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### **Title**

Field evaluation of the versatile aerosol concentration enrichment system (VACES) particle concentrator coupled to the rapid single-particle mass spectrometer (RSMS-3)

## **Permalink**

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## **Journal**

Journal of Geophysical Research-Atmospheres, 110(D7)

## **ISSN**

0148-0227

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## **Publication Date**

2005-02-01

## DOI

10.1029/2004|D004644

Peer reviewed

## Field evaluation of the versatile aerosol concentration

## enrichment system (VACES) particle concentrator coupled

- 4 to the rapid single-particle mass spectrometer (RSMS-3)
- 5 Y. Zhao, K. J. Bein, A. S. Wexler, C. Misra, P. M. Fine, and C. Sioutas
- 6 Received 13 February 2004; revised 28 July 2004; accepted 3 August 2004; published XX Month 2005.
- 7 [1] A field evaluation of versatile aerosol concentration enrichment system (VACES)
- 8 coupled to a rapid single-particle mass spectrometer (RSMS-3) was conducted as part
- of the U.S. Environmental Protection Agency Supersite program in Pittsburgh during
- March 2002. RSMS-3 hit rate increases were measured, and possible particle
- composition changes introduced by the VACES were examined in the single-particle
- mass spectra. The hit rates increased by 5-20 times at particle sizes ranging from 40 to
- 13 640 nm. VACES only enhances the hit rate by about a factor of 2 for large
- particle sizes because the RSMS-3 flow rates for these particles did not match the
- optimum operating condition of VACES. During the 3 days of measurements most of
- the particles were a mixture of carbonaceous material and ammonium nitrate with a
- variation across the spectrum from particles that were mostly carbonaceous to particles
- 18 that were mostly ammonium nitrate. Both ambient and concentrated carbonaceous and
- ammonium nitrate composition distributions were indistinguishable with RSMS-3,
- 20 suggesting that VACES introduces an insignificant artifact for those particles.
- 21 Citation: Zhao, Y., K. J. Bein, A. S. Wexler, C. Misra, P. M. Fine, and C. Sioutas (2005), Field evaluation of the versatile aerosol
- 22 concentration enrichment system (VACES) particle concentrator coupled to the rapid single-particle mass spectrometer (RSMS-3),
- 23 J. Geophys. Res., 110, D07S02, doi:10.1029/2004JD004644.

## 1. Introduction

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- [2] Atmospheric ultrafine particles are either formed by gas-to-particle conversion processes, in which hot and supersaturated vapors undergo condensation upon being cooled to ambient temperatures, or directly emitted as products of incomplete combustion processes [Finlayson-Pitts and Pitts, 1986]. Although the mass fraction of the ultrafine mode is negligible, this size range contains the highest number of ambient particles as well as the highest total surface area. Because of their increased number and surface area, ultrafine particles are particularly important in atmospheric chemistry and environmental health.
- [3] Recently, increasing toxicological and epidemiological evidence supports the link between respiratory health effects and exposures to ultrafine particles. Recent epidemiological studies [Heyder et al., 1996; Peters et al., 1997] demonstrate a stronger association between health effects and exposures to ultrafine particles compared to accumulation mode or coarse particles. Toxicological stud-

ies by *Donaldson et al.* [1998] indicate that ultrafine 44 particles exerted a stronger physiological effect than 45 the same mass of coarse or fine particles. A recent study 46 by *Li et al.* [2003] indicates that ultrafine particulate matter 47 (PM) is most potent in inducing cellular heme oxygenase-1 48 (HO-1) expression and depleting intracellular glutathione, 49 both sensitive markers for oxidative stress, compared to 50 concurrently collected accumulation and coarse mode PM. 51 The same study showed that ultrafine particles, and to a 52 lesser extent fine particles, localize in mitochondria where 53 they induce major structural damage.

[4] A rapid single-particle mass spectrometer (RSMS) 55 was developed at the University of California, Davis, 56 and the University of Delaware [Phares et al., 2002] 57 for measuring the size and chemical composition of indi- 58 vidual atmospheric fine and ultrafine single particles. The 59 second generation of the single-particle mass spectrometer, 60 RSMS-2, was deployed at the U.S. Environmental Protec- 61 tion Agency (EPA) Supersite in Atlanta in August 1999 62 [Rhoads et al., 2003] and in Houston from 23 August to 18 63 September 2000 [Phares et al., 2003]. Over 15,000 indi- 64 vidual particles were recorded covering 14-1300 nm size 65 range and composing of 70 compound classes in the Atlanta 66 Supersite experiment. In Houston, transient plumes of 67 ultrafine particles that were present at the site for short 68 duration were detected because of the instrument's fine 69 temporal resolution and its ability to run continuously for 70 a period of time. The RSMS-2 was modified to its third 71 generation, RSMS-3, in 2001. In comparison with RSMS-2, 72 there are two major improvements in RSMS-3: (1) both 73

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positive and negative ions can be detected and (2) two of the four digitizer channels are used to record each ion polarity signal to increase the spectral dynamic range (one channel is set at low sensitivity for strong signals and the other at high sensitivity for weak signals) and then signals in the two channels are combined by a computer program. One RSMS-3 was installed at the U.S. EPA Supersite in Pittsburgh. A quarter of million single-particle mass spectra were analyzed over a 1 year period from September 2001 to October 2002. Results indicate that a rich array of multicomponent ultrafine particles were present [Bein et al., 2005]. Another RSMS-3 was installed at the U.S. EPA Supersite in Baltimore for semicontinuous operation over 9 months from 2001 to 2002 [Lake et al., 2003; Tolocka et al., 2005], where the characteristics of specific chemical components, such as metals [Tolocka et al., 2004a], sulfate [Lake et al., 2004], and nitrate [Tolocka et al., 2004b], and association among multiple components in the same particle were examined. A disadvantage of the RSMS-3 single-particle instrument is its insufficient hit rate for all but polluted urban conditions. Laboratory tests showed that the detection efficiency of RSMS was about one in a million and varied with particle size, shape, and composition [Kane and Johnston, 2000; Phares et al., 2002]. In order to increase the RSMS hit rate for cleaner conditions and therefore to broaden its applicability, several methods are under consideration. The goal is to either increase the sampling efficiency without changing its sizing ability or concentrate particles before they enter the instrument. Theoretical work shows that the hit rate may be increased by more than 10 times using a new inlet system with capped cone structure [Middha and Wexler, 2003]. Another way to increase the hit rate is to introduce a particle concentrator to the sampling inlet of the RSMS-3 mass spectrometer, which is the topic of this presentation.

[5] A particle concentrator (versatile aerosol concentration enrichment system, VACES) has been developed at the University of Southern California and deployed in many field experiments [Sioutas et al., 1999; Kim et al., 2001a, 2001b]. In its optimum configuration, VACES concentrates fine particles, including the ultrafine mode, by a factor up to 30, depending on the ratio of total-to-minor flow rates of the virtual impactor [Sioutas et al., 1999; Kim et al., 2001a, 2001b]. Evaluation of the VACES was previously performed in both laboratory and field and the results are described in a great detail by Kim et al. [2000, 2001a, 2001b] and Geller et al. [2002]. The ability of the VACES to concentrate particles has been laboratory tested using different type of particles, including polystyrene latex (PSL), silica beads, ammonia sulfate, and ammonia nitrate, in the size range from 50 to 1900 nm and at three minor flow rates of 7, 10, and 20 liters per minute (lpm) with the major intake flow rate of 220 lpm. TSI Condensation Particle Counter (CPC) was used to measure the number concentration of the original aerosols at upstream and concentrated aerosols at the downstream of the VACES. TSI Scanning Mobil Particle Sizer (SMPS) was used to measure the size distribution of those aerosols. The resulting enrichment factors (ratio of downstream aerosol number concentration to upstream) were very close to the ideal values (ratio of total-to-minor flow rate) and the aerosol size distribution was fairly well preserved during the concentration enrichment process. Hygroscopic aerosols, such as 136 ammonium sulfate and ammonia nitrate were concentrated 137 as efficiently as hydrophobic PSL particles [Kim et al., 138 2001a]. Field evaluations of the VACES were conducted 139 outdoors in Southern California [Geller et al., 2002; Kim et 140 al., 2001b], where measurements of concentration-enriched 141 aerosols were compared to direct ambient measurements 142 made with micro-orifice uniform deposit impactor 143 (MOUDI). Downstream and upstream measurements 144 showed very good agreement (correlation coefficient  $r^2 = 145$ 0.80 for coarse particles, 0.66 for PM<sub>2.5</sub> nitrate, 0.84 for PM 146 2.5 sulfate, and 0.94 for ultrafine elemental carbon). Aver- 147 aged concentration enrichment of those aerosols was very 148 close to the ideal values. These experimental results indi- 149 cated that the concentrator does not distort the size distribution of the original ultrafine aerosols on the basis of bulk 151 measurements of particle chemical composition. Compar- 152 isons between the VACES and a reference monitor for 153 ammonium nitrate, the Harvard/EPA Annular Denuder 154 System, HEADS [Koutrakis et al., 1988], shows excellent 155 agreement between the nitrate concentrations between 156 HEADS and VACES [Kim et al., 2000].

[6] A field evaluation of the VACES concentrator cou- 158 pled to the RSMS-3 ultrafine single-particle mass spectrom- 159 eter was conducted at the Pittsburgh EPA Supersite in 160 March 2002 to determine the hit rate increase and elucidate 161 possible particle composition changes introduced by the 162 concentrator, on the basis of single-particle mass spectra.

#### 2. Methods

#### 2.1. Instrumental Setup

165 [7] Figure 1 shows a schematic diagram of rapid single- 166 particle mass spectrometer, RSMS-3. The principle of 167 RSMS-3 is nearly the same as RSMS-2, which has been 168 described in detail previously [*Phares et al.*, 2002], so only 169 a brief description is given here. RSMS-3 consists of a 170 Nafion dryer (PD-750-12SS, Perma Pure Inc., Toms River, 171 New Jersey), a rotary valve orifice bank, an inlet system, 172 two liners jointed with a source region, two Microchannel 173 Plate (MCP) detectors (25 mm BiPolar Time-of-Flight, 174 Burle Opto-Electronics Inc., Sturbridge, Massachusetts), 175 and an UV ArF Excimer laser (EX10, GAM Laser Inc., 176 Orlando, Florida). Sample air with particles passes through 177 a dryer and a rotary valve orifice bank and then arrives at 178 the inlet system composed of an aerodynamic lens with four 179 vacuum stages. A 3 lpm carrying dry air passes through the 180 dryer and removes primarily water vapor from the sample. 181 The orifice bank controls inlet pressure. The inlet system 182 creates a particle beam with a narrow particle size range and 183 skims off most of the gas. The optimum particle size that is 184 focused depends on the upstream pressure, which is con- 185 trolled by the rotary valve orifice bank. A 193 nm, pulsed 186 UV laser beam from an ArF Excimer laser is aligned 187 coaxially with the particle beam by a 45° folding-aligning 188 mirror and focused at the source region by a lens. The laser 189 emits laser pulses at 50 or 100 Hz and the laser energy is 190 between 5 and 8 mJ. When the laser beam hits a particle in 191 the source region, the particle is ablated and ionized. 192 Positive ions are accelerated by an electric field and fly 193 inside the liner to the MCP detector on the positive side of 194 the instrument. Negative ions fly in the opposite direction 195

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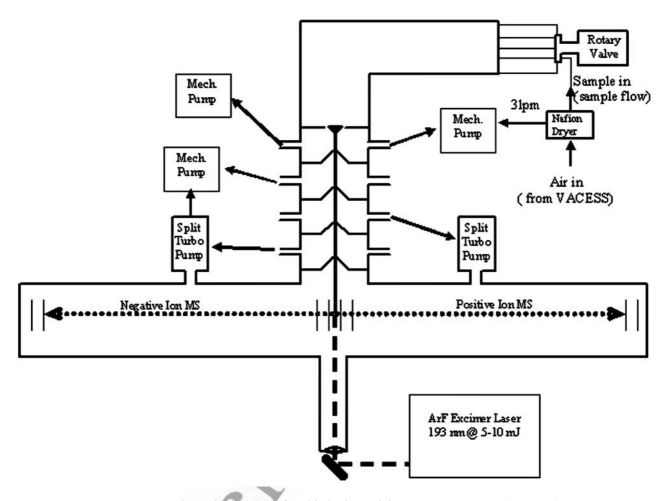


Figure 1. Schematic diagram of rapid single-particle mass spectrometer (RSMS-3).

and are detected by another MCP on the negative side. Signals from the MCP detectors are digitized by a four-channel A/D converter (two A/D channels for each MCP signal to increase dynamic range) and recorded by a computer. There are nine orifices in the orifice bank, so RSMS-3 can measure nine particle sizes. Table 1 lists the inlet pressure, flow rates, and particle sizes (Stokes number = 1.14) at each orifice. The 3 lpm air drawn from the Nafion dryer was not included in Table 1, but shown in Figure 1.

[8] Figure 2 is a schematic diagram of VACES fine plus ultrafine particle concentrator. The VACES consists of a sample line, a saturation-condensation system, a virtual impactor, and a diffusion dryer (Model 3062, TSI Inc. Shoreview, Minnesota). Sample air is drawn through the sample line into a 35°C saturation chamber above a warm DI water bath where particles and air are humidified. This warm saturated aerosol is then introduced into a section cooled by 10°C, thereby supersaturating the air and causing rapid condensation and particle growth. A virtual impactor concentrates the particles in its minor flow, which is then dehydrated so the particles return to their original sizes by means of a series of diffusion dryers. Particle enrichment by the VACES concentrator depends on the ratio of the virtual impactor's total-to-minor flow rates [Sioutas et al., 1999; Kim et al., 2001a]. The principle of a virtual impactor is similar to that of an inertial impactor [e.g., Willeke and Baron, 1993]: both

methods use particulate inertia to separate particles from 222 gases. A jet of particle-laden air is injected at a collection 223 medium, which causes an abrupt deflection of the air stream- 224 lines. Particles larger than a certain size (the so-called cut 225 point of the impactor) cross the air streamlines and, in the case 226 of an inertial impactor, are collected on the medium, while 227 particles smaller than a certain size follow the deflected 228 streamlines. The main difference between an inertial and a 229 virtual impactor is that in the latter, particles are injected into a 230 collection probe rather than onto a collection medium. To 231 separate larger particles continuously from the collection 232

Table 1. RSMS-3 Sizing Ability and Sample Flow Rates

Orifice	Orifice ID, inches/mm	Inlet Pressure, torr	Sample Flow Rate, <sup>a</sup> lpm	Sizing, nm	t1.2
1	0.063/1.600	151.7	15.4	1100	t1.3
2	0.035/0.889	60.5	4.96	640	t1.4
3	0.024/0.610	30.2	2.82	354	t1.5
4	0.017/0.432	15.2	1.44	184	t1.6
5	0.015/0.381	9.6	1.09	117	t1.7
6	0.013/0.330	9.0	0.74	109	t1.8
7	0.011/0.280	6.0	0.55	73	t1.9
8	0.008/0.203	3.2	0.23	40	t1.1
9	0.006/0.152	1.5	0.10	18	t1.1

<sup>a</sup>Here, lpm, liters per minute.

t1.12

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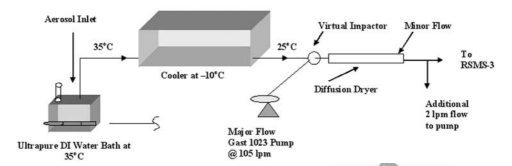


Figure 2. Schematic diagram of VACES fine plus ultrafine particle concentrator.

probe, a fraction of the total flow, referred to as the minor flow (typically 10-20% of the total flow), is allowed to pass through the probe, leaving particles larger than the cut point contained in a small fraction of the initial gases. The cut point of the virtual impactor used in VACES was 2.5  $\mu$ m.

[9] In this experiment, the minor flow of the VACES concentrator was coupled to the RSMS-3 instrument; that is, the minor flow of the concentrator is directly connected to the sample port of the RSMS-3 mass spectrometer. The two instruments were operated at their original configurations, as described in the previous paragraphs in this section, without any modifications.

### 2.2. Data Analysis

[10] After firing each laser pulse, 5000 data points of the digitizer signals were acquired and examined by a computer. A single-particle mass spectrum was recorded when the height of any peak in the selected spectral region was greater than the predefined threshold value. Afterward, the data were transmitted from Pittsburgh Supersite to the UC Davis campus for storage and postprocessing. The singleparticle mass spectra were first mass calibrated (converted from time to mass coordinates) covering the spectral range from m/z = -256 to m/z = +256. Spectra with a broad peak centering at m/z = 149 were caused by instrument emission and therefore were considered as background. After quality control to remove the background spectra, the calibrated spectra were integrated and normalized at integer m/z values and finally classified using the Adaptive Resonance Theory-2a (Art-2a) algorithm, which was first introduced by Hopke and Song [1997] for mass spectra analysis.

[11] The Art-2a algorithm uses the vector dot product as its similarity metric to classify the particles and is controlled by two parameters. The vigilance factor sets the similarity condition and the learning rate determines the rate at which the parameters adjust. The algorithm first selects each normalized spectrum in a random order and compares it to an existing set of weight vectors. If a winning weight vector is found to have the largest degree of similarity with the selected spectrum and its dot product with the corresponding particle vector is greater than the predefined vigilance factor, the selected spectrum is considered to belong to the class that the winning weight vector represents and then it is incorporated into the winning weight vector. In this case, the weight vector components are shifted toward the added spectrum's by the learning rate. If no weight vector satisfies the vigilance criterion with the selected spectrum, the particle vector becomes a new weight vector

and is then added to the set of existing weight vectors. The 280 first selected spectrum must be a new weight vector because 281 there are no existing weight vectors to compare at that time. 282 Once all spectra were selected, the whole procedure was 283 repeated with a set of weight vectors produced in the 284 previous iteration. Phares et al. [2001] validated the appli-285 cation of Art-2a algorithm in the analysis of single-particle 286 spectra generated in laboratory with aerosols composing of 287 single and mixed know chemical components. It was shown 288 that a higher vigilance factor tended to overclassify (more 289 classes than the real number) while a lower vigilance factor 290 tended underclassify (less classes than the real) these 291 laboratory-generated mass spectra. A vigilance factor of 292 0.6 was recommended to produce a class number that was 293 very close to the real number [Phares et al., 2001]; on this 294 basis, the vigilance factor was predefined as 0.6 in this 295 study. The final weight vectors are presented with equal- 296 weighted averages of all spectra belonging to that class, in 297 order to calculate the standard deviations in the peak heights 298 for each class. Therefore no learning rate was used in this 299 data analysis.

[12] Although RSMS-3 is a bipolar mass spectrometer 301 measuring both positive and negative ions simultaneously, 302 only positive spectra were analyzed with Art-2a algorithm 303 and presented in this study. Previous research shows that 304 most of negative spectra were contributed by sulfate, which 305 is very difficult to ablate and detect in the fine and ultrafine 306 particles sampled here [Kane and Johnston, 2001; Lake et 307 al., 2004]. Because of the large uncertainty in the detection 308 of negative ion spectra, they were excluded in this study.

[13] The RSMS-3 hit rates were defined as the nonbackground particle hits divided by the corresponding measurement interval. Enhancements of the hit rate by VACESS 312
concentrator were determined by ratios of the hit rate of 313
concentrated air to that of ambient air. Ideal enrichments of 314
particle concentration by VACES were predicted by the 315
ratio of impactor's total-to-minor flow rate. The detection 316
efficiency of RSMS-3 relative to the real aerosol concentration in the atmosphere will not be discussed in this work. 318
Previously studies showed that the efficiency of RSMS 319
instrument was about 10<sup>-6</sup> and varied with particle size, 320
shape, and chemical composition [Kane and Johnston, 321
2000; Phares et al., 2002].

## 3. Results and Discussion

[14] Experiments were performed during two sampling 325 periods in spring of 2002. The first, 5 March, provided 326

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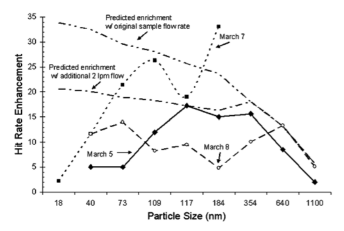


Figure 3. Enhancement of RSMS-3 hit rate by VACES particle concentrator. Predicted particle concentration enrichments were calculated using the ratio of VACES total-to-minor flow rate, where the 3 lpm flow rate passing through the RSMS-3 Nafion dryer was taken into account.

preliminary data and understanding of the use of the instruments together. The second, 7-8 March, was a more thorough test of the coupled system.

#### 3.1. Data of 5 March 2002

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[15] RSMS-3 was scheduled to measure one sample of unconcentrated ambient air followed by a concentrated sample at each particle size. The measurement time for each orifice (each particle size bin) was terminated by a maximum of 30 particle hits or 5 minutes whichever came first. Each measurement was repeated 2-3 times. The VACES concentrator was operated with a fixed major flow of 105 lpm. In total, 342 mass spectra from ambient air sample and 462 from sample with concentrator were recorded on 5 March 2002 after removal of the background

[16] Figure 3 shows the enhancement of RSMS-3 hit rates by VACES concentrator at different particle sizes observed in this study. Enrichments of particle concentration by VACES were also predicted on the basis of the ratios of total-to-minor flow rates of the impactor. Since the minor flow of the VACES concentrator was directly coupled to the RSMS-3 sample port, the minor flow of VACES was equal to the RSMS-3 sample flow listed in Table 1 plus the 3 lpm flow of the Nafion dryer which is not listed in Table 1. Diamonds connected with a solid line represent the 5 March results. The hit rate enhancement of RSMS-3 varies with particle size for a number of reasons. Since the VACES minor flow rate changed when RSMS-3 was sampling different particle sizes (see Table 1), while its major flow rate was fixed, the ratio of total-to-minor flow rate, and therefore the VACES concentration enrichment, changed with particle sizes, which is consistent with the predicted enrichment of particle concentration by VACES at large particle sizes and can readily be seen in Figure 3. With the VACES concentrator, particle hit rates of the RSMS-3 were increased by 10-17 times for the particles with sizes ranging from 109 to 354 nm. From size 640 nm to 1100 nm, the hit rate enhancements became smaller as the

RSMS-3 sampling flow rate, and therefore the minor 365 flow rate of the concentrator, increased. For the particles 366 of 1100 nm, the enhancement of hit rates was only 2. The 367 predicted enrichment was 5 at this point. Table 1 indicates 368 that the flow rate at this particle size was greater than 369 15 lpm, which plus the 3 lpm of dryer flow was too high for 370 the VACES concentrator to operate in its optimum range 371 [Sioutas et al., 1999].

[17] At the small particle side, the hit rate enhancements 373 also decreased as the flow rate decreased which is contrary 374 to the predicted particle concentration enrichment that 375 increased. At these small particle sizes, the flow rates (about 376 3 lpm) into RSMS-3 were lower than the design conditions 377 for the concentrator, so particle losses to the walls of the 378 diffusion dryer would be substantial. On the other hand, the 379 enrichment of particle concentration by VACES is affected 380 to a large extent by the actual minor flow ratio and deviates 381 from its ideal value as this ratio becomes smaller (i.e., less 382 than about 5%). This is because as this minor flow 383 decreases, particle losses mostly on the collection nozzle 384 of the virtual impactor increase, thereby decreasing the 385 overall enrichment [Marple and Chien, 1980; Sioutas et 386 al., 1994]. This is the case also with the virtual impactors 387 used in the VACES to concentrate the grown ultrafine 388 particles [Sioutas et al., 1999]. Despite these losses, the 389 hit rate enhancements at 40 and 73 nm were about 5. 390 Accuracy of the real flow rate measurement is another 391 factor in difference between the predicted VACES concen- 392 tration enrichment and RSMS-3 hit rate enhancement. The 393 ablation ability of RSMS-3 for different particle sizes [Kane 394 and Johnston, 2000] would be another cause for the 395 difference. It should be emphasized that this experiment 396 was conducted with the RSMS-3 and VACES in their 397 original configurations and the main premise of this re- 398 search was to find out problems in coupling the two 399 instruments together and ways to fix them, but no efforts 400 were made to achieve the maximum enhancement of 401 RSMS-3 hit rates.

[18] All single-particle mass spectra obtained on 5, 7, and 403 8 March were divided into two groups, with and without the 404 concentrator, to calculate the hit rate enhancements and 405

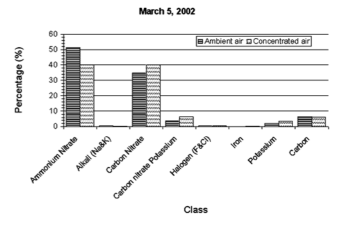
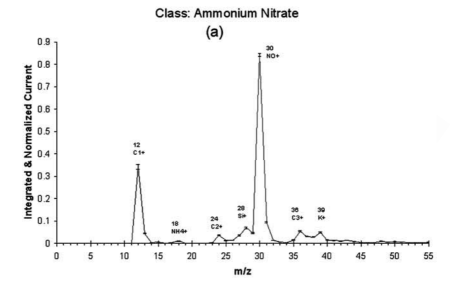
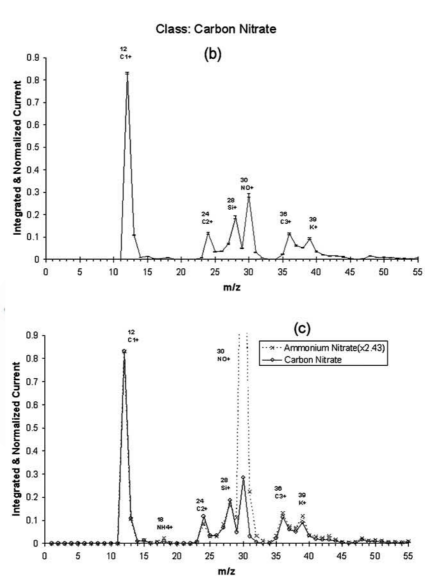


Figure 4. Comparison of chemical classes between ambient air and concentrated air measured on 5 March 2002.





**Figure 5.** Average spectra of (a) ammonium nitrate and (b) carbon nitrate and (c) comparison between the two average spectra. The ammonium nitrate spectrum has been scaled up by 2.43 in Figure 5c to show the similarity between the two spectra.

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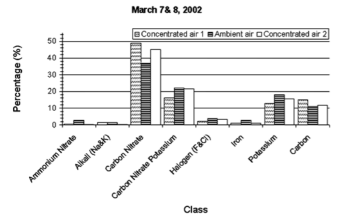
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**Figure 6.** Same as Figure 4 but for 7–8 March 2002. The concentrated air was sampled immediately before (concentrated air 1) and after (concentrated air 2) the ambient air.

classified using Art-2a algorithm. All spectra were partitioned into 8 composition classes. Each of the eight classes represented more than 10 particles and those classes containing less 10 particles were considered minor and not presented. Figure 4 shows the fractions of total hits for each class observed on 5 March. The two major classes were ammonium nitrate (nitrate peak is domain in the spectra) and carbon nitrate (carbon peaks are domain in the spectra with a small nitrate peak), whose average spectra are shown in Figures 5a and 5b. Without the concentrator, about 51% of the particles were in the ammonium nitrate class and about 35% particles were in the carbon nitrate class, while their values were about 40% and 40%, respectively, when the concentrator was used. Figure 5c compares the average spectrum of the ammonium nitrate class with the one in the carbon nitrate class. The two spectra are very similar except for the  $NO^+$  peak at m/z = 30 where the peak in the carbon nitrate class was much lower than in the ammonium nitrate class. Thus it appears that the concentrator might introduce a compositional change in a fraction of the particles, primarily shifting nitrate from particles in the ammonium nitrate class to those in the carbon nitrate class. This shift could be also due to a change in atmospheric composition during the experiment, so experiments on 7 and 8 March were designed to address this potential ambiguity. Including the effects of changes in chemical position of ambient air, the statistical uncertainty in RSMS-3 measurements, and the coupling with particle concentrator, in total about 8% particles were shifted from the ammonium nitrate class to the carbon nitrate class during the experiments on 5 March. More discussion regarding to this shift will be made in the following sections.

### 3.2. Data of 7 and 8 March 2002

[19] The RSMS-3 operating schedule was adjusted to measure one sample with concentrated air, one without, and again one with, at each particle size to separate changes in particle composition due to ambient conditions from those due to the concentrator. In order to obtain a more statistically significant sample, the measurement time was terminated at 30 particle hits at each particle size without setting a time limit. On 8 March, an additional 2 lpm was 446 drawn from the minor flow port through diffusion dryer of 447 the concentrator when RSMS-3 was sampling particles at 448 sizes from 40 to 184 nm, in order to keep the changes in the 449 ratio of total-to-minor flow rates small and minimize small 450 particle losses. In total, 227 particles were collected without 451 the concentrator and 701 with after removal of the back- 452 ground spectra.

[20] The hit rate enhancements are shown in Figure 3 for 454 7 March (closed squares) and 8 March (open circles). 455 Decrease of the hit rate enhancement for small particles 456 can still be seen on 7 March, but this dependence on RSMS- 457 3 flow rate was partially corrected by pulling out the 458 additional 2 lpm. On 7 March the hit rate enhancement of 459 RSMS-3 at 184 nm was even higher than the predicted 460 VACES enrichment of particle concentration, which is 461 likely not true and might be caused by the removal of the 462 background spectra. The causes for day-to-day hit rate 463 variation, evident in Figure 3, may be due to variation in 464 ambient conditions, RSMS-3 operating conditions such as 465 laser intensity and laser beam alignment, or imprecise 466 removal of background spectra.

[21] Class comparisons between samples collected with 468 and without the concentrator on 7 and 8 March are shown in 469 Figure 6. The four major classes were carbon nitrate (as 470 shown in Figure 5b), carbon nitrate potassium, potassium, 471 and carbon. Particles in the ammonium nitrate class were 472 not observed on these days. The experiments of 7 and 473 8 March were designed to sample concentrated particles 474 immediately before and after ambient particles to identify 475 whether or not a shift in particle composition occurred 476 during the sampling period. There is a 4% difference in 477 the particle fractions between the two concentrated samples 478 that could be due to either change in ambient air or 479 instrument drift. A 10% difference is also seen in the carbon 480 nitrate class between the ambient air and the average of the 481 two concentrated samples indicating a possible class shift 482 from other nitrate containing classes to carbon nitrate.

#### 3.3. Causes of the Class Shift

[22] Figures 4 and 6 indicate that about 8–10% particles 485 shifted from the ammonium nitrate class (or other nitrate 486 containing classes) to the carbon nitrate class during the 487 experiments on 5, 7, and 8 March when the RSMS-3 was 488 coupled with the VACES particle concentrator. As dis- 489 cussed, changes in the ambient air, operational conditions 490 of the RSMS-3, and artifacts caused by the VACES particle 491 concentrator are the possible sources for the observed 492 differences. This section will address more possible causes 493 for the class shift.

[23] Figures 7a-7d compare the chemical classes be- 495 tween concentrated particles and ambient particles at differ- 496 ent particle sizes measured on 5, 7, and 8 March 2002. 497 Although the RSMS-3 can measure nine sizes of particles 498 as given in Table 1, only two size ranges, a small size (40 – 499 117 nm) representing ultrafine particles and a large size 500 (184–1100 nm) for the fine particles minus ultrafines, are 501 grouped and shown in these figures in order to present 502 statistically significant results. On 5 March the class shift 503 from ammonium nitrate to carbon nitrate of ultrafine par- 504 ticles (Figure 7a) is the same pattern as the average of all 505 sizes (Figure 4) but the shift of fine particles (Figure 7b) is 506

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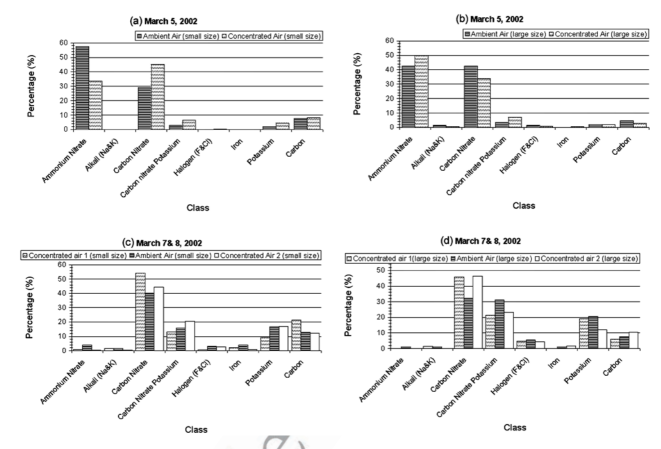
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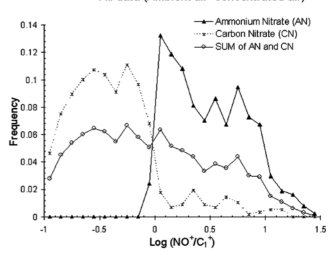


**Figure 7.** Same as Figure 4 but (a) for small size (ultrafine) particles of 5 March 2002 between 40 and 117 nm, (b) for large size (fine) particles of 5 March 2002 between 184 and 1100 nm, (c) for small size (ultrafine) particles of 7–8 March between 40 and 117 nm, and (d) for large size (fine) particles of 7–8 March between 184 and 1100 nm.

opposite. By comparison, on 7 and 8 March, the shifts at both fine and ultrafine particles (Figures 7c and 7d) are the same direction as that of the average. Observation by RSMS-3 on these days did not indicate any dependence of the class shift on particle sizes because the shift directions were random. From this point of view, it is thus unlikely that the VACES particle concentrator introduced the shift of particles from one class to another. If the concentrator did introduce such a shift, the patterns of the class shift should not change day-by-day.

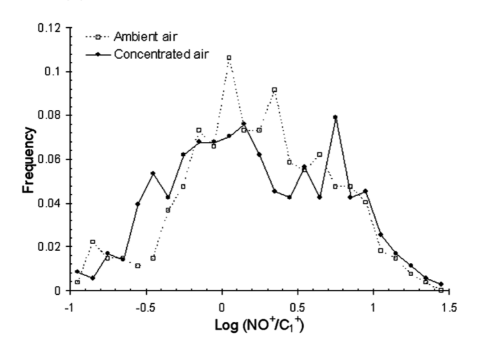
[24] Figure 5c shows that the average spectra for the ammonium nitrate and carbon nitrate classes are identical when scaled to the  $C_1^+$  peak heights, except for the height of the NO<sup>+</sup> peaks. Therefore it appears that the two classes have similar underlying carbonaceous cores with varying amounts of ammonium nitrate condensed, presumably due to varying particle age. Figure 8 shows the frequency distribution of particles from both classes as a function of the logarithm of the NO<sup>+</sup>/C<sub>1</sub><sup>+</sup> peak ratio. Areas under each curve in Figure 8 are normalized to 1. It is seen that the frequency of ammonium nitrate class has a maximum at  $NO^+/C_1^+ > 1$ , while the maximum of carbon nitrate class is located at  $NO^+/C_1^+ \le 1$ . There is a valley between the two maxima near  $NO^{+}/C_{1}^{+} = 1$ . The Art-2a algorithm breaks the two classes near this valley. Since the valley is not very pronounced and in fact the distribution between the ammonium nitrate and carbon nitrate classes is more of a

#### All data (Ambient air+concentrated air)

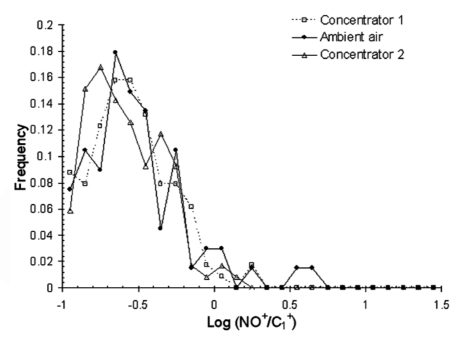


**Figure 8.** Frequency of the ratios of  $NO^+$  peak to  $C_1^+$  peak of ammonium nitrate class, carbon nitrate class, and sum of the two classes. Areas under each curve are normalized to 1.

## (a) Sum of Ammonium nitrate and Carbon nitrate, March 5



# (b) Sum of Ammonium nitrate and Carbon nitrate, March 7&8



**Figure 9.** Comparisons of the frequency between ambient air and concentrated air measured on (a) 5 March 2002 and (b) 7-8 March 2002. Each curve is the sum of the ammonium nitrate class and the carbon nitrate class. Areas under each curve are normalized to 1.

continuous distribution, subtle differences between ammonium nitrate class and carbon nitrate class near  $NO^+/C_1^+ = 1$ can move particle spectra from one class to the other. The atmospheric conditions, instrument operating conditions, and the Art-2a initial conditions could cause this movement. 538 It can also be seen in Figure 8 that some of the high  $NO^+/C_1^+$  539 particles were classified as carbon nitrate, which may be due 540 to other peak information, since when the Art-2a algorithm 541

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compares a selected spectrum with the existing set of weight vector, all peaks in the spectrum make contribution to the vector dot product. Therefore, when the contribution of other peaks to the dot product becomes more significant, the Art-2a may shift the class from one to another.

[25] Comparisons of these frequency distributions for ambient air and concentrated air are shown in Figures 9a and 9b for 5 March and 7 and 8 March, respectively. On 7 and 8 March the frequency distribution was skewed toward the  $NO^+/C_1^+ < 1$  (see Figure 9b) indicating that on that day carbon nitrate particles (carbonaceous particles with a small amount of nitrate condensed on them) were observed much more frequently than ammonium nitrate particles (carbonaceous particles with a lot of nitrate condensed on them). Figure 6 shows the same results. Since both ammonium nitrate and carbon nitrate particle classes were observed in more equal proportions on 5 March (see Figure 4), the frequency distribution appears on both sides of  $NO^+/C_1^+ = 1$  as shown in Figure 9a. There appear to be no significant difference in the frequency distributions between ambient and concentrated samples on both days and that the class shift from ammonium to carbon nitrate was not due to the particle concentrator.

### **Summary and Conclusions**

- [26] Our field evaluation of coupling VACES and RSMS-3 resulted in the following conclusions:
- [27] 1. By coupling with the VACES concentrator, hit rates of the RSMS-3 single-particle mass spectrometer increased by 5-20 times except when RSMS-3 sampled the smallest and largest particle sizes where its flow rate was off the optimum configuration of the VACES
- [28] 2. Small differences in chemical composition were observed between samples with and without the VACES particle concentrator. The shift of 8–10% particles from one class to another could be caused by the changes in the composition of ambient air, or due to statistical variation in RSMS-3 measurements, or spectrum classification. There was no evidence showing that the VACES particle concentrator introduced the particle shift.
- [29] 3. The minor flow rates of the VACES concentrator must be in its optimum range to effectively couple it to RSMS-3. Outside this range, the minor flow is either too high, resulting in a low concentration enhancement, or too low, causing ultrafine particle loss. Therefore it is necessary to control the minor flow rate of VACES in the coupling system when RSMS-3 samples different sizes of particles.
- [30] 4. Since Art-2a judges class membership on the basis of a single vigilance factor, it broke a carbonaceous particle distribution with a wide range of ammonium nitrate content into two composition classes. Better algorithms should be developed that can identify distributions of composition within a class.
- 598 [31] Acknowledgments. This work was supported by EPA and DOE funding for the Pittsburgh Supersite (Y.Z., K.J.B., and A.S.W.) and the 599 Southern California Particle Center and Supersite (SCPCS) funded by the U.S. EPA (STAR award R82735201) (C.M., P.M.F., and C.S.). This 601 602 manuscript has not been subjected to the EPA peer and policy review

and therefore does not necessarily reflect the views of the Agencies. No 603 official endorsement should be inferred.

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