 2013)”
P22180 L15-16 Re-ordered to read “(Iinuma et al., 2010; Kitanovski et al., 2012; Lauraguais et al., 2014)”
P22181 L5-6 Re-ordered to read “(Cabañas et al., 2005; Villanueva et al., 2007)”
P22182 L21-22 Re-ordered to read “(Lobert et al., 1991; Schade and Crutzen, 1995; Ma and Hays et al., 2008; Barnes et al., 2010; Ge et al., 2011).”
P22184 L7 Re-ordered to read “(Friedli et al., 2001; Meinardi et al., 2003; Akagi et al., 2011; Simpson et al., 2011)”
P22185 L11 Re-ordered to read “(Holzinger et al., 1999; Christian et al., 2004)”
P22186 L11-12 Re-ordered to read “(Christian et al., 2003; Akagi et al., 2011; Yokelson et al., 2013; St. Clair et al., 2014)”
P22187 L10 Re-ordered to read “(Simpson et al., 2011; Akagi et al., 2013)

Tables Table 2 - Line 2 of the caption - Suggest adding the word type after fuel
The table caption now reads “Table 2a. Emission ratios to benzene, phenol, and furan for aromatic hydrocarbons, phenolic compounds, and substituted furans in lumped fuel-type categories.

Figures Figure 1 - Suggest labeling each plot as a, b, or c since they are referred to this way in the text
Thanks for the suggestion and we have labeled the three sections of the figure a, b, and c.

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This has been addressed both in the text and in the figure caption as initially prompted by Anonymous Referee 1.

Figure 5 - Line 3 of the caption - Suggest changing considered the fuel average to considered in the fuel average
We have changed the caption to read “Figure 5. (a) The distribution in average fuel EF for several phenolic compounds, where compound specific contributions are indicated by color. The EFs for compounds additionally analyzed a single time for select fires are included but are not a true average. (b) The linear correlation of select phenolic compounds with phenol during an organic hay burn (Fire 119).”

Voluntary changes:

P22190, L11: “was” changed to “were” – L12: changed “regions biofuel” to “regions where biofuel” - L18:
changed “chlorine organic gases were not readily detectable” to “chlorine containing organic gases were not readily observed”

We changed “Ringed Aromatics” to “Aromatics” on the y axis label of Figure 4.

We eliminated m/z 81 from the Supplementary Tables because it is a fragment considered in the calculation of total carbon, but not a primary emission.

We added the EFs for the tire fire to the supplementary tables based on using HCOOH as an internal standard.
We’ve also added several rows of additional species to Table S3. We have also extended the column averages in the supplemental tables for fuel types to include extended species even if there was no standard deviation.

We added an extra row for MCE in Tables 2a and 2b.

At the end of the acknowledgements we added: “We thank C. Geron for providing a sample of NC peat.”


In Figure 6’s caption we’ve changed “furan derivatives” to “substituted furans”. We’ve also clarified which compounds are not averaged: The caption now reads “Figure 6. (a) The distribution in average fuel EF for furan and substituted furans, where individual contributions are indicated by color. The EFs for substituted furans additionally analyzed a single time are not true averages. (b) The linear correlation of furan with select substituted furans for an African grass fire (Fire 49).”

In Figure 4’s caption we’ve changed “The EF for p-Cymene is only calculated for select burns and should not be considered an average for each particular fuel type.” To: “The EFs for p-Cymene are only calculated for select fires and should not be considered a true average.”


Characterization of biomass burning smoke from cooking fires, peat, crop residue and other fuels with high resolution proton-transfer-reaction time-of-flight mass spectrometry

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Abstract

We deployed a high-resolution proton-transfer-reaction time-of-flight mass spectrometer (PTR-TOF-MS) to measure biomass burning emissions from peat, crop-residue, cooking fires, and many other fire types during the fourth Fire Lab at Missoula Experiment (FLAME-4) laboratory campaign. A combination of gas standards calibrations and composition sensitive, mass dependent calibration curves were applied to quantify gas-phase non-methane organic compounds (NMOCs) observed in the complex mixture of fire emissions. We used several approaches to assign best identities to most major “exact masses” including many high molecular mass species. Using these methods approximately 80-96% of the total NMOC mass detected by PTR-TOF-MS and Fourier transform infrared (FTIR) spectroscopy was positively or tentatively identified for major fuel types. We report data for many rarely measured or previously unmeasured emissions in several compound classes including aromatic hydrocarbons, phenolic compounds, and furans; many of which are suspected secondary organic aerosol precursors. A large set of new emission factors (EFs) for a range of globally significant biomass fuels is presented. Measurements show that oxygenated NMOCs accounted for the largest fraction of emissions of all compound classes. In a brief study of various traditional and advanced cooking methods, the EFs for these emissions groups were greatest for open 3-stone cooking in comparison to their more advanced counterparts. Several little-studied nitrogen-containing organic compounds were detected from many fuel types that together accounted for 0.1-8.7% of the fuel nitrogen and some may play a role in new particle formation.

1 Introduction

Biomass burning (BB) injects large amounts of primary, fine carbonaceous particles and trace gases into the global atmosphere and significantly impacts its physical and chemical properties (Crutzen and Andreae, 1990; Bond et al., 2004, 2013). While BB emissions are recognized as the second largest global atmospheric source of gas-phase non-methane organic compounds (NMOCs) after biogenic emissions, a significant portion of the higher molecular weight species remain unidentified (Christian et al., 2003; Warneke et al., 2011; Yokelson et al., 2013). It is widely
accepted that the addition of large amounts of these highly reactive species into the atmosphere alters chemistry on local to global scales (Andreae and Merlet, 2001; Andreae et al., 2001; Karl et al., 2007). NMOCs particularly impact smoke evolution by rapid formation of secondary organic aerosols (SOA) and secondary gases including photochemical ozone (O_3) (Reid et al., 1998; Trentmann et al., 2005; Alvarado and Prinn, 2009; Reid et al., 1998; Trentmann et al., 2005; Yokelson et al., 2009; Vakkari et al., 2014).

The many unknowns and initial gas-phase variability of BB emissions limit our ability to accurately model the atmospheric impacts of fire at all scales (Trentmann et al., 2005; Mason et al., 2006; Alvarado and Prinn, 2009; Alvarado et al., 2009; Wiedinmyer et al., 2011). Estimating or modeling the potential of smoke photochemistry to generate secondary aerosols or O_3 requires realistic estimates of NMOC emissions in fresh smoke and knowledge of the chemical processing environment. Measurements capable of identifying and quantifying rarely measured and presently unidentified emissions of NMOCs, in particular the chemically complex low volatility fraction, are vital to advance current understanding of the BB impact on air quality and climate.

Proton-transfer-reaction time-of-flight mass spectrometry (PTR-TOF-MS) is an emerging technique that simultaneously detects most NMOCs present in air samples including: oxygenated organics, aromatics, alkenes, and nitrogen-containing species at parts per trillion detection limits (pptv) (Jordan et al., 2009; Graus et al., 2010). The instrument uses H_3O^+ reagent ions to ionize NMOCs via proton-transfer-reactions to obtain high resolution mass spectra of protonated NMOCs with a low degree of molecular fragmentation at a mass accuracy sufficient enough to determine molecular formulas (C_nH_mN_oO_p).

Although there are many advantages to PTR-TOF-MS over conventional PTR quadrupole mass spectrometers (increased mass range, high measurement frequency, and high mass resolution) there remain several difficulties involving PTR technology including (1) detection is limited to molecules with a proton affinity greater than water, (2) complicated spectra due to parent ion fragmentation or cluster ion formation, and (3) the inability of the method to isolate isomers. Despite the limitations of this technology, PTR-TOF-MS is ideal for studying complex gaseous mixtures such as those present in BB smoke.

This study was carried out as part of a large scale experiment to characterize the initial properties and aging of gas- and particle-phase emissions in smoke from globally significant fuels. Experiments were conducted from October to November of 2012 during the fourth Fire Lab at Missoula Experiment (FLAME-4) as detailed by Stockwell et al. (2014). A major goal of the study focused on the identification and quantification of highly reactive NMOCs in order to: (1) better characterize the overall chemical and physical properties of fresh BB emissions, (2) better understand the distribution of emitted carbon across a range of volatilities in fresh and aged smoke, and (3) improve the capability of current photochemical models to simulate the climatic, radiative, chemical, and ecological impacts of smoke on local to global scales. In a companion paper the FLAME-4 emissions were compared extensively to field measurements of fire emissions and they were shown to be representative of “real-world” biomass burning either as is or after straightforward adjustment procedures detailed therein (Stockwell et al., 2014). In this work, we describe, to our knowledge, the first application of PTR-TOF-MS technology to laboratory biomass burning smoke.
to characterize emissions from a variety of authentic globally significant fuels. In this work, we perform, to our knowledge, the first application of PTR-TOF-MS technology to laboratory biomass burning smoke to characterize emissions from a variety of authentic globally significant fuels. We report on several new or rarely measured gases and present a large set of useful emission ratios (ERs) and emission factors (EFs) for major fuel types that can inform/update current atmospheric models.

2 Experimental details

2.1 Missoula fire sciences laboratory

The US Forest Service Fire Sciences Laboratory (FSL) in Missoula, MT houses a large indoor combustion room described in detail elsewhere (Christian et al., 2003; Burling et al., 2010; Stockwell et al., 2014). Briefly, fuels are burned on a bed located directly below a 1.6 m diameter exhaust stack. The room is slightly pressurized by outdoor air that generates a large flow that entrains the fire emissions up through the stack. Emissions are drawn into sampling lines fixed in the stack at a platform height 17 m above the fuel bed. Past studies demonstrated that temperature and mixing ratios are constant across the width of the stack at the platform height, confirming well-mixed emissions (Christian et al., 2004).

Burns were conducted using two separate configurations as described in Stockwell et al. (2014). In this paper we will focus on 125 of the 157 burns. During these fires, well mixed fresh smoke was sampled directly from the combustion stack by PTR-TOF-MS, roughly 5 s after emission. Results obtained during the remaining burns investigating photochemically processed smoke composition in dual smog chambers with a suite of state-of-the-art instrumentation are presented elsewhere (Tkacik et al., 2014).

2.2 Biomass fuels

Descriptions and ignition methods of each fuel type burned during FLAME-4 are detailed in Stockwell et al. (2014). Authentic globally significant fuels were collected including: African savanna grasses; US grasses; US and Asian crop-residue; Indonesian, temperate, and boreal peat; temperate and boreal coniferous canopy fuels; woods in traditional and advanced cooking stoves; shredded tires; and trash. The range of fuel loading was chosen to simulate real-world conditions for the investigated fuel types with global examples of biomass consumption shown in Akagi et al. (2011).

2.3 Proton-transfer-reaction time-of-flight mass spectrometer

Real-time analysis of NMOCs was performed using a commercial PTR-TOF-MS 8000 instrument from Ionicon Analytik GmbH (Innsbruck, Austria) that is described in detail by Jordan et al. (2009). The PTR-TOF-MS sampled continuously at a frequency of 0.2 Hz through heated PEEK tubing (0.0003 m o.d., 80°C) positioned facing upward to limit particulate uptake. The instrument was configured with a mass resolution (m/Δm) in the range of 4000 to 5000 at m/z 21 and a typical mass range from m/z 10-600. The drift tube was operated at 600 V with a pressure of 2.30 mbar at 80 °C (E/N ~ 136Td, with E as the electric field strength and N as the concentration of neutral gas; 1
Calibration factors (CFs) were determined using the relationship established using the oxygenated species. Cyclic compounds were considered collectively and together they more closely follow the trend of the oxygenated species. Thus, substantially higher trasmission efficiency (Fig. 1b and 1c) can be expected for hydrocarbon species (green labels) as this compound is known to be susceptible to substantial fragmentation in the drift tube. Sulfur and nitrogen-containing compounds were considered collectively and together they more closely follow the trend of the oxygenated species. Thus, in cases where a compound contains a non-oxygen heteroatom (such as methanethol), the mass dependent calibration factor was determined using the relationship established using the oxygenated species. Calibration factors were then determined according to the exact mass for all peaks where the chemical formula has been determined. Our
approach does not yet account for the potential for ions to fragment and/or cluster, however, we expect this impacts
less than 30% of NMOC and usually to a small degree for any individual species. These latter issues change the
mass distribution of observed carbon, but should not have a large effect on the total observed carbon.

It is difficult to assess the overall error introduced using this method of calibration factor approximation, as only a
limited number of comparable measurements of calibration factors are available. The deviation of measured
calibration factors for species contained in the gas standard from the linear approximation yields a range of errors
(21 ± 19%) with a maximum of 50% observed in all cases (excluding α-pinene, for reasons detailed above). While
PTR-TOF-MS is typically known as a soft ionization method, fragmentation is common among higher molecular
weight species and therefore needs to be considered as a limitation of this technique. For the individual species
identified it would be misleading to give a set error based on this limited analysis, however, in the absence of any
known molecular fragmentation a maximum error of 50% is prescribed, but with larger errors possible for
compounds with N and S heteroatoms. Better methods for the calculation of mass dependent calibration factors by
compound class should be developed in the near future to improve the accuracy of volatile organic compound
(VOC) measurements using PTR-TOF-MS.

2.4 OP-FTIR

To enhance application of the MS data, emission ratios to carbon monoxide (CO) were calculated where possible
using measurements from an open-path Fourier transform infrared (OP-FTIR) spectrometer described elsewhere
(Stockwell et al., 2014). The system includes a Bruker Matrix-M IR Cube spectrometer with an open White cell that
was positioned to span the width of the stack to sample the continuously rising emissions. The spectral resolution
was set to 0.67 cm⁻¹ and spectra were collected every 1.5 s with a duty cycle greater than 95%. Other gas-phase
species quantified by this method included carbon dioxide (CO₂), methane (CH₄), ethyne (C₂H₂), ethene (C₂H₄),
propylene (C₃H₆), formaldehyde (HCHO), formic acid (HCOOH), methanol (CH₃OH), acetic acid (CH₃COOH),
glycolaldehyde (C₃H₆O₂), furan (C₄H₄O), water (H₂O), nitric oxide (NO), nitrogen dioxide (NO₂), nitrous acid
(HONO), ammonia (NH₃), hydrogen cyanide (HCN), hydrogen chloride (HCl), and sulfur dioxide (SO₂) and were
obtained by multi-component fits to selected regions of the mid-IR transmission spectra with a synthetic calibration
non-linear least-squares method (Griffith, 1996; Yokelson et al., 2007).

The OP-FTIR system had the highest time resolution with no sampling line, storage, fragmentation, or clustering
artifacts; thus, for species in common with PTR-TOF-MS, the OP-FTIR data was used as the primary data. The
results from the inter-comparison (for methanol) of OP-FTIR and PTR-TOF-MS show excellent agreement using an
orthogonal distance regression to determine slope (0.995 ± 0.008) and the R² coefficient (0.789). The agreement
between these two measurements is within the uncertainties of both instruments. This result is consistent with the
good agreement for several species measured by both PTR-MS and OP-FTIR observed in numerous past studies of
laboratory biomass burning emissions (Christian et al., 2004; Karl et al., 2007; Veres et al. 2010; Warneke et al.,
2011).

2.5 Emission ratio and emission factor determination
Excess mixing ratios (denoted $\Delta X$ for each species “X”) were calculated by applying an interpolated background correction (determined from the pre and post fire concentrations). The molar emission ratio (ER) for each species “X” relative to CH$_2$OH ($\Delta X$/CH$_2$OH) is the ratio between the integral of $\Delta X$ over the entire fire relative to the integral of CH$_2$OH over the entire fire. We selected CH$_2$OH as the species in common with the OP-FTIR to serve as an internal standard for the calculation of the fire-integrated ERs of each species X to CO (Supplement Table S1). We do this by multiplying the MS-derived ER ($\Delta X$/CH$_2$OH) by the FTIR-derived ER ($\Delta$CH$_2$OH/$\Delta$CO), which minimizes error due to occasional reagent ion depletion or the different sampling frequencies between instruments that would impact calculating $\Delta X$ to $\Delta$CO directly. Several fires have been excluded from this calculation as data was either not collected by OP-FTIR and/or PTR-TOF-MS or alternatively, methanol data could not be applied for the conversion because (1) the mixing ratios remained below the detection limit or (2) methanol was used to assist ignition purposes during a few fires. As discussed in Sect. 2.3.1, ~50 additional masses were analyzed for selected fires and the ERs (to CO) for these fires are included in the bottom panels of Table S1. The combined ERs to CO from the FTIR and PTR-TOF were then used to calculate emission factors (EFs, g kg$^{-1}$ dry biomass burned) by the carbon mass-balance method (CMB), based on the assumption that all of the burned carbon is volatilized and that all of the major carbon-containing species have been measured (Ward and Radke, 1993; Yokelson et al., 1996, 1999; Burling et al., 2010). EFs were previously calculated solely from FLAME-4 OP-FTIR data as described in Stockwell et al. (2014) and a new larger set of EFs, which include more carbon-containing species quantified by PTR-TOF-MS, are now shown in Supplement Table S2. With the additional carbon compounds quantified by PTR-TOF-MS, the EFs calculated by CMB decreased ~1-2% for most major fuels with respect to the previous EFs reported in Stockwell et al. (2014). In the case of peat and sugar cane fires, the OP-FTIR derived EFs are now reduced by a range of ~2-5% and 3.5-7.5%, respectively. Along with these small reductions, this work now provides EFs for many additional species that were unavailable in Stockwell et al. (2014). Finally, the EFs reported in Supplement Table S3 were adjusted (when needed) according to procedures established in Stockwell et al. (2014) to improve laboratory representation of real-world biomass burning emissions. This table contains the EF we recommend other workers use and it appears in the Supplement only because of its large size. In addition to the comparisons considered in Stockwell et al. (2014), we find that our EFs in Table S3 are consistent (for the limited number of overlap species) with additional, recent field studies including Kudo et al. (2014) for Chinese crop residue fires and Geron and Hays (2013) for NC peat fires.

Fire emissions are partially dependent on naturally changing combustion processes. To estimate the relative amount of smoldering and flaming combustion that occurred over the course of each fire, the modified combustion efficiency (MCE) is calculated by (Yokelson et al., 1996):

$$MCE = \frac{\Delta CO_2}{\Delta CO_2 + \Delta CO} = \frac{1}{1 + (\frac{\Delta CO}{\Delta CO_2})}$$
Though flaming and smoldering combustion often occur simultaneously, a higher MCE value (approaching 0.99) designates relatively pure flaming combustion (more complete oxidation), a lower MCE (0.75-0.84) designates pure smoldering combustion and, thus, an MCE of ~0.90 represents roughly equal amounts of flaming and smoldering. Each fire-integrated MCE is reported in Tables S1-S2S3.

3 Results

3.1 Peak assignment

As exemplified by a typical PTR-TOF-MS spectrum of diluted smoke (Fig. 2a), the complexity of BB smoke emissions presents challenges to mass spectral interpretation and ultimately emissions characterization. Figure 2b shows a smaller mass range of the smoke sample shown in Fig. 2a on a linear scale to illustrate the typical relative importance of the masses (note the intensity of acetaldehyde (m/z 45) and acetic acid plus glycolaldehyde (m/z 61), which together account for almost 25% of the total signal). Although the spectra are very complex, systematic treatment of the burn data, assisted at some m/z by extensive published “off-line” analyses can generate reasonable assignments for many major peaks and result in useful emissions quantification.

As described earlier, the PTR-TOF-MS scans have sufficiently high resolution to assign molecular formulas \((C_nH_{2n}N_{2}O_{n})\) to specific ion peaks by matching the measured exact mass with possible formula candidates for the protonated compound. Specific compound identification for formula candidates can be unambiguous if only one species is structurally plausible or explicit identification of the compound had previously been confirmed by BB smoke analysis (Akagi et al., 2011; Yokelson et al., 2013; Akagi et al., 2014, etc.). Supplement Table S4 lists every mass and formula assignment for observable peaks up to m/z 165 and categorizes each mass as a confirmed identity, a tentative (most likely) species assignment, or an unknown compound. For several confirmed identities, the most abundant species at that exact mass is listed with likely contributions to the total signal from the secondary species listed in column 5. Most of the tentatively identified species have, to our knowledge, typically not been directly observed in BB smoke, but have been frequently verified with off-line techniques as major products in the extensive literature describing biomass pyrolysis experiments of various fuel types (Liu et al., 2012; Li et al., 2013; Pittman Jr. et al., 2012; Li et al., 2013; more citations in Table S4). Several tentative assignments are supported by off-line analyses being published elsewhere (Hatch et al., 2014), for example, simultaneous grab samples analyzed by two dimensional gas chromatography (2D-GC) support tentative assignments for furan methanol, salicylaldehyde, and benzofuran. In the case of nitrogen-containing formulas, the suggested compounds have been observed in the atmosphere, tobacco smoke, or lab fire smoke at moderate levels (Lobert, 1991; Ge et al., 2011; Lobert, 1991, etc.). Select studies supporting these assignments are referenced in the mass table with alternative possibilities also listed. An exhaustive list of all the many papers supporting the assignments is beyond the scope of this work. Several remaining compounds are also classified as tentative assignments as the identities designated are thought to be the most structurally likely. We anticipate that some or even many of the tentative assignments (and a few of the confirmed assignments) will be refined in future years as the results of more studies become available. We offer the
tentative assignments here as a realistic starting point that improves model input compared to an approach in which these species are simply ignored.

3.2 Unidentified compounds

The identities of several compounds remain unknown, especially at increasing mass where numerous structural and functional combinations are feasible. However, compared to earlier work at unit mass resolution (Warneke et al., 2011; Yokelson et al., 2013), the high-resolution capability of the PTR-TOF-MS has enhanced our ability to assign mass peaks while always identifying atomic composition. With unit mass resolution spectrometers, FTIR, and GC-MS grab samples, Yokelson et al. (2013) estimated that ~31% to ~72% of the gas-phase NMOC mass remained unidentified for several fuel types. For similar, commonly burned biomass fuels (chaparral, grasses, crop residue, etc.), considering a PTR-TOF range up to m/z 165, we estimate that ~7% of the detected NMOC mass remains unidentified, while ~12% is tentatively assigned using selection criteria described in Sect. 3.1. The compounds considered in this study cover a smaller mass range (up to m/z 165 rather than m/z 214) than in the earlier study, but in that earlier study, the compounds in the range m/z 165-214 accounted for only ~1.5% of the NMOC mass (Yokelson et al., 2013). Thus, the molecular formula assignments from the PTR-TOF aided in positive and tentative identification and quantification resulting in a reduction of the estimate of unidentified NMOCs from ~31% down to ~7%.

Calculations of unidentified and tentatively assigned emissions relative to overall NMOC emissions (including FTIR species) for several lumped fuel groups are summarized in Table 1. Estimates of total intermediate and semivolatile gas-phase organic compounds (IVOC + SVOC, estimated as the sum of species at or above the mass of toluene) are also included as these less volatile compounds are likely to generate SOA via oxidation and/or cooling. Similar to previous organic soil fire data, the percentages of unidentified and tentatively identified NMOCs for peat burns are significantly larger than for other fuel types (sum ~37%) and they could be a major source of impacts and uncertainty during El-Niño years when peat combustion is a major global emission source (Page et al., 2002; Akagi et al., 2011).

4 Discussion

For all fuel types, there is noticeable variability concerning which compounds have the most significant emissions. Figure 3 includes both FTIR and PTR emissions grouped into the following categories: non-methane hydrocarbons, oxygenates containing only one oxygen, oxygenates containing two oxygen atoms, and oxygenates containing three oxygen atoms. Within these categories, the contribution from aromatics, phenolic compounds, and furans are further indicated. As shown in Fig. 3, oxygenated compounds account for the majority of the emissions for all fuels where EF calculations were possible (several fuels are excluded including tires and plastic bags due to insufficient FTIR methanol data). Oxygenated compounds containing only a single oxygen atom accounted for ~50% of the total raw mass signal (> m/z 28, excluding m/z 37) on average and normally had greater emissions than oxygenated compounds containing two oxygen atoms or hydrocarbons. Sugar cane has the highest emissions of oxygenated compounds as was noted earlier in the FTIR data (Stockwell et al., 2014) and is one of the few fuels where the
emissions of compounds containing two oxygens are the largest. To facilitate discussion we grouped many of the assigned (or tentatively assigned) mass peak features into categories including: aromatic hydrocarbons; phenolic compounds; furans and its derivatives; nitrogen-containing compounds; and sulfur-containing compounds. These categories do not account for the majority of the emitted NMOC mass, but account for most of the rarely-measured species reported in this work. We then also discuss miscellaneous compounds at increasing m/z.

### 4.1 Aromatic hydrocarbons

Aromatic hydrocarbons contributed most significantly to the emissions for several major fuel types including ponderosa pine, peat, and black spruce. The identities of these ringed structures are more confidently assigned due to the small H to C ratio at high masses. The aromatics confidently identified in this study include benzene (m/z 79), toluene (m/z 93), phenylacetylene (m/z 103), styrene (m/z 105), xylenes/ethylbenzene (m/z 107), 1,3,5-trimethylbenzene (m/z 121), and naphthalene (m/z 129), while masses more tentatively assigned include dihydronaphthalene (m/z 131), p-cymene (m/z 135), and methylnaphthalenes (m/z 143). All masses are likely to have minor contributions from other hydrocarbon species. The EFs for aromatic species quantified during all fires are averaged by fuel type and shown in Fig. 4a. The EF for p-cymene was only calculated for select burns and has been included in Fig. 4a for comprehensiveness.

Aromatic structures are susceptible to multiple oxidation pathways and readily drive complex chemical reactions in the atmosphere that are highly dependent on hydroxyl radical (OH) reactivity (Phousongphouang and Arey, 2002; Ziemann and Atkinson, 2012; Phousongphouang and Arey, 2002). Ultimately these gas-phase aromatic species have high yields for SOA as their physical and chemical evolution lead to lower volatility species that condense into the particle phase. SOA yields from these parent aromatic HCs have been shown to strongly vary depending on environmental parameters including relative humidity, temperature, aerosol mass concentration, and particularly the level of nitrogen oxides (NOx) and availability of RO2 radicals, further adding to the complexity in modeling the behavior and fate of these compounds (Ng et al., 2007; Song et al., 2007; Henze et al., 2008; Chhabra et al., 2010, 2011; Im et al., 2014).

Biofuel and biomass burning together comprise the largest global atmospheric source of benzene (Andreae and Merlet, 2001; Henze et al., 2008; Andreae and Merlet, 2004), thus not surprisingly benzene is a significant aromatic in our dataset. The ERs relative to benzene for the aromatics listed above are shown in Table 2 and are positively correlated with benzene as demonstrated by Fig. 4b. Henze et al. (2008) outlines how ERs to CO of major aromatics (benzene, xylene, and toluene) can be implemented as a part of a model to predict SOA formation. An identical or similar approach that incorporates the additional aromatics detected by PTR-TOF-MS in this work may be useful to predict the contribution of aromatics from BB to global SOA by various reaction pathways.

Toluene, another major emission, often serves as a model compound to study the formation of SOA from other small ringed volatile organic compounds (Hildebrandt et al., 2009). Black spruce yielded the greatest toluene ER (to benzene) during FLAME-4 (3.24 ± 0.42) and has been linked to significant OA enhancement during chamber photo-
oxidation aging experiments investigating open biomass burning emissions during FLAME-III, though not significant enough to account for all of the observed SOA (Hennigan et al., 2011).

Naphthalene is the simplest species in a class of carcinogenic and neurotoxic compounds known as polycyclic aromatic hydrocarbons (PAH) and was detected from all fuels. The rapid rate of photo-oxidation of these smaller-ringed gas-phase PAHs (including naphthalene and methylnaphthalenes) can have important impacts on the amount and properties of SOA formed and yields significantly more SOA over shorter timespans in comparison to lighter aromatics (Chan et al., 2009). Under low-NOx conditions (BB events generate NOx though at lower ratios to NMOC and/or CO than those present in urban environments) the SOA yield for benzene, toluene, and m-xylene was ~30% (Ng et al., 2007), while naphthalene yielded enhancements as great as 73% (Chan et al., 2009).

In summary, many of the species identified and detected during FLAME-4 are associated with aerosol formation under diverse ambient conditions (Fisseha et al., 2004; Na et al., 2006; Ng et al., 2007; Chan et al., 2009). We present here initial emissions for a variety of aromatics from major global fuels. A more focused study to probe the extent and significance of SOA formation in BB plumes by these aromatic precursors was performed by chamber oxidation during the FLAME-4 campaign and will be presented in Tkacik et al. (2014).

4.2 Phenolic compounds

Phenol is detected at m/z 95. Earlier studies burning a variety of biomass fuels found that OP-FTIR measurements of phenol accounted for the observed PTR-MS signal at this mass even at unit mass resolution, though small contributions from other species such as vinyl furan were possible, but not detected (Christian et al., 2004). 2D-GC grab samples in FLAME-4 find that other species with the same formula (only vinyl furan) are present at levels less than ~2% of phenol (Hatch et al., 2014). Thus, we assume that within experimental uncertainty m/z 95 is a phenol measurement in this study and find that phenol is one of the most abundant oxygenated aromatic compounds detected. Several phenol derivatives were speciated for every fire and included catechol (m/z 111), vinylphenol (m/z 121), salicylaldehyde (m/z 123), xylenol (m/z 123), and guaiacol (m/z 125) (Fig. 5a). Several additional species were quantified for selected fires and included cresol (m/z 109), creosol (m/z 139), 3-methoxycatechol (m/z 141), 4-vinylguaiacol (m/z 151), and syringol (m/z 155). The EFs for these additional phenolic compounds were calculated for select burns and are included in Fig. 5a with the regularly analyzed compounds. Significant emissions of these compounds are reported in Table 2 relative to phenol and the selected compounds shown in Fig. 5b demonstrate the tight correlation between these derivatives and phenol.

Phenol, methoxyphenols (guaiacols), dimethoxyphenols (syringol), and their derivatives are formed during the pyrolysis of lignin (Simoneit et al., 1993) and can readily react with OH radicals leading to SOA formation (Coeur-Tourneur et al, 2010; Lauraguais et al., 2014). Hawthorne et al. (1989,1992) found that phenols and guaiacols accounted for 21% and 45% of aerosol mass from wood smoke, while Yee et al. (2013) noted large SOA yields for phenol (24-44%), guaiacol (44-50%), and syringol (25-37%) by photo-oxidation chamber experiments under low-NOx conditions (<10 ppb).
Softwoods are considered lignin-rich and are associated predominately with guaiacyl units (Shafizadeh, 1982). Thus not surprisingly, guaiacol emissions were significant for ponderosa pine. Peat, an accumulation of decomposing vegetation (moss, herbaceous, woody materials), has varying degrees of lignin-content depending on the extent of decomposition, sampling depth, water table levels, etc. (Williams et al., 2003). The peat burns all emitted significant amounts of phenolic compounds, with noticeable compound specific variability between regions (Indonesia, Canada, and North Carolina). It is also noteworthy that sugar cane, which also produced highly oxygenated emissions based on FTIR and PTR-TOF-MS results, had the greatest total emissions of phenolic compounds.

The photochemical formation of nitrophenols and nitroguaiacols by atmospheric oxidation of phenols and substituted phenols via OH radicals in the presence of NOx is a potential reaction pathway for these compounds (Atkinson et al., 1992; Olariu et al., 2002; Harrison et al., 2005; Lauraguais et al., 2014). Nitration of phenol in either the gas or aerosol phase is anticipated to account for a large portion of nitrophenols in the environment. Higher nitrophenol levels are correlated with increased plant damage (Hinkel et al., 1989; Natangelo et al., 1999) and consequently are linked to forest decline in central Europe and North America (Rippen et al., 1987). Nitrophenols are also important components of brown carbon and can contribute to SOA formation in biomass burning plumes (Kitanovski et al., 2012; Desyaterik et al., 2013; Mohr et al., 2013; Zhang et al., 2013; Kitanovski et al., 2012). Nitratated phenols including nitroguaiacols and methyl-nitrocatechols are suggested as suitable BB molecular tracers for secondary BB aerosol considering their reactivity with atmospheric oxidants is limited (Iinuma et al., 2010; Kitanovski et al., 2012; Lauraguais et al., 2014; Iinuma et al., 2010). The oxidation products from the phenolic compounds detected in fresh smoke here have not been directly examined and would require a more focused study beyond the scope of this paper.

As with the aromatic compounds, the ERs provided in Table 2 can be used to estimate initial BB emissions of phenolic species, both rarely measured or previously unmeasured, from a variety of fuels in order to improve atmospheric modeling of SOA and nitrophenol formation.

### 4.3 Furans

Other significant oxygenated compounds include furan and substituted furans which arise from the pyrolysis of cellulose and hemicellulose. The substituted furans regularly quantified included 2-methylfuran (m/z 83), 2-furanone (m/z 85), furfural (m/z 97), furfuryl alcohol (m/z 99), methylfurfural (m/z 111), benzo[5]furan (m/z 119), and hydroxymethylfurfural (m/z 127), while 2,5-dimethylfuran (m/z 97) and methylbenzofurans (m/z 133) were occasionally quantified. The ERs to furan for these compounds are summarized in Table 2 and Fig. 6a shows the average EF for the regularly quantified masses and the individual fire EFs for the occasionally quantified compounds.

Furan and substituted furans are oxidized in the atmosphere primarily by OH (Bierbach et al., 1995), but also by NO3 (Berndt et al., 1997) or Cl atoms (Cabañas et al., 2005; Villanueva et al., 2007; Cabañas et al., 2005). Photo-oxidation of furan, 2-methylfuran, and 3- methylfuran produce butenedial, 4-oxo-2-pentenal, and 2-methylbutenedial (Bierbach et al 1994, 1995). These products are highly reactive and can lead to free radical
In fact, aerosol formation from photo-oxidation chamber experiments has been observed for furans and their reactive intermediates listed above (Gomez Alvarez et al., 2009; Strollo and Ziemann, 2013). Even less is known concerning SOA yields from furans with oxygenated functional groups, which comprise the majority of the furan emissions in this study. Alvarado and Prinn (2009) added reaction rates for furans based on 2-methylfuran and butenedial values (Bierbach et al., 1994, 1995) to model O$_3$ formation in an aging savanna smoke plume. Although a slight increase in O$_3$ was observed after 60 min, it was not large enough to account for the observed O$_3$ concentrations in the plume. The furan and substituted furan ERs compiled here may help explain a portion of the SOA and O$_3$ produced from fires that cannot be accounted for based upon previously implemented precursors (Grieshop et al., 2009).

Furfural was generally the dominant emission in this grouping consistent with concurrent 2D-GC measurements (Hatch et al., 2014) while emissions from 2-furanone and furan also contributed significantly. Friedli et al. (2001) observed that ERs of alkyl furans linearly correlated with furan and concluded that these alkylated compounds likely break down to furan. Our expanded substituted furan list includes a variety of functionality ranging from oxygenated substituents to those fused with benzene rings for diverse fuel types. Similar to the behavior observed for alkylated furans, the emissions of our substituted furans linearly correlate with furan as shown in Fig. 6b. As noted for phenolic compounds, sugar cane produced the largest emissions of furans excluding Canadian peat, supporting sugar cane as an important emitter of oxygenated compounds. The emissions from furan, phenol, and their derivatives reflect variability in cellulose and lignin composition of different fuel types. Cellulose and hemicellulose compose ~75% of wood while lignin only accounts for ~25% on average (Sjöström, 1993). Accordingly the Σfurans/Σphenols for initially analyzed compounds indicate that furans are dominant in nearly every fuel type.

4.4 Nitrogen-containing compounds

Many nitrogen (N)-containing peaks were not originally selected for post-acquisition analysis in every fire. However, the additional analysis of selected fires included a suite of N-containing organic compounds to investigate their potential contribution to the N-budget and new particle formation (NPF). Even at our mass resolution of ~5000, the mass peak from N-compounds can sometimes be overlapped by broadened $^{13}$C “isotope” peaks of major carbon containing emissions. This interference was not significant for the following species that we were able to quantify in the standard or added analysis: C$_2$H$_3$N (acetonitrile, calibrated), C$_2$H$_5$N (dimethylamine; ethylamine), C$_2$H$_6$NO (acetamide), C$_3$H$_7$N (trimethylamine), C$_4$H$_8$NO (assorted amides), C$_5$H$_5$N (benzonitrile). As illustrated by the multiple possibilities for some formulas, several quantified nitrogen-containing species were observed but explicit single identities or relative contributions could not be confirmed. The logical candidates we propose are based upon atmospheric observations and include classes of amines and amides shown in Table S4 (Ge et al., 2011; Lobert et al., 1991; Schade and Crutzen, 1995; Ma and Hays et al., 2008; Barnes et al., 2010; Ge et al., 2011 Ma and Hays et al., 2008). Additional N-containing compounds were clearly observed in the mass spectra such as acrylonitrile, propanenitrile, pyrrole, and pyridine, but they were often overlapped with isotopic peaks of major carbon compounds, thus a time-intensive analysis would be necessary to provide quantitative data.
For the species in this category, quantification was possible for select fires by 2D-GC-MS and they are reported by Hatch et al. (2014) for the FLAME-4 campaign.

We present in Supplement Table S5 the abundance of each N-containing gas quantified by PTR-TOF-MS and FTIR relative to NH₃ for selected fires. The additional nitrogen-containing organic gases detected by PTR-TOF-MS for these 29 fires summed to roughly 22 ± 23% of NH₃ on average and accounted for a range of 0.1-8.7% of the fuel nitrogen. These compounds contributed most significantly to fuel N for peat and this varied by sampling location. This is not surprising since environmental conditions and field sampling depths varied considerably. Stockwell et al. (2014) reported large differences for N-containing compounds quantified by FTIR between FLAME-4 and earlier laboratory studies of emissions from peat burns. In any case, the additional NMOCs (including N-containing compounds) speciated by PTR-TOF-MS substantially increases the amount of information currently available on peat emissions.

The relevance of the N-containing organics to climate and the N-cycle is briefly summarized next. Aerosol particles acting as cloud condensation nuclei (CCN) critically impact climate by production and modification of clouds and precipitation (Novakov and Penner, 1993). NPF, the formation of new stable nuclei, is suspected to be a major contributor to the amount of CCN in the atmosphere (Kerminen et al., 2005; Laaksonen et al., 2005; Sotiropoulou et al., 2006). Numerous studies have suggested that organic compounds containing nitrogen can play an important role in the formation and growth of new particles (Smith et al., 2008; Kirkby et al., 2011; Yu and Luo, 2014). The primary pathways to new particle formation include (1) reaction of organic compounds with each other or atmospheric oxidants to form higher molecular weight, lower volatility compounds that subsequently partition into the aerosol phase or (2) rapid acid/base reactions forming organic salts. The observation of significant emissions of N-containing organic gases in FLAME-4 could improve understanding of the compounds, properties, and source strengths contributing to new particle formation and enhance model predictions on local to global scales. The identities and amounts of these additional nitrogen containing emissions produced by peat and other BB fuels are also important in rigorous analysis of the atmospheric nitrogen budget.

### 4.5 Sulfur, phosphorous, and chlorine-containing compounds

Sulfur emissions are important for their contribution to acid deposition and climate effects due to aerosol formation. Several S-containing gases have been detected in BB emissions including SO₂, carbonyl sulfide (OCS), dimethylsulphide (DMS), and dimethyl disulphide (DMDS), where DMS is one of the most significant organosulfur compounds emitted by BB and is quantified by PTR-TOF-MS in our primary dataset (Friedli et al., 2001; Meinardi et al., 2003; Akagi et al., 2011; Meinardi et al., 2003; Friedli et al., 2001; Simpson et al., 2011). The signal at m/z 49 had a significant mass defect and is attributed to methanethiol (methyl mercaptan, CH₃SH), which to our knowledge has not been previously reported in real-world BB smoke though it has been observed in cigarette smoke (Dong et al., 2010) and in emissions from pulp and paper plants (Toda et al., 2010). Like DMS, the photochemical oxidation of CH₃SH leads to SO₂ formation (Shon and Kim, 2006), which can be further oxidized to sulfate or sulfuric acid and contribute to the aerosol phase. The emissions of CH₃SH are dependent on the fuel S-content and are
negatively-correlated with MCE. The greatest EF(CH$_3$SH) in our additional analyses arose from organic alfalfa, which had the highest S-content of the selected fuels and also produced significant emissions of SO$_2$ detected by FTIR.

Other organic gases containing chlorine and phosphorous were expected to be readily detectable because of their large, unique mass defects and possible enhancement by pesticides and fertilizers in crop residue fuels. However, they were not detected in significant amounts by our full mass scans. Fuel P and Cl may have been emitted primarily as aerosol, ash, low proton affinity gases, or as a suite of gases that were evidently below our detection limit.

4.6 Miscellaneous (order of increasing m/z)

$m/z$ 41: The assignment of propyne is reinforced by previous observations in BB fires, and it is of some interest as a BB marker even though it has a relatively short lifetime of ~2 days (Simpson et al., 2011; Akagi et al., 2013; Yokelson et al., 2013). Considering that propyne was not detected in every fuel type, a level of uncertainty is added to any use of this compound as a BB tracer and in general, the use of multiple tracers is preferred when possible.

$m/z$ 43: The high-resolution capabilities of the PTR-TOF-MS allowed propylene to be distinguished from ketene fragments at $m/z$ 43. The propylene concentrations are superseded in our present dataset by FTIR measurements, however, the two techniques agree well.

$m/z$ 45: PTR technology has already been reported as a reliable way to measure acetaldehyde in BB smoke (Holzinger et al., 1999; Christian et al., 2004; Holzinger et al., 1999). Photolysis of acetaldehyde can play an important role in radical formation and is the main precursor of peroxy acetyl nitrate (PAN) (Trentmann et al., 2003). A wide range in EF(acetaldehyde) (0.13-4.3 g kg$^{-1}$) is observed during FLAME-4 and reflects variability in fuel type. The detailed emissions from a range of fuels in this dataset can aid in modeling and interpretation of PAN formation in aging BB plumes of various regions (Alvarado et al., 2010, 2013). Crop-residue fuels regularly had the greatest emissions of acetaldehyde, which is important considering many crop-residue fires evade detection and are considered both regionally and globally underestimated. Sugar cane burning had the largest acetaldehyde EF (4.3 ± 1.4 g kg$^{-1}$) and had significant emissions of oxygenated and N-containing compounds, consequently it is likely to form a significant amount of PAN.

$m/z$ 57: The signal at $m/z$ 57 using unit-mass resolution GC-PTR-MS was observed to be primarily acrolein with minor contributions from alkenes (Karl et al., 2007). In the PTR-TOF-MS, the two peaks at $m/z$ 57 (C$_3$H$_5$O$^+$ and C$_4$H$_9^+$) are clearly distinguished and acrolein is often the dominant peak during the fire with the highest emissions from ponderosa pine and sugar cane.

$m/z$ 69: The high resolution of the PTR-TOF-MS allowed three peaks to be distinguished at $m/z$ 69, identities attributed to carbon suboxide (C$_3$O$_2$), furan (C$_4$H$_4$O), and mostly isoprene (C$_5$H$_8$) (Fig. 7). Distinguishing between isoprene and furan is an important capability of the PTR-TOF-MS. The atmospheric abundance and relevance of carbon suboxide is fairly uncertain and with an atmospheric lifetime of ~10 days (Kessel et al., 2013) the reactivity...
and transport of CO$_2$ emitted by fires could have critical regional impacts. The emissions of CO$_2$ by BB will be interpreted in detail at a later date (S. Kessel, personal communication, 2014).

$m/z$ 75: Hydroxyacetone emissions have been reported from both field and laboratory fires (Christian et al., 2003; Akagi et al., 2011; Yokelson et al., 2013; St. Clair et al., 2014; Yokelson et al., 2013; Akagi et al., 2011; Christian et al., 2003). Christian et al. (2003) first reported BB emissions of hydroxyacetone, and noted very large quantities from burning rice straw. The EF(C$_3$H$_6$O$_2$) for rice straw was noticeably high (1.10 g kg$^{-1}$) in the FLAME-4 dataset and only sugar cane had greater emissions.

$m/z$ 85, 87: The largest peak at $m/z$ 85 was assigned as pentenone as it was monitored/confirmed by PIT-MS/GC-MS in an earlier BB study (Yokelson et al., 2013). Pentenone was a substantial emission from several fuels with ponderosa pine having the greatest EF. By similar evidence the minor peak at $m/z$ 87 was assigned to pentanone but was only detected in a few of the fires in the second set of analyses with the most significant emissions arising from Indonesian peat.

$m/z$ 107: Benzaldehyde has the same unit mass as xylenes, but is clearly separated by the TOF-MS. Greenberg et al. (2006) observed benzaldehyde during low temperature pyrolysis experiments with the greatest emissions from ponderosa needles (ponderosa pine produced the greatest EF in our dataset, range 0.10-0.28 g kg$^{-1}$). Benzaldehyde emissions were additionally quantified by GC-MS during a laboratory BB campaign and produced comparable EF to that of xylenes (Yokelson et al., 2013). During FLAME-4 the $\text{EF(benzaldehyde)}$ was comparable to $\text{EF(xylenes)}$ calibrated as p-xylene as seen earlier except for peat burns where p-xylene was significantly higher.

$m/z$ 137: At unit mass resolution the peak at $m/z$ 137 is commonly recognized as monoterpenes which can further be speciated by GC-MS. However, as shown in Fig. 8 there can be up to three additional peaks at this mass that presently remain unidentified oxygenated compounds. As anticipated, the hydrocarbon monoterpene peak is significant for coniferous fuels such as ponderosa pine but much smaller for grasses. In this work we calibrated for $\alpha$-pinene, which has been reported as a major monoterpene emission from fresh smoke (Akagi et al., 2013; Simpson et al., 2011; Akagi et al., 2013).

4.7 Cookstoves

Trace gas emissions were measured for four cookstoves including: a traditional 3-stone cooking fire, the most widely used stove design worldwide; two “rocket” type designs (Envirotif G3300 and Ezy stove); and a “gasifier” stove (Philips HD4012). Several studies focus on fuel efficiency of cookstove technology (Jetter et al., 2012), while the detailed emissions of many rarely measured and previously unmeasured gases are reported here and in Stockwell et al. (2014) for FLAME-4 burns. For cooking fires, ~3-6% of the NMOC mass remained unidentified, with the Envirotif rocket stove design generating the smallest percentage in the study. To improve the representativeness of our laboratory open cooking emissions, the EFs of smoldering compounds reported for 3-stone cooking fires were adjusted by multiplying the mass ratio of each species “X” to CH$_4$ by the literature-average field $\text{EF(CH}_4)$ for open
cooking in Akagi et al. (2011). Flaming compounds were adjusted by a similar procedure based on their ratios to CO₂. The preferred values are reported in Table S3. With these adjustments, the emissions of aromatic hydrocarbons (Fig. 9a), phenolic compounds (Fig. 9b), and furans (Fig. 9c) distinctively increased with the primitiveness of design, thus, 3-stone cooking fires produced the greatest emissions. The advancement in emissions characterization for these sources will be used to upgrade models of exposure to household air pollution and the ERs/EFs should be factored in to chemical-transport models to assess atmospheric impacts.

BB is an important source of reactive nitrogen in the atmosphere producing significant emissions of NOₓ and NH₃ while non-reactive HCN and CH₃CN are commonly used as BB marker compounds (Yokelson et al., 1996, 2007; Goode et al., 1999; de Gouw et al., 2003). The FTIR used in FLAME-4 provided the first detection of HCN emissions from cooking fires and the HCN/CO ER was about a factor of 5 lower than most other BB fuels burned (Stockwell et al., 2014). Similarly, acetonitrile emissions were measured for the first time for cooking fires by PTR-TOF-MS in this study and the CH₃CN/CO ERs from cooking fires are much lower (on average a factor of ~15) than those from other fuels. This should be considered when using CH₃CN/CO ERs to drive source apportionment in areas with substantial emissions from biofuel cooking sources.

5 Conclusions

We investigated the primary BB NMOC emissions from laboratory simulated burns of globally significant fuels using a PTR-TOF-MS instrument. In this first PTR-TOF-MS deployment dedicated to fires we encountered some specific challenges. The fast change in concentration necessitated a fast acquisition rate, which decreases the signal to noise for the emissions above background. The large dynamic concentration range necessitated dilution to minimize reagent ion depletion at peak emissions and the dilution further reduced the signal to noise ratio. Positive identification of some species by co-deployed grab sampling techniques will be explored further in a separate paper, but is challenged by the difficulty of transmitting some important fire emissions through GC columns (Hatch et al., 2014). We attempted to enhance compound identification by switching reagent ions (O₂⁺ and NO⁺), however, this approach with two broadly sensitive ions in a complex mixture resulted in spectra with complexity whose comparative analysis is beyond the scope of the present effort. Future experiments might consider instead using a less broadly sensitive reagent ion such as NH₃⁺ as the alternate reagent ion. We were limited to our pre-chosen calibration mixture based primarily on gases previously observed in smoke. For future experiments we suggest adding more standards to generate more accurate calibration factors, specifically including major species such as furan and phenol and more compounds with S and N heteroatoms. In addition, measuring the fragmentation, if any, of more of the species identified in this work would be of great value. Despite these practical limitations, the experiment produced a great deal of useful new information.

The PTR-TOF-MS obtains full mass scans of NMOCs with high enough resolution to distinguish multiple peaks at the same nominal mass and high enough accuracy to assign chemical formulas from the “exact” masses. This aided in compound identification and more than 100 species were categorized as a confirmed identity, a tentative (most likely) assignment, or unidentified but with a chemical formula. Chemical identification was aided by observations
of compounds reported in smoke emissions, pyrolysis experiments, and those species at relevant concentrations in the atmosphere. This allowed the identification of more masses up to m/z 165 than in earlier work at unit mass resolution though an estimated range of 12-37% of the total mass still remains unidentified and tentatively identified. The analysis provides a new set of emission factors for ~68 compounds in all fires plus ~50 more in select fires, in addition to species previously quantified by FTIR (Stockwell et al., 2014) and other techniques during FLAME-4 (Hatch et al., 2014). While significant variability was observed between fuels, oxygenated compounds collectively accounted for the majority of emissions in all fuels with sugar cane producing the highest EF of oxygenated species on average possibly due to its high sugar content.

We also report emission ratios to benzene, phenol, or furan for the aromatic hydrocarbons, phenolic compounds, and substituted furans, respectively. Reporting emissions of previously unmeasured or rarely measured compounds relative to these more regularly measured compounds facilitates adding several new compounds to fire emissions models. To our knowledge this is the first on-line, real-time characterization of several compounds within these “families” for biomass burning. Emissions were observed to vary considerably between fuel types. Several example compounds within each class (i.e. toluene, guaiacol, methylfuran, etc.) have been shown, by chamber experiments, to be highly reactive with atmospheric oxidants and contribute significantly to SOA formation. The ERs and EFs characterized by PTR-TOF-MS of fresh BB smoke are presented in Tables S1-S3 and (especially the recommended values in Table S3) should aid model predictions of O₃ and SOA formation in BB smoke and the subsequent effects on air quality and climate on local-global scales.

A large number of organic nitrogen-containing species were detected with several identities speculated as amines or amides. These N-containing organic gases may play an important role in new particle formation by physical, chemical, and photochemical processes, though a more focused study is necessary to measure NPF yields from these compounds and processes. The additional N-containing gases detected here account for a range of 1-87% of NH₃ dependent on fuel type with the most significant contribution of additional N-species to fuel N arising from peat burns. The ERs of acetonitrile to CO for cooking fires were significantly lower than other fuels and should be factored into source apportionment models in regions where biofuel use is prevalent if CH₃CN is used as a tracer.

The S-containing compounds detected by PTR-TOF-MS included dimethyl sulfide and methanethiol, where methanethiol was detected for the first time in BB smoke to our knowledge. These compounds may play a role in acid deposition and aerosol formation though to what extent has yet to be extensively studied. Phosphorous and chlorine organic gases were not readily detected in our dataset, which may reflect that these species were below our detection limit.

Using full mass scans from a high resolution PTR-TOF-MS to characterize fresh smoke has aided in identification of several compounds and provided the chemical formula of other organic trace gases. The additional NMOCs identified in this work are important in understanding fresh BB emissions and will improve our understanding of BB atmospheric impacts. The subsequent oxidation products of these gases are the focus of a companion paper probing
BB aging. Taken together, this work should improve BB representation in atmospheric models, particularly the formation of ozone and secondary organic aerosol at multiple scales.

Acknowledgements

FLAME-4, rental of PTR-TOF-MS, C. S. and R. Y. were supported primarily by NSF grant ATM-0936321. FSL operational costs were supported by NASA Earth Science Division Award NNX12AH17G to S. Kreidenweis, P. DeMott, and G. McMeeking whose collaboration in organizing and executing FLAME-4 is gratefully acknowledged. The collaboration of A. Robinson in organizing FLAME-4, and the cooking fires is also gratefully acknowledged. We thank C. Geron for providing a sample of NC peat.

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**Figure 1.** (a) The normalized response of calibration factors (“CF,” ncps/ppbv) versus mass (calibrated species labeled by name) overlaid with the linearly fitted mass-dependent transmission curve (black markers and dotted line). Separate linear approximations (b) oxygenated (blue) and (c) hydrocarbon (green) species used to calculate approximate calibration factors for all observed masses where explicit calibrations were not available.

**Figure 2.** A typical full mass scan of biomass burning smoke from the PTR-TOF-MS on a logarithmic (a) and a smaller range linear (b) scale. The internal standard (1,3-diiodobenzene) accounts for the major peaks ~m/z 331 and fragments at peaks near m/z 204 and 205.

**Figure 3.** The emission factors (g kg$^{-1}$) of total observed hydrocarbons and total observed species oxygenated to different degrees averaged for each fire type based on a synthesis of PTR-TOF-MS and OP-FTIR data. The patterned sections indicate the contribution to each of the above categories by selected functionalities discussed in the text (aromatic hydrocarbons, phenolics, furans). The parenthetical expressions indicate how many oxygen atoms are present. The distribution of oxygenated and hydrocarbon emission factors averaged for each fuel type where FTIR data were available. EFs are included for FTIR compounds and the 68 masses initially analyzed by PTR-TOF-MS. The patterned sections indicate the contributions from various “families” based on functionality and their oxygen content.

**Figure 4.** (a) The EFs of the aromatics analyzed in all fires averaged and shown by fuel type. Individual contributions from benzene and other aromatics are indicated by color. The EFs for p-Cymene are only calculated for select burns and should not be considered an average for each particular fuel type. (b) The correlation plots of selected aromatics with benzene during a black spruce fire (Fire 74). Similar behavior was observed for all other fuel types.

**Figure 5.** (a) The distribution in average fuel EF for several phenolic compounds, where compound specific contributions are indicated by color. The EFs for compounds additionally analyzed are only calculated for select fires and should not be considered the fuel average. (b) The linear correlation of select phenolic compounds with phenol during an organic hay burn (Fire 119).

**Figure 6.** (a) The distribution in average fuel EF for furan and substituted furans, where individual contributions are indicated by color. The EFs for substituted furans additionally analyzed are not true averages. The EFs for additionally analyzed substituted furans are also included but should not be considered fuel averages. (b) The linear correlation of furan with select furan derivatives for an African grass fire (Fire 49).

**Figure 7.** Expanded view of the PTR-TOF-MS spectrum at m/z 69 demonstrating the advantage over unit mass resolution instruments of distinguishing multiple peaks, in this instance separating carbon suboxide (C$_2$O$_2$), furan (C$_4$H$_4$O), and mostly isoprene (C$_5$H$_8$) in ponderosa pine smoke (fire 70).

**Figure 8.** Expanded view of the PTR-TOF-MS spectrum of NC peat (fire 61) at m/z 137 showing multiple peaks.

**Figure 9.** Emission factors (g kg$^{-1}$) of aromatic hydrocarbons (a), phenolic compounds (b), and furans (c), for traditional and advanced cookstoves. The EF for traditional stoves were adjusted from original lab data (Sect. 4.7).
Table 1. Quantities for various categories of compounds (g kg\(^{-1}\)) and calculation of mass ratios and/or percentages for several fuel types.

<table>
<thead>
<tr>
<th>Quantity or Ratio</th>
<th>Chaparral</th>
<th>Coniferous Canopy</th>
<th>Peat</th>
<th>Grasses</th>
<th>Cooking Fires</th>
<th>Crop Residue</th>
<th>Trash</th>
</tr>
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<tr>
<td>Σ NMOCs</td>
<td>13.1</td>
<td>23.9</td>
<td>40.5</td>
<td>5.17</td>
<td>8.16</td>
<td>29.6</td>
<td>7.13</td>
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<tr>
<td>Σ I/SVOCs(^a)</td>
<td>3.49</td>
<td>7.13</td>
<td>14.6</td>
<td>1.38</td>
<td>1.33</td>
<td>7.21</td>
<td>1.83</td>
</tr>
<tr>
<td>Σ Tentatively assigned NMOCs</td>
<td>1.43</td>
<td>2.77</td>
<td>7.01</td>
<td>0.72</td>
<td>0.72</td>
<td>4.38</td>
<td>0.51</td>
</tr>
<tr>
<td>Σ Unidentified NMOCs</td>
<td>1.23</td>
<td>1.79</td>
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<td>0.39</td>
<td>0.33</td>
<td>2.10</td>
<td>0.41</td>
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<tr>
<td>Σ (I/SVOCs) / Σ NMOC</td>
<td>0.21</td>
<td>0.28</td>
<td>0.37</td>
<td>0.26</td>
<td>0.15</td>
<td>0.24</td>
<td>0.26</td>
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<tr>
<td>Percent NMOCs Tentatively assigned</td>
<td>8.35</td>
<td>9.74</td>
<td>17.5</td>
<td>13.9</td>
<td>8.19</td>
<td>14.0</td>
<td>7.20</td>
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<td>Percent NMOCs Unidentified</td>
<td>7.24</td>
<td>6.75</td>
<td>19.5</td>
<td>7.19</td>
<td>3.77</td>
<td>6.90</td>
<td>5.75</td>
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<td>Percent NMOCs Tentatively + Unidentified</td>
<td>16</td>
<td>16</td>
<td>37</td>
<td>21</td>
<td>12</td>
<td>21</td>
<td>13</td>
</tr>
</tbody>
</table>

\(^a\) See section 3.2 for definition
<table>
<thead>
<tr>
<th>ER/Benzene</th>
<th>Fuel Type (# burns)</th>
<th>Grasses (42)</th>
<th>Coniferous Canopy (14)</th>
<th>Chaparral (8)</th>
<th>Peat (6)</th>
<th>Crop Residue (food, 19)</th>
<th>Crop Residue (feed, 9)</th>
<th>Open 3-Stone Cooking (3)</th>
<th>Rocket Cookstoves (5)</th>
<th>Gasifier Cookstove (1)</th>
<th>Trash (2)</th>
<th>Tires (1)</th>
<th>Plastic Bags (1)</th>
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<tr>
<td>Toluene</td>
<td>C6H5</td>
<td>0.44(0.26)</td>
<td>2.19(0.84)</td>
<td>0.49(0.17)</td>
<td>0.53(0.17)</td>
<td>0.70(0.22)</td>
<td>1.00(0.44)</td>
<td>0.95(0.029)</td>
<td>0.98(1.39)</td>
<td>0.24</td>
<td>0.41(0.20)</td>
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<td>Phenylacetylene</td>
<td>C6H5</td>
<td>0.094(0.022)</td>
<td>0.13</td>
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<td>-</td>
<td>0.65(0.45)</td>
<td>0.14(0.09)</td>
<td>0.10(0.05)</td>
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<td>0.11(0.02)</td>
<td>0.074(0.020)</td>
<td>0.087(0.027)</td>
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<td>0.11(0.05)</td>
<td>0.10(0.05)</td>
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<td>0.95(0.017)</td>
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<td>0.32(0.16)</td>
<td>0.20(0.08)</td>
<td>0.24(0.11)</td>
<td>0.052(0.034)</td>
<td>0.10(0.05)</td>
<td>0.048</td>
<td>0.86(0.16)</td>
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<td>0.11(0.03)</td>
<td>0.043(0.023)</td>
<td>0.17(0.08)</td>
<td>0.11(0.05)</td>
<td>0.11(0.06)</td>
<td>0.014(0.007)</td>
<td>0.05(0.048)</td>
<td>0.026</td>
<td>0.033(0.016)</td>
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<td>0.18(0.16)</td>
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<td>0.10(0.03)</td>
<td>0.15(0.09)</td>
<td>0.20(0.17)</td>
<td>0.18(0.11)</td>
<td>0.21(0.05)</td>
<td>0.30(0.17)</td>
<td>0.12</td>
<td>0.10</td>
<td>0.19</td>
<td>0.059</td>
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<td>Dihydronaphthalene</td>
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<td>0.034(0.016)</td>
<td>0.020(0.010)</td>
<td>0.050(0.019)</td>
<td>0.059(0.028)</td>
<td>0.051(0.021)</td>
<td>0.019(0.006)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>9.81E-03</td>
<td>-</td>
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<td>p-Cymene</td>
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<td>0.11(0.01)</td>
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<td>nm</td>
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<td>nm</td>
<td>nm</td>
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<td>Methyl Naphthalenes</td>
<td>C6H10,10</td>
<td>0.032(0.009)</td>
<td>0.053(0.005)</td>
<td>0.033(0.007)</td>
<td>-</td>
<td>0.19(0.09)</td>
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<td>-</td>
<td>-</td>
<td>nm</td>
<td>-</td>
<td>0.031</td>
<td>-</td>
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</table>

**Note:** “nm” indicates not measured; blank indicates species remained below the detection limits; values in parenthesis indicate one standard deviation.

* Significant contributions from both methylfurfural and catechole reported in pyrolysis reference papers, thus there is no indication which species is the major contributor at this mass.

### Table 2. Emission ratios to benzene, phenol, and furan for aromatic hydrocarbons, phenolic compounds, and substituted furans in lumped fuel-type categories.
Figure 1. (a) The normalized response of calibration factors ("CF," ncp/ppbv) versus mass (calibrated species labeled by name) overlaid with the linearly fitted mass-dependent transmission curve (black markers and dotted line). Separate linear approximations (b) oxygenated (blue) and (c) hydrocarbon (green) species used to calculate approximate calibration factors for all observed masses where explicit calibrations were not available.
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Figure 4. (a) The EFs of the aromatics analyzed in all fires averaged and shown by fuel type. Individual contributions from benzene and other aromatics are indicated by color. The EFs for p-Cymene are only calculated for select burns and should not be considered a true average. (b) The correlation plots of selected aromatics with benzene during a black spruce fire (Fire 74). Similar behavior was observed for all other fuel types.
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Figure 6. (a) The distribution in average fuel EF for furan and substituted furans, where individual contributions are indicated by color. The EFs for substituted furans additionally analyzed a single time are not true averages. (b) The linear correlation of furan with select substituted furans for an African grass fire (Fire 49).
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