

¹ Microfluidic Electrochemical Sensor for On-Line Monitoring of ² Aerosol Oxidative Activity

³ Yupaporn Sameenoi,[†] Kirsten Koehler,[‡] Jeff Shapiro,[‡] Kanokporn Boonsong,[†] Yele Sun,^{§,||} ⁴ Jeffrey Collett, Jr.,[§] John Volckens,[‡] and Charles S. Henry^{*,[†]}

⁵ Department of Chemistry, [‡]Department of Environmental and Radiological Health Sciences, and [§]Department of Atmospheric
 ⁶ Science, Colorado State University, Fort Collins, Colorado 80523, United States

7 Supporting Information

ABSTRACT: Particulate matter (PM) air pollution has a 8 significant impact on human morbidity and mortality; 9 however, the mechanisms of PM-induced toxicity are poorly 10 defined. A leading hypothesis states that airborne PM induces 11 harm by generating reactive oxygen species in and around 12 human tissues, leading to oxidative stress. We report here a 13 system employing a microfluidic electrochemical sensor 14 coupled directly to a particle-into-liquid sampler (PILS) 15 system to measure aerosol oxidative activity in an on-line 16 format. The oxidative activity measurement is based on the 17



dithiothreitol (DTT) assay, where, after being oxidized by PM, the remaining reduced DTT is analyzed by the microfluidic sensor. The sensor consists of an array of working, reference, and auxiliary electrodes fabricated in a poly(dimethylsiloxane)based microfluidic device. Cobalt(II) phthalocyanine-modified carbon paste was used as the working electrode material, allowing selective detection of reduced DTT. The electrochemical sensor was validated off-line against the traditional DTT assay using filter samples taken from urban environments and biomass burning events. After off-line characterization, the sensor was coupled to a PILS to enable on-line sampling/analysis of aerosol oxidative activity. Urban dust and industrial incinerator ash samples were

- 24 aerosolized in an aerosol chamber and analyzed for their oxidative activity. The on-line sensor reported DTT consumption rates
- (oxidative activity) in good correlation with aerosol concentration (R^2 from 0.86 to 0.97) with a time resolution of approximately 3 min.

27 INTRODUCTION

28 Airborne particulate matter (PM) is a prime candidate for the 29 generation of biological oxidative stress.^{1,2} Epidemiological and 30 clinical research has demonstrated strong links between 31 atmospheric aerosols and adverse health effects, including 32 premature deaths,³ impaired pulmonary function,⁴ neuro-33 degenerative disorders,⁵ and respiratory and cardiovascular 34 diseases.⁶ Chemical compounds in ambient PM, including 35 aromatic compounds and transition metals such as Fe, V, Cr, 36 Mn, Co, Ni, Cu, Zn, and Ti, may contribute to these effects 37 through the generation of reactive oxygen species (ROS).^{4,7,8} 38 The exact mechanism by which PM causes oxidative stress is 39 not completely understood; however, PM-associated ROS can 40 cause damage to lipids, proteins, and DNA, and these species 41 have been implicated in pro-inflammatory effects in living 42 tissues.^{4,5,7–12} In normal biological systems, generation of ROS 43 as a result of natural aerobic metabolism is balanced by 44 endogenous antioxidants.¹³ When ROS levels exceed cellular 45 antioxidant capacity, the redox status of the cell and its 46 surrounding environment changes, thereby triggering a cascade 47 of events associated with inflammation and, at higher 48 concentrations, significant cellular damage.^{14,15}

⁴⁹ Various approaches for measuring the oxidative activity of ⁵⁰ PM have been developed to study PM-induced oxidative stress.^{16–19} Chemical assays offer the best potential for analysis 51 of effective ROS dose in a format that can support 52 epidemiological research,^{1,20} and many different types of 53 chemical assays have been developed for assessing PM oxidative 54 activity.^{1,16–18,20–22} The oldest of these assays focused on 55 measuring redox-specific chemicals such as transition metals 56 and polyaromatic hydrocarbons (PAHs).²² Other methods 57 have been proposed that make use of chromatography,^{18,23} 58 electron paramagnetic resonance (EPR),²⁴ and fluores- 59 cence.^{25,26} The dithiothreitol (DTT)-based chemical activity 60 assay is currently the most widely reported technique used to 61 assess the capacity of PM to catalyze ROS generation.^{27,28} In 62 this assay, reduced DTT is oxidized to its disulfide in the 63 presence of ROS generated by PM. After the reaction, the 64 remaining reduced DTT is reacted with Ellman's reagent (5,5'- 65 dithiobis(2-nitrobenzoic acid, DTNB) to produce a chromo- 66 phore that absorbs light at 412 nm (Scheme 1). Thus, the rate 67 s1 of DTT consumption is proportional to the oxidative activity of 68 the PM sample.² Using this assay, redox-active quinones have 69 been shown to catalyze the transfer of electrons from DTT to 70 oxygen, generating superoxide.^{27,29} Furthermore, levels of PM 71

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Scheme 1. Analysis of Aerosol Oxidative Activity Using the DTT Assay with Traditional (Yellow Box) and Microfluidic Electrochemical Detection (Blue Box) Methods



72 oxidative activity measured by this assay have been correlated 73 with increased levels of biological oxidative stress in vitro.²⁸ 74 The traditional assay, however, requires the use of both 75 quenching and developing agents, which results in sample 76 dilution and a higher detection limit. Another major limitation 77 of all current assays for PM oxidative activity is that they rely on 78 a classic filter-based collection of PM. These methods require 79 long (up to 110 h) aerosol sampling durations to capture 80 sufficient mass for detection.³⁰ The long sampling times not 81 only reduce the temporal resolution of the measurement but ⁸² also increase the potential for collected species to react and ⁸³ change composition prior to analysis.^{19,21} These methods also 84 require analysis using laboratory-based instrumentation that is 85 not readily integrated into portable, field measurement systems. To overcome problems with filter collection and off-line 86 ⁸⁷ laboratory analysis, an on-line analysis system was developed by ⁸⁸ the Hopke group.^{25,26,31} Their system uses the particle-into-⁸⁹ liquid sampler (PILS)^{32,33} for aerosol sampling and a 90 dichlorofluorescein (DCFH)-based assay to determine par-91 ticle-bound ROS activity. The PILS offers the potential for 92 direct, real-time measurement of aerosol-bound ROS and 93 represents the first step toward on-line measurement of aerosol 94 oxidative activity. However, the stability of the DCHF reagent, 95 due to photobleaching and photo-oxidation, proved problem-96 atic during analysis, resulting in larger than desired 97 variability.^{34,35} In addition, an internal standard was not used to account for sample dilution by the PILS system.^{25,31,35} 98 Temporal resolution of this system was also limited (>20 min/ 99 100 sample) by the long sampling periods needed for sufficient 101 mass capture and subsequent sample flushing/rinsing periods to ensure proper detector performance.³⁵ 102

Here we present a microfluidic electrochemical sensor for 103 on-line monitoring of aerosol oxidative activity that is smaller, 104 105 less expensive, and more portable than previously reported 106 systems. Microfluidic devices can handle small sample volumes efficiently, and thus, they are attractive for field-based 107 measurements.^{36,37} They can also be multiplexed to carry out 108 109 multiple types of chemistry at the same time.³⁸⁻⁴⁰ Electro-110 chemical sensing is also well-suited for microfluidics because of 111 the ease of integration and low cost.^{41,42} By the choice of 112 detection potential and/or electrode modification, electro-113 chemistry also provides high sensitivity and high selectivity 114 even when working with low analyte masses.⁴³⁻⁴⁵ The sensor 115 reported here is based on the existing DTT assay, with several

simplifying modifications. A schematic of the procedure for 116 both the traditional and new DTT assays is shown in Scheme 1. 117 Following the reaction of DTT with PM, the remaining DTT is 118 analyzed directly by the sensor, eliminating the need for 119 quenching and developing reagents associated with UV–vis 120 detection. The electrochemical sensor is highly sensitive and 121 capable of detecting small changes in the DTT electrochemical 122 signal following reaction with a small amount of PM. Reducing 123 the required sample mass also increases temporal resolution of 124 the instrument, as less mass is needed for each individual 125 measurement. Reducing the number and quantity of reagents 126 also simplifies the system, making it more portable. 127

To create an electrochemical sensing device for DTT 128 detection, a cobalt(II) phthalocyanine (CoPC)-modified 129 carbon paste electrode (CPE) was used as an electrode 130 material. CoPC-CPE has shown good selectivity for the 131 catalytic oxidation of thiol compounds such as DTT, is stable 132 for long periods of time, and can be fabricated in a microfluidic 133 device.^{46,47} The electrode design and fabrication is based on our 134 prior work and utilizes a poly(dimethylsiloxane) (PDMS)- 135 containing binder to generate a CPE with high physical stability 136 and good electron transfer properties. The electrode 137 composition and system operating parameters were optimized 138 using cyclic voltammetry. System performance was then 139 characterized off-line using flow injection analysis and 140 amperometric detection to establish the linear range, detection 141 limit, and sensitivity of the electrode toward DTT. Then to 142 characterize the DTT assay, the working range and sensitivity 143 of the assay chemistry were then determined using a model 144 oxidant, 1,4-naphthoquinone (1,4-NQ). Reactions of DTT and 145 1,4-NQ were performed off-line, and the remaining DTT was 146 directly measured by the sensor. The sensor performance was 147 found to depend on the starting concentration of DTT, with 148 lower concentrations giving higher sensitivity but a lower 149 working range. As a final off-line validation step, 14 extracted 150 filtered samples of ambient urban PM and biomass burning 151 aerosols were analyzed. The results showed no significant 152 difference in the oxidative activity measured by the sensor 153 versus the traditional method. Finally, to demonstrate that the 154 sensor can be applied for the measurement of aerosol oxidative 155 activity in situ, we connected the sensor to an on-line aerosol 156 sampling system (Scheme 2). PM collected by the PILS reacted 157 s2 with DTT in sample transfer lines, and the remaining reduced 158 DTT was analyzed directly. Lithium fluoride was used as an 159

Scheme 2. Automated Sampling/Analysis System for Aerosol Oxidative Activity a



"Aerosol was collected by the PILS and mixed with DTT reagent and lithium fluoride (internal standard LiF). Following reaction within the sample line, the remaining DTT was analyzed directly by a microfluidic electrochemical sensor.

¹⁶⁰ internal standard to account for aerosol dilution by the PILS. A ¹⁶¹ strong linear correlation between aerosol concentration and the ¹⁶² measured oxidative activity (DTT consumption rate) was ¹⁶³ observed at concentrations similar to those found in polluted ¹⁶⁴ air (4–120 μ g m⁻³). High temporal resolution was obtained; at ¹⁶⁵ least three aerosol samples were analyzed every 10 min. To the ¹⁶⁶ best of our knowledge, the system gives the fastest time ¹⁶⁷ information on the aerosol oxidative activity available, which ¹⁶⁸ can greatly contribute to the future understanding of how ¹⁶⁹ aerosols affect human health during short-term exposure events.

170 RESULTS AND DISCUSSION

Electrode Composition and System Optimization. 172 Carbon paste electrodes, a mixture of graphite and binders 173 (mineral oils, nonconducting polymers, etc.), have shown 174 potential as electrochemical sensors in microchip devices^{48,49} 175 because of their ease of fabrication and the ability to modify the 176 electrode with a range of chemically selective dopants.⁵⁰ 177 Various methods have been reported for carbon paste electrode 178 fabrication on microfluidic devices, including the insertion of

tube sleeves into the device and screen printing.^{49,51} Of these 179 methods, screen printing is particularly attractive because it can 180 be performed directly on-chip with electrode dimensions 181 controlled by screens or channels on the device itself. In this 182 work, an electrode fabrication method analogous to screen- 183 printing was used to produce on-chip electrodes using carbon 184 paste with a custom-designed binder (details for electrode 185 fabrication provided in the Supporting Information, Figure S- 186 1).⁴⁴ We found that these electrode systems provided 187 robustness and good electrochemical properties. The electro- 188 chemical sensor can be reused over a month by being rinsed 189 with deionized water daily. They were also characterized with 190 catecholamines that have limited fouling potential. DTT 191 requires the addition of a catalyst, however, to reduce the 192 oxidation potential and reduce fouling.^{55,56} Cobalt phthalocya- 193 nine (CoPC) is one of the more common electrocatalytic 194 agents used and acts as a redox mediator that lowers the 195 overpotential for thiols. $^{52-54}$ The two-step electrocatalytic 196 mechanism starts with the electrochemical oxidation of 197 cobalt(II) phthalocyanine to cobalt(III) phthalocyanine, 198 followed by the chemical oxidation of DTT and regeneration 199 of the cobalt(II) phthalocyanine.⁵² Since both the solution pH 200 and CoPC composition impact DTT detection, the signal for 1 201 mM DTT as a function of the CoPC concentration (%) and 202 solution pH was studied using cyclic voltammetry. 52,55-57 A 203 two-variable experimental design was used for this optimization 204 study.⁵⁸ Optimal values giving the highest signal of anodic 205 current (scan range -0.1 to +1 V vs unmodified CPE) were 206 obtained at a CoPC concentration of 12% (w/w) and a 207 solution pH of 7 (detailed discussion provided in the 208 Supporting Information, Figure S-2). Therefore, this compo- 209 sition of CoPC was used for CPE modification, and a solution 210 pH of 7 was used as the running buffer for all subsequent 211 experiments. 212

As a first step to test the performance of the sensor, a flow 213 injection analysis system was designed for off-line measure- 214 ments of aerosol oxidative activity. The voltammetric behavior 215 of the systems was established first (Figure 1A). The 216 fl voltammogram shape is different from those of most 217 hydrodynamic voltammograms where the current plateaus at 218 higher potentials because of mass transport. The unusual 219 behavior shown here can be attributed to many factors such as 220 additional oxidation and decomposition of the phthalocyanine 221 ring at higher potentials and irreversible complexation of the 222



Figure 1. Selectivity of the microfluidic electrochemical sensor for DTT. (A) Hydrodynamic voltammogram plotted as the signal-to-noise ratio as a function of the applied potential from 100 μ M DTT injection (n = 3). (B) Flow profiles from injections of DTT and extracted aerosol samples.

223 Co(III) center.^{57,59,60} While the highest signal-to-noise ratio $_{224}$ (S/N) was observed at +0.5 V, we selected a potential of +0.2 V 225 for selective DTT detection to avoid potential interference (i.e., 226 oxidation signal) from other redox-active species typically 227 present in ambient aerosols. These species include metals such 228 as Fe, Cr, V, and Ni and a broad spectrum of organic 229 compounds such as PAHs (for example, pyrene, fluoranthene, 230 chrysene), redox cycling agents (hydroquinones), olefins, 231 aldehydes, ketones, and nitro compounds.^{61,62} Although some 232 chemicals can be oxidized at 0.2 V (according to standard 233 reduction potentials), our electrochemical sensor is chemically 234 modified as discussed above for selective detection of DTT .⁶³ For each measurement, we also injected an aerosol sample 235 extract in the absence of DTT (i.e., as a negative control) to 236 ensure the sample did not contribute to the electrochemical 237 signal. Since aerosol composition is highly variable, 14 different 238 239 filter samples were employed to test for interferences. These samples included biomass burning aerosol and urban aerosols 240 collected during both summer and winter seasons. All samples 241 showed negligible inferences at the DTT detection potential 242 (data not shown). Example results (Figure 1B) show a high 243 244 signal for DTT (20 μ M) and no signal for the extracted aerosol 245 sample.

Analytical Figures of Merit. After determination of the 246 247 optimal electrochemical conditions, the figures of merit for 248 DTT were determined to ensure that the experimental 249 conditions provide effective analysis for the remaining reduced 250 DTT using flow injection analysis for DTT concentrations of 251 10–100 μ M. A plot of the average peak current (nA) (n = 3) as 252 a function of the DTT concentration (μ M) gave a linear 253 calibration curve from 10 to 100 μ M (y = 0.037x - 0.43, $R^2 =$ 254 0.997) (see the Supporting Information, Figure S-3). The 255 relative standard deviation from 10 consecutive injections of 256 100 μ M DTT was 7.0%, and electrode fouling was not 257 observed (data not shown). The limit of detection for DTT defined as the concentration that gives a signal $3 \times$ larger than 258 259 the baseline noise was 2.49 \pm 0.20 μ M (n = 5) (equivalent to 260 24.9 pmol for a 10 μ L injection), which is comparable to those of similar microfluidic electrochemical sensors. 57,64 261

262 **Sensor Performance Study.** Following calibration, the 263 sensor was used to measure PM oxidative activity. The effect of 264 the DTT starting concentration on the assay dose–response 265 curve using 1,4-NQ as a model oxidant was studied for 266 sensitivity and working range. The results shown in Figure 2 267 demonstrate that, at low starting DTT (25 nmol), the signal



Figure 2. Impact of the initial DTT amount on the assay dose-response (n = 3).

dropped quickly with increasing 1,4-NQ concentration, 268 providing the highest sensitivity of all three conditions tested 269 (sensitivities of -1.50%, -0.75%, and -0.50% DTT remaining/ 270 ng of 1,4-NQ for 25, 50, and 75 nmol, respectively). The 271 decrease in signal at higher starting DTT levels (75 nmol) is 272 more gradual but provides a larger assay working range. These 273 results indicated that the assay sensitivity and working range 274 can be tuned according to the levels of DTT present in 275 solution. In all remaining off-line assays, 25 nmol (50 μ L of 0.5 276 mM) of DTT was used to provide high sensitivity at the low 277 oxidative activity of our PM samples.

Finally, the new sensor was compared to the traditional DTT 279 assay for aerosol oxidative activity using 14 representative 280 aerosol filter samples (Supporting Information, Table S-1). The 281 1,4-NQ equivalent values obtained by the two methods were 282 compared using a paired t test and plotted for correlation 283 (Figure 3). There was no significant difference ($t_{obsd} = 1.621$, 284 f3



Figure 3. Comparison of PM oxidative activity (1,4-NQ equivalent unit, $ng_{NQ}/\mu g_{PM}$) between the traditional DTT assay and the microfluidic electrochemical sensor (off-line). The data represent aqueous extracts of 14 different aerosol samples.

 $t_{\rm critical} = 2.179$, p = 0.05) in the 1,4-NQ equivalent values 285 obtained using the electrochemical DTT sensor and the 286 traditional DTT assay. A good correlation for the 1,4-NQ 287 equivalent values determined by the two methods was observed 288 (R^2 = 0.96). The equivalence between the two methods 289 demonstrates that the new electrochemical DTT assay is 290 suitable for the measurement of oxidative activity from PM 291 samples collected on filters. Moreover, the electrochemical 292 assay for filter samples requires 100 times less sample for 293 detection when compared to the traditional assay (10 μ L vs 294 1000 μ L).^{27–29,65} This reduction allows for a commensurate 295 reduction in field sampling duration, representing a significant 296 advantage over the traditional DTT assay. The inclusion of 297 several aerosol types (biomass burning smokes, urban winter 298 aerosols, urban summer aerosols) indicates that this finding is 299 not restricted to a small class of aerosol types. 300

On-Line Aerosol Oxidative Activity Measurement. 301 After the electrochemical microfluidic sensor was validated 302 for measurement of DTT consumption by PM from filters, the 303 sensor was connected to a PILS to create an on-line aerosol 304 oxidative activity analysis system (Scheme 2). 305

As a first step, the system was tested to show the ability to 306 detect reduced DTT in the presence of nonoxidizing aerosols. 307 Sodium chloride (NaCl) aerosol, which has no oxidative 308 activity, was created in the chamber at various concentrations 309 and the DTT signal measured. The results shown in Figure 4A 310 f4

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Figure 4. Initial study of the on-line aerosol oxidative activity system. (A) DTT (black) and fluoride (green) response curve as a function of the salt aerosol concentration without oxidative activity. DTT levels were measured using the electrochemical sensor. F^- levels were measured using ion chromatography. (B) DTT signal response as standard oxidant (1,4-NQ) was added to react on-line without aerosol delivered.



Figure 5. Correlation of the DTT consumption rate with the aerosol concentration for (A) standard reference material urban dust and (B) standard reference fly ash. The top panels show the DTT consumption rate (black y axis) and aerosol concentration (green y axis) as a function of the experiment time. The bottom panels plot the DTT consumption rate as a function of the aerosol concentration and show the resultant correlation coefficient.

311 demonstrate a decreasing DTT signal for increasing NaCl 312 aerosol concentration (and thus increased water) delivered to 313 the PILS impaction plate, which is indicative of sample dilution. 314 The internal standard was used to account for this dilution 315 effect and to correct the measurement of aerosol oxidative 316 activity. A decrease in measured fluoride concentration, 317 proportional to the salt aerosol concentration, is also shown 318 in Figure 4A and demonstrates our ability to account for this 319 phenomenon.

In the on-line system, various experimental conditions were different from those of our validated off-line assay, including temperature (\sim 37 to \sim 28 °C) and the chemical mixing environment. The on-line DTT assay was therefore tested using 1,4-NQ as a model oxidant. For this test the PILS was allowed to sample only filtered, particle-free air. To simulate exposure to an oxidant under the conditions of the PILS, 1,4NQ was injected through a T-valve at the entrance to the PILS, 327 which is above the impaction plate. The decreasing DTT signal 328 for higher concentrations of 1,4-NQ demonstrates DTT 329 consumption by a standard oxidant under on-line operation 330 (Figure 4B). The amount of 1,4-NQ used was equivalent to 331 what was used in the off-line system, and the DTT 332 consumption of the standard oxidant was of the same 333 magnitude as that observed in the off-line system, indicating 334 the viability of the on-line DTT assay. 335

To demonstrate on-line performance more fully, standard 336 reference samples of urban dust and fly ash (industrial 337 incinerator ash) were aerosolized, sampled, and analyzed for 338 their oxidative activity directly. These aerosols were selected 339 because of their varying chemical composition and because they 340 represent typical toxicants found in outdoor air. For each 341 sample, aerosol concentrations generated in the chamber were 342

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343 in the range of those found in the urban atmosphere $(4-120 \mu g$ $_{344}$ m⁻³). The results shown in the top panel of Figure 5A show 345 DTT consumption corresponding to urban dust aeroso 346 concentrations. As the aerosol concentration increased, th 347 DTT consumption rate increased. Furthermore, the system 348 provided high temporal resolution, reporting an independen 349 measurement approximately once every 3 min. To the best of 350 our knowledge, this is the highest temporal resolution for a 351 aerosol oxidative activity measurement system that has bee 352 reported.^{25,26,31} Higher temporal resolution could ultimately b 353 obtained by reducing the volume of the injection loop and 354 increasing the buffer flow rate through the system. In terms of 355 aerosol mass, the on-line system required between 7 and 214 n 356 of particle mass per injection to observe quantifiable DTT consumption. This range was calculated using the PILS air 357 sampling rate (12.5 L min⁻¹), aerosol concentrations measured 358 359 in the chamber (4–120 μ gm⁻³), and a 10 μ L injection loop. 360 This mass range is approximately 3 orders of magnitude lower 361 than the amount required for the traditional DTT assay (5-40) $_{362}$ µg).^{27–29,65} A correlation plot between DTT consumption rate 363 and aerosol concentration was constructed, and a strong linear 364 correlation coefficient was obtained ($R^2 = 0.97$) (Figure 5A, 365 bottom panel). The performance of the system was further 366 confirmed with a fly ash test aerosol. In this example, the fly ash aerosol concentration was varied faster and the DTT 367 consumption rate was analyzed (Figure 5B, top panel) 368 continuously. The on-line system was able to measure DTT 369 370 consumption rates that were strongly correlated with the aerosol concentration ($R^2 = 0.86$), even during periods of rapid 371 concentration change. The oxidative activities of the urban dust 372 and fly ash samples were comparable. The oxidative activity of 373 the fly ash aerosol is believed to result from the transition-metal 374 content of the sample.^{66,67} ROS generated from urban dust 375 376 aerosol might be attributed to PAHs and nitro-PAHs, which are major components of such a sample.⁶⁸ 377

378 CONCLUSIONS

379 We present here for the first time a high temporal resolution 380 on-line sampling/analysis system for aerosol oxidative activity 381 using a microfluidic electrochemical sensor coupled with an on-382 line aerosol collection system. The determination of aerosol 383 oxidative activity was based on the widely reported DTT assay 384 but used electrochemical detection instead of photometric detection. The sensor was validated off-line for its performance 385 386 in aerosol oxidative activity measurement. No significant differences for the aerosol oxidative activity expressed as the 387 388 1,4-NQ equivalent were observed between the traditional assay 389 and the sensor for 14 extracted ambient aerosol and biomass 390 burning smoke filter samples. Using on-line monitoring of 391 aerosol oxidative activity, high correlations between aerosol 392 concentration and DTT consumption rate were observed for 393 two representative test aerosols. The on-line system developed 394 here shows promise as an eventual tool for field studies of 395 aerosol oxidative activity. Such studies may lead to a better 396 understanding of how PM can affect human and environmental 397 health.

398 **ASSOCIATED CONTENT**

399 Supporting Information

400 Experimental details, electrode fabrication diagram, electrode 401 composition optimization results, DTT flow injection profiles 402 and standard calibration, and extracted aerosol sample 414

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information for the method validation test. This material is 403 available free of charge via the Internet at http://pubs.acs.org. 404

AUTHOR INFORMATION	405
Corresponding Author	406
chuck.henry@colostate.edu	407
Present Address	408
^{II} State Key Laboratory of Atmospheric Boundary Layer and	. 409
Atmospheric Chemistry Institute of Atmospheric Physics,	410
Chinese Academy of Sciences, Beijing, China.	411
Notes	412
The authors declare no competing financial interest.	413

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