

This discussion paper is/has been under review for the journal Atmospheric Chemistry and Physics (ACP). Please refer to the corresponding final paper in ACP if available.

Investigations of primary and secondary particulate matter of different wood combustion appliances with a high-resolution time-of-flight aerosol mass spectrometer

M. F. Heringa¹, P. F. DeCarlo^{1,*}, R. Chirico^{1,**}, T. Tritscher¹, J. Dommen¹, E. Weingartner¹, R. Richter¹, G. Wehrle¹, A. S. H. Prevot¹, and U. Baltensperger¹

¹Laboratory of Atmospheric Chemistry, Paul Scherrer Institut, 5232 Villigen PSI, Switzerland now at: AAAS Science and Technology Policy Fellow, US EPA, Washington, DC, USA now at: Italian National Agency for New Technologies, Energy and Sustainable Economic Development (ENEA), UTAPRAD-DIM, Via E. Fermi 45, 00044 Frascati, Italy

Received: 18 February 2011 - Accepted: 27 February 2011 - Published: 9 March 2011

Correspondence to: A. S. H. Prevot (andre.prevot@psi.ch)

Published by Copernicus Publications on behalf of the European Geosciences Union.

Discussion Paper

Discussion

Paper

Discussion Paper



Back

Printer-friendly Version

Interactive Discussion



11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

ACPD

M. F. Heringa et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures

Close

Full Screen / Esc

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



A series of photo-oxidation smog chamber experiments were performed to investigate the primary emissions and secondary aerosol formation from two different log wood burners and a residential pellet burner under different burning conditions: starting and flaming phase. Emissions were sampled from the chimney and injected into the smog chamber leading to primary organic aerosol (POA) concentrations comparable to ambient levels. The composition of the aerosol was measured by an Aerodyne high resolution time-of-flight aerosol mass spectrometer (HR-TOF-AMS) and black carbon (BC) instrumentation. The primary emissions were then exposed to xenon light to initiate photo-chemistry and subsequent secondary organic aerosol (SOA) production. After correcting for wall losses, the average increase in organic matter (OM) concentrations by SOA formation for the starting and flaming phase experiments with the two logwood burners was found to be a factor of 4.1 ± 1.4 after five hours of aging. No SOA formation was observed for the stable burning phase of the pellet burner. The startup emissions of the pellet burner showed an increase in OM concentration by a factor of 3.3. Average emission factors of BC + POA + SOA, calculated from CO₂ emission, were found to be in the range of 0.04 to 3.9 g kg⁻¹ wood for the stable burning pellet burner and an old log wood burner during startup respectively. SOA contributed significantly to the ion $C_2H_4O_2^+$ at mass to charge ratio m/z 60, a commonly used marker for primary emissions of wood burning. The primary organic emissions from the three different burners showed a wide range in O/C atomic ratio (0.19-0.60) for the starting and flaming conditions, which also increased during aging. Primary wood burning emissions have a rather low relative contribution at m/z 43 (f43) to the total organic mass spectrum. The non-oxidized fragment C₃H₇⁺ has a considerable contribution at m/z 43 for the fresh OA with an increasing contribution of the oxygenated ion $C_2H_3O^+$ during aging. After five hours of aging, the OA has a rather low C₂H₃O⁺ signal for a given CO₂ fraction, possibly indicating a higher ratio of acid to non-acid oxygenated compounds in wood burning OA compared to other OOA.

ACPD

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures







Back

Wood burning for domestic heating and cooking is very common in many parts of the world, and, because of its importance as a source of renewable energy, there is currently an increasing interest in its usage. By estimation, three billion people use small-scale appliances (three-stone fires or cooking stoves) that are both inefficient and highly polluting (World Energy Council, 2007). Wood combustion emits a wide range of volatile organic compounds (VOC) and particulate matter (PM) which consists mainly of organics and black carbon (BC) (McDonald et al., 2000; Schauer et al., 2001). On a global scale, biomass burning is estimated to contribute up to 90% of the combustion generated primary particulate organic carbon (OC) (Bond et al., 2004). Wood burning is responsible for more than 30% of the particulate carbonaceous matter in Europe (Simpson et al., 2007) and was shown to be the dominant source of PM in the Swiss Alpine valleys in winter (Sandradewi et al., 2008). These gas phase and PM emissions contribute to various adverse effects on human health (Nel, 2005; Pope and Dockery, 2006).

Organic aerosol (OA) is an important fraction of the non-refractory PM_1 (PM with an aerodynamic diameter $d_a < 1 \mu m$) in many parts of the world (Zhang et al., 2007). The OA mass spectra from the Aerodyne aerosol mass spectrometer (AMS) can be deconvolved to give information about sources contributing to the total OA mass (Lanz et al., 2007). Hydrocarbon-like OA (HOA) from traffic, biomass burning OA (BBOA), and oxygenated OA (OOA) were distinguished in many datasets where secondary organic aerosol (SOA) is assumed to be the main contributor to OOA (Lanz et al., 2010; Jimenez et al., 2009). OOA can be further separated into a low volatility fraction (LVOOA), which is more aged and exhibits a higher O/C ratio, and a semi-volatile fraction (SV-OOA) which represents fresher OOA with a lower O/C ratio (Lanz et al., 2007; Ng et al., 2010).

Levoglucosan was found to be a substantial constituent of primary organic aerosol (POA) particles from biomass burning (Fraser and Lakshmanan, 2000). The AMS

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I₫

►I

Close

- 4

•

Back

Full Screen / Esc

Printer-friendly Version



11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I ← ▶I

← Back Close

Printer-friendly Version

Full Screen / Esc

Interactive Discussion

© **()**

mass spectrum of nebulized levoglucosan is highly correlated with ambient measurements at a wood burning dominated site in Roveredo, Switzerland. Three fragments, m/z 60, 73, and 137, have been suggested as marker fragments for wood burning generated aerosols (Alfarra et al., 2007). The fragment at m/z 60 was observed for open burning (Lee et al., 2010) as well as for residential wood burners where the markers were found to depend on burning conditions and wood type (Weimer et al., 2008).

Emissions from a small wood burning appliance have been shown to form significant amounts of SOA during irradiation with UV light (Grieshop et al., 2009). Such an aged OA closely resembles the OOA spectrum, which makes it difficult to extract the total contribution of wood burning in ambient OA. Smog chamber experiments are suitable to give insight into the properties of POA and to characterize the SOA from wood burning exhaust.

In this paper, we describe the POA, the SOA production, and OA aging processes from three different residential wood burning appliances during the startup and flaming phases using measurements from a high resolution time-of-flight aerosol mass spectrometer (HR-TOF-AMS). SOA production and estimated emission factors for BC, POA, and SOA are reported for all three burners separated by starting phase and flaming phase. We present a correlation between O/C ratio and m/z 44 obtained from unit mass resolution (UMR) mass spectra, which can be used to estimate the O/C ratio of wood burning aerosols, POA and SOA, from UMR data. In addition, results from pure SOA experiments, where only the gas phase emissions are injected and processed, are presented.

2 Materials and methods

2.1 Experimental setup

The experiments were performed in the smog chamber of the Paul Scherrer Institut (PSI). The PSI chamber is a 27-m³ Teflon bag suspended in a temperature controlled

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

ACPD

M. F. Heringa et al.

Title Page **Abstract** Introduction Conclusions

References

Figures Tables







Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



housing. The smog chamber was operated at 20°C and a relative humidity (RH) of ~50% during all experiments described here. Four xenon arc lamps of 4 kW each were used to simulate the atmospheric light spectrum and intensity during a typical Swiss winter day at noon. The smog chamber housing is covered with a reflecting coating to increase light intensity and light diffusion. A more detailed description of the PSI smog chamber and its standard instrumentation can be found elsewhere (Paulsen et al., 2005).

The experiments were performed with two different residential log wood burners and one residential pellet burner. The first burner is an old style log wood burner (WESO Duplex, built around 1960) with a small (~0.018 m³) combustion chamber. The second burner is a modern log wood burner (Attika Avant, 2009, combustion chamber ~0.037 m³) with well defined air flows to reach optimal combustion efficiencies. The wood used for these burners, 100% beech, was obtained from a local distributor, cut into ~400 g pieces and stored in a dry place. The logs were positioned on top of the kindling and lit from below. Emissions were injected into the smog chamber either during starting, flaming or smoldering phases of the burning cycle. For the starting phase experiments the kindling was lit using a small gas torch immediately before chamber injection. Experiments on the gas phase only emissions were performed by filtering the hot emissions through a heated filter (150°C). These experiments are referred to as pure SOA experiments throughout the rest of this paper and are a combination of starting and flaming phase emissions. The pellet burner (Rüegg KEA, 2005, Pmax 9 kW) was operated at 80% of its maximum power using wood pellets. The burners were connected to a chimney with a diameter of 0.16 m and a height of 3.5 m.

The introduction of the wood burning emissions into the chamber was done from a pickoff point in the middle of the chimney at a height of approximately 2 m. To avoid particle coagulation, the emissions were diluted by a factor of ~7 using a heated ejector diluter (Dekati Ltd., Tampere, Finland). The preheated dilution air was provided by a pure air generator (737-250 series, AADCO Instruments, Inc., USA). The dilution system, dilution air, and all sampling lines were usually operated at 150 °C to prevent

Printer-friendly Version

condensation of semi-volatile organic compounds (SVOC's). Finally, the emissions experienced a second dilution (~220 times) and cooling to 20°C when entering the smog chamber. A schematic representation of the inlet system and the smog chamber setup is shown in Fig. 1. The filling time and/or the flow into the smog chamber were varied to achieve atmospherically relevant concentrations (1-30 µg m⁻³) of POA. The smoldering phase emissions generally produced low POA emissions and therefore only one smoldering experiment, which showed a high organic emission, is described in this paper.

After injection and a 15-min homogenization/mixing period the primary emissions were measured for 30-60 min before the xenon lamps were switched on to start photooxidation. After each experiment the smog chamber was cleaned by the addition of several ppm of ozone for ~3 h and flushing the chamber with zero air for at least 36 h. Blank experiments were performed to make sure that the organic matter (OM) produced during the experiments is not significantly influenced by background impurities in the smog chamber. An overview of the experiments described in this paper can be found in Table 1.

Instrumentation

High resolution time of flight aerosol mass spectrometer

An Aerodyne high resolution time-of-flight aerosol mass spectrometer (HR-TOF-AMS) was used for the on-line quantification of the submicron (PM₁) non-refractory aerosol components. The term "non-refractory" is assigned to those species that evaporate rapidly at 600 °C under vacuum conditions, e.g. OM, NH₄NO₃ and (NH₄)₂SO₄ (Allan et al., 2004). Potassium salts like KNO₃ or K₂SO₄ can not be quantitatively measured with the AMS. A detailed description can be found elsewhere (DeCarlo et al., 2006), however a short description is given here.

The aerosol particles are sampled through a 100-µm critical orifice into an aerodynamic lens which focuses particles in the size range ~35 nm to 1.5 µm into a narrow **ACPD**

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back

Full Screen / Esc

Interactive Discussion

particle beam (Zhang et al., 2004). The expansion at the exit of the aerodynamic lens into the first vacuum chamber accelerates the particles towards a heated tungsten element. The particles are vaporized on impact and ionized by electron ionization (EI) at 70 eV. The produced ions are extracted from the ionization region into the TOF (H-TOF 5 series, Tofwerk, Thun, Switzerland) followed by the orthogonal pulsed mass analysis.

The high mass resolution and accuracy of the HR-TOF-AMS made it possible to determine the elemental composition of the ions up to approximately m/z 100 (DeCarlo et al., 2006). Integration of the organic ions with the elemental composition $C_x H_v O_z N_D^+$ gives information on the elemental ratios of O/C, H/C, N/C as well as the OM/OC ratio (Aiken et al., 2007; Aiken et al., 2008). Based on their elemental composition, ions were grouped into four different families: C_xH_v ($x \ge 1$ and $y \ge 0$), $C_xH_vO_z$, $C_xH_vN_p$ and $C_xH_yO_zN_p$ $(x \ge 1, y \ge 0, z \ge 1 \text{ and } p \ge 1)$. These families will be referred to as CH, CHÓ, CHN and CHON from hereon.

Data analysis was performed using Igor Pro 6 (Wavemetrics, Lake Oswego, OR) with the Squirrel TOF analysis toolkit v1.48 or later and the TOF HR analysis toolkit v1.07 or later. During every experiment, after filling and before lights on, HEPA filtered air was sampled from the smog chamber to measure the gas phase contributions to the mass spectra. The AMS fragmentation table (Allan et al., 2004) was modified accordingly for m/z 16, 18, 29 and 44. In addition, the improved fragmentation table (Aiken et al., 2008) was used with the following fragmentation patterns in relation to the measured CO_2^+ signal: $CO_2^+ = 100\%$, $H_2O_2^+ = 22.5\%$, $OH_2^+ = 5.625\%$ (25% of $H_2O_2^+$), $O^+ = 0.90\%$ (4% of H_2O^+). The contributions of these ions are added to the CHO family. A particle collection efficiency (CE) of 1 was used to estimate the non-refractory aerosol mass concentration. Primary emissions from a complete burning cycle of a log wood burner were sampled during a separate measurement campaign. A CE of one was found based on the comparison of the sum of the AMS species and BC with a tapered element oscillating microbalance (TEOM, Series 1400a, Thermo Scientific).

ACPD

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

> **Figures Tables**

Close

Back

BC was measured using an aethalometer (AE31, Magee Scientific Company, Berkeley CA) at 880 nm using a specific attenuation cross section of $16.6\,\mathrm{m}^2\,\mathrm{g}^{-1}$ and an empirical correction for the shadowing effect (Weingartner et al., 2003) during the first five experiments (Table 1). For the other experiments BC was measured with a multi angle absorption photometer (MAAP) (Model 5012, Thermo Fisher Scientific). BC concentrations were calculated at a one-minute time resolution using a mass specific absorption cross section of $6.6\,\mathrm{m}^2\,\mathrm{g}^{-1}$ at the wavelength 630 nm.

2.2.3 Gas phase instrumentation

The evolution of several gas phase species was measured during the experiments. Carbon dioxide (CO_2) was measured using a differential, non-dispersive, infrared (NDIR) gas analyzer (LI-7000, Li-Cor Biosciences) operated with a time resolution of one second. Carbon monoxide (CO) was measured with an ultra fast fluorescence analyzer (AL5002, Aero-laser GmbH). A chemiluminescence NO_x analyzer (Monitor Labs model ML9841A) was used to measure NO_x and NO_x ($NO + NO_2$) and ozone (O_3) was measured with an ozone analyzer (Monitor Labs Inc. Model 8810).

3 Results and discussion

3.1 Primary emissions

The starting phase and flaming phase emissions from the old log wood burner were dominated by BC with an average OM/BC ratio of 0.21 ± 0.06 . The OM/BC ratio for the modern log wood burner changed significantly between the starting phase and the flaming phase, with values of 1.2 ± 0.4 and 0.12 ± 0.04 , respectively. The emissions from the pellet burner showed an average ratio of 2.5 ± 0.9 for the starting phase and during stable burning conditions. OM/BC ratios were found to be lower than published

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

[4

-1

Back

Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



8088

Abstract Conclusions

Introduction References

ACPD

11, 8081-8113, 2011

Primary and

secondary aerosol from wood

combustion

M. F. Heringa et al.

Title Page

Figures Tables

Close

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



values for off-line methods (Szidat et al., 2006 and the references therein) and on the lower end of previously published smog chamber data (Grieshop et al., 2009). Two possible explanations can be given. Our experiments were at lower concentrations (1- $30 \,\mu g \, m^{-3}$ as compared to $\geq 40 \,\mu g \, m^{-3}$), which affects the partitioning of organics between the gas and particle phase (Lipsky and Robinson, 2006), and the modified combustion efficiency (MCE, defined as [CO₂]/([CO₂]+[CO])) was higher (Table 1) which has been shown to lead to lower OM/BC ratios (Grieshop et al., 2009).

Secondary organic aerosol production

Figure 2 shows the evolution of OM, NO₃, SO₄ and BC for a typical experiment (No. 15 Table 1) with log wood burner emissions. After the injection, 15 min mixing time and the characterization of the primary emissions, the smog chamber lights were switched on. During the dark phase the concentrations of BC and OM decrease due to wall loss. The total PM mass as measured by the AMS was dominated by OM for all experiments. On average, the highest OM fractions were measured for the starting phase of the logwood burner (93%), followed by the starting phase of the pellet burner (86%), the flaming phase of the log wood burners (76%) and the stable burning phase of the pellet burner (53%). After the lights were switched on, the production rate of OM is greater than the wall loss rate, resulting in a rapid OM concentration increase from SOA formation in the first hour. In the second hour of lights on, the OM production rate slows down and OM reaches its maximum concentration once the SOA production rate equals the wall loss rate. After two hours, the wall loss rate dominates, resulting in a slow decrease in the measured OM concentration.

The wall loss rate was determined using BC as an inert tracer. Based on the assumption that the aerosol is internally mixed and that particles deposited at the walls are in equilibrium with the suspended material, the wall loss rate of BC can be used to correct the concentrations of other particulate species. The wall loss corrected concentration of OM (OM_{WLC}) can be derived using the equation:

where $OM_{meas}(t)$ refers to the concentration of OM measured at time t. $BC(t_0)$ and BC(t) are the concentrations of BC when lights were switched on and at time t, respectively.

During some experiments, as shown in Fig. 2, an apparent increase in the BC concentration was seen after lights were switched on and OM increased substantially. This observation due to increased light absorption by BC with thicker coatings is in agreement with findings in the literature. An increase in light absorption efficiency due to an organic coating up to a factor of ~2 has previously been reported (Schnaiter et al., 2005; Shiraiwa et al., 2010). In similar experiments using diesel exhaust in the PSI smog chamber this increase of black carbon concentration was not observed, likely due to a thinner coating (Chirico et al., 2010). This increase in absorption leads to an underestimation of the wall loss rate and consequent underestimation of the SOA production if the BC concentration calculated from the absorption measurement is used. To get a representative wall loss rate the BC data was fit for the time period prior to lights on using an exponential fit to zero. Equation (1) can be modified to:

$$OM_{WLC}(t) = OM_{meas}(t) \times \left[\frac{BC_{fit}(t_0)}{BC_{fit}(t)} \right]$$
(2)

where $\mathrm{BC}_{\mathrm{fit}}(t_0)$ and $\mathrm{BC}_{\mathrm{fit}}(t)$ are the concentrations of BC derived from the exponential fit when lights were switched on and at time t respectively. The pure SOA experiments were not used to quantify the SOA formation and therefore were not corrected for wall losses.

The relative increase of OM in relation to the concentration of POA is expressed as the OM enhancement ratio (OM_{ER}) . Taking the wall loss correction as shown in Eq. (2) into account, the OM_{ER} at time t can be calculated by:

OM_{ER}(t) =
$$\left[\frac{OM_{\text{meas}}(t)}{OM_{\text{meas}}(t_0)} \right] \times \left[\frac{BC_{\text{fit}}(t_0)}{BC_{\text{fit}}(t)} \right]$$
8090

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

4

Back Close

Full Screen / Esc

Printer-friendly Version



All log wood burner experiments show a substantial increase in OM for both starting and flaming phase emissions. The average OM_{ER} for these experiments after 5 h of photo oxidation was found to be 4.1 ± 1.4 . During injection for experiment No. 1, the fire stopped flaming and started smoldering which may be the reason for the lower OM_{ER} of 1.6. In contrast, the high emission smoldering event (No. 6) showed an OM loss of 54% (after wall loss correction) within the first hour and a slow increase during the following hours. The mass loss during this experiment is discussed in detail in Sect. 3.5. The pellet burner produced an average OM_{ER} of 3.3 ± 0.6 during the starting phase whereas no SOA formation was observed for the flaming phase, which is explained by the very stable and controlled burning conditions for the pellet burner in contrast to the log-wood ovens.

3.3 Emission factors

To perform smog chamber experiments at atmospherically relevant OM concentrations it was necessary to adjust the filling time and dilution factor to the type of burner and burning condition to stay below $40\,\mu g\,m^{-3}$ primary OM concentration. Therefore the concentrations of the measured aerosol species need scaling before they can be compared or averaged. Two abundant gas phase products from wood burning, CO_2 and CO, were found to be useful scalars for dilution effects and emission factors. The average concentrations of BC, POA and SOA, normalized by the increase in the CO_2 concentration, for the different burners and burning conditions are shown in Fig. 4.

Since there was no information on the mass of wood burned during injection, an estimate of the emission factors were calculated based on the increase in CO₂ concentration. The mass of wood burned was calculated using the following formula

$$m_{\rm WB} = \frac{P \times V}{R \times T} \times \Delta \rm{CO}_2 \times M_W C \times \frac{1}{f_C}$$
 (4)

where $m_{\rm WB}$ is the mass of wood burned (g), $\Delta {\rm CO_2}$ the increase in concentration (ppm), 8091

ACPD

Discussion Paper

Discussion Paper

Discussion Paper

Discussion Paper

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.









Printer-friendly Version



 $M_{\rm W}$ C the molecular mass of carbon, and $f_{\rm C}$ the carbon fraction of wood. P, V, R and T are the pressure (Pa), volume (m³), universal gas constant, and temperature (K) respectively. Using a carbon fraction of 50% for beech (Joosten et al., 2004), a 1ppm increase in CO₂ per cubic meter air in the smog chamber corresponds to 1 mg burned wood. These estimated emissions factors, in g kg⁻¹ wood burnt, are shown on a second axis in Fig. 4.

All of the burners tested showed higher emission factors for BC, POA and SOA during the startup phase compared to the flaming or stable phases. The old log wood burner produced the highest total emissions during the starting phase and flaming phase followed by the new log wood burner and then the pellet burner. All values from Fig. 4 as well as their standard deviations are presented in Table 2.

The old logwood burner showed a similar BC production during the starting and flaming phase. Total OM (POA + SOA) showed a relatively small decrease in emission factor during the flaming phase. The similarity between starting and flaming phases may be explained by the small combustion chamber, which leads to relatively inefficient combustion even under flaming conditions. The modern log wood burner showed similar BC emission factors during the starting phase and flaming phase, but was roughly a factor of 3-4 lower than the old wood stove. The emission factor of OM for the modern log wood burner was significantly lower during the flaming phase. This indicates that the new design with a large combustion chamber and sophisticated airflows decreases the emission factors for OM once the burner reaches operating temperatures. The improved design of the new log wood burner thus influences both the primary emissions and secondary formation of organic aerosol. The emission factors of the pellet burner were dominated by OM and were considerably lower during stable burning compared to the starting phase. A wide range of POA/CO ratios (3.9-48 µg m⁻³ ppm⁻¹) were observed for the different burners and burning conditions (Table 1). This range is in agreement with the average OM/CO ratio of 38 that was found for the wood burning dominated site in Roveredo, Switzerland (Gaeggeler et al., 2008) and with values found for other smog chamber studies (Grieshop et al., 2009).

ACPD

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back

Discussion Paper

11, 8081–8113, 2011

Emissions from wood burning are highly variable in quantity and properties as shown above. An example of the family spectra of the POA and of the aged organic aerosol is shown in Fig. 5. Experiment No. 15 was chosen as it is in the middle of the observed range in SOA production and O/C ratio. POA is dominated by CHO (51.9%) followed by CH (43.6%), CHON (3.2%) and CHN (1.3%). After 5 h of aging, CHO showed an increase to 63.3% whereas CH decreased to 31.1%. CHN increased from 3.2% to 4.4% and CHON stayed constant at 1.3%. The POA spectrum shows a strong signal for the ion series at m/z 27, 41, 55, 69, 83 and m/z 29, 43, 57, 71 which are often assigned to cycloalkanes or alkenes ($C_nH_{2n+1}^+$) and normal or branched alkanes ($C_nH_{2n+1}^+$) respectively (Alfarra et al., 2004; Canagaratna et al., 2004; Weimer et al., 2008). However, the high resolution data shows that, except for m/z 27 and 41, the signals are dominated by oxygen containing ions and cannot be assigned to either of these hydrocarbon series alone.

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Discussion Paper

Discussion Paper

Conclusions Tables

Abstract

≯I

Introduction

References

Figures

•

Close

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

© BY

Levoglucosan, a product from wood burning and biomass burning, has been used as molecular marker in ambient organic aerosol (Fraser and Lakshmanan, 2000; Simoneit et al., 1999). The ion at m/z 60, a dominant fragment in the mass spectrum of levoglucosan, has been used as a wood burning marker for AMS measurements in order to estimate the wood burning contribution to the ambient organic aerosol (Alfarra et al., 2007; DeCarlo et al., 2008). Recent studies showed that levoglucosan is not stable under atmospheric conditions (Hennigan et al., 2010; Hoffmann et al., 2010). AMS reference spectra of levoglucosan (\geq 98%, Fluka) showed that the signal at m/z 60 corresponds to the ion $C_2H_4O_2^+$ fit at m/z 60.0211. The wall loss corrected evolution of this

Conclusions

Figures Tables

ACPD

11, 8081-8113, 2011

Primary and

secondary aerosol

from wood

combustion

M. F. Heringa et al.

Title Page

Abstract



Introduction

References



Close

Back

Full Screen / Esc

Printer-friendly Version

Interactive Discussion



ion for all smog chamber experiments is shown in Fig. 6. A decrease in C₂H₄O₂⁺ was observed for the pellet burner experiments and the smoldering experiments. One log wood burner experiment showed a decrease, whereas the remaining 11 experiments showed an increase, in $C_2H_4O_2^+$. This increase in the signal must result from SOA products that contribute to the same $C_2H_4O_2^+$ fragment as levoglucosan. These SOA products depend on the burning conditions and were not observed during the aging experiments performed with the pellet burner. The contribution of SOA to the $C_2H_4O_2^+$ ion at m/z 60 can overcompensate for the degradation of levoglucosan, thus leading to an overestimation of the contribution of either wood burning or biomass burning to the total OM if this ion is used as a primary tracer for wood burning or biomass burning aerosol in ambient measurements. Concurrent measurements of levoglucosan and aerosol mass spectra were recently performed in our chamber and will be analyzed and published in the near future.

Elemental ratios

From the high resolution AMS spectra the average elemental composition and different atomic ratios can be derived (Aiken et al., 2007). The bulk organic aerosol properties can change by condensation of SOA and/or heterogeneous chemistry. During oxidation OA becomes more oxygenated, less volatile, and more hygroscopic such that the O/C ratio can be used as an approximation for the degree of oxidation of the organic aerosol (Jimenez et al., 2009).

Figure 7 shows the evolution of the O/C ratio for all smog chamber experiments. The O/C ratios of the freshly emitted organic aerosol were highly variable and spanned the range of 0.19-0.58 for the different starting and flaming phase experiments. There was no separation in O/C ratio possible between the starting and flaming phases, nor between the different burners. The smoldering experiment showed a significantly higher O/C ratio, of 0.87, compared to the starting and flaming phases. All experiments show an increase in the O/C ratio after lights on, with the exception of the smoldering phase experiment. Different rates for the increase in the O/C ratio were found for the first two

Interactive Discussion

hours after lights on compared to the last period (3-5 h). For the log wood burners the O/C ratio increased initially by 0.055 ± 0.020 and 0.036 ± 0.012 per hour for the starting phase and flaming phase respectively. This is 3-5 times faster than for the latter part of the experiments where it increased by 0.016 ± 0.004 and 0.007 ± 0.010 per hour for the starting phase and flaming phase respectively. The O/C ratio of the start-up emissions from the pellet burner increased by 0.086 ± 0.010 during the first two hours and 0.007 ± 0.010 during the last two hours of the experiment. The pure SOA experiments did not show an increase during the first two hours (-0.03 ± 0.03) and no significant increase during the last two hours (0.003 ± 0.005), indicating that the condensing OM has a similar O/C ratio during the entire experiment.

During the smoldering experiment No. 6, an increase in the O/C ratio was observed during the first 1.5 h, which was then followed by a gradual decrease, while the OM had exactly the opposite trend. This peculiar behavior could potentially be explained by the fact that the POA seems to have a fraction of highly oxygenated OM and a smaller fraction of less oxygenated compounds. The loss of mass in the first hour of the experiment (Fig. 6) suggest that these less oxygenated species fragment upon oxidation and thus less likely partition into the OA (Jimenez et al., 2009). In the later phase semi-volatile species with a lower O/C ratio than the existing OM condense on the aerosol resulting in a decrease in the O/C ratio (Fig. 7).

The fraction of m/z 44 at unit mass resolution can be used to estimate the O/C ratio when their relationship is known. This relationship has been published for an analysis of ambient datasets (Aiken et al., 2008), but not for primary and secondary aerosols from domestic wood burning. Figure 8 shows the data for all starting and flaming phase experiments as well as from the pure SOA experiments. The slope of the separately fit starting and flaming phase experiments were very similar with values of 0.033 and 0.030 respectively. The intercept found for the starting phase experiments is higher than the intercept for the flaming phase experiments and therefore more sensitive to the number of experiments fit for each condition. The fitting shown in Fig. 8 is based on the average slope and intercept from the separate fittings of the starting and flaming

ACPD

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Back Close Full Screen / Esc

Printer-friendly Version

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page Abstract Introduction Conclusions References Tables Figures I ◆ ▶I ◆ Back Close Full Screen / Esc

phase experiments. It shows a slightly lower slope and higher intercept than reported for the ambient data.

Recently a new mass spectral diagnostic tool was developed to follow the aging of the OOA component in the atmosphere (Ng et al., 2010). The fractions of two major masses, m/z 44 (CO₂⁺) and m/z 43 (mostly C₂H₃O⁺) were used to create a triangular space f44 vs. f43 in which OOA moves towards the apex during the aging process. In Fig. 9a the POA at lights on (t = 0 h) and the bulk aerosol after five hours of aging is plotted in this two-dimensional space. Most data points are within this triangle close to the left margin whereby ageing moves the data to higher f44 for most experiments, whereas only a small increase or decrease was observed in the f43. However, in contrast to OOA, the POA of wood burning has a significant contribution from the nonoxygenated ion $C_3H_7^+$ at m/z 43, increasing the m/z 43/44 ratio. Using the HR data, it is possible to plot the fractions of the particle CO_2^+ (pCO_2^+) and $C_2H_3O^+$ ions (Fig. 9b) which represent the OA functionalities better than the UMR data. The figure shows that the POA from all wood burning experiments have a characteristic low C₂H₃O⁺ signal for a given pCO_2^+ fraction compared to ambient OOA. During the SOA formation, the pCO₂⁺ and C₂H₃O⁺ fraction of the bulk aerosol increases, resulting in a shift in this space up and to the right. After 5 h, about one third of the experiments were found at the lower edge of the $C_2H_3O^+$ vs. pCO_2^+ triangle found for ambient OOA, while the others were still outside the triangle.

4 Conclusions

In this study, we present the results from photo-oxidation smog chamber experiments on two different log wood burners and a residential pellet burner under different burning conditions. OM increased by SOA formation on average by a factor of 4.1 ± 1.4 for the starting and flaming phases of the two log wood burners. The startup phase of the pellet burner showed an increase by a factor of 3.3 ± 0.6 , whereas no measureable SOA formation was observed for the stable burning phase. This implies that the gas

Printer-friendly Version

Interactive Discussion

phase emissions from log wood burners play an import role in the total contribution of OM from residential wood burning appliances to ambient OM and should be considered to be included in future legislations.

The primary organic emissions from the three different burners showed a wide range ₅ in the O/C atomic ratio of OA (0.19–0.60) for all burning conditions and showed an increase during aging. The increase in the O/C ratio was significantly faster during the first two hours compared to 3-5 h after lights on. After five hours of aging, the O/C ratio of the bulk aerosol reached values >0.6 for several experiments, which is close to the O/C ratio of LV-OOA found for aged ambient OA.

The O/C ratio of the pure SOA experiments stayed relatively stable during the entire experiment. One smoldering event with high emissions was captured, which showed a decrease of the OM concentration accompanied by an increase in the O/C ratio (0.87 to 0.95) during the first 1.5 h, followed by an increase of the OM concentration and a decrease in the O/C ratio. This exemplifies the complex combination of processes that are involved in aging (partitioning including condensation and evaporation, and oxidation coupled to either fragmentation or functionalization).

The wood burning marker at m/z 60 ($C_2H_4O_2^+$ fragment of levoglucosan) showed an increase in concentration for more than 90% of the log wood burner experiments for the starting and flaming phases. This increase is attributed to SOA species also contributing to this fragment. The formation of the precursor leading to this SOA contribution depends on the burning conditions, and is not observed during the pellet burner experiments. The fractions of C₂H₃O⁺ and pCO₂⁺, when projected onto the two-dimensional space described by Ng et al. (2010), was found to initially be outside of the triangle found for ambient OOA, but on aging moved up and to the right, towards the triangle. After five hours of aging, the OA has a rather low C₂H₃O⁺ signal for a given CO₂⁺ fraction, possibly indicating a higher ratio of acid to non-acid oxygenated compounds in wood burning OA compared to other OOA.

ACPD

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Close

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I**∢** ►I

◆ •

Back Close

Full Screen / Esc

Printer-friendly Version

Interactive Discussion

© BY

Supplementary material related to this article is available online at: http://www.atmos-chem-phys-discuss.net/11/8081/2011/acpd-11-8081-2011-supplement.pdf.

Acknowledgements. This work was supported by the IMBALANCE project of the Competence Center Environment and Sustainability of the ETH Domain (CCES), the Bundesamt für Umwelt (BAFU), the Bundesamt für Energie (BFE), as well as the Swiss National Science Foundation. PFD is grateful for the postdoctoral support from the US-NSF (IRFP# 0701013). Attika Feuer AG is kindly acknowledged for the provided wood stove.

References

 Aiken, A. C., DeCarlo, P. F., and Jimenez, J. L.: Elemental analysis of organic species with electron ionization high-resolution mass spectrometry, Anal. Chem., 79, 8350–8358, 2007.

Aiken, A. C., Decarlo, P. F., Kroll, J. H., Worsnop, D. R., Huffman, J. A., Docherty, K. S., Ulbrich, I. M., Mohr, C., Kimmel, J. R., Sueper, D., Sun, Y., Zhang, Q., Trimborn, A., Northway, M., Ziemann, P. J., Canagaratna, M. R., Onasch, T. B., Alfarra, M. R., Prevot, A. S. H., Dommen, J., Duplissy, J., Metzger, A., Baltensperger, U., and Jimenez, J. L.: O/C and OM/OC ratios of primary, secondary, and ambient organic aerosols with high-resolution time-of-flight aerosol mass spectrometry, Environ. Sci. Technol., 42, 4478–4485, 2008.

Alfarra, M. R., Coe, H., Allan, J. D., Bower, K. N., Boudries, H., Canagaratna, M. R., Jimenez, J. L., Jayne, J. T., Garforth, A. A., Li, S. M., and Worsnop, D. R.: Characterization of urban and rural organic particulate in the lower Fraser valley using two aerodyne aerosol mass spectrometers, Atmos. Environ., 38, 5745–5758, 2004.

Alfarra, M. R., Prevot, A. S. H., Szidat, S., Sandradewi, J., Weimer, S., Lanz, V. A., Schreiber, D., Mohr, M., and Baltensperger, U.: Identification of the mass spectral signature of organic aerosols from wood burning emissions, Environ. Sci. Technol., 41, 5770–5777, 2007.

Allan, J. D., Delia, A. E., Coe, H., Bower, K. N., Alfarra, M. R., Jimenez, J. L., Middlebrook, A. M., Drewnick, F., Onasch, T. B., Canagaratna, M. R., Jayne, J. T., and Worsnopf, D. R.: A generalised method for the extraction of chemically resolved mass spectra from Aerodyne aerosol mass spectrometer data, J. Aerosol Sci., 35, 909–922, 2004.

Interactive Discussion

Bond, T. C., Streets, D. G., Yarber, K. F., Nelson, S. M., Woo, J. H., and Klimont, Z.: A technology-based global inventory of black and organic carbon emissions from combustion, J. Geophys. Res-Atmos., 109, D14203, doi:10.1029/2003JD003697, 2004.

Canagaratna, M. R., Jayne, J. T., Ghertner, D. A., Herndon, S., Shi, Q., Jimenez, J. L., Silva, P. J., Williams, P., Lanni, T., Drewnick, F., Demerjian, K. L., Kolb, C. E., and Worsnop, D. R.: Chase studies of particulate emissions from in-use New York City vehicles, Aerosol. Sci. Technol., 38, 555-573, 2004.

Chirico, R., DeCarlo, P. F., Heringa, M. F., Tritscher, T., Richter, R., Prvt, A. S. H., Dommen, J., Weingartner, E., Wehrle, G., Gysel, M., Laborde, M., and Baltensperger, U.: Impact of aftertreatment devices on primary emissions and secondary organic aerosol formation potential from in-use diesel vehicles: results from smog chamber experiments, Atmos. Chem. Phys., 10, 11545-11563, doi:10.5194/acp-10-11545-2010, 2010.

DeCarlo, P. F., Kimmel, J. R., Trimborn, A., Northway, M. J., Jayne, J. T., Aiken, A. C., Gonin, M., Fuhrer, K., Horvath, T., Docherty, K. S., Worsnop, D. R., and Jimenez, J. L.: Field-deployable, high-resolution, time-of-flight aerosol mass spectrometer, Anal. Chem., 78. 8281-8289, doi:10.1021/ac061249n, 2006.

DeCarlo, P. F., Dunlea, E. J., Kimmel, J. R., Aiken, A. C., Sueper, D., Crounse, J., Wennberg, P. O., Emmons, L., Shinozuka, Y., Clarke, A., Zhou, J., Tomlinson, J., Collins, D. R., Knapp, D., Weinheimer, A. J., Montzka, D. D., Campos, T., and Jimenez, J. L.: Fast airborne aerosol size and chemistry measurements above Mexico City and Central Mexico during the MILAGRO campaign, Atmos. Chem. Phys., 8, 4027-4048, doi:10.5194/acp-8-4027-2008, 2008.

20

Fraser, M. P. and Lakshmanan, K.: Using levoglucosan as a molecular marker for the longrange transport of biomass combustion aerosols, Environ. Sci. Technol., 34, 4560-4564, 2000.

Gaeggeler, K., Prevot, A. S. H., Dommen, J., Legreid, G., Reimann, S., and Baltensperger. U.: Residential wood burning in an Alpine valley as a source for oxygenated volatile organic compounds, hydrocarbons and organic acids, Atmos. Environ., 42, 8278-8287, 2008.

Grieshop, A. P., Logue, J. M., Donahue, N. M., and Robinson, A. L.: Laboratory investigation of photochemical oxidation of organic aerosol from wood fires 1: measurement and simulation of organic aerosol evolution, Atmos. Chem. Phys., 9, 1263-1277, doi:10.5194/acp-9-1263-2009, 2009.

Hennigan, C. J., Sullivan, A. P., Collett, J. L., and Robinson, A. L.: Levoglucosan stability in biomass burning particles exposed to hydroxyl radicals, Geophys. Res. Lett., 37, L09806, **ACPD**

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Introduction **Abstract**

Conclusions References

Tables Figures

Close

Close

Interactive Discussion

doi:10.1029/2010GL043088, 2010.

20

Hoffmann, D., Tilgner, A., linuma, Y., and Herrmann, H.: Atmospheric stability of levoglucosan: A detailed laboratory and modeling study, Environ. Sci. Technol., 44, 694–699, 2010.

Jimenez, J. L., Canagaratna, M. R., Donahue, N. M., Prevot, A. S. H., Zhang, Q., Kroll, J. H., DeCarlo, P. F., Allan, J. D., Coe, H., Ng, N. L., Aiken, A. C., Docherty, K. S., Ulbrich, I. M., Grieshop, A. P., Robinson, A. L., Duplissy, J., Smith, J. D., Wilson, K. R., Lanz, V. A., Hueglin, C., Sun, Y. L., Tian, J., Laaksonen, A., Raatikainen, T., Rautiainen, J., Vaattovaara, P., Ehn, M., Kulmala, M., Tomlinson, J. M., Collins, D. R., Cubison, M. J., Dunlea, E. J., Huffman, J. A., Onasch, T. B., Alfarra, M. R., Williams, P. I., Bower, K., Kondo, Y., Schneider, J., Drewnick, F., Borrmann, S., Weimer, S., Demerjian, K., Salcedo, D., Cottrell, L., Griffin, R., Takami, A., Miyoshi, T., Hatakeyama, S., Shimono, A., Sun, J. Y., Zhang, Y. M., Dzepina, K., Kimmel, J. R., Sueper, D., Jayne, J. T., Herndon, S. C., Trimborn, A. M., Williams, L. R., Wood, E. C., Middlebrook, A. M., Kolb, C. E., Baltensperger, U., and Worsnop, D. R.: Evolution of organic aerosols in the atmosphere. Science, 326, 1525-1529, 2009.

Joosten, R., Schumacher, J., Wirth, C., and Schulte, A.: Evaluating tree carbon predictions for beech (Fagus sylvatica L.) in western Germany, Forest Ecol. Manag., 189, 87-96, 2004.

Lanz, V. A., Alfarra, M. R., Baltensperger, U., Buchmann, B., Hueglin, C., and Prvt, A. S. H.: Source apportionment of submicron organic aerosols at an urban site by factor analytical modelling of aerosol mass spectra, Atmos. Chem. Phys., 7, 1503-1522, doi:10.5194/acp-7-1503-2007, 2007.

Lanz, V. A., Prévôt, A. S. H., Alfarra, M. R., Weimer, S., Mohr, C., DeCarlo, P. F., Gianini, M. F. D., Hueglin, C., Schneider, J., Favez, O., D'Anna, B., George, C., and Baltensperger, U.: Characterization of aerosol chemical composition with aerosol mass spectrometry in Central Europe: an overview, Atmos. Chem. Phys., 10, 10453-10471, doi:10.5194/acp-10-10453-2010, 2010.

Lee, T., Sullivan, A. P., Mack, L., Jimenez, J. L., Kreidenweis, S. M., Onasch, T. B., Worsnop, D. R., Malm, W., Wold, C. E., Hao, W. M., and Collett, J. L.: Chemical smoke marker emissions during flaming and smoldering phases of laboratory open burning of wildland fuels, Aerosol. Sci. Technol., 44, I-V, 2010.

Lipsky, E. M. and Robinson, A. L.: Effects of dilution on fine particle mass and partitioning of semivolatile organics in diesel exhaust and wood smoke, Environ. Sci. Technol., 40, 155-162, 2006.

McDonald, J. D., Zielinska, B., Fujita, E. M., Sagebiel, J. C., Chow, J. C., and Watson, J. G.:

Paper

Abstract

Introduction

Conclusions

References **Figures**

Tables

ACPD

11, 8081–8113, 2011

Primary and

secondary aerosol

from wood

combustion

M. F. Heringa et al.

Title Page





Interactive Discussion

Fine particle and gaseous emission rates from residential wood combustion, Environ. Sci. Technol., 34, 2080-2091, 2000.

- Nel, A.: Air pollution-related illness: Effects of particles, Science, 308, 804-806, 2005.
- Ng, N. L., Canagaratna, M. R., Zhang, Q., Jimenez, J. L., Tian, J., Ulbrich, I. M., Kroll, J. H., Docherty, K. S., Chhabra, P. S., Bahreini, R., Murphy, S. M., Seinfeld, J. H., Hildebrandt, L., Donahue, N. M., DeCarlo, P. F., Lanz, V. A., Prévôt, A. S. H., Dinar, E., Rudich, Y., and Worsnop, D. R.: Organic aerosol components observed in Northern Hemispheric datasets from Aerosol Mass Spectrometry, Atmos. Chem. Phys., 10, 4625-4641, doi:10.5194/acp-10-4625-2010, 2010,
- Paulsen, D., Dommen, J., Kalberer, M., Prevot, A. S. H., Richter, R., Sax, M., Steinbacher, M., Weingartner, E., and Baltensperger, U.: Secondary organic aerosol formation by irradiation of 1,3,5-trimethylbenzene-NO_v-H₂O in a new reaction chamber for atmospheric chemistry and physics, Environ. Sci. Technol., 39, 2668-2678, 2005.
 - Pope, C. A. and Dockery, D. W.: Health effects of fine particulate air pollution: Lines that connect, J. Air Waste Manage., 56, 709-742, 2006.
 - Sandradewi, J., Prevot, A. S. H., Szidat, S., Perron, N., Alfarra, M. R., Lanz, V. A., Weingartner, E., and Baltensperger, U.: Using aerosol light absorption measurements for the quantitative determination of wood burning and traffic emission contributions to particulate matter, Environ. Sci. Technol., 42, 3316–3323, 2008.
- Schauer, J. J., Kleeman, M. J., Cass, G. R., and Simoneit, B. R. T.: Measurement of emissions from air pollution sources. 3. C-1-C-29 organic compounds from fireplace combustion of wood, Environ. Sci. Technol., 35, 1716-1728, 2001.
 - Schnaiter, M., Linke, C., Mohler, O., Naumann, K. H., Saathoff, H., Wagner, R., Schurath, U., and Wehner, B.: Absorption amplification of black carbon internally mixed with secondary organic aerosol, J. Geophys. Res-Atmos., 110, D19204, doi:10.1029/2005JD006046, 2005.
 - Shiraiwa, M., Kondo, Y., Iwamoto, T., and Kita, K.: Amplification of light absorption of black carbon by organic coating, Aerosol. Sci. Technol., 44, 46–54, 2010.
- Simoneit, B. R. T., Schauer, J. J., Nolte, C. G., Oros, D. R., Elias, V. O., Fraser, M. P., Rogge, W. F., and Cass, G. R.: Levoglucosan, a tracer for cellulose in biomass burning and atmospheric particles, Atmos. Environ., 33, 173-182, 1999.
- Simpson, D., Yttri, K. E., Klimont, Z., Kupiainen, K., Caseiro, A., Gelencser, A., Pio, C., Puxbaum, H., and Legrand, M.: Modeling carbonaceous aerosol over Europe: Analysis of the CARBOSOL and EMEP EC/OC campaigns, J. Geophys. Res.-Atmos., 112, D23S14,

ACPD

11, 8081–8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Introduction **Abstract**

Conclusions References

> **Tables Figures**

Close **Back**

Full Screen / Esc

doi:10.1029/2006JD008158, 2007.

25

- Szidat, S., Jenk, T. M., Synal, H.-A., Kalberer, M., Wacker, L., Hajdas, I., Kasper-Giebl, A., and Baltensperger, U.: Contributions of fossil fuel, biomass-burning, and biogenic emissions to carbonaceous aerosols in Zurich as traced by ¹⁴C, J. Geophys. Res., 111, D07206, doi:10.1029/2005JD006590, 2006.
- Weimer, S., Alfarra, M. R., Schreiber, D., Mohr, M., Prévôt, A. S. H., and Baltensperger, U.: Organic aerosol mass spectral signatures from wood-burning emissions: Influence of burning conditions and wood type, J. Geophys. Res., 113, D10304, doi:10.1029/2007JD009309, 2008.
- Weingartner, E., Saathoff, H., Schnaiter, M., Streit, N., Bitnar, B., and Baltensperger, U.: Absorption of light by soot particles: determination of the absorption coefficient by means of aethalometers, J. Aerosol Sci., 34, 1445–1463, 2003.
 - World Energy Council: 2007 Survey of energy resources, available at: http://www.worldenergy.org/publications/survey_of_energy_resources_2007/, 2007.
- Zhang, Q., Jimenez, J. L., Canagaratna, M. R., Allan, J. D., Coe, H., Ulbrich, I., Alfarra, M. R., Takami, A., Middlebrook, A. M., Sun, Y. L., Dzepina, K., Dunlea, E., Docherty, K., De-Carlo, P. F., Salcedo, D., Onasch, T., Jayne, J. T., Miyoshi, T., Shimono, A., Hatakeyama, S., Takegawa, N., Kondo, Y., Schneider, J., Drewnick, F., Borrmann, S., Weimer, S., Demerjian, K., Williams, P., Bower, K., Bahreini, R., Cottrell, L., Griffin, R. J., Rautiainen, J., Sun, J. Y., Zhang, Y. M., and Worsnop, D. R.: Ubiquity and dominance of oxygenated species in organic aerosols in anthropogenically-influenced Northern Hemisphere midlatitudes, Geophys. Res. Lett., 34, L13801, doi:10.1029/2007GL029979, 2007.
- Zhang, X. F., Smith, K. A., Worsnop, D. R., Jimenez, J. L., Jayne, J. T., Kolb, C. E., Morris, J., and Davidovits, P.: Numerical characterization of particle beam collimation: Part II Integrated aerodynamic-lens-nozzle system, Aerosol. Sci. Technol., 38, 619–638, 2004.

ACPD

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I∢

◆

Close

Full Screen / Esc

Back

Printer-friendly Version



Table 1. Overview of several parameters for the different smog chamber experiments.

Experiment	POA (μg m ⁻³)	OA _{5 h} /POA (WLC)	POA O/C	OA _{5 h} O/C	POA/CO $(\mu g m^{-3} ppm^{-1})$	MCE ^a	$\Delta O_3 t_{5h} - t_{0h}$ (ppb)
Old log wood burner							
1 flaming + smoldering	3.8	1.6	0.51	0.58	3.9	0.986	32
2 flaming	4.4	3.8	0.38	0.50	n/a	n/a	59
3 flaming	28	5.1	0.19	0.36	11	0.960	111
4 starting	27	6.9	0.32	0.42	7.6	0.943	52
5 starting	6.5	5.7	0.23	0.46	6.4	0.959	65
6 smoldering	17	0.7	0.87	0.79	14	0.909	16
7 flaming	7.6	3.8	0.25	0.36	38	0.994	10
Pellet burner							
8 stable burning	4.2	1.0	0.23	0.50	48	0.999	n/a
9 starting	3.6	2.8	0.52	0.72	29	0.992	26
10 starting	3.5	3.7	0.59	0.84	22	0.988	24
Modern log wood burner							
11 flaming	1.4	3.6	0.48	0.62	2.5	0.993	
12 flaming	4.9	5.3	0.37	0.50	3.1	0.987	79
13 flaming	6.1	2.7	0.35	0.41	11	0.995	3
14 gas-phase only ^b	_	_	_	0.50	_	0.989	73
15 starting	31	4.2	0.37	0.56	20	0.958	23
16 starting	3.9	3.4	0.41	0.62	28	0.979	12
17 starting	20	2.3	0.33	0.51	16	0.966	25
18 gas-phase only ^b	_	_	_	0.36	_	0.994	10
19 gas-phase only ^b	_	_	_	0.39	_	0.993	25
20 flaming	2.0	2.1	0.50	0.54	5.8	0.995	2

^a Modified combustion efficiency defined as [CO₂]/([CO₂]+[CO]).

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I◀

►I

- ▼

Back

Close

Full Screen / Esc

Printer-friendly Version



^b Gas-phase only experiments were performed on a mixture of starting and flaming phase emissions.

Table 2. Emission factors of BC, POA and SOA normalized to the amount of burnt wood (g kg⁻¹) and their relative contribution to the total PM emissions (OM + BC) after 5 h of aging.

Experiment ^{a,b}	BC (g kg ⁻¹)	BC (%)	POA (g kg ⁻¹)	POA (%)	SOA (g kg ⁻¹)	SOA (%)
Old start	1.51 ± 0.11	39	0.36 ± 0.13	10	2.0 ± 1.0	51
Old flaming	1.3 ± 1.2	53	0.25 ± 0.19	10	0.88 ± 0.59	37
Modern start	0.53 ± 0.02	16	0.73 ± 0.19	22	2.1 ± 0.9	62
Modern flaming	0.34 ± 0.05	71	0.03 ± 0.01	7	0.10 ± 0.11	22
Pellet start	0.10 ± 0.03	10	0.25 ± 0.03	27	0.57 ± 0.23	63
Pellet stable	0.014	33	0.027	67	0	0

^a Old = old log wood burner;

8104

Abstract

Introduction References

Figures

Tables

I⋖

Conclusions

ACPD

11, 8081-8113, 2011

Primary and secondary aerosol from wood

combustion

M. F. Heringa et al.

Title Page









^b Modern = modern log wood burner.

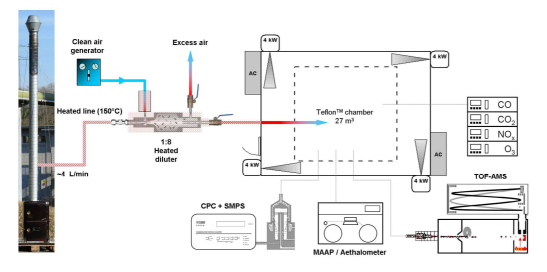


Fig. 1. Schematic representation of the inlet system and smog chamber setup.

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract

Introduction

Conclusions

References

Tables

Figures













Full Screen / Esc

Printer-friendly Version



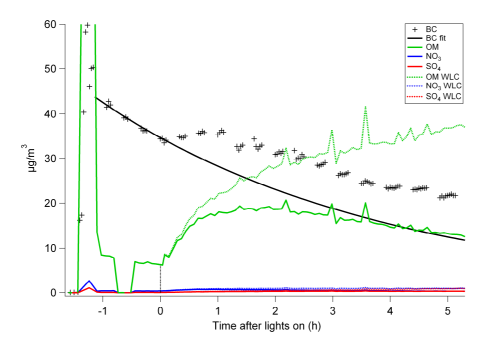


Fig. 2. Temporal evolution of the AMS species and BC (aethalometer) for the starting phase emissions of the modern log wood burner (experiment No. 15). BC data was fitted for the time period prior to lights on. Wall loss corrected (WLC) time series are calculated according to Eq. (2).

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

Close

Full Screen / Esc

Back

Printer-friendly Version





11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

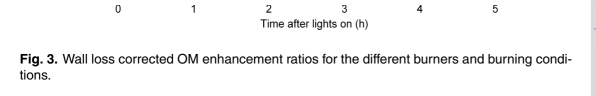
ACPD

M. F. Heringa et al.



Printer-friendly Version

Interactive Discussion



Starting phase Flaming phase

Log wood burner

--- Pellet burner

6

5

2 -

0

OM enhancement ratio

Flaming + smoldering phase Smoldering phase



11, 8081-8113, 2011

ACPD

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.



Printer-friendly Version

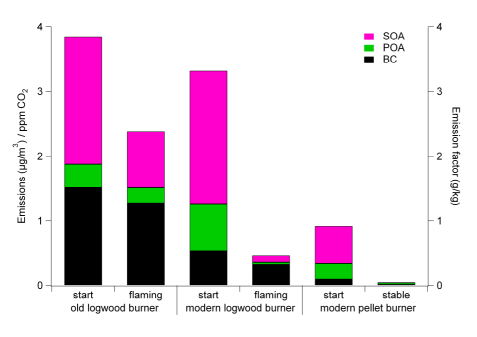


Fig. 4. Average concentrations of BC, POA and SOA after 5h of aging normalized by the increase in CO₂ concentration. The right axis shows the estimated emission factor in g kg⁻¹ wood burnt.

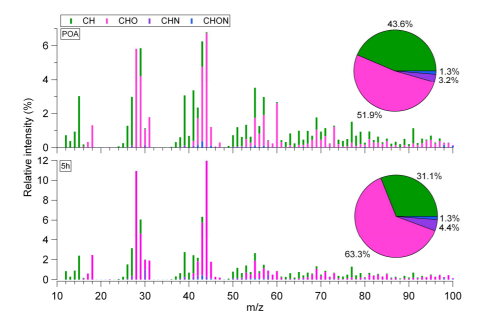


Fig. 5. Mass spectra of the starting phase log wood burner experiment (No. 15). The top panel shows the POA and the lower panel the organic aerosol after 5 h of aging. Ions are grouped into the families CH, CHO, CHN and CHON based on their elemental composition. The pie charts show the relative contributions of the families.

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Printer-friendly Version





11, 8081-8113, 2011

ACPD

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.



Printer-friendly Version

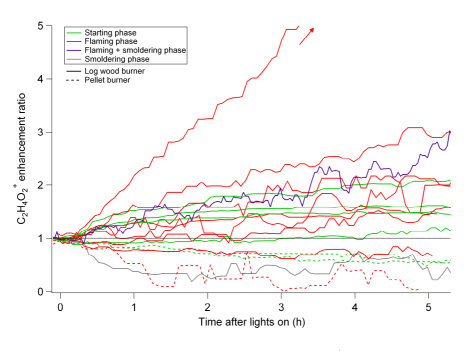


Fig. 6. Wall loss corrected enhancement ratios of the $C_2H_4O_2^+$ ion at m/z 60.0211 for the different burners and burning conditions.

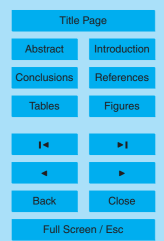


11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

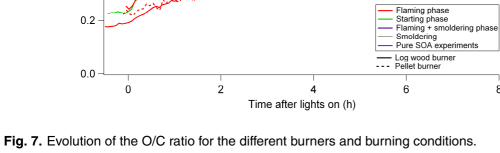
ACPD

M. F. Heringa et al.



Printer-friendly Version

Interactive Discussion



1.0 -

0.8

0.6

0.4

O/C ratio

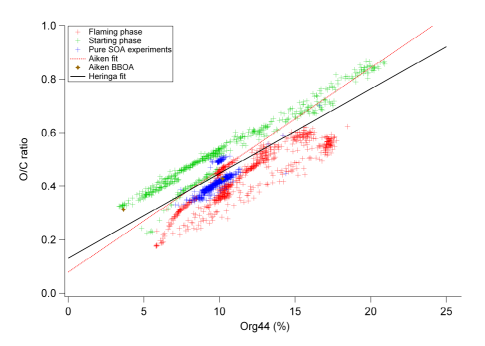


Fig. 8. Scatter plot of the O/C ratio vs. the percentage m/z 44 of the normalized organic spectrum. The red line shows the correlation found for different ambient studies (Aiken et al., 2008). The black line shows the average of the fittings of the starting and flaming phase experiments $(0.0316 \pm 0.0005x + 0.132 \pm 0.006)$.

11, 8081-8113, 2011

Primary and secondary aerosol from wood combustion

M. F. Heringa et al.

Title Page

Abstract Introduction

Conclusions References

Tables Figures

I**⊲** ►I

4

Back Close

Full Screen / Esc

Printer-friendly Version







Full Screen / Esc

Back

Abstract

Conclusions

Tables

Printer-friendly Version

ACPD

11, 8081-8113, 2011

Primary and

secondary aerosol

from wood

combustion

M. F. Heringa et al.

Title Page

Introduction

References

Figures

Close



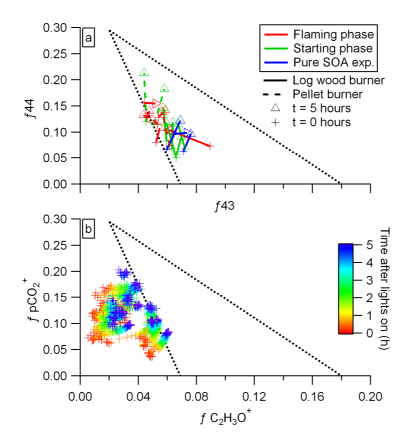


Fig. 9. Fraction plot of (a) mass f_{44} vs. f_{43} and (b) high resolution fit $f pCO_2^+$ vs. $f C_2H_3O^+$. The triangular space found by Ng et al. (2010) to accommodate the OOA component of ambient organic aerosol is shown as well.