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1 Long-term evaluation of air sensor technology under ambient conditions in Denver, Colorado

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ABSTRACT

14 Air pollution sensors are quickly proliferating for use in a wide variety of applications, with a low price 15 point that supports use in high density networks, citizen science, and individual consumer use. This 16 emerging technology motivates the assessment under real-world conditions, including varying pollution 17 levels and environmental conditions. A seven-month, systematic field evaluation of low-cost air 18 pollution sensors was performed in Denver, Colorado over 2015-2016; the location chosen to evaluate 19 the sensors in a high altitude, cool, and dry climate. A suite of particulate matter (PM), Ozone (O₃), and 20 nitrogen dioxide (NO2) sensors were deployed in triplicate, and were collocated with Federal Equivalent 21 Method (FEM) monitors at an urban regulatory site. Sensors were evaluated for their data 22 completeness, correlation with reference monitors, and ability to reproduce trends in pollution data, 23 such as daily concentration values and wind-direction patterns. Most sensors showed high data 24 completeness when data loggers were functioning properly. The sensors displayed a range of 25 correlations with reference instruments, from poor to very high (e.g. hourly-average PM Pearson 26 correlations with reference measurements varied from 0.01 to 0.86). Some sensors showed a change in 27 response to laboratory audits/testing from before the sampling campaign to afterwards, such as the 28 Aeroqual, where the O₃ response slope changed from about 1.2 to 0.6. Some PM sensors measured 29 wind-direction and time of day trends similar to those measured by reference monitors, while others did 30 not. This study showed different results for sensor performance than previous studies performed by the 31 U.S. EPA and others, which could be due to different geographic location, meteorology, and aerosol 32 properties. These results imply that continued field testing is necessary to understand emerging air 33 sensing technology.

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1. INTRODUCTION

Next generation air monitoring (NGAM) is a quickly evolving and expanding field. Low-cost air pollution

36 sensors have improved the access for both citizens and researchers to obtain pollutant concentration

37 data in more locations. Many new sensors are now sold and marketed to consumers, and come with

38 messaging on implications for health. In addition to improving the accessibility of measurement data, air

pollution sensors have been used to supplement ambient air monitoring by providing high spatial

40 density and high time-resolution measurements (Mead et al., 2013; Snyder et al., 2013; Kaufman et al.,

41 2017). Low-cost air pollution sensors have the potential to be important enablers of smart cities and the

42 Internet of things (IoT), especially in terms of forecasting and health messaging in megacities with

43 significant variability in microenvironments (Mead et al., 2013; Kumar et al., 2015; Ramaswami et al,

2016). Sensors also enable new techniques for mobile monitoring. (McKercher and Vanos 2017;

45 Woodall et al., 2017). However, without a proper understanding of sensor data quality and calibration,

46 low-cost sensors have the potential to mislead interested community and research groups (Rai et al.,

47 2017). Evaluating how well these sensors perform in both laboratory and field environments is critical

48 for understanding their possible uses in research, citizen science, and consumer use, for individual

49 exposure assessment.

50 Low-cost air pollution sensors, with purchase prices ranging from the low hundreds to the low

51 thousands of dollars per pollutant, have been developed for both particulate and gas phase pollutants,

52 including ozone (O₃) and nitrogen dioxide (NO₂). Particulate matter (PM) sensors typically measure

53 particle counts using light scattering principles. By using light scattering to measure an ensemble of

54 particles, sensors can be produced that are miniaturized, lower cost, and provide real-time data.

However, this detection approach can result in bias and inaccuracy from measurement artifacts (Gao et

al., 2015; Holstius et al., 2014). Some sensors, such as the OPC-N2 (Alphasense) measure single particles

and allocate them into size bins. This approach is subject to measurement artifacts due to humidity

58 effects, potential particle coincidence, and assumes particles are spherical and of a homogenous density

59 (Mukherjee et al., 2017). Gas phase sensors produce a signal through the reaction of the target gases

60 with electrochemical or metal oxide sensors. However, the reactive agents used in these types of

61 sensors may degrade over time, and measurement artifacts may also exist, such as cross-interferences

and impacts of temperature (Rai et al., 2017). Therefore, it is necessary to evaluate sensor performance

63 in long-term, real-world study conditions (Alastair Lewis and Peter Edwards, 2016; Williams et al., 2014).

64 The evaluation of low-cost air pollution sensors and their performance is continually evolving

65 (McKercher et al., 2017b). Many sensors are evaluated in laboratory settings by exposure to known

66 concentrations of gasses and PM, with PM often being evaluated by well-defined aerosol, such as

67 polystyrene latex, in controlled conditions (Wang et al., 2015; Lewis et al., 2016; Manikonda et al.,

68 2016). In outdoor, field settings, sensors are often evaluated to determine their performance in

69 comparison with reference methods (Borrego et al. 2016; Jiao et al., 2017; Crilley et al., 2017;

70 Mukherjee et al., 2017; Hagan et al; 2018). Correlations of low-cost sensors have been found to vary

71 from study to study, spanning from negligible to high correlations. Recent studies have shown the

72 correlation between sensors and reference measurements can be improved by the application of

73 correction factors for environmental conditions such as relative humidity (Crilley et al., 2017) or

74 multivariate models and machine learning (Cross et al., 2017; Zimmerman et al, 2018; Hagan et al.,

75 2018).

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76 There are relatively few efforts that exist to systematically examine air pollution sensor technology 77 performance, that test a variety of replicate sensor types against reference monitors in a real-world 78 environment. In the United States, the U.S. EPA and the South Coast Air Quality Management District 79 (SCAQMD) have developed field- and laboratory-testing programs for both gas and particulate matter 80 sensors. These efforts represent specific geographic locations and concentration ranges (U.S. EPA, 2017; 81 SCAQMD, 2017). For example, EPA's Community Air Sensor Network (CAIRSENSE) project tested a 82 variety of gas-phase and particulate matter sensors in Atlanta, GA, under conditions that were high 83 temperature, high humidity, and fairly low ambient concentrations (e.g., hourly PM_{2.5} ranging 0 to 40 84 μg/m³) (Jiao et al., 2016). The SCAQMD AQ-SPEC program similarly conducts field testing of sensor 85 technology in Diamond Bar, California, at a near-road location nominally two months. Evaluation of 86 identical sensors by the EPA and SCAQMD has revealed that the sensor performance may vary by 87 geographical region. For example, Jiao et al., (2016) found Airbeam sensor correlations to be moderate 88 $(r^2 \approx 0.43)$, SCAQMD (2017) reported much stronger correlations $(r^2 \approx 0.74)$. This might be a result of 89 from both different concentration ranges as well as the optical properties of the aerosol being 90 measured.

The Community Air Sensor Network (CAIRSENSE) project was a multi-year, multi-location project that focused on evaluating performance characteristics and limitations of low-costs sensors. A prior CAIRSENSE study in Atlanta, Georgia was conducted in 2014 and early 2015 (Jiao, et al., 2016). Atlanta was chosen to test the sensors' performance in the face of higher temperatures and humidity. For the second part of the CAIRSENSE study, Denver Colorado was chosen to test the sensors' performance under conditions of high altitude, dry and lower temperature conditions. Beyond assessing sensor performance through correlation with a reference monitor, this study also investigates the degree to which data from sensors is able to produce similar temporal, wind-direction, and transient event trends in comparison to a high time-resolution reference monitors.

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2. METHODS

102 Sensors for this study were selected based on cost, commercial availability, market prevalence, 103 capability, and applicability to EPA research objectives. Table 1 lists the sensors chosen for this study, 104 pollutants measured by each sensor, and the measurement principle used by each sensor. Cost 105 information for these sensors are published on the EPA's Air Sensor Toolbox (U.S. EPA 2017). Two 106 different Dylos units were used for this study. Unit 1 was a Dylos DC1100, while units 2 and 3 were Dylos 107 DC1100 Pro models, where the Pro models are advertised to have increased sensitivity for smaller 108 particles. The Shinyei, Dylos, Airbeam, Aeroqual, and Cairclip sensors were used in both the Denver and 109 Atlanta studies (Jiao et al., 2016). Additionally, several of these sensors have been evaluated in 110 laboratory or short term ambient settings (Air Sensor Toolbox reference; Sousan et al., 2016; SCAQMD 111 2017; etc.). 112 Air pollution sensors were acquired and deployed in triplicate. Before deployment, laboratory sensor 113 response audits were performed for all of the available sensors. PM sensors were zero-checked in a 114 clean room environment, all reporting <2 µg/m³ values under those conditions, except for the Air

Assure. The software for the Air Assure performs its own zeroing, therefore they were operated 'as-is'. A

116 pre-deployment sensor response audit was not performed for the TZOA as it was received shortly

before deployment. Sensor output was not adjusted based on the calibration audits in order to reflect

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Table 1 Sensors used during the CAIRSENSE-Denver Study

Pollutant(s) Measured	Principle of Operation		
O ₃	Electrochemical Sensor		
PM	Light Scattering		
PM	Light Scattering		
NO ₂ + O ₃	Electrochemical Sensor		
PM	Laser particle counter		
PM	Laser particle counter		
PM	Light Scattering		
PM	Light Scattering		
PM	Laser particle counter		
	O ₃ PM PM NO ₂ + O ₃ PM PM PM PM		

their 'out of the box' performance. Sensor responses were also audited by either challenge concentrations (Aeroqual and CairClip sensors), or in a clean air environment (PM sensors) after the end of the measurement period, to evaluate possible sensor drift. Laboratory audit results are presented in the supplemental information.

Sensors were deployed at the downtown Denver Continuous Ambient Monitoring Program (CAMP) regulatory monitoring site (Latitude: 39.751184; Longitude: -104.987625) from September 2015 to March 2016. The CAMP site was operated by the state of Colorado for the duration of the study. Sensors were placed in a ventilated, multi-level shelter designed to allow ambient air circulation and prevent intrusion from precipitation, as shown in Fig. 1. A full description of the shelter has been previously reported (Jiao, 2016). The sensors were connected to data loggers stored in weatherproof enclosures attached to the bottom of the shelter. Most of the sensors were connected to Arduino (single-board) microprocessors with either Ethernet (IEEE 802.3 standard) or Recommended Standard 232 (RS-232) serial communication cables. The OPC-N2 and Speck sensor data were logged using laptops, and the TZOA data was stored internally on secure digital (SD) cards. To comply with EPA data security requirements, the cloud based storage capability of the Air Assure sensors was disabled, and these units reported data locally via the Arduino microprocessors with onboard memory. The Cairclip sensor measures the combined signal from NO₂ and O₃. Therefore, both NO₂ and O₃ measurements from the Cairclip were determined by subtracting the opposite (collocated) reference measurement. The Dylos units also measure multiple particle size fractions. In this study, the "small" particle size fraction, as described by the manufacturer, was used for PM_{2.5} comparisons. TZOA sensors did not have a real-time

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Figure 1 Sensor deployment shelter

clock, and only measured time as the elapsed number of milliseconds since the device was powered on. Therefore, field operators were required to accurately record start and end times as a means of establishing the sensor response time series.

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A total of four Arduino microprocessors and three laptops were used simultaneously for data logging. Between the data loggers, laptops and onboard data storage, there were many different sensor data output formats. Separate data scripts were developed to process each different data format into similarly formatted files for each air pollution sensor type. Once data collections were initiated in September 2015, the sensors were operated with little or no intervention through the entirety of the study. Noted interventions included restarting data systems when they 'locked up', or removing windblown snow from the shelves housing the sensors during a major winter snowstorm.

Federal Equivalent Method (FEM) measurements at the Denver monitoring site were collected using a Tolodyna 4005 Or monitor. Tolodyna 2005 I. NO. analyzer and a GRIMM FDM 180 Dust monitor which

Federal Equivalent Method (FEM) measurements at the Denver monitoring site were collected using a
Teledyne 400E O₃ monitor, Teledyne 200EU NO₂ analyzer and a GRIMM EDM 180 Dust monitor, which
measured PM_{2.5} and PM₁₀ mass at one-minute intervals using optical detection. All sensors and
monitors collected pollutant data at one minute intervals or less. One minute values were used to
generate concentrations at multiple time intervals, with primarily one hour averages used for data

generate concentrations at multiple time intervals, with primarily one hour averages used for data analysis. All averaging and other data processing was performed using the following software: RStudio

version 0.98.1103, R version 3.2.2, and the ggplot2, scales, plyr, lattice, corrplot, and 'data.table'

158 (extension of 'data.frame') packages.

159 Sensor data were recovered from the connected laptops and SD cards connected to the data loggers.

160 Most sensors reported data in one-minute intervals. The Alphasense OPC-N2 units recorded

161 concentrations every ten seconds. These measurements were used to calculate one-minute averages.

162 The TZOA sensors reported data based on time elapsed from turning on each unit. The start times for

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each unit and total elapsed time for each measurement were combined to generate 5-second time stamps for the TZOA measurements. These values were then used to calculate one-minute averages.

In order to best replicate actual use by non-experts and avoid biasing the results towards a positive direction, minimal screening of data was performed. Quality assurance screening consisted primarily of removing data where there was a clear malfunction of the sensor, such as non-numeric data output, or when a sensor (e.g., Cairclip unit 1) became 'stuck', reporting a repeated, value (value = 255) for long time spans. These types of errors had previously been identified for the output of this sensor type. The Aeroqual units had significant numbers of measurements that, for some reason, were reported as zero.

171 These were possibly due to the inability of the sensor to detect trace concentrations, and were

therefore not screened out of the data.

Timestamps for all sensors except the TZOA were recorded in Mountain Standard Time. As previously mentioned, TZOA timestamps were generated by combining the initial recording time and the elapsed time reported by the sensors. One-minute measurements and averages were used to calculate 5-minute and hourly averages. Hourly averages were further used to calculate 12-hour and daily averages. FEM measurements from the State of Colorado instruments were also recorded at one-minute intervals and averaged in the same manner as the sensor data. Data from all sensors and reference instruments were stored in separate data files and combined based on timestamps for analyses using 'R' scripts.

Sensors were also investigated for how well they replicated different trends in the regulatory monitor measurement data. The trends analyzed included average sensor responses based on time of day and wind direction. In order to evaluate these trends, different normalized sensor responses were used. The normalized average sensor response for the diel (daily, 24-hour) patterns was calculated as the average concentration for a given hour divided by the average concentration for the hour beginning at 12:00 PM. The normalized average sensor response for wind direction data was defined as the mean concentration for each 10-degree wind 'bin', divided by the average concentration of the 170 to 180-degree bin. The sensor response times were also analyzed by calculating the average one-minute relative sensor response, as defined by the distribution of the one-minute concentration differences divided by the average sensor response.

3. RESULTS & DISCUSSION

Table 2 shows a summary of data completeness from the air pollution sensors, including the total percentage of minutes measured, percentage of measurements missed, by not logging data, and the percentage of completely missing data. The majority of missing data was due to events where the sensor and data loggers were inoperative. The most significant of these events was due to snow intrusion into the monitoring platform in December 2015, which caused units to shut down. Most sensors had a very high data capture rate throughout the study when the units were on (and operational). The Cairclip units had significant amounts of missing data, likely due to data transmission errors from the universal asynchronous receiver-transmitter (UART) serial communication system. In the previous Atlanta study as well as in a Newark-based citizen science study (Kaufman et al., 2017), Cairclip units with identical sensors but different universal serial bus (USB) data connections were used and did not have significant amounts of missing data.

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Table 2 Sensor Data Completeness

	Measurement	Sensor on and	Completely	
Sensor	%	not Logging %	Missing %	Comments
	82%	0%	18%	45% of logged values were 0
	73%	0%	27%	42% of logged values were 0
Aeroqual	81%	5%	13%	32% of logged values were 0
	87%	0%	13%	
Air Assure	87%	0%	13%	
	87%	0%	13%	
	74%	0%	25%	
Airbeam	62%	6%	32%	
	62%	6%	32%	
	29%	53%	18%	56% of logged values were 255
Cairclip	63%	13%	24%	No data before 10/8/15
	63%	23%	13%	
	82%	0%	18%	
Dylos	82%	0%	18%	
	72%	1%	27%	
	82%	0%	18%	
Shinyei	73%	0%	27%	
	87%	0%	13%	

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Measurements from air pollution sensors and regulatory monitors were time-averaged at multiple intervals for comparison. The time intervals included 5 minute, hourly, 12-hour, and daily averages. For each set of time averaging, regressions were calculated to evaluate sensor correlation and bias when compared to regulatory measurements. Additionally, intercomparisons were made between sensors of the same pollutant type (e.g., correlations between PM sensors). Table 3 displays a summary of regression statistics for sensors when compared to regulatory measurements for 1-hour time averages. In general, correlations were greatest at the 1-hour time average. Correlations in general improved slightly with increasing length of the averaging period up to hourly averages. Reduced correlations for most sensors at the 12-hour and daily averages may be a result of a lower number of data points. In contrast to most other measurements, sensors that reported data for coarse PM (Dylos) or PM₁₀ (OPC-N2) showed improved correlations with increasing averaging time for those measurements. The correlations for all the time averaging periods can be found in the supplemental information. Figure 2 shows a Pearson correlation (R) plot for 1-hour average measurements of the PM sensors. The PM units show high correlation among sensors of the same model, except for when one sensor in a group had significant issues. Of the PM_{2.5} sensors, the Air Assure, Airbeam, and Dylos (R = 0.73 to 0.86) units exhibited the highest correlation with reference measurements. Dylos unit 1 had the highest linearity, however it had the lowest particle count response, both of which are likely explained by not detecting the smallest particles as effectively as other units. Cairclip unit 1 rarely properly transmitted data throughout the study, leading to its low correlations. Cairclip units 2 and 3 had more sporadic data transmission issues. All Cairclip units recovered data properly once returned to the lab after the field campaign where their internal data storage was used. The response from Shinyei unit 3 changed in mid-

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Table 3 Regression Results for CAIRSENSE sensors (1-hour time averaged)

Sensor	Pollutant	Reference Average Concentration ¹	Slope	Intercept	Pearson Correlation, R	Number of Hourly Measurements
3611301	Tonutune	Concentration	0.56	-0.004	0.93	3325
Aeroqual SM- 50	O₃, ppb	18.8 ppb	0.58	-0.004	0.92	2963
	C ₃ , pp5	20.0 pps	0.38	-0.004	0.96	3279
			1.14	2.64	0.8	3486
TSI Air Assure	PM, μg/m³	7.8 μg/m³	1.13	-0.04	0.78	3486
			1.19	-1.38	0.78	3486
			1.19	-1.56	0.81	
AirCasting AirBeam	Particle Count, hundreds of particles per cubic foot (hppcf)	7.8 μg/m³	273	-323	0.82	3028
			278	-124	0.84	2539
			322	-352	0.82	2532
Cairpol		18.8 ppb	NA ²	NA ²	NA ²	738
Cairclip	O ₃ , ppb		-0.04	-23.6	-0.06	2831
			1.03	-39.0	0.46	2852
Cairpol		26.8 ppb	NA^2	NA ²	NA ²	738
Cairclip	NO ₂ , ppb		0.65	-10	0.87	2831
			0.67	-15	0.84	2852
Dylos DC1100/DC11 00 Pro	"Small" Particle Count, hppcf	7.8 μg/m³	64	-152	0.86	3324
			428	-1182	0.78	3324
			431	-941	0.73	2937
Dylos DC1100/DC11 00 Pro	"Large" Particle Count, hppcf	12.0 μg/m³	1.3	5.5	0.40	3324
			5.7	73	0.33	3324
			4.9	84	0.27	2937
	PM _{2.5} , μg/m ³	7.8 μg/m³	0.4	-0.30	0.45	2969
Alphasense OPC-N2			0.49	-1.66	0.34	2939
			0.07	0.60	0.11	2735
Alphasense OPC-N2	PM ₁₀ , μg/m³	19.6 μg/m³	0.45	2.98	0.47	2969
			0.54	-1.06	0.68	2939
			0.12	2.86	0.20	2735
Shinyei PMS- SYS-1	PM _{2.5} , μg/m ³	7.8 µg/m³	0.58	0.24	0.71	3325
			0.54	0.8	0.72	2963
			0.42	4.35	0.01ª	3486
AirViz Speck	PM _{2.5} , μg/m ³	7.8 μg/m³	0.76	13	0.24	3557
			0.74	15	0.40	3584
			0.62	10	0.35	3971
TZOA PM Research Sensor	Particle Count, hppcf	7.8 μg/m³	NA ²	NA ²	NA ²	2341
			6.68	1.37	0.66	1838
			6.75	2.16	0.72	1836

¹Average Concentration calculated for hours with valid sampling data.

²Correlation results not shown due to large amount of missing or invalid data

^a Shinyei Unit 3's correlation improved to 0.84 when only considering data from October 16 and later

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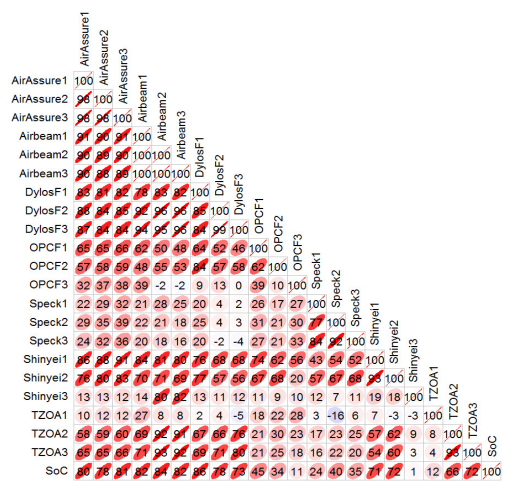


Figure 2 Correlation (r*100) plot for sensors measuring fine PM. Ellipses represent the overall scatter of the data (1-hour averaged measurements)

October. The correlation between the unit and the reference monitor was initially 0.01, then increased to 0.84 when comparing only the data starting October 16 and later.

Several sensor models were used in both the Atlanta and Denver CAIRSENSE evaluation campaigns. Both studies deployed the Airbeam, Dylos, and Shinyei PM sensors. In all cases except for Shinyei unit 3, these sensors showed greater linearity in Denver than in Atlanta, when comparing 12-hour averages. When only considering data after October 16, Shinyei unit 3 also had higher correlation in Denver than in Atlanta. This may be due to less noise caused by lower humidity in Denver than in Atlanta. Aeroqual and Cairclip air pollution sensors were also deployed in both Atlanta and Denver. O_3 measured by the Aeroqual units showed similar correlations in both locations ($R^2 = 0.82$ to 0.94 in Atlanta, $R^2 = 0.85$ to 0.92 in Denver). O_3 measured by Cairclip units 2 and 3 in Denver showed poorer correlations than the Cairclip units used in Atlanta ($R^2 = 0.00$ to 0.21 in Denver versus $R^2 = 0.68$ to 0.88 in Atlanta). However,

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 NO_2 measured by Cairclip units 2 and 3 in Denver was more highly correlated than in Atlanta ($R^2 = 0.71$ to 0.76 in Denver versus 0.57 in Atlanta).

While Denver is not necessarily known for high humidity, humidity artifacts were observed in some sensors. Figure 3a shows the $PM_{2.5}$ concentrations measured by one of the OPC-N2 against relative humidity. At RH around 90%, the PM concentration spikes significantly, suggesting that humidity is interfering with the sensor response measurement. This behavior is similar to that observed by Sousan et al., (2016). Some other instruments also had different responses based on humidity. Figure 3b shows hourly particle counts measured by an Airbeam sensor against $PM_{2.5}$ concentration measured by the reference instrument, stratified by relative humidity. There appear to be two separate relations between reference measured concentrations and sensor measured particle counts, with a greater particle count response occurring more at higher humidity. This relationship was observed in each of the Airbeam sensors. An example of humidity relationships from each sensor type can be found in the supplemental information.

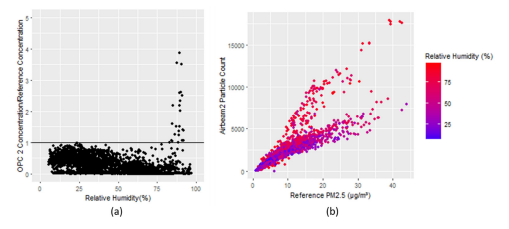


Figure 3 OPC 2 PM_{2.5} and Relative Humidity (a) and Hourly Average FRM PM_{2.5} concentration and Airbeam Particle count stratified by Relative Humidity (b)

In addition to understanding the precision of air pollution sensors and how well they correlate with reference measurements, it is also important to understand how well a sensor can capture trends and distributions of pollutant concentrations. There are many ways to examine these trends and distributions. Figure 4 shows the diel patterns of $PM_{2.5}$ (a) and O_3 (b) reference and sensor measurements respectively. The results, for each sensor, represent the measurements of the best performing unit for each sensor type/model, as determined by R-squared values. The various PM air pollution sensors have a wide range of comparisons to the reference monitor. Two sensors (TZOA and Airbeam) show similar patterns throughout the day, while some other sensors do not reflect the reference diel pattern at all (e.g., OPC, Speck, etc.). The Aeroqual sensor diel pattern was similar to that of the reference O_3 monitor. The nature of the calculation of O_3 and NO_2 by subtraction, and missing data from the Cairclip sensors, prevented this analysis from providing meaningful results.

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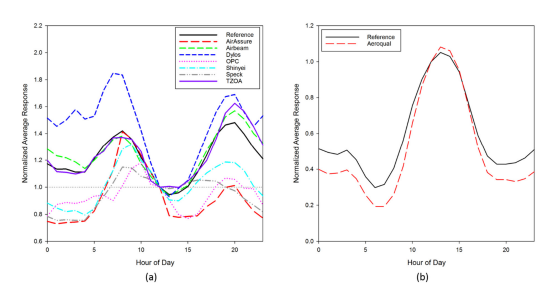


Figure 4 Diel patterns for PM_{2.5} (a) O₃ (b) sensor and reference measurements.

Air quality measurements are also known to be dependent on wind direction, and it is important to know if these differences were reflected in the sensor measurements. Figure 5 shows the normalized average sensor response $PM_{2.5}$ (a) and O_3 (b) response of the sensors and the reference monitors respectively. The reference monitor response is represented by the black line. The sensors generally compared more favorably with the reference monitors when examining the wind direction dependence of concentration. This is most apparent in the OPC-N2 sensor, where the sensor trends track the trends measured by the reference monitor. This increases the confidence that sensors may be useful in studies that pair wind direction with concentration to determine potential bearings or locations of pollution sources to supplement source apportionment and receptor modeling. It also raises questions as to why an air pollution sensor would be able to reproduce wind direction trends but not necessarily reproduce daily concentration measurement patterns. We undertook exploration of this perplexing result, but were not able to determine a clearly identifiable cause.

The high-time resolution data collected for this study allowed for the examination of air pollution sensor response trends compared to that of regulatory air pollution monitors. Figure 6 shows a cumulative distribution function (CDF) for the

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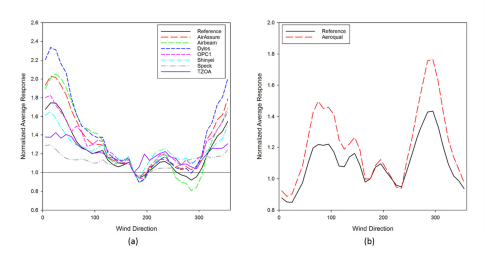


Figure 5 Wind direction patterns for PM_{2.5} (a) O₃ (b) sensor and reference measurements.

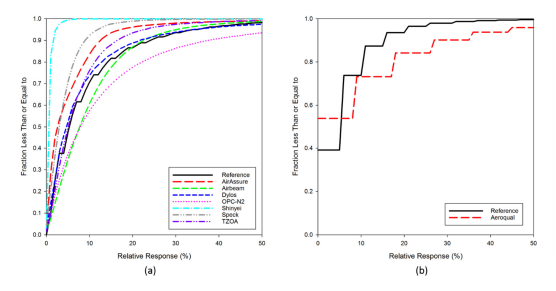


Figure 6 Cumulative distribution functions for 1-minute response differences for $PM_{2.5}$ (a) O_3 (b) sensor and reference measurements.

relative change in sensor and regulatory monitor response between 1-minute measurements for $PM_{2.5}$ (a) and O_3 (b) sensor and reference measurements respectively. The relative response was calculated as the absolute value of the difference between consecutive one-minute measurements divided by the mean measurement for each sensor/monitor. If the reference monitor were considered a perfect measurement, sensor curves to the left and above the reference monitor line would be slower to respond to changes in concentration, while curves below and to the right of the monitor line would signify high levels of measurement noise. Most PM monitors exhibited a slower response to changes in

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concentration than the reference monitor. The OPC-N2 and Airbeam sensors were the only ones with curves to the right of the reference monitor, suggesting that they may have more noise in their measurements. The Aeroqual sensor showed more O₃ measurement noise when compared to the reference measurement.

4. Conclusions

Nine different air pollution sensor devices were deployed in triplicate with collocated air pollution reference monitors in Denver, CO over an extended operational timeline of longer than six months. The sensors showed a wide range of correlations with reference measurements, but tended to have high correlation with sensors of the same model. PM sensors deployed in both Denver and Atlanta had higher correlations with reference monitors in Denver than in Atlanta. This is likely due to less humidity related response in Denver. Aeroqual O₃ measurements in Denver showed similar linearity to those measured in Atlanta. Cairclip O₃ correlations were lower in Denver than in Atlanta, but NO₂ correlations were higher. Sensors that have also been evaluated by the South Coast Air Quality Management District (SCAQMD) tended to show similar results in terms of correlation (SCAQMD, 2017). However, in all cases, sensors' performance in this long-term field deployment was less than that of laboratory based comparisons performed in this study and others (U.S. EPA, 2017). This study demonstrates the need for long-term, real-world evaluation studies for current and future air pollution sensors, that will need to be performed in locations with different air pollutant concentration ranges and aerosol characteristics.

Several air pollution sensors were able to capture variations in important trends, such as diel patterns and wind direction dependence on concentration. However, the OPC-N2 units showed similar results as reference monitor measurement data when analyzing the wind direction trends, but not when analyzing 'time-of-day' trends. These promising results show that sensors have the possibility for supplementing measurement research capabilities when interested in air pollution trends such as those dependent on wind direction. Analyses of wind direction based air pollutant trends could be useful for possible identification of source locations or regions, especially with the use of a sensor-based network.

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326	Data Availability
327 328 329	The CAIRSENSE dataset will be available at the EPA environmental dataset gateway (https://edg.epa.gov) (EPA 2018) where the dataset can be retrieved by searching for "CAIRSENSE Denver." Project data can also be requested from the corresponding author.
330	
331	Author Contribution
332 333 334 335 336 337 338	Stephen Feinberg performed the data analysis and prepared the manuscript with contribution from all co-authors. Ron Williams was the Principal Investigator and advised data analysis and manuscript preparation. Gayle Hagler was Co-Investigator and advised data analysis and manuscript preparation. Joshua Rickard, Ryan Brown, Daniel Garver, and Robert Judge provided discussion and assistance in the establishment of the study design and execution. Greg Harshfield, Phillip Stauffer, and Erick Mattson carried out day-to-day operations. Sam Garvey was in charge of the establishment of the initial site, data recovery, and principal operation of the sensors.
339	
340	Disclaimer
341 342 343 344 345	This article was supported by Jacobs Technology, Inc. (contract EP-C-15-008) for the National Exposure Research Laboratory, U.S. Environmental Protection Agency (U.S. EPA), Research Triangle Park, NC. It has been reviewed by the U.S. EPA and approved for publication. Mention of trade names or commercial products does not constitute endorsement or recommendation for use.

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