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*Supplement of*

## **Free tropospheric aerosols at the Mt. Bachelor Observatory: more oxidized and higher sulfate content compared to boundary layer aerosols**

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11      **Section 1. Distinguish Free Tropospheric Air from Boundary Layer-Influenced Air**

12      Extensive work has been done at MBO to differentiate free troposphere air (FT) from boundary layer (BL)-  
13 influenced air. Several water vapor ( $H_2O_{(g)}$ ) mixing ratio-based segregation techniques have been used to date. Weiss-  
14 Penzias et al. (2006) and Fischer et al. (2010) used percentiles of water vapor mixing ratio to identify FT and BL-  
15 influences air masses. These same studies also used water vapor calculated from the 0 and 12 UTC National Weather  
16 Service soundings from Medford, Oregon and Salem, Oregon to determine a representative altitude for the air masses  
17 sampled at MBO. Reidmiller et al. (2010) used chairlift soundings of water vapor mixing ratios at Mt. Bachelor  
18 coupled with high-resolution  $NO_x$  data from MBO to determine a time of day when the BL influence began at the  
19 mountain summit. While all these studies focus on segregating FT and BL-influenced air masses during Spring,  
20 Ambrose et al. (2011) used sounding data from Medford to compare with  $H_2O_{(g)}$  distributions at MBO and obtained a  
21 seasonal mean  $H_2O_{(g)}$  mixing ratios for MBO FT data, which for winter, spring, summer, and fall seasons corresponded  
22 with WV < 3.28, < 3.28, < 5.4, and < 4.12 g kg<sup>-1</sup>. Based on the same work by Ambrose et al. (2011), McClure et al.  
23 (2016) interpolated between the seasonal values into a monthly criterion. Zhang and Jaffe (2017) contributed to a more  
24 accurate monthly FT/BL-influence isolation at MBO based on comparison of MBO  $H_2O_{(g)}$  distributions to the  $H_2O_{(g)}$   
25 soundings from Medford and Salem, Oregon, at equivalent pressure level.  $H_2O_{(g)}$  criteria for FT air masses at MBO  
26 were referred to the cut points when equivalent monthly averages of the retained drier portion of the MBO data and  
27 the soundings were obtained. The FT/BL-influenced water vapor criterion for each month is 3.26, 2.64, 2.46, 2.55,  
28 3.06, 4.25, 5.14, 5.23, 4.60, 4.36, 3.44, 2.97 (from January to December).

29      Venzac et al. (2009) derived the boundary layer height of the puy de Dome using the ECMWF IFS model. Model  
30 results showed that the site (1465 a.s.l.) lied above the BL all times in winter while in summer it was strongly  
31 influenced by BL during the day but was above the BL at night. Collaud Coen et al. (2011) estimated BL influence at  
32 the Jungfraujoch research site (JFJ; 3580 a.s.l) by classifying daily synoptic weather situations and concluded that JFJ  
33 was largely influenced by the BL at night in summer during the subsidence weather type.

34      We derived the BL height at MBO during the study period based on the HYSPLIT model using the 40 km resolution  
35 US Eta Data Assimilation System (EDAS) meteorological data. The  $H_2O_{(g)}$  mixing ratio at MBO showed a clear  
36 average diurnal variation similar to that of the BL height (Fig. 2). This suggest that it is valid to use  $H_2O_{(g)}$  mixing  
37 ratio to distinguish FL and BL-influenced air masses in this study. Comparing to the estimated BL height, measured  
38  $H_2O_{(g)}$  mixing ratio provide more details in its temporal variations. For example, as seen in Fig. S14, from the evening  
39 of 7/25 to the morning of 7/26, the estimated mixing depth at MBO was below 300 m which was expected for a typical  
40 evening. However,  $H_2O_{(g)}$  remained high over the course (~ 6 g kg<sup>-1</sup>), suggestive of BL air masses. Indeed, the  
41 sampling site experienced relatively high pressure, temperature, and RH during this period, indicative of different  
42 synoptic conditions. In addition, nighttime  $H_2O_{(g)}$  mixing ratios in August appeared to be higher than those in July in  
43 general (Fig. S14). This may also associate with synoptic weather. Based on these observations, we believe that  $H_2O_{(g)}$   
44 is a better criterion for differentiating FL and BL-influenced air masses than modeled BL height. We based on Zhang  
45 and Jaffe (2017) and classify periods with observed  $H_2O_{(g)}$  above the minimum monthly water vapor criteria value,  
46 2.5 g kg<sup>-1</sup>, and CO > 80 ppb as those dominated by “BL-influenced air” and the rest by “FT air”.

**Table S1.** A summary of mass concentration and composition of NR-PM<sub>1</sub> species measured at different high-altitude locations around the world

Location <sup>1</sup>	Altitude (m a.s.l.)	Sampling Time	Air mass type	Instrument <sup>2</sup>	NR-PM <sub>1</sub> ( $\mu\text{g m}^{-3}$ )	Mass fraction of NR-PM <sub>1</sub> (%)					Reference
						Org	SO <sub>4</sub>	NO <sub>3</sub>	NH <sub>4</sub>	Chl	
MBO	2763	Jul. - Aug. 2013	Mixed	HR-AMS	3.16	85	11	1	3	0	this study
MBO	2763	Jul. - Aug. 2013	FT	HR-AMS	0.85	54	39	1	6	0	this study
NE Pac.	> 3000	May, 2006	FT	HR-AMS	0.7	27	53	2	18	0	Dunlea et al. (2009)
Whistler	2182	Apr. - May, 2006	Mixed	HR-AMS	1.9	55	30	3	12	0	Sun et al. (2009)
Sierra	1315	Aug. - Sept., 2007	BL	HR-AMS	4.1	73	15	5	7	0	Worton et al. (2011)
SPL	3210	Apr., 2004	Mixed	Q-AMS	2.11	42	28	10	19	1	Zhang et al. (2007)
RMNP	2740	Aug., 2010	Mixed	HR-AMS	5.13	75	17	4	4	0	Schurman et al. (2015)
WF. Mt.	600	Jul. - Aug., 2002	Mixed	Q-AMS	1.41	67	27	2	4	0	Zhang et al. (2007)
Montsec	1570	Jul. - Sept., 2011	Mixed	ACSM	7.3	4.0	1.9	0.5	0.9	0.04	Ripoll et al. (2015)
PdD	1450	Feb. 2012	Mixed	C-AMS	11.2	35	19	31	15	0	(Freney et al., 2016)
PdD	1450	Feb. 2012	FT	C-AMS	1.52	33	50	8	9	0	Freney et al. (2016)
JFJ	3580	Summer, 2013	Mixed	ACSM	0.8	57	22	8	13	0	Fröhlich et al. (2015)
JFJ	3580	Jul. 2012 - Oct. 2013	FT	ACSM	0.5	42	44	4	10	0	Fröhlich et al. (2015)
Mt. CM	2165	Jun. - Jul., 2012	Mixed	HR-AMS	4.5	63	21	7	9	0	Rinaldi et al. (2015)
Nam Co	4730	Jun. - Jul., 2015	Mixed	HR-AMS	1.84	74	16	2	8	0	Xu et al. (2018)
Mt. WZ	958	Mar. - Apr., 2015	mixed	HR-AMS	10.2	47	33	5	15	0	Zhu et al. (2016)
E. Asia	1-3 km	Mar. - May, 2001	mixed	Q-AMS	7.8	45	59	7	19	0	Bahreini et al. (2003)

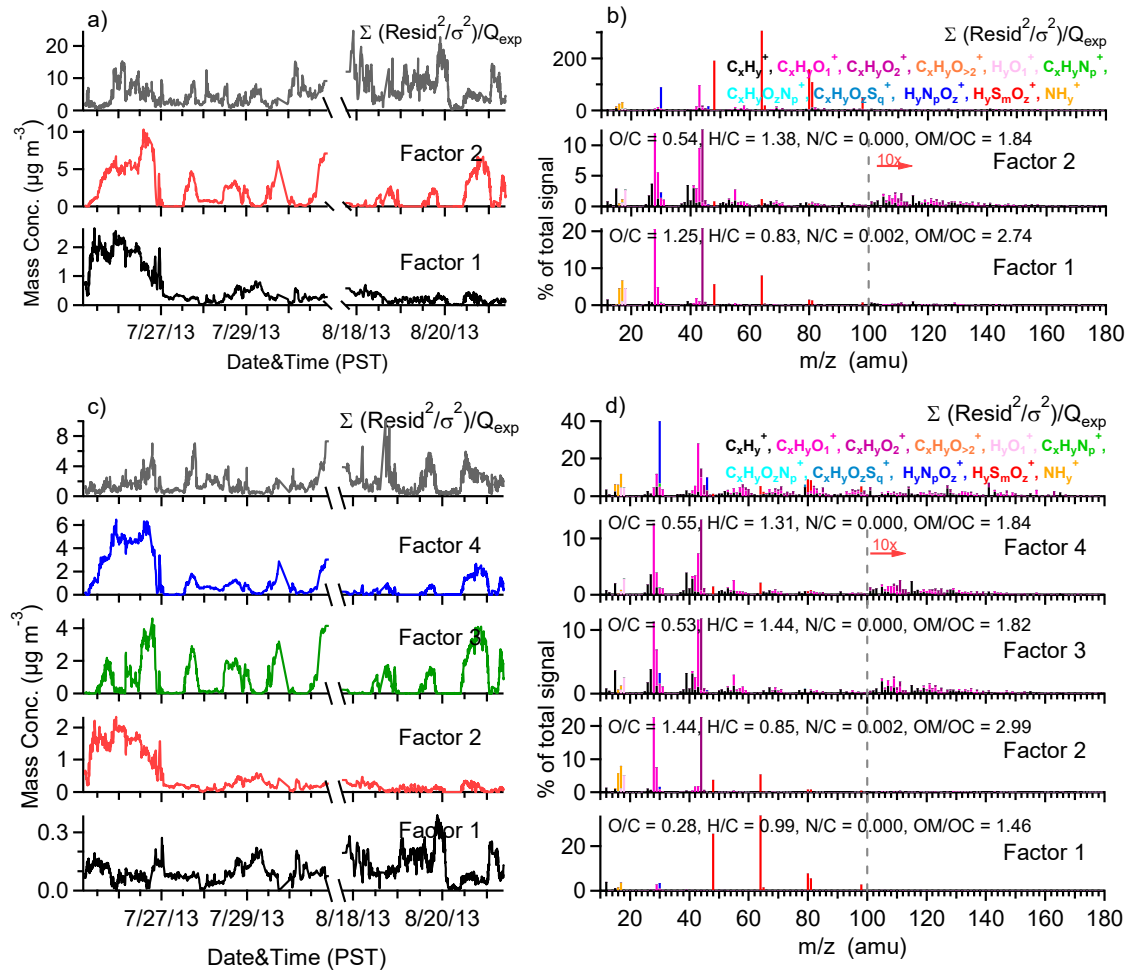
<sup>1</sup>Locations:

MBO: Mt. Bachelor Observatory, OR, USA

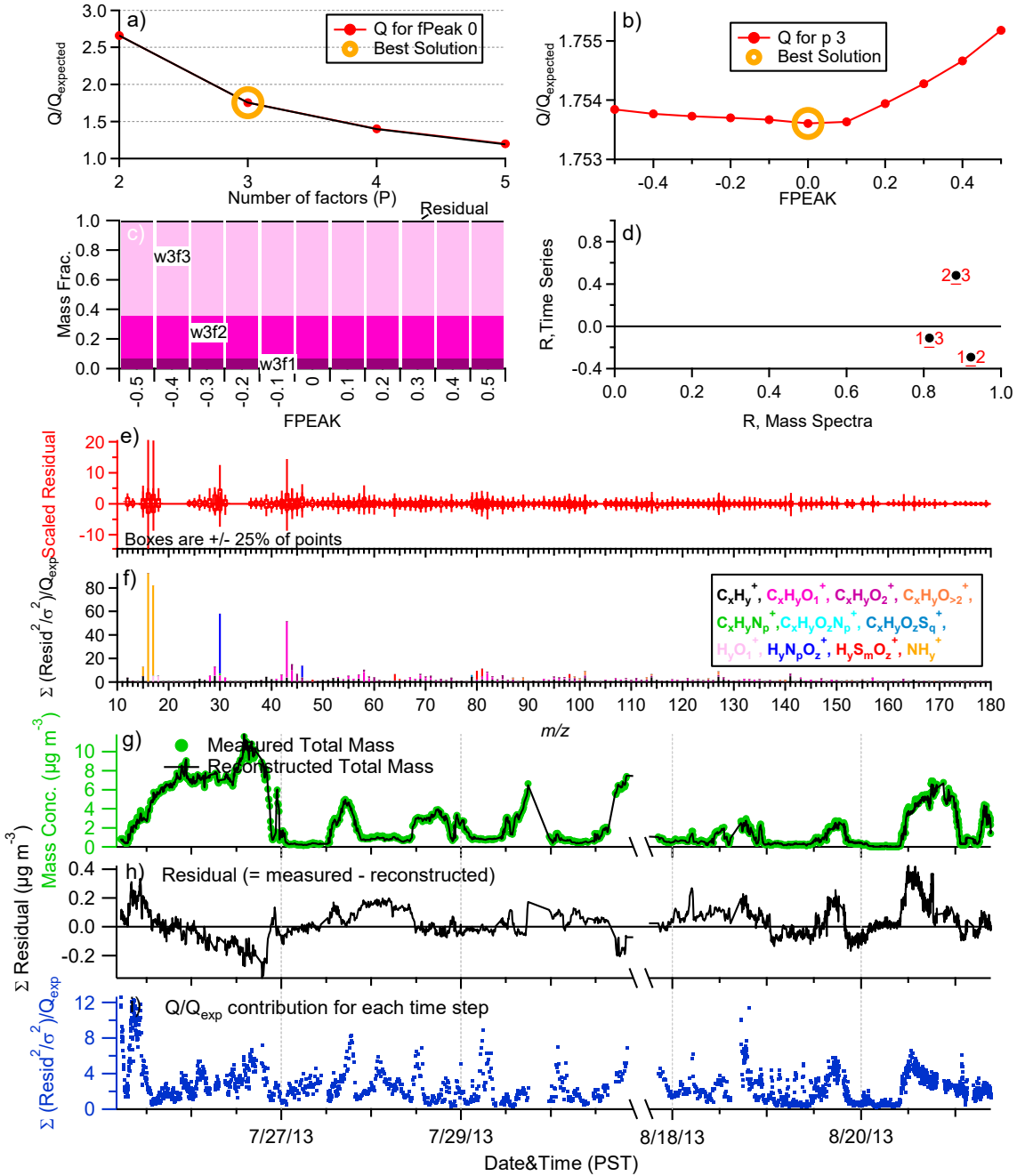
NE Pac.: Northeast Pacific  
Whistler: Whistler Mountain, Canada  
Sierra: Sierra Nevada, USA  
SPL: Storm Peak Laboratory, USA  
RMNP: Rocky Mountains National Park, USA  
WF. Mt.: Whiteface Mountain, USA  
Montsec: Montsec d'Ares mountain, Spain  
PdD: Puy-de-Dome, France  
JFJ: Jungfraujoch, Switzerland  
Mt. CM: Mt. Cimone, Italy  
Tibet: Tibet, China  
Mt. WZ: Mt. Wuzhi, China  
E. Asia: East Asia

<sup>2</sup>Instruments:

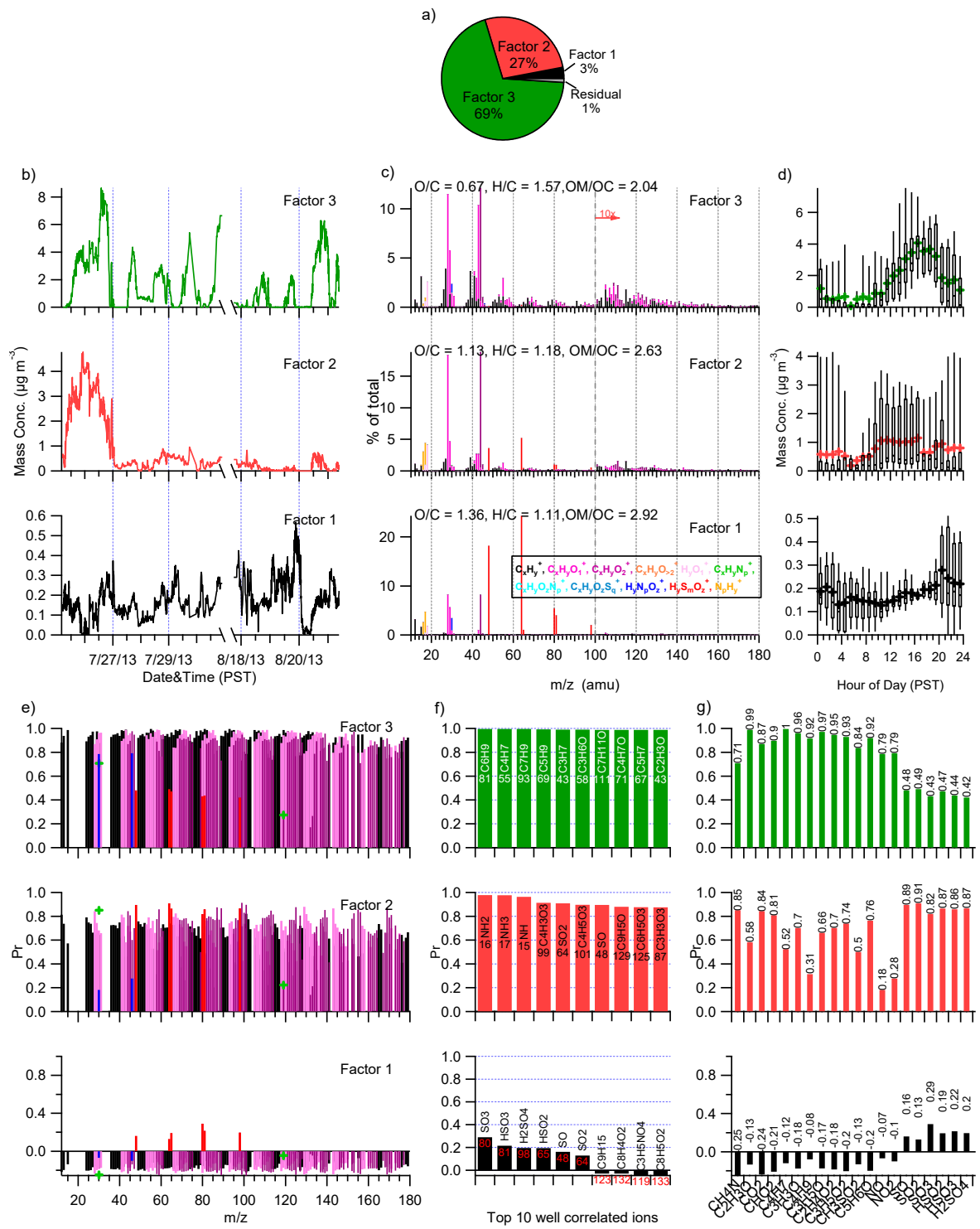
HR-AMS: High Resolution Time-of-Flight Aerosol Mass Spectrometer  
Q-AMS: Quadrupole Aerosol Mass Spectrometer  
C-AMS: Compact Time-of-Flight Aerosol Mass Spectrometer  
ACSM: Aerosol Chemical Speciation Monitor



**Fig. S1.** Time series and mass spectral profiles of total Q/Q<sub>exp</sub> and each OA factor for the 2-factor (a, b) and 4-factor (c, d) solution.

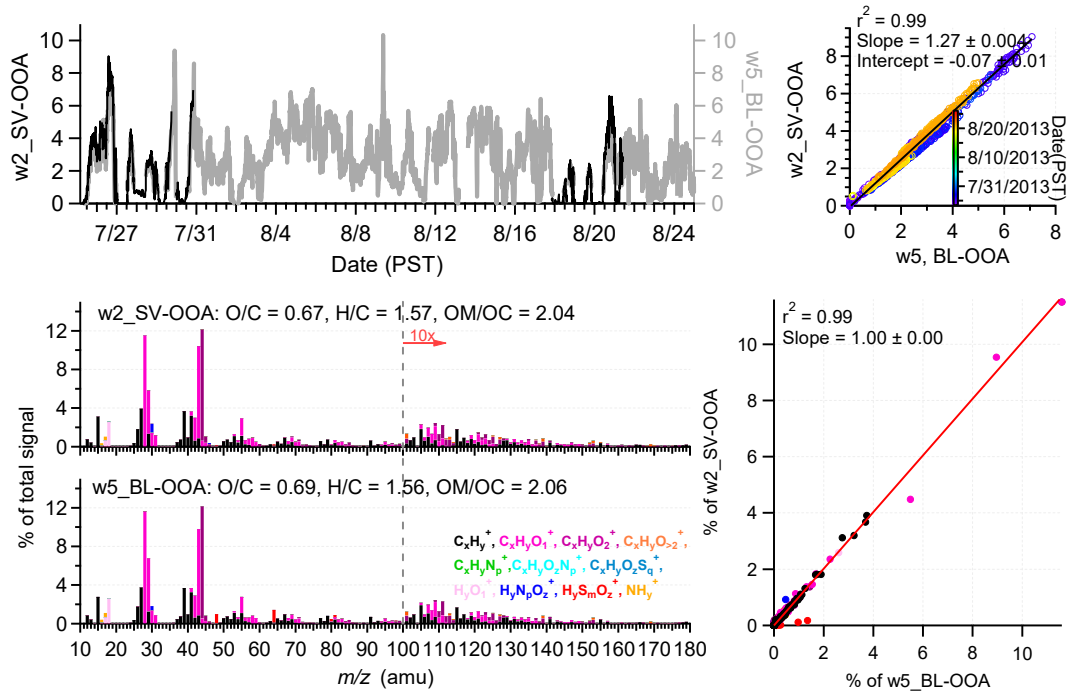


**Fig. S2.** Summary of the evaluation of PMF results for the 3-factor solution: (a)  $Q/Q_{\text{exp}}$  as a function of number of factors ( $P$ ), (b)  $Q/Q_{\text{exp}}$  as a function of fPeak values, (c) fractions of PMF factors as a function of fPeak values, (d) correlations among the 3 factors in terms of mass spectrum and time series, (e) box plot of the scaled residuals for each ion, (f)  $Q/Q_{\text{exp}}$  values for each ion, (g) time series of the measured and reconstructed NR-PM<sub>1</sub> mass concentration, (h) residual time series, and (i)  $Q/Q_{\text{exp}}$  time series.

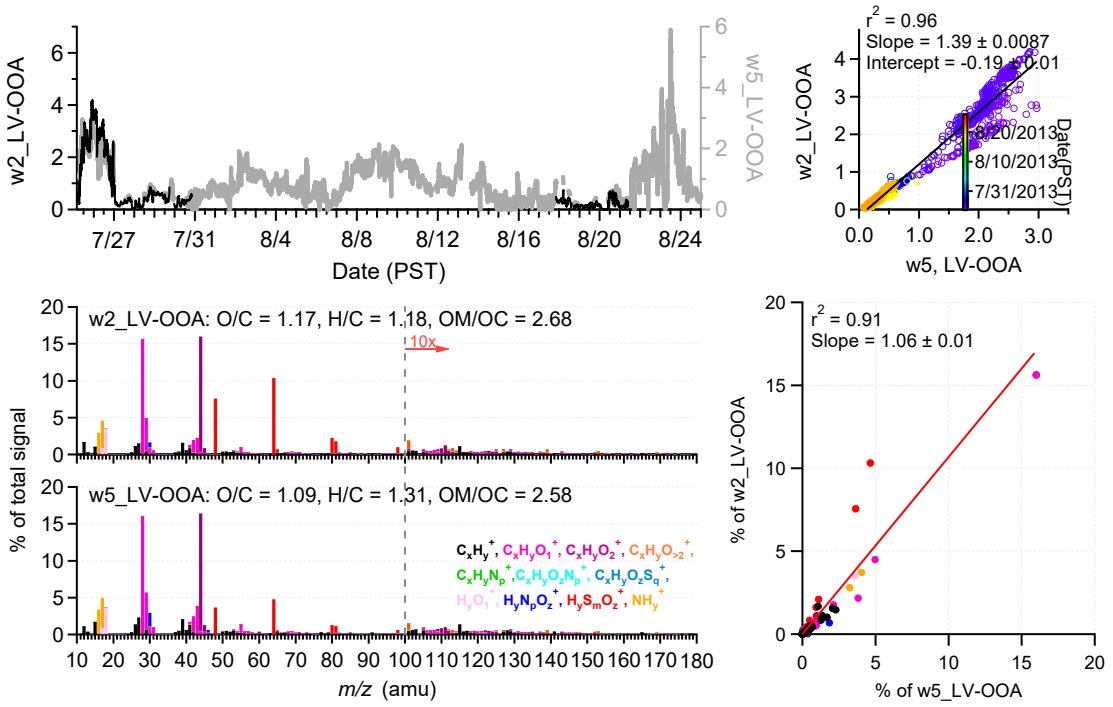


**Fig. S3.** PMF results for the 3-factor solution: (a) time series, (b) high-resolution mass spectrum (HRMS) colored by eight ion families, and (c) diurnal variations of each factor. The elemental ratios (IA method) of each OA factor are shown in the legends of (c). Correlations (Pearson's  $r$ ) between each OA factor and (e) the HRMS ions (share color code in (c)), (f) top 10 HRMS ions that showed the tightest correlation, and (g) characteristic ions. The nominal

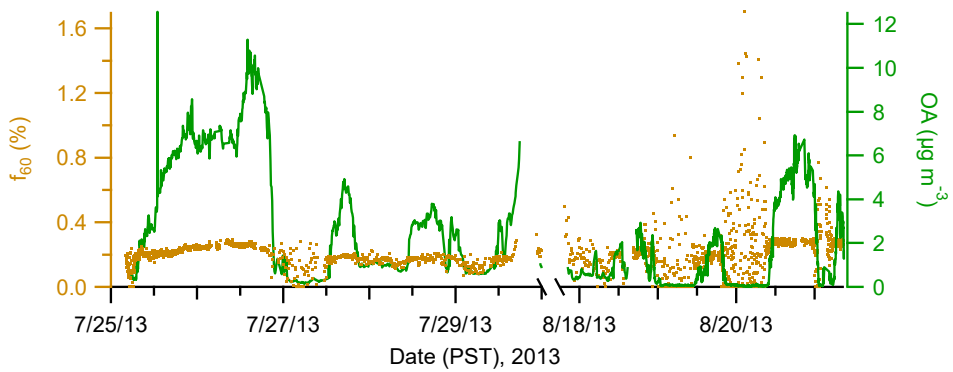
mass and identity of the top 10 HRMS ions are shown in (f). The Pearson's r number between each OA factor and the characteristic ions are shown in (g).



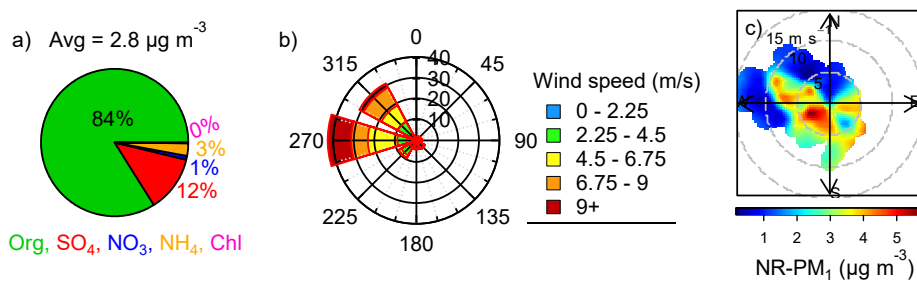
**Fig. S4.** Comparison of the time series (upper panel) and mass spectrum (lower panels) between the SV-OOA factor in the reconstructed 2-factor solution determined in this study (w2\_BL-OOA) and the BL-OOA factor from the 5-factor solution determined in Zhou et al. (2017) (w5\_BL-OOA).



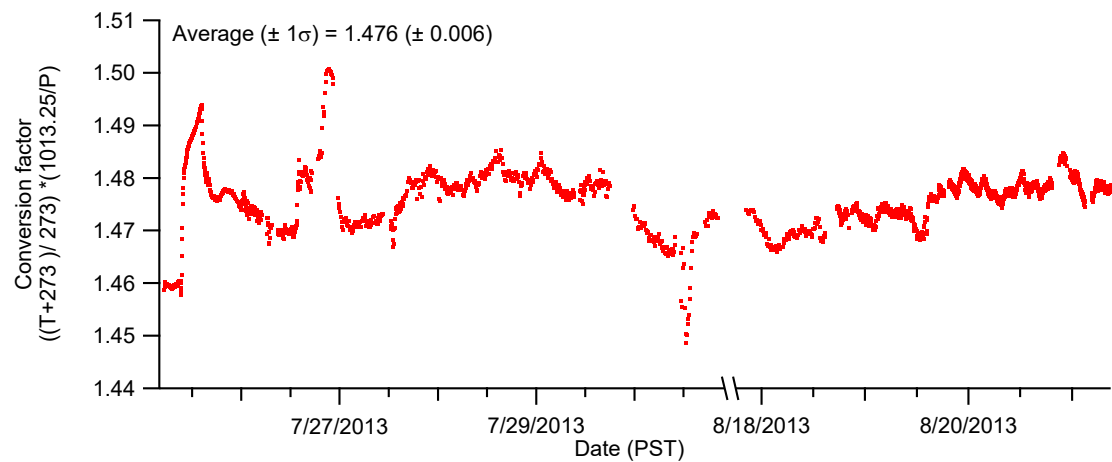
**Fig. S5.** Comparison of the time series (upper panel) and mass spectrum (lower panel) between the LV-OOA factor in the reconstructed 2-factor solution determined in this study (w2\_FT-OOA) and the LV-OOA factor from the 5-factor solution determined in Zhou et al. (2017) (w5\_LV-OOA).



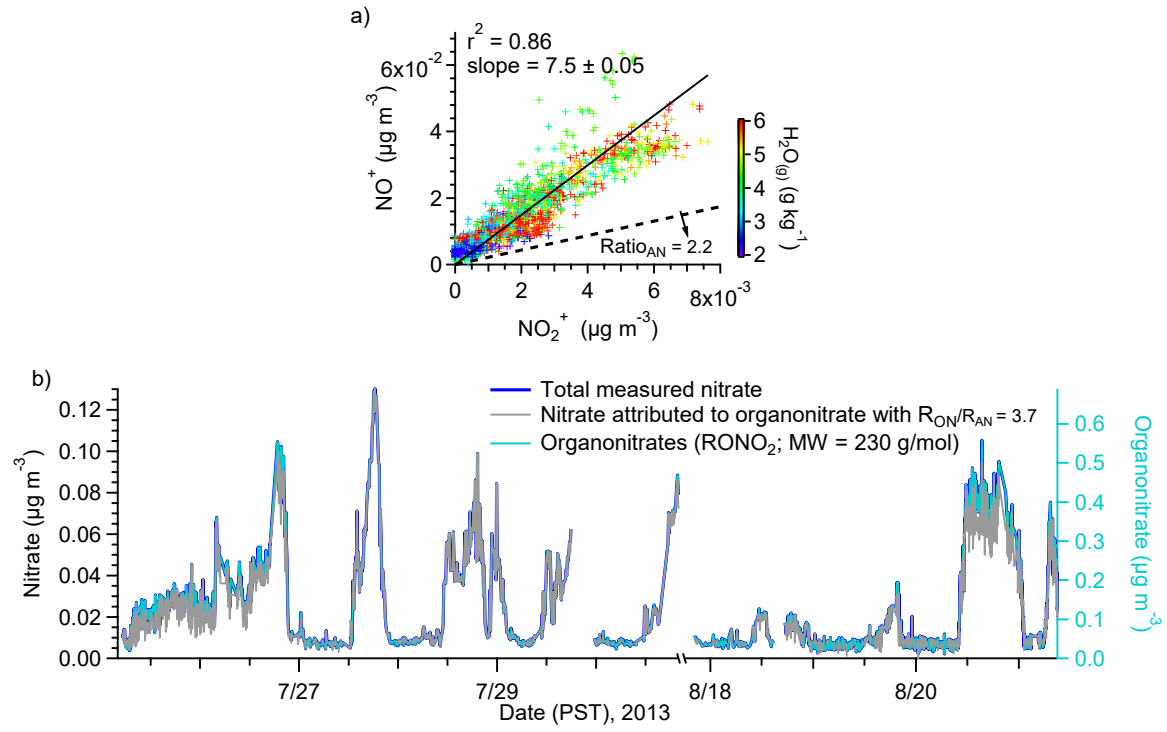
**Fig. S6.** Time series of  $f_{60}$ , the fraction of  $\text{C}_2\text{H}_4\text{O}_2^+$  ( $m/z = 60.021$ ) signal over total OA, and total OA mass concentration during the clean periods.



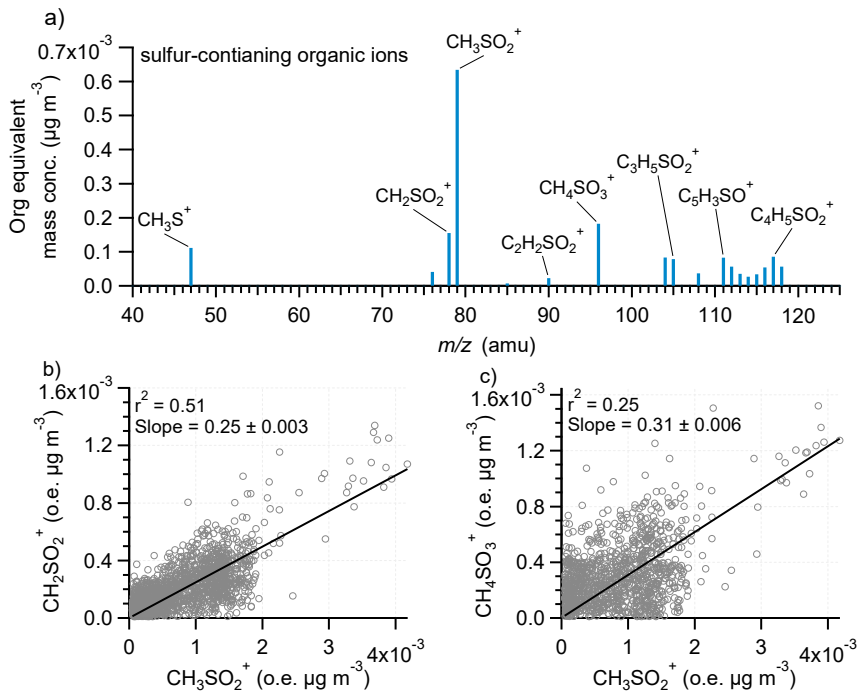
**Fig. S7.** Average (a) NR-PM<sub>1</sub> composition, (b) wind rose, and (c) bivariate polar plots of NR-PM<sub>1</sub> concentrations during the clean periods.



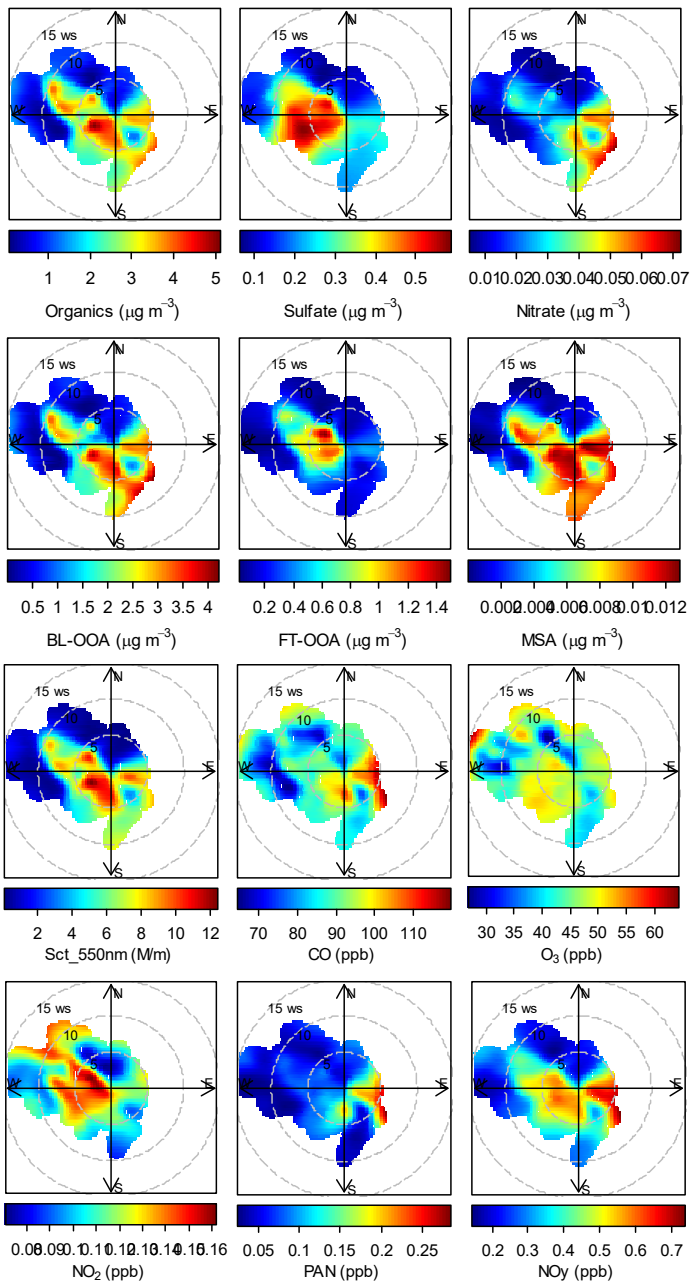
**Fig. S8.** Time series of the factor converting aerosol mass concentration under ambient temperature and pressure at MBO to that under standard temperature (K = 273 K) and pressure (P = 1013.25 hPa).



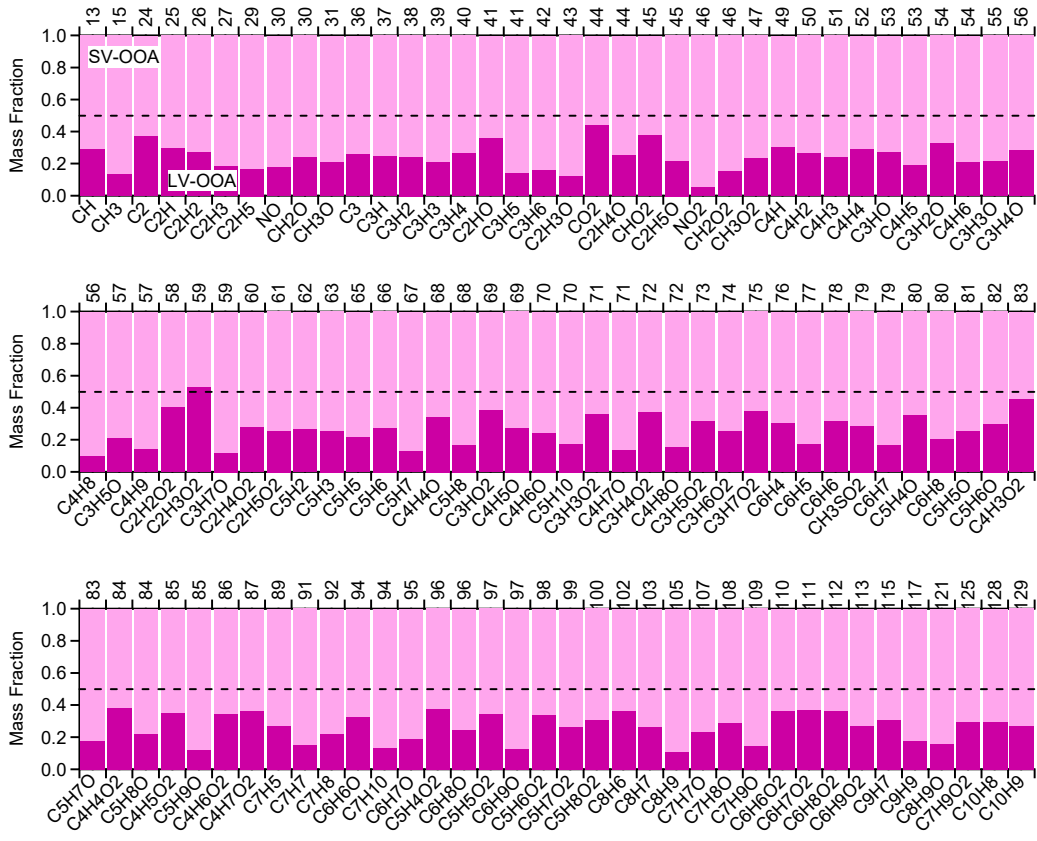
**Fig. S9.** (a) Scatter plot of  $\text{NO}^+$  vs.  $\text{NO}_2^+$  colored by water vapor mixing ratio. Data fitting was performed using the orthogonal distance regression (ODR) forced through origin. (b) Time series of total measured nitrate and estimated nitrate attributed to organonitrates on the left axis, and estimated organonitrate mass concentration on the right axis.



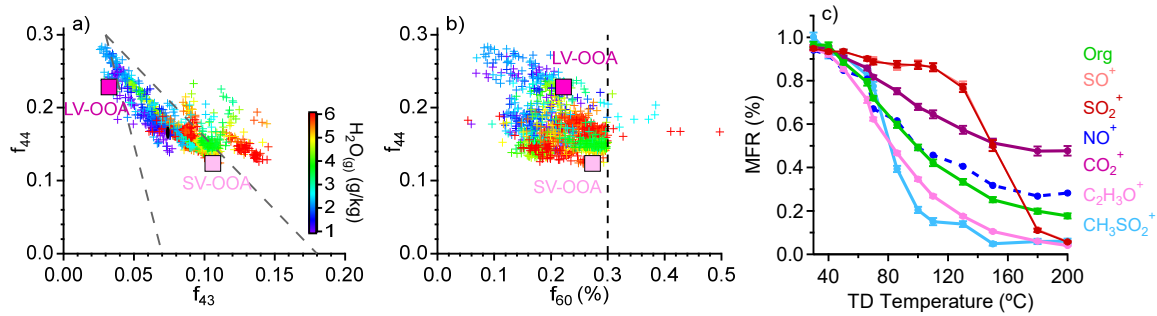
**Fig. S10.** (a) Average high-resolution aerosol mass spectrum of the organosulfur ions detected at MBO. Scatter plots of (b)  $\text{CH}_2\text{SO}_2^+$  and (c)  $\text{CH}_4\text{SO}_3^+$  vs.  $\text{CH}_3\text{SO}_2^+$  in organic-equivalent mass concentration. Data fitting was performed using the orthogonal distance regression (ODR) forced through origin.



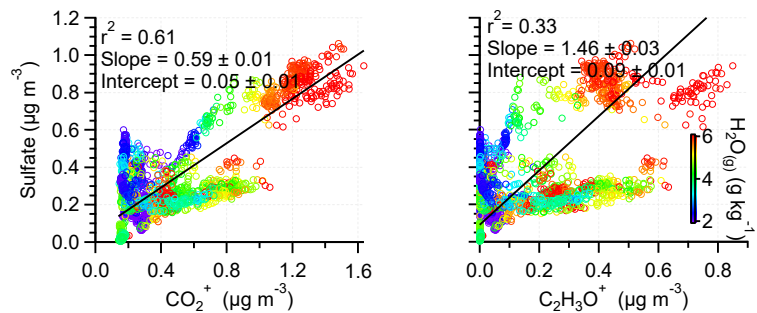
**Fig. S11.** Bivariate polar plots of aerosol and gas species measured at MBO during the clean periods.



**Fig. S12.** Ion signal distribution between two PMF factors for additional ions that were not presented in Fig.4.  
Nominal masses of individual ions are shown on the top axes.



**Fig. S13.** (a)  $f_{44}$  versus  $f_{43}$  (b)  $f_{44}$  versus  $f_{60}$  for extracted OA factors (square) and the measured ambient OA during the clean periods colored by water vapor mixing ratio (cross). The black dashed lines in (a) show the boundaries set by Ng et al. (2010). The vertical dashed line in (b) indicates the background line ( $f_{60} = 0.3\%$ ) from Cubison et al. (2011) (c) Mass fraction remaining (MFR) of total organics and selected signature ions as a function of thermodenuder (TD) temperature during the clean periods.



**Fig. S14.** Scatter plots of sulfate vs.  $\text{CO}_2^+$  and  $\text{C}_2\text{H}_3\text{O}^+$  colored by water vapor mixing ratio. Data fitting was performed using the orthogonal distance regression (ODR).  $\text{CO}_2^+$  and  $\text{C}_2\text{H}_3\text{O}^+$  are in organic-equivalent mass concentration.

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