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Executive Summary

The purpose of the Arizona Department of Environmental Quality (ADEQ) Annual Ambient Air Assessment Report is to serve as an informational and technical document for use within ADEQ, other state and local agencies, and health organizations such as the American Lung Association. This report summarizes the results of air quality monitoring conducted by ADEQ in 2017. Monitor locations, purpose of monitoring, and monitoring methods are discussed. Also included are area/monitor compliance values based on air monitoring conducted by ADEQ in 2017, an overview of long-term monitoring concentrations, and Air Quality Index (AQI) values.

Data were collected from 36 ADEQ air monitoring sites located throughout Arizona in 2017. Many of the sites have multiple instruments measuring a variety of gaseous, particulate, meteorological, and visibility parameters.

Criteria Pollutants

The majority of the ADEQ's air quality measurements are for criteria pollutants (carbon monoxide (CO), nitrogen dioxide (NO₂), ozone (O₃), sulfur dioxide (SO₂), particulate matter (PM₁₀ and PM_{2.5}), and lead (Pb)) classified as State and Local Air Monitoring Stations (SLAMS) used for regulatory compliance. To show regulatory compliance, the U.S. Environmental Protection Agency (EPA) has established National Ambient Air Quality Standards (NAAQS). These standards are the concentrations at which each pollutant becomes harmful to human health and are unique to each pollutant.

Compliance statuses are presented in this document for informational purposes only and are considered preliminary. For each of the criteria pollutants the preliminary compliance status and a summary of the 2017 AQI values are provided below:

CO

- In compliance with the NAAQS.
- AQI values for 2017:

Site	"Good" Days	Missing Days
Alamo Lake	300	65
JLG Supersite	283	82

• NO₂

- o In compliance with the NAAQS.
- o AQI values for 2017:

Site	"Good" Days	"Moderate" Days	Missing Days
JLG Supersite	305	5	55

O₃

o Four sites in violation of the NAAQS: JLG Supersite, Queen Valley, Tonto National Monument, and Yuma Supersite.

o AQI values for 2017:

Site	"Good" Days	"Moderate " Days	"Unhealthy for Sensitive Groups" Days	"Unhealthy" Days	Missing Days
Alamo Lake	275	77	2	0	11
Flagstaff Middle School	284	78	1	0	2
JLG Supersite	237	109	13	1	5
Prescott Pioneer Park	219	103	5	0	38
Queen Valley	233	109	12	1	10
Tonto National Monument	261	92	8	1	3
Yuma Supersite	293	48	10	0	14

• SO₂

o Four sites in violation of the NAAQS: Hayden Old Jail, Miami Jones Ranch, Miami Ridgeline, and Miami Townsite.

o AQI values for 2017:

Site	"Good" Days	"Moderate" Days	"Unhealthy for Sensitive Groups" Days	"Unhealthy" Days	Missing Days
Hayden Old Jail	133	122	95	15	0
JLG Supersite	306	0	0	0	59
Miami Jones Ranch	262	47	31	5	20
Miami Ridgeline*	188	27	16	0	37
Miami Townsite	295	34	18	1	17

^{*268} possible monitoring days.

• PM₁₀

Three sites in violation of the NAAQS (preliminary, based on incomplete data): Douglas Red Cross,
 Rillito, and Yuma Supersite.

o AQI values for 2017:

Site	"Good" Days	"Moderate" Days	"Unhealthy for Sensitive Groups" Days	"Unhealthy" Days	"Very Unhealthy" Days	Missing Days
Ajo	356	9	0	0	0	0
Alamo Lake	347	5	0	0	0	13
Bullhead City	344	6	0	0	0	15
Douglas Red Cross	329	27	2	0	0	7
Hayden Old Jail	341	23	0	0	0	1
JLG Supersite	323	31	0	0	0	11
Miami Golf Course	358	5	0	0	0	2
Nogales Post Office	301	63	1	0	0	0
Paul Spur Chemical Lime Plant	356	3	0	0	0	6
Payson Well Site	358	4	0	0	0	3
Rillito	239	124	2	0	0	0
Yuma Supersite	296	65	2	1	1	0

PM_{2.5}

- o In compliance with the NAAQS.
- o AQI values for 2017:

Site	"Good" Days	"Moderate" Days	"Unhealthy for Sensitive Groups" Days	"Unhealthy" Days	Missing Days
Alamo Lake	345	3	0	0	17
Douglas Red Cross	263	10	0	0	92
JLG Supersite	307	44	1	0	13
Nogales Post Office	276	60	2	2	25
Yuma Supersite	315	48	2	0	0

Pb

- Two sites in violation of the NAAQS: Globe Highway and Hillcrest.
- An AQI is not established for Pb.

Non-Criteria Pollutants

In addition to sampling for criteria pollutants, ADEQ conducts manual sampling of O₃ forming compounds, speciated particulate matter, and hazardous air pollutant sampling under the Photochemical Assessment Monitoring Stations (PAMS), Chemical Speciation Network (CSN), National Air Toxics Trends Stations (NATTS), and National Core (NCore) Monitoring Network. Brief summaries from the CSN, NATTS, and PAMS networks are provided below:

CSN

- The 2017 annual average percent compositions for speciated PM_{2.5} major elements at JLG Supersite included the following: 31% Organic Carbon, 26% Other, 18% Crustal Component, 9% Elemental Carbon, 9% Sulfate, 6% Nitrate, and 1% Ammonium.
- 23% increasing trend to 57% decreasing trend in these major elements (excluding 'Other') from 2000-2017.

NATTS

- Air toxics concentrations at JLG Supersite and South Phoenix continue to show many decreasing trends, ranging from a 9% increasing trend to a 106% decreasing trend.
- Manganese may be of local concern as the 2017 average exceeded the 2014 national average by a factor of 2.4.

PAMS

 Samples for Total Non-Methane Organic Compound (TNMOC) concentrations were not collected in 2017.

Visibility

Further monitoring includes special continuous monitoring for the optical/visual characteristics of the atmosphere under the urban visibility network required by House Bill 2538 of the Forty-fifth AZ State Legislature. A description of the trends observed at the Phoenix visibility instrument is provided below:

• Phoenix Transmissometer

- o 48% increase in the mean visual range for all hours from 1995-2017.
- o 40% increase in the mean visual range in the morning hours from 1995-2017.
- o Between 16% and 40% increase in the mean visual range for winter, spring, and fall from 2006-2017; 22% decrease in the summer months from 2013-2017, and 40% increase from 2006-2011.

Additionally, ADEQ serves as an operator for the Interagency Monitoring of Protected Visual Environments (IMPROVE) which tracks visual conditions in and around national parks and monuments, as well as in some urban areas.

Conclusion

Overall, monitoring data collected by ADEQ in 2017 showed NAAQS compliance with CO, NO_2 , and $PM_{2.5}$. SO_2 , O_3 , PM_{10} , and Pb were found to be in violation of the NAAQS. PM_{10} and O_3 have been long-standing pollutants of concern in Arizona and SO_2 and Pb continue to be monitored at local areas near copper smelting operations. Despite these violations, the total average of the AQI values for days with valid monitoring data operated by ADEQ indicated that 85% of the days in 2017 were "Good" days. A summary of the percentage of "Good" days for each pollutant is listed below:

- 100% for the CO monitors;
- 98% for the NO₂ monitor;
- 73% for the O₃ monitors;
- 76% for the SO₂ monitors (100% for JLG Supersite, 81% for Miami area, 36% Hayden Old Jail);
- 91% for the PM₁₀ monitors; and
- 90% of the PM_{2.5} monitors.

Similar to the criteria pollutants, many decreasing trends were also observed for speciated particles and air toxics. With reductions in air pollutant concentrations over the years, visibility measurements made by ADEQ showed great improvements with increased visual range.

Section I – Introduction to ADEQ's Ambient Air Monitoring Program

1.0 Purpose and Background

The ADEQ Annual Ambient Air Assessment Report summarizes the air monitoring activities and the results of air quality monitoring conducted by the Arizona Department of Environmental Quality (ADEQ) for the calendar year 2017. This report shows monitor locations, purpose of monitoring, and monitoring methods. Also included is an overview of long-term monitoring concentrations, Air Quality Index (AQI) values, and area/monitor compliance values based on air monitoring conducted by ADEQ in 2017. The purpose of this report is to serve as an informational and technical document intended for use within ADEQ, other state and local agencies, and health organizations such as the American Lung Association.

Data from 36 sites located throughout Arizona are included in this report. Many of the sites have multiple instruments measuring a variety of gaseous, particulate, meteorological, and visibility parameters. The majority of the air quality measurements are for criteria pollutants (carbon monoxide (CO), nitrogen dioxide (NO₂), ozone (O₃), sulfur dioxide (SO₂), particulate matter (PM₁₀ and PM_{2.5}), and lead (Pb)). The monitors used to measure these pollutants are classified as State and Local Air Monitoring Stations (SLAMS) used for regulatory compliance. Per requirements in the Clean Air Act, the U.S. Environmental Protection Agency (EPA) established the National Ambient Air Quality Standards (NAAQS) (40 CFR Part 50). These standards are the concentrations at which each criteria pollutant becomes harmful to human health and the environment, and are unique to each pollutant.

In addition to sampling for criteria pollutants, ADEQ performs special continuous monitoring for the optical/visual characteristics of the atmosphere under the urban visibility network required by House Bill 2538 of the Forty-fifth AZ State Legislature. Further monitoring includes manual sampling of O₃ forming compounds and other hazardous air pollutants, and speciated particulate matter sampling under the Photochemical Assessment Monitoring Stations (PAMS), Chemical Speciation Network (CSN), National Air Toxics Trends Stations (NATTS), and National Core (NCore) Monitoring Network. ADEQ also oversees industrial networks to determine the effects of their emissions on local air quality and how well pollution control technologies are working. Additionally, ADEQ serves as an operator for the Interagency Monitoring of Protected Visual Environments (IMPROVE) network which tracks visual conditions in and around national parks and monuments, as well as in some urban areas. Lastly, ADEQ operates a network of portable particulate monitors for public awareness to provide information regarding pollutant levels from wildfires and wood burning.

Air monitoring is commonly classified by networks based on individual pollutants or by a group of related pollutants. Monitoring networks for ambient air quality are established to sample pollution in a variety of settings representative of different populations and geographical areas, to assess health and welfare effects, and to assist in determining air pollution sources. The ambient monitoring networks cover both

urban and rural areas of the state and are composed of one or more monitoring sites whose data are compared to the NAAQS for compliance and statistically analyzed for trends analysis. ADEQ also tracks data recovery, quality control, and quality assurance parameters for the instruments operated at its various sites. Most of the sites within each network also measure meteorological variables. ADEQ networks monitor a wide variety of pollutant and atmospheric characteristics including urban, industrial, rural, transport, and background surveillance.

2.0 Standards and Guidelines

The Federal Clean Air Act (CAA) of 1970 requires the EPA to assist states and local agencies in establishing ambient air quality monitoring networks to characterize human health exposure and public welfare effects from criteria pollutants. The way public welfare is measured is by analyzing ambient air conditions using a variety of instruments specifically designed to measure a certain pollutant. These instruments show pollution concentrations for a given time period, and then the data is analyzed to identify certain concentrations which can affect human health. Because different concentrations of pollutants affect human health at different levels, an Air Quality Index (AQI) is used for showing when a specific concentration can be bad for human health.

For data completeness, EPA requires 75 percent valid data recovery over a set time period for values to be considered valid. Depending on the pollutant, this can apply to the scales of hourly, daily, quarterly, and yearly, for which each averaging period must be 75 percent complete. For regulatory purposes, if the data completeness is greater than 75 percent for the specified time period, then completeness criteria are met and data may be used for area designations.

2.1 Air Quality Index

The AQI is an index for reporting daily air quality to the general population. It indicates how clean or polluted the air is, and what associated health effects might be of concern for that day. The AQI focuses on health effects that may be experienced within a few hours or days after breathing polluted air. The EPA calculates the AQI for the criteria air pollutants regulated by the CAA: ground-level O₃, PM₁₀, PM_{2.5}, CO, SO₂, Pb, and NO₂. For each of these pollutants, EPA has established national air quality standards to protect public health. When the AQI reaches 100, this indicates that a concentration has exceeded the standard set forth by the EPA. Generally, ground-level O₃ and airborne PM are the two pollutant types that pose the greatest threat to human health in this country. EPA's AQI website AIRNow is found at www.airnow.gov.



Figure 1 - Chart of AQI Levels

Each category in Figure 1 corresponds to a different level of health concern. The six levels of health concern are:

- "Good" AQI is 0 50. Air quality is considered satisfactory, and air pollution poses little or no risk to human health.
- "Moderate" AQI is 51 100. Air quality is acceptable; however, for some pollutants there may be a moderate health concern for a very small number of people. For example, people who are unusually sensitive to O₃ may experience respiratory symptoms.
- "Unhealthy for Sensitive Groups" AQI is 101 150. Although general public is not likely to be affected at this AQI range, people with lung disease, older adults, and children are at a greater risk from exposure to O₃, whereas persons with heart and lung disease, older adults, and children are at greater risk from the presence of particles in the air.
- "Unhealthy" AQI is 151 200. Everyone may begin to experience some adverse health effects, and members of the sensitive groups may experience more serious effects.
- "Very Unhealthy" AQI is 201 300. This would trigger a health alert signifying that everyone may experience more serious health effects.
- "Hazardous" AQI is greater than 300. This would trigger a health warning of emergency conditions. The entire population is more likely to be affected.

2.2 National Ambient Air Quality Standards

The NAAQS were first established by the EPA per the CAA in 1970 and are continuously evaluated and updated based on current scientific research on the effects of pollution exposure to the population. Focus is placed on those groups who are sensitive to air pollution. Table 1 shows the current NAAQS.

Table 1 – Current NAAQS (Source: USEPA TTN NAAQS)

Pollu	tant	Primary/ Secondary	Averaging Time	Level	Form
Carbon M	onoxide	nrimanı	8-hour	9 ppm	Not to be exceeded more
(CO)		primary	1-hour	35 ppm	than once per year
Lead (Pb)		primary and secondary	Rolling 3 month average	0.15 μg/m ³	Not to be exceeded
Nitrogen Dioxide		primary	1-hour	100 ppb	98 th percentile, averaged over 3 years
(NO ₂)		primary and secondary	Annual	53 ppb	Annual Mean
Ozone (O ₃	Ozone (O ₃) primary and secondary 8-hour		0.070 ppm	Annual fourth-highest daily maximum 8-hr concentration, averaged over 3 years	
		primary	Annual	12 μg/m³	annual mean, averaged over 3 years
	PM _{2.5}	secondary	Annual	15 μg/m³	annual mean, averaged over 3 years
Particle Pollution		primary and secondary	24-hour	35 μg/m ³	98 th percentile, averaged over 3 years
	PM ₁₀	primary and secondary	24-hour	150 μg/m³	Not to be exceeded more than once per year on average over 3 years
Sulfur Dio	xide	primary	1-hour	75 ppb	99 th percentile of 1-hour daily maximum concentrations, averaged over 3 years
		secondary	3-hour	0.5 ppm	Not to be exceeded more than once per year

Ambient concentrations to be compared to the NAAQS are defined under the "Form" column in Table 1 and are calculated based on the averaging time stated. For many of the pollutants, there is a primary standard and a secondary standard. Primary standards provide public health protection, including protecting the health of "sensitive" populations such as asthmatics, children, and the elderly. Secondary standards provide public welfare protection, including protection against decreased visibility and damage to animals, crops, vegetation, and buildings.

An area may be designated as nonattainment, attainment, or unclassifiable based on exceedances of the standards. Area designations occur after a revision to the NAAQS occurs, or at other times the EPA deems appropriate.

- A nonattainment area is one in which either the primary or secondary standard has been violated, and both the local agency and the EPA have acknowledged the area as being in nonattainment. The EPA can designate that area as nonattainment with certain sanctions or penalties being placed in order to bring that area into attainment. When EPA designates an area as nonattainment, a State Implementation Plan (SIP) is put in place. A SIP outlines the actions the state will take to improve air quality in the area. This can include instituting lower maximum pollutant allowances for industrial processes, paving of roads, replacing aging equipment, and other control strategies. If the controls outlined in the SIP do not achieve the standard or are inadequate, the EPA has the option to develop a Federal Implementation Plan (FIP) for the area.
- An attainment area is one which is achieving the standards and no additional measures are necessary.
- An unclassifiable area is an area for which data are incomplete and do not support a designation
 of attainment or nonattainment. Data may be incomplete due to lack of monitoring or
 completeness criteria for the year.

It is important to understand the difference between an exceedance and a violation of a standard in order to define area designation. An exceedance occurs when the pollutant concentration reaches a level where the AQI is at or above 100 for a given time period. This is above the acceptable level that is defined in the NAAQS, but does not necessarily indicate that the NAAQS have been violated.

For example, the 24-hour PM_{2.5} NAAQS level of exceedance is 35 micrograms per cubic meter ($\mu g/m^3$). If the concentration on a given day is above this level, an exceedance occurs, but a violation of standard does not necessarily occur. The form by which the PM_{2.5} standard is calculated is the 98th percentile of samples averaged over three years. If the 24-hour average for a day goes above the 35 $\mu g/m^3$ level, but the 98th percentile averaged over three years is not above 35 $\mu g/m^3$, then there is an exceedance for that day, but there is no violation of the standard.

2.3 Monitoring Objectives

There are three basic monitoring objectives as outlined below, each containing multiple aspects and purposes: NAAQS comparison for regulatory compliance, research, and public information. There may be certain monitors which have more than one purpose due to crossover between different networks. For example, a NAAQS comparison monitor may also be used for research purposes in some circumstances.

A. NAAQS Comparison

The majority of ADEQ's monitoring falls under the NAAQS comparison category. This monitoring is governed by the CAA and is explained under the Code of Federal Regulations (CFR) 40 Part 58. This monitoring is required based on rules spelled out in the CFR, including: total emissions in an area, population of an area, attainment/nonattainment areas, population weighted emissions, traffic counts, and pollutant point sources. ADEQ designates these monitors as SLAMS. These are long term monitors that operate for the primary purpose of comparison to the NAAQS and are used for regulatory decision making. They may also support compliance with air quality standards and emissions strategy development, and provide air pollution data to the general public in a timely manner. The SLAMS network includes monitors at stations with network affiliations such as NCore, PAMS, or Speciation, but does not include Special Purpose Monitors (SPM) and other monitors used for non-regulatory or industrial monitoring purposes. Once monitoring occurs, EPA designates areas as attainment or nonattainment based on the comparison of observed concentrations with the NAAQS.

B. Research

ADEQ operates a range of monitors used mainly for research purposes. These instruments may not be regulated by the EPA as part of the NAAQS, but represent various precursor and toxic pollutants. The programs that fall under research include PAMS, NATTS, CSN, IMPROVE, urban air toxics monitoring program (UATMP), and NCore. PAMS, NATTS, CSN, and IMPROVE will be discussed at length in this report (Sections III and IV). These programs are required by the EPA as part of a national network of instruments, but many of the pollutants measured are not specifically regulated. Due to the cross over between NCore and criteria pollutants, a brief description of NCore is provided below. Crossover also exists between UATMP and NATTS and is also described below.

NCore is a multi-pollutant network throughout the whole nation that integrates several advanced measurement systems for particles, pollutant gases, and meteorology. The stations are equipped with several measurement systems to monitor particulate matter (PM_{2.5}, PM₁₀, and PM_{10-2.5}), O₃, CO, SO₂, total reactive oxides of nitrogen (NO_γ), and basic meteorological parameters (temperature, wind speed, wind direction, and relative humidity). The NCore stations should be perceived as developing a representative report card on air quality across the nation, capable of delineating differences among geographic and climatological regions. There are approximately 83 NCore stations nationwide. NCore meets a number of important data needs: improved flow and timely reporting of data to the public, including supporting air quality forecasting and information systems such as AIRNow; continued determination of NAAQS compliance; improved development of emissions control strategies; enhanced accountability for the effectiveness of emission control programs; and more complete information for scientific, public health, and ecosystem assessments.

Emphasis of NCore is placed on high sensitivity instruments with the capability to detect low levels of the precursor gases CO, SO_2 , and NO_y . These gases play important roles in the formation of atmospheric O_3 , air toxics, and PM, which are linked to human health issues. This interconnection among distinct air quality issues requires an integrated multiple pollutant air quality monitoring and management approach which NCore successfully incorporates. ADEQ supports the NCore network by monitoring all required pollutants at its Phoenix JLG Supersite. Concentration data will be shown and analyzed in Section II of this document.

O UATMP is a subset of the Air Toxics program, which includes NATTS, school air toxics monitoring initiative (SATMI), and community scale air toxics ambient monitoring (CSATAM) projects. All programs that are a part of Air Toxics utilize similar monitoring methods and are therefore directly comparable. All have similar monitoring objectives of characterizing toxics in an area of interest. UATMP specifically looks at toxics trends in urban areas. Concentration data will be shown and analyzed in Section III of this document.

C. <u>Public Information</u>

Data generated by ADEQ's SLAMS monitors for regulatory compliance are reported to AIRNow on an hourly basis to show real-time conditions to the public. AIRNow data are used to report the AQI only, not to show regulatory status of an area.

ADEQ also operates networks that do not have EPA approved monitoring techniques, for the purpose of providing public health and visibility information to the general public. Although these networks do not use approved monitoring techniques, they give the public a general understanding of current air quality in their area. These networks do not report data to AIRNow.

For public health, a network of temporary PM_{2.5} monitors without an EPA method code are used to provide air quality conditions to the public in areas that do not require monitoring under the CAA or per the CFR. These areas are mainly located in the northeastern part of Arizona and are often impacted by summer wildfires. These monitors can be easily deployed around a wildfire to describe the impacts that it has on air pollution for the local population. There are also monitors located in areas where wood burning is the main source of heating in the winter. Wildfires and wood burning create small particulates that are harmful to human health. Information regarding these portable particulate monitors can be found at http://phoenixvis.net/PPMmain.aspx.

In the Phoenix metropolitan area, ADEQ operates a variety of instruments used to determine the visibility on a daily basis. Data and pictures for visibility will be shown and analyzed under Section IV of this document.

3.0 Quality Assurance

ADEQ sustains a quality system as required by EPA to ensure high quality data are produced that meet the users' needs. The EPA primarily specifies the quality assurance (QA) requirements for operating SLAMS, SPM, CSN, NCore, NATTS, PAMS, and prevention of significant deterioration (PSD) air monitors in 40 CFR Part 58 Appendix A, the Quality Assurance Handbook for Air Pollution Measurement Systems: Volume II: Ambient Air Quality Monitoring Program, Technical Assistance Documents (TADs), and other supporting guidance documents. In response, ADEQ develops quality assurance plans for air monitoring networks, which provide detailed information regarding the specifics of each air monitoring network and how data will be managed. Components of ADEQ's quality system include, but are not limited to:

- ADEQ being established as the primary quality assurance organization (PQAO) for the criteria and non-criteria pollutant air monitoring data collected and reported to EPA's Air Quality System (AQS) database.
- An agency-level Quality Management Plan (QMP), which is an "umbrella" document that details, in broad terms, the strategies used to carry out QA/QC in environmental data collection activities.
- Division-level quality assurance program and project plans (QAPPs) for each major, ongoing air monitoring network. Each QAPP describes:
 - o purpose for operating the monitoring station or network
 - data quality objectives (DQOs) and/or measurement quality objectives (MQOs) along with data quality indicators (DQIs) that specify the amount of tolerable error in the data using statistical metrics
 - o variety of regularly occurring quality control (QC) checks along with pass/fail criteria
 - o types of QA assessments and reports needed from the network
 - o data validation processes and data reporting requirements
- Standard operating procedures (SOPs) that document procedures to assure that work products
 are reliable, reproducible, and consistent in quality. SOPs also serve to clearly communicate any
 process customizations in-use, providing a means of attesting that work products are credible,
 legally defensible, and meet or exceed our customers' and/or stakeholders' needs or
 requirements. Additional forms of standardization are used to document procedures for
 accuracy.

ADEQ uses a multi-tiered approach to data validation to ensure consistent quality. It requires all data to move through different levels of QA by separate reviewers. Data have five different levels associated with each tier level.

Raw – Original unchanged data recorded by the sampler or produced by laboratory analysis.

- Level 1 Data are reviewed programmatically and automatically using software written to flag
 data upon receipt to ADEQ's database. The data are flagged valid or invalid based on certain
 instrumentation parameters.
- Level 2 Data are reviewed manually on a weekly basis by an initial data reviewer to flag any
 discrepancies found. This gives the data a preliminary validation decision and identifies outliers,
 anomalous data, and instrumentation/laboratory issues.
- Level 3 Data are reviewed manually on a monthly basis by the final data reviewer by looking at the data spatially and temporally. QC measures are incorporated, environmental events are identified, and a final determination on the validity of data is made.
- Certified Data are uploaded to AQS quarterly and are certified annually by ADEQ by ensuring compliance with programmatic goals of data completeness, data precision, and data bias.

Periodically, EPA publishes reports for some of the criteria pollutant networks, and potentially non-criteria pollutant networks, that rate and/or rank monitoring organizations' performance over a three- year period. The QA Team, as well as other personnel in the data management and quality assurance (DM&QA) unit, reviews these reports to gauge how well ADEQ's networks are performing with those across the nation. If needed, corrective actions are taken to ensure data of the highest quality possible are collected.

4.0 Monitoring Location Summary

Table 2 contains some of ADEQ's site associated metadata including the site number from EPA's AQS database, the city in which the site is located, the main monitoring objectives for the site, the site coordinate location, and which parameters were recorded at the site in 2017.

Table 2 – Site Index

Site Name	Air Quality System ID	Location	Monitoring Objective(s)	Lat. Long. (Deg.)	Parameters Recorded
ADEQ Building	None	Phoenix, AZ	Visibility	33.4483, -112.088	Digital High Resolution Image
Ajo	04-019-0001	Ajo, AZ	NAAQS Comparison	32.3820, -112.858	PM ₁₀ , Wind, Temp/RH
Alamo Lake	04-012-8000	Alamo Lake State Park, AZ	NAAQS Comparison	34.2439, -113.559	O ₃ , CO, PM ₁₀ , PM _{2.5} , Wind, Temp/RH
Banner Mesa Medical Center	None	Mesa, AZ	Visibility	33.4335, -111.843	Digital High Resolution Image
Bullhead City	04-015-1003	Bullhead City, AZ	NAAQS Comparison/ Research	35.1539, -114.566	PM ₁₀ , PM _{2.5} (EBAM)
Douglas Red Cross	04-003-1005	Douglas, AZ	NAAQS Comparison	31.3492, -109.54	PM ₁₀ , PM _{2.5} , Temp/RH, Wind,
Estrella Mountain Community College	None	Avondale, AZ	Visibility	33.4836, -112.350	Digital High Resolution Image
Flagstaff Middle School	04-005-1008	Flagstaff, AZ	NAAQS Comparison/ Public Information	35.2061, -111.653	O ₃ , PM _{2.5} (EBAM)
Globe Highway	04-007-1002	Winkelman, AZ	NAAQS Comparison	33.002, -110.765	Pb, Temp/RH, Wind
Hayden Old Jail	04-007-1001	Hayden, AZ	NAAQS Comparison	33.0062, -110.786	SO ₂ , PM ₁₀ , Temp/RH, Wind
Hillcrest	04-007-1003	Hayden, AZ	NAQQS Comparison	33.004 - 110.782	Pb
JLG Supersite	04-013-9997	Phoenix, AZ	NAAQS Comparison/ Research	33.5038, -112.096	CO, NO ₂ , NO ₇ , O ₃ , SO ₂ , Carbonyl, VOC, SVOC, Pb- PM ₁₀ , / PM ₁₀ metals speciation, PM ₁₀ , PM _{10-2.5} , PM _{2.5} , PM _{2.5} speciation, Temp/RH, Wind, IMPROVE, Delta Temp, Horizontal Solar Radiation, Ultraviolet Solar Radiation
Meadview	04-015-9000	Meadview, AZ	Visibility	36.0193, -114.068	IMPROVE
Miami Golf Course	04-007-8000	Miami, AZ	NAAQS Comparison	33.4190, -110.83	Pb, PM ₁₀ , Temp/RH, Wind
Miami Jones Ranch	04-007-0011	Miami, AZ	NAAQS Comparison	33.3853, -110.867	SO ₂
Miami Ridgeline	04-007-0009	Miami, AZ	NAAQS Comparison	33.3992, -110.859	SO ₂
Miami Townsite	04-007-0012	Miami, AZ	NAAQS Comparison	33.3973, -110.874	SO ₂
Nogales Post Office	04-023-0004	Nogales, AZ	NAAQS Comparison	31.3372, -110.937	PM ₁₀ , PM _{2.5} , Temp/RH, Wind, IMPROVE
North Mountain Summit	None	Phoenix, AZ	Visibility	33.5855, -112.072	Digital High Resolution Image
Organ Pipe National Monument	04-019-005	Ajo, AZ	Visibility	31.950, -112.80	IMPROVE
Paul Spur Chemical Lime Plant	04-003-0011	Paul Spur, AZ	NAAQS Comparison	31.366, -109.73	PM ₁₀ , Temp/RH, Wind

Site Name	Air Quality System ID	Location	Monitoring Objective(s)	Lat. Long. (Deg.)	Parameters Recorded
Payson Well Site	04-007-0008	Payson, AZ	NAAQS Comparison/	34.230,	PM ₁₀ , PM _{2.5} (EBAM),
			Public Information	-111.33	Temp/RH, Wind
Phoenix Transmissometer Receiver	None	Phoenix, AZ	Visibility	33.490, -112.08	Bext, Temp/RH
Phoenix Transmissometer Transmitter	None	Phoenix, AZ	Visibility	33.525, -112.10	Bext
Prescott Pioneer Park	04-025-8034	Prescott, AZ	NAAQS Comparison/ Public Information	31.612, -112.463	O ₃ , PM _{2.5} (EBAM)
Queen Valley	04-021-8001	Queen Valley, AZ	NAAQS Comparison	33.294, -111.29	O ₃ , Temp/RH, Wind
Rillito	04-019-0020	Rillito, AZ	NAAQS Comparison	32.414, -111.16	PM ₁₀ , Temp/RH, Wind
Saguaro Nation Park West	04-019-9000	Tucson, AZ	Visibility	32.249, -111.22	IMPROVE (Protocol)
San Luis Rio Colorado Well 10	80-026-8012	San Luis Rio Colorado, Mexico	Research	32.467, -114.769	O ₃ , Temp/RH, Wind
Sedona Fire Station AQD	None	Sedona, AZ	Public Information	34.868, -111.763	PM _{2.5} (EBAM)
Show Low	None	Show Low, AZ	Public Information	34.2524, -110.0364	PM _{2.5} (EBAM)
South Phoenix	04-013-4003	Phoenix, AZ	Research	33.403, 112.08	VOC
Springerville	None	Springerville, AZ	Public Information	34.1284, -109.2892	PM _{2.5} (EBAM)
Tonto National Monument	04-007-0010	Roosevelt, AZ	NAAQS Comparison	33.655, -111.11	O ₃ , IMPROVE
Verde Ranger Station	None	Camp Verde, AZ	Public Information	34.5490, -111.8472	PM _{2.5} (EBAM)
Yuma Supersite	04-027-8011	Yuma, AZ	NAAQS Comparison	32.690, -114.62	O ₃ , PM ₁₀ , PM _{2.5} , Temp/RH, Wind

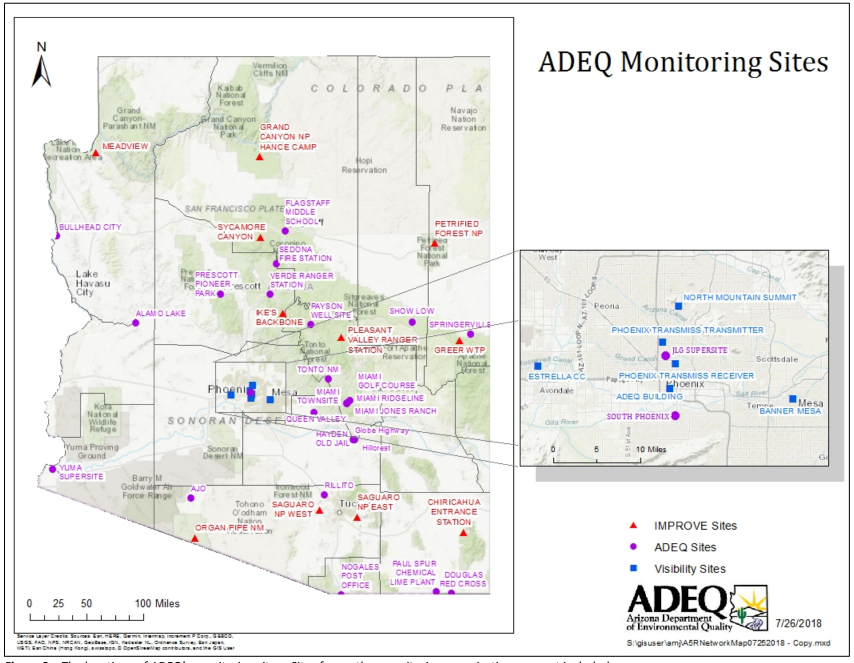


Figure 2 – The locations of ADEQ's monitoring sites. Sites from other monitoring organizations are not included.

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Section II - Criteria Pollutants

The seven criteria pollutants as outlined in the CAA are discussed at length in this Section. The health effects of each pollutant, the specific background of the pollutants in Arizona, and an explanation of current monitoring techniques are provided. Also included is a discussion of 2017 data, which shows a history of the pollutant standards and compliance with the current standards. Additionally, a trend analysis provides a history of the monitors operating in Arizona. The trend analysis includes a summary of the quantitative increase/decrease of the pollutant over a period of 5 years, 10 years, and length of record. Lastly, a summary of the AQI values in 2017 for each pollutant is presented.

1.0 Carbon Monoxide (CO)

Carbon Monoxide is a colorless, odorless, tasteless gas that is produced by the incomplete combustion of fuels. It has a variety of adverse health effects that arise from its ability to chemically bind to blood hemoglobin. Carbon Monoxide successfully competes with oxygen for binding with hemoglobin and thereby impairs oxygen transport. This impaired transport leads to several central nervous system effects, such as headache, fatigue, and dizziness. Chronic CO exposure also contributes to or exacerbates arteriosclerotic heart disease. Chronic exposure to low levels of CO can lead to depression, confusion, and memory loss.

1.1 Background

According to the 2014 National Emissions Inventory (NEI), in Arizona, 52 percent of CO emissions come from mobile sources including on-road motor vehicles, off-road motor vehicles, construction equipment, and lawn and garden equipment; 24 percent from biogenic sources; 19 percent from fires; 3 percent from miscellaneous sources; and 2 percent from fuel combustion. This pollutant has background levels, with the concentrations next to busy streets, and elevated concentrations in locations with significant amounts of emissions transported from upwind areas. Concentrations peak from November to January because emissions are highest in cold weather. Automotive emissions of CO are greatest when engines operate in open loop. This occurs for longer periods of time in cold weather as the engine attempts to warm up and because the mixed layer of the atmosphere is most shallow in wintertime



Figure 3 – Map of ADEQ's CO sites

due to decreased solar heating. Hourly concentrations tend to be at their maximum during the morning rush hour and between 6 p.m. and midnight.

Emission controls have reduced overall CO emissions, and the standards have been achieved in the metropolitan Phoenix area since 1996, in stark contrast to the first half of the 1980s when more than 100 exceedances were recorded each year. Similar improvements have occurred in Tucson, where the last eight-hour exceedances were recorded in 1988. Equipping vehicles with catalytic converters and electronic ignition systems was the most effective control, but significant reductions can also be attributed to the vehicle emissions inspection program (beginning in 1976) and oxygenated fuels (beginning in 1989). In the early 2000's, the EPA re-designated the Phoenix and Tucson metropolitan areas to attainment for the NAAQS for CO, and approved the attainment demonstration and maintenance plan. Each area is now in its second maintenance plan period.

ADEQ operated two CO monitors in 2017: one at its NCore station and a second added in September at Alamo Lake as a background SPM monitor. Additional CO monitors are operated by other State and Local agencies as required, but will not be discussed in this report.

1.2 Monitoring Methods

Carbon Monoxide is monitored continuously with a nondispersive infrared (IR) instrument. The IR light passes through a gas filter correlation wheel that alternately uses a CO filled chamber and a chamber with no CO present. The light path travels through a sample cell following the correlation wheel and continues on toward an IR light detector which converts the light into an electrical signal. Carbon Monoxide absorbs a specific wavelength of IR light and the energy loss through the sample cell is compared with the zero reference provided by the gas filter correlation wheel to produce an electrical signal that is proportional to concentration. Raw data readings are retrieved by a data collection system and stored in a database. Readings are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis.

1.3 Compliance/Summary of Design Values

On April 30, 1971, the EPA promulgated NAAQS for CO based on a criteria document published by the U.S. Department of Health, Education and Welfare in March 1970. Identical primary (health-based) and secondary (welfare-based) NAAQS for CO were set at 35 parts per million (ppm), one-hour average, and at 9 ppm, eight-hour average, neither to be exceeded more than once a year. After the most recent review of the CO NAAQS, on August 31, 2011, EPA proposed to retain the current primary standards. After review of the air quality criteria, EPA further concluded that no secondary standard should be set for CO at this time. Table 3 summarizes the history of the NAAQS for CO during the period 1971-2011. At present there are two primary standards for CO. The one-hour standard is 35 ppm and the eight-hour standard is 9 ppm.

Table 3 – History of the National Ambient Air Quality Standards for CO during the period 1971-2011 (Source: USEPA TTN NAAOS)

(Source: OSETATTI	Source. Usera TTN NAAQS)						
Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form		
1971			1-hour	25	Maximum, not to be exceeded		
	Primary and	60	period	35 ppm	more than once in a year		
36 FR 8186	Secondary	CO	8-hour	0	Maximum, not to be exceeded		
Apr 30, 1971			period	9 ppm	more than once in a year		
1985							
	Primary standards retained, without revision; secondary standards revoked.						
50 FR 37484	Primary	stanuarus re	rtained, witho	out revision	; secondary standards revoked.		
Sept 13, 1985							
1994							
		Drima	ry standards	rotained u	vith out rovision		
59 FR 38906		PIIIII	iry Stariuarus	retained, w	vithout revision.		
Aug 1, 1994							
2011							
		Drima	ry standards	rotained w	vithout revision.		
76 FR 54294		PIIIII	iry Stariuarus	retained, w	VILLIOUL LEVISION.		
Aug 31, 2011							

According to 40 CFR Part 50, compliance for both standards is determined by having no more than one exceedance per year. EPA determines attainment of the standard by evaluating two calendar years of data from each site. The highest of the annual second-highest values in a two-year period must not exceed the standard of 35 ppm for the one-hour standard or 9 ppm for the eight-hour standard.

Table 4 and Table 5 below show the 1st maximum and 2nd maximum values for both the current primary standards for the years 2016 and 2017. No exceedances of the one-hour or eight-hour standards were recorded in 2016 and 2017. The CO monitor at Alamo Lake and JLG Supersite are considered to be in compliance for the year 2016 and 2017.

Table 4 - CO One-Hour Compliance Summary

Table 4 Co one floar compliance summary								
2016 to 2017 One-Hour CO Compliance Summary (in ppm)								
(NAAQS primary one-hour standard 35 pp	(NAAQS primary one-hour standard 35 ppm)							
2016 2017 Compliance								
Site Name	1 st Max	2 nd Max	1 st Max	2 nd Max	Compliance			
	Value	Value	Value	Value	Value			
La Paz County								
Alamo Lake ¹	0.5#	0.4#	0.3	0.3	0.4*			
Maricopa County								
JLG Supersite	2.3#	2.2#	2.6	2.4	2.4*			
Number of Sites in Violation of the NAAQS					0			

¹Monitor began operation in September 2016.

[#]Indicates the data do not satisfy EPA's summary criteria.

^{*}Preliminary Design Value since data completeness was not satisfied.

Table 5 – CO Eight-Hour Compliance Summary

2016 to 2017 Eight-Hour CO Compliance Summary (in ppm) (NAAQS primary eight-hour standard 9 ppm)						
2016 2017 Compliance						
Site Name	1 st Max Value	2 nd Max Value	1 st Max Value	2 nd Max Value	Compliance Value	
La Paz County						
Alamo Lake ¹	0.3#	0.3#	0.3	0.3	0.3*	
Maricopa County						
JLG Supersite	1.9#	1.6#	1.8	1.8	1.8*	
Number of Sites in Violation of the NAAQS					0	

¹Monitor began operation in September 2016.

1.4 Trends

Monitoring of CO throughout the state of Arizona contains the longest history of all the criteria pollutants. Most of this long-term monitoring was located in the highly urbanized areas of Phoenix and Tucson, and several of these CO sites contain monitoring records dating back to the 1970s. For the purpose of this report, the examination of CO trends will include the ADEQ monitors only.

Figure 4 and Figure 5 illustrate the temporal variability of CO for a period of nineteen years from 1999 to 2017. ADEQ has monitored CO at JLG Supersite since 1999 and recently Alamo Lake starting in 2016. ADEQ started trace-level monitoring of CO at JLG supersite in 2010 as part of the NCore program. Most of the improvements can be attributed to emission control programs as stated in section 1.1. The average trends for a period of 5 years, 10 years, and length of record for JLG Supersite are provided in Table 6 below.

^{*}Indicates the data do not satisfy EPA's summary criteria.

^{*}Preliminary Design Value since data completeness was not satisfied.

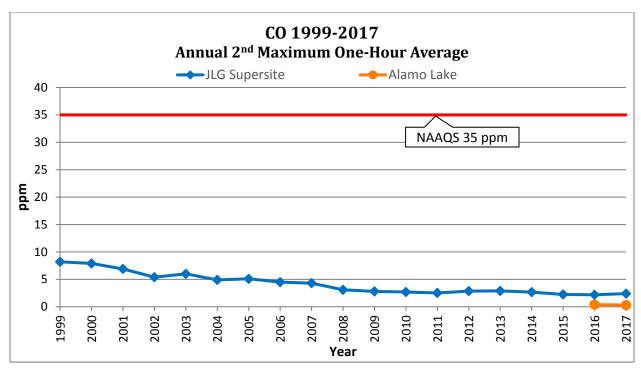


Figure 4 – CO One-Hour Average Trend

Note: Some years might not satisfy completeness criteria.

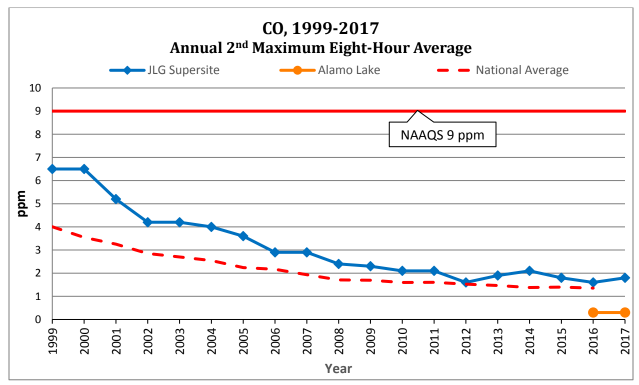


Figure 5 – CO Eight-Hour Average Trend

Link to National Averages (https://www.epa.gov/air-trends/carbon-monoxide-trends). 2017 National Averages were not available.

Note: Some years might not satisfy completeness criteria.

Table 6 – CO One-Hour & Eight-Hour Average Trend

	JLG Supersite			
Trend Length	2 ND MAX 1 HR	2 ND MAX 8 HR		
5 Year (2013 - 2017)	21% ↓ Decrease	14% ↓ Decrease		
10 Year (2008 - 2017)	23% ↓ Decrease	27% ↓ Decrease		
Length of Record	82% ↓ Decrease 1999-2017	85% ↓ Decrease 1999-2017		

1.5 Air Quality Index (AQI)

The CO daily AQI values for 2017 are categorized into the different levels of health concerns shown in Table 7. Background levels of CO are near zero in areas without manmade sources. Levels of CO in more populous areas are low and stable due to significant emissions controls placed on mobile sources (cars, trucks, etc.). Emissions are highest in the winter months due to the increased time it takes engines to warm up. Wintertime meteorology in Arizona is typified by stable conditions, therefore increasing AQI values. A graphical representation of the CO Daily AQI values for 2017 is shown in Figure 6.

AQI Values	Levels of Health Concern	Number	of Days
AQI values	Levels of Health Concern	JLG Supersite	Alamo Lake
0 - 50	Good	283	300
51 - 100	Moderate	0	0
101 - 150	Unhealthy for Sensitive Groups	0	0
151 - 200	Unhealthy	0	0
201 - 300	Very Unhealthy	0	0
301 - 500	Hazardous	0	0
	Missing	82	65
	Total Days	365	365

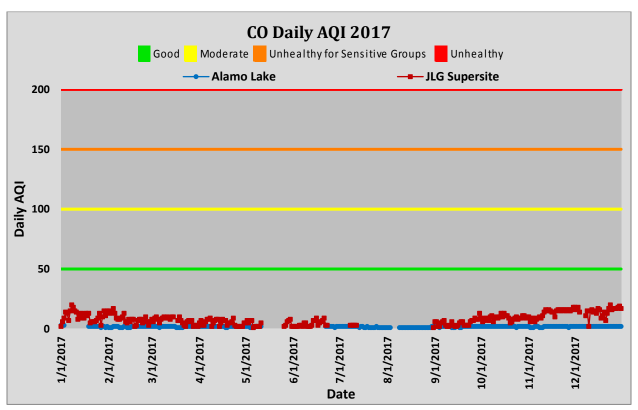


Figure 6 - CO Daily AQI 2017

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data.

2.0 Nitrogen Dioxide (NO₂)

 NO_2 is a reddish-brown gas that is formed by the oxidation of nitric oxide (NO) - a byproduct of combustion. Adverse health effects associated with NO_2 include the risk of respiratory illness in children and vary depending on the level of NO_2 and exposure time. Short exposure to low levels may result in changes to airway responsiveness and decreased lung function in individuals with pre-existing conditions. Irreversible changes may occur to lungs due to long-term exposure to higher levels. This pollutant is also of concern in its reduction of visibility and its contributory role in the photochemical formation of ground level O_3 and acid rain.

2.1 Background

Combustion emissions of nitrogen oxides (NO_x) are 95 percent NO and five percent NO₂. NO₂ concentrations often serve as the indicator for the larger group of nitrogen oxides since NO rapidly oxidizes to NO₂. In the 2014 NEI, Arizona NOx emissions were led by the transportation sector with 72 percent of the emissions from mobile sources such as cars and trucks; 18 percent came from fuel combustion processes such as utility power plants; and 6 percent from biogenics, 2 percent from fires, 2 percent from processes industrial and 1 percent miscellaneous. NO and NO₂ concentrations are highest near major roadways. NO concentrations decrease rapidly with distance from the roadway, whereas NO₂ concentrations are more evenly distributed because of their formation through oxidation and their subsequent transport. Concentrations of NO₂ are highest in the late afternoon and early evening of winter, when rush hour emissions of NO are converted to NO2 under

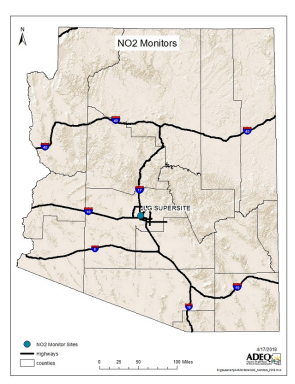


Figure 7 – Map of ADEQ's NO₂ sites

relatively stable atmospheric conditions. Because NO reacts rapidly with O_3 , nocturnal O_3 concentrations in cities are often reduced to near zero levels, while concentrations at background sites remain higher.

NO emissions have been reduced over time using several different techniques. NO emissions from motor vehicles have been reduced through retardation of spark timing, lowering the compression ratio, exhaust gas recirculation systems, diesel retrofits, Ozone Action Days, and three-way catalysts. Also, the vehicle inspection program's NO_x test for light-duty gasoline vehicles age 1981 and newer (in Phoenix only) has helped reduce emissions. Controls on electricity generating units include catalytic reductions

and natural gas re-burn. Reformulated gasolines also decrease NO emissions: The Phoenix Metropolitan area adopted "Arizona Cleaner Burning Gasoline" in 1998 to reduce vehicular NO emissions.

One NO₂ monitor was operated by ADEQ in 2017 and is located in a local neighborhood which represents a typical Phoenix area community. This monitor is part of the PAMS network at JLG Supersite. Additionally, one total NO_y monitor was operated by ADEQ, which was operated as part of the NCore station and PAM network at JLG supersite. However, NO_y is not a criteria pollutant and is not further evaluated in this section.

2.2 Monitoring Methods

 NO_2 is monitored continuously with chemiluminescence instruments which operate on the principle that when two chemicals combine, a unique wavelength of light is emitted. This wavelength of light is detected using a light sensor, and the intensity of that light is a direct correlation to the concentration of the target chemical species (NO_2). An NO_2 analyzer is based on the chemiluminescence of an excited NO_2 molecule which determines NO and NO_x (the sum of NO_2 and NO) concentrations. Readings from all instruments are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis. Readings are retrieved by a data collection system and stored in a database.

2.3 Compliance/Summary of Design Values

The NAAQS for NO_2 was reviewed in 2010 and the original 1971 primary NAAQS of 53 parts per billion (ppb) for the annual mean was retained. However, a new primary one-hour NO_2 standard of 100 ppb was added. In 2012, the annual secondary standard of 53 ppb was reviewed and again retained. The annual standard is attained when the annual arithmetic mean concentration in a calendar year is less than or equal to 53 ppb. The one-hour standard is attained when the three-year average of the 98^{th} percentile of the yearly distribution of one-hour daily maximum NO_2 concentrations is below 100 ppb. A history of the NAAQS for NO_2 is provided in Table 8.

Table 8 − History of the National Ambient Air Quality Standards for NO₂ during the period 1971-2010 (Source: USEPA TTN NAAQS)

Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form
1971 36 FR 8186 Apr 30, 1971	Primary and Secondary	NO ₂	Annual	53 ppb	Annual arithmetic average
1985 50 FR 25532 Jun 19, 1985	Priı	mary and sec	condary NO ₂ s	standards ret	ained, without revision.

Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form
1996					
61 FR 52852 Oct 8, 1996	Priı	mary and sec	condary NO ₂ s	standards ret	ained, without revision.
2010		NO ₂	1-hour	100 ppb	98 th percentile, averaged over 3
	Primary	1402	1-11001	100 bbp	years
75 FR 6474	Fillialy	Drin	mary annual N	IO standard	retained, without revision.
Feb 9, 2010		PIII	ilaly allitual i	NO ₂ Standard	retained, without revision.
2012					
77 FR 20218 Apr 3, 2012	Secondary	Existing so	econdary NO ₂	2 standard (ar	nnual) retained, without revision.

Table 9 below shows the 2017 annual mean for the 53 ppb primary standard. The annual mean of JLG Supersite is significantly below this primary standard, and is in compliance with the NAAQS.

Table 9 - NO₂ Annual Mean Compliance Summary

2017 NO₂ Annual Mean (in ppb) (NAAQS Annual Mean 53 ppb)	
Site Name	2017 Annual Mean
Maricopa County	
Maricopa County	
JLG Supersite	15.28*

^{*}Preliminary Design Value since data completeness was not satisfied.

The NO_2 three-year average of the one-hour averages at the 98th percentile was approximately half of the 100 ppb standard at JLG Supersite and is in compliance with the NAAQS. Refer to Table 10 for the 2017 three-year average.

Table 10 – NO₂ One-Hour Compliance Summary

2015 to 2017 One-Hour Average NO₂ Compliance (in ppb) (NAAQS One-Hour Average 100 ppb)				
Site Name	98 th Percentile Samples			Three-Year Average
	2015	2016	2017	
Maricopa County				
JLG Supersite	50.0#	51.2	51.6#	51*
Number of Sites in Violation of the NAAQS			0	

^{*}Indicates the data do not satisfy EPA's summary criteria.

^{*}Preliminary Design Value since data completeness was not satisfied.

2.4 Trends

ADEQ began monitoring for NO_2 in Phoenix at JLG Supersite in 1993. However, data are not readily available in AQS prior to 1999. As a result, the assessment of trends in NO_2 uses a period of nineteen years from 1999 to 2017.

Figure 8 and 9 illustrate the temporal variability of JLG Supersite over the 1999 to 2017 period in the form of annual means and one-hour averages at the 98th percentile. The average trends for a period of 5 years, 10 years, and length of record for JLG Supersite are provided in Table 11 below.

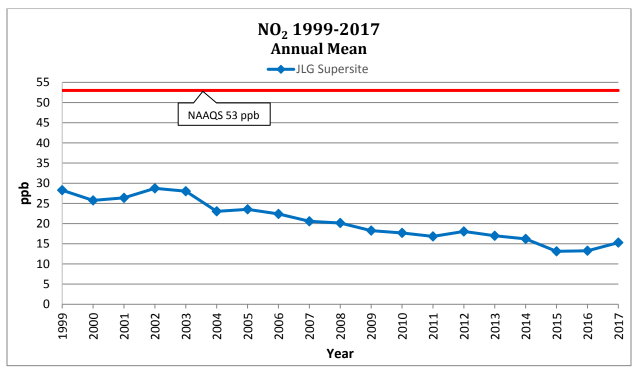


Figure 8 - NO₂ Annual Mean Trend

Note: Some years might not satisfy completeness criteria.

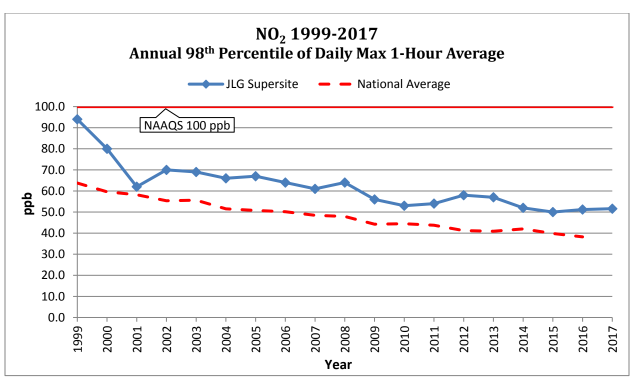


Figure 9 – NO₂ One-Hour Average Trend

Link to National Averages (https://www.epa.gov/air-trends/nitrogen-dioxide-trends). 2017 National Averages were not available.

Note: Some years might not satisfy completeness criteria.

Table 11 - NO₂ One-Hour Average Trend

	3	
	Site Name	
Trend Length	JLG Supersite	
5 Year	8% ↓	
(2013 - 2017)	Decrease	
10 Year	15% ↓	
(2008 - 2017)	Decrease	
	39% ↓	
Length of Record	Decrease	
	1999-2017	

2.5 Air Quality Index (AQI)

The NO_2 daily AQI values for 2017 are categorized into the different levels of health concerns in Table 12. Background levels of NO_2 are near zero in areas without manmade sources. Levels of NO_2 in more populous areas are low and stable due to significant emissions controls placed on mobile sources (cars, trucks, etc.). Emissions are highest in the winter months due to the NO_2 from mobile sources in stable atmospheric conditions, which can elevate AQI values. A graphical representation of the NO_2 Daily AQI values for 2017 is shown in Figure 10.

Table 12 - NO₂ Daily AQI Count 2017

AQI Values	Levels of Health Concern	Number of Days
AQI Values	Levels of Health Concern	JLG Supersite
0 - 50	Good	305
51 - 100	Moderate	5
101 - 150	Unhealthy for Sensitive Groups	0
151 - 200	Unhealthy	0
201 - 300	Very Unhealthy	0
301 - 500	Hazardous	0
	Missing	55
	Total Days	365

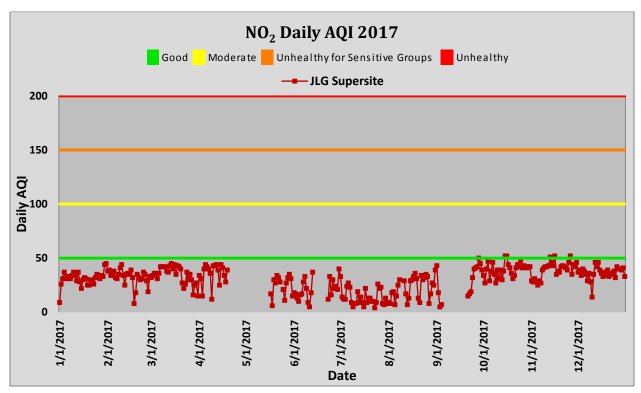


Figure 10 - NO₂ Daily AQI 2017

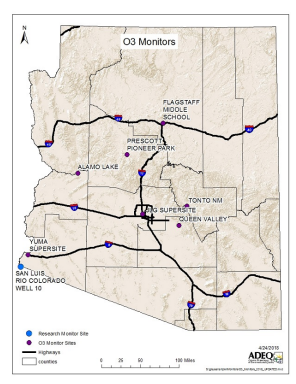
Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data.

$3.0 \, Ozone \, (O_3)$

Ozone is a colorless, slightly odorous gas that is both a natural component of the upper atmosphere and a key air contaminant in the lower atmosphere. In the stratosphere, O₃ blocks harmful ultraviolet radiation. In the lower atmosphere, its photochemical formation by the reaction of volatile organic compounds (VOC), hydrocarbons (HC), and Nitrogen Oxides (NO_x), leads to concentrations harmful to people, animals, plants, and materials (plastics, tires, o-rings). In both animals and humans, O₃ causes significant physiological and pathological changes at concentrations present in many urban environments. Short-term (one to two hours) exposure to concentrations in the range of 100 ppb to 400 ppb induce changes in lung function, including increased respiratory rates, increased pulmonary resistance, decreased volume of air, and changes in lung mechanics. Symptomatic responses in exercising adults include throat dryness, chest tightness, substernal pain, coughing, wheezing, pain on deep inspiration, shortness of breath, and headache. These symptoms also have been observed at lower concentrations for longer exposure times. Evidence suggests that O₃ exposure makes the respiratory airways more susceptible to other bronchoconstrictive challenges and interferes with or inhibits the immune system. Ozone at ambient concentrations also injures the stomata of plants, which are the cells that regulate plant respiration, resulting in flecks on the upper leaf surfaces of dichotomous plants and the death of the tips of coniferous needles. Plant scientists consider O₃ to be the most important of all of the phytotoxic air pollutants, causing over 90 percent of all plant injury from air pollution on a global basis.

3.1 Background

High O₃ concentrations are a phenomenon caused when sunlight, emissions from plant life, and evaporating fuel emissions peak. Fuel emissions consist of VOCs and NOx and are two of the main components of O₃ formation. According to the 2014 NEI, VOC emissions in Arizona are: 89 percent biogenic emissions (grass, shrubs, trees); 4 percent mobile sources (cars and trucks, off-road vehicles and equipment such as lawn mowers); 3 percent fires (wildfire and wood burning); 3 percent industrial solvent processes; and 1 percent miscellaneous. Nitrogen oxides are: 72 percent mobile sources; 18 percent fuel combustion (power plants, industrial processes); 6 percent biogenic emissions; 2 percent industrial processes; 2 percent fires; and 1 percent miscellaneous. Ozone has relatively high background levels, with the daily maximum in remote areas being about one-half to three-quarters of the daily maximum in urban areas. In these areas, the highest O₃ concentrations tend to occur on the downwind edge, although high concentrations do occur less Figure 11 - Map of ADEQ's O3 sites



frequently in the central city. Urban O_3 concentrations are low to near zero at night and rise rapidly through the morning and peak in the afternoon.

Controls to reduce the precursors of O₃, VOC, and NO_x have been successfully implemented for years. Nitrogen oxides and VOC from vehicular exhaust have been reduced through engine modifications and three-way catalytic converters. Reformulated gasoline standards, ethanol blends, ozone education days, and alternative fuel programs all help reduce VOC emission. Evaporative HC from vehicles have been reduced through better engineered fuel tanks and auxiliary plumbing combined with carbon absorption canisters. Additional reductions of vehicular VOCs have come through ADEQ's vehicle emissions inspection program, which tests all gasoline fueled vehicles for HC (in Phoenix and Tucson), through vapor capturing equipment for gasoline tankers, vapor recovery systems at retail gas stations (Phoenix area only), and cleaner burning gasoline (Phoenix area only). Stationary sources of HC have been reduced through a variety of better control equipment required by stricter regulations.

In 2017, ADEQ operated a network of eight O_3 monitors to support a variety of monitoring objectives; chief among them is for NAAQS compliance. Other monitoring objectives include support for the NCore station, PAMS, and to show O_3 transport coming from across Arizona's borders.

3.2 Monitoring Methods

Continuous monitoring for O_3 is done with an ultraviolet absorption instrument. A specific ultraviolet wavelength of light which O_3 absorbs is passed through a sample cell. A drop in light intensity is detected by a light sensor and that drop is a direct correlation to the concentration of O_3 in the sample cell. This results in accurate readings of O_3 concentrations continuously. Readings are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis. Readings are retrieved by a data collection system and stored in a database.

3.3 Compliance/Summary of Design Values

On Oct. 1, 2015, the U.S. Environmental Protection Agency (EPA) strengthened the National Ambient Air Quality Standards (NAAQS) for ground-level ozone based on EPA's review of the air quality criteria for ozone (O_3) and related photochemical oxidants and for O_3 . Further, based on extensive scientific evidence about ozone's effects on public health and welfare, EPA revised the primary and secondary ozone standard levels to 0.070 parts per million (ppm), and retained their indicators (O_3), forms (fourth-highest daily maximum, averaged across three consecutive years) and averaging times (eight hours). The designation rule finalized the nonattainment area boundaries for areas that do not meet the 0.070 ppm standard. The standard is met when the 4^{th} highest rolling eight-hour average for the year, averaged over three years is less than the 0.070 ppm standard. Thus, an exceedance above the 0.070 ppm standard for a given year may not cause a violation of the standard, if the three-year average is still below 0.070 ppm. The designations are based on air quality monitoring data and the history of the NAAQS for O_3 is provided in Table 13 below.

 $\textbf{Table 13} - \text{History of the National Ambient Air Quality Standards for } O_3 \text{ during the period } 1971-2015$

(Source: USEPA TTN NAAOS)

Final	Primary/	Indicator	Averaging Time	Level	Form
Rule/Decision	Secondary				
1971 36 FR 8186 Apr 30, 1971	Primary and Secondary	Total photochemical oxidants	1 hour	0.08 ppm	Not to be exceeded more than one hour per year
1979 44 FR 8202 Feb 8, 1979	Primary and Secondary	O ₃	1 hour	0.12 ppm	Attainment is defined when the expected number of days per calendar year, with maximum hourly average concentration greater than 0.12 ppm, is equal to or less than 1
1993					
58 FR 13008 Mar 9, 1993	EPA de	cided that revision	ons to the sta	indards wer	e not warranted at the time
1997 62 FR 38856 Jul 18, 1997	Primary and Secondary	O ₃	8 hours	0.08 ppm	Annual fourth-highest daily maximum 8-hr concentration, averaged over 3 years
2008 73 FR 16483 Mar 27, 2008	Primary and Secondary	O ₃	8 hours	0.075 ppm	Annual fourth-highest daily maximum 8-hr concentration, averaged over 3 years
2015 80 FR 65292 Oct 26, 2015	Primary and Secondary	O ₃	8 hours	0.070 ppm	Annual fourth-highest daily maximum 8 hour average concentration, averaged over 3 years

The data in Table 14 are from the sites in operation from 2015 to 2017 and have been evaluated based on the 2015 O_3 standard (0.070 ppm). Three sites met the revised standard: Flagstaff Middle School, Alamo Lake and Prescott Pioneer Park. The other four sites were in violation of the current 0.070 ppm O_3 NAAQS.

Table 14 – O₃ Compliance Summary

2015 to 2017 Eight-Hour O₃ Compliance (in ppm) (NAAQS eight-hour 0.070 ppm)									
	Fourt	th-Highest \	/alue	Three-					
Site Name	2015	2016	2017	Year Average					
Coconino County									
Flagstaff Middle School ¹	0.070	0.064	0.066	0.066					
Gila County									
Tonto NM ¹	0.073	0.070	0.076	0.073					
La Paz County									
Alamo Lake ¹	0.070	0.067	0.069	0.068					
Maricopa County									
JLG Supersite	0.075	0.075	0.078	0.076					
Pinal County									
Queen Valley ¹	0.074	0.072	0.075	0.073					
Yavapai County ²									
Prescott College AQD ¹	0.067	0.064	N/A	0.067					
Prescott Pioneer Park	N/A	N/A	0.072	0.067					
Yuma County									
Yuma Supersite ¹	0.077	0.067	0.074	0.072					
Number of Sites in Violation of the NAAQS				4					

¹Seasonal monitor in 2015, operational during March 1st to December 31st.

Bold denotes exceedances and sites in violation of the 2015 NAAQS of 0.070 ppm.

²For compliance purposes, Prescott College and Prescott Pioneer Park site data are combined to calculate the Three-Year Average Design Value.

3.4 Trends

Monitoring for O₃ has occurred for many years in Arizona, mainly in the urbanized centers of Phoenix and Tucson. Figure 12 illustrates the temporal variability in the O₃ Annual 4th Maximum of Daily Max Eight-Hour averages for the period of thirteen years from 2005 to 2017. The examination of O₃ trends includes the monitors that have been run by ADEQ only for the years 2005 to 2017, which includes Alamo Lake, JLG Supersite, Queen Valley, and Tonto National Park. Additionally, trends were analyzed for the years 2008 to 2017 for Flagstaff Middle School, Prescott College/Pioneer Park, and Yuma Supersite. In 2017, the Prescott College site was relocated to the Prescott Pioneer Park site. The locations of both sites are close enough that they are considered the same site for the purpose of trends analysis. The average trends for a period of 5 years, 10 years, and length of record for each site are provided in Tables 15 and 16 below.

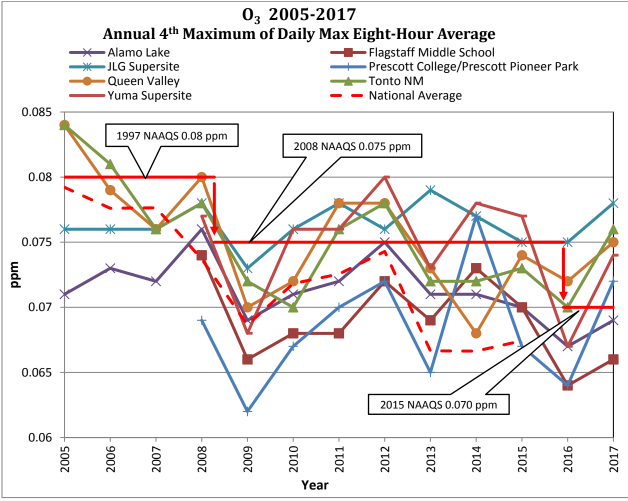


Figure 12 − O₃ Annual 4th Maximum of Daily Max Eight-Hour Average
Link to National Averages (https://www.epa.gov/air-trends/sulfur-dioxide-trends). 2017 National Averages were not available.

Note: Some years might not satisfy completeness criteria.

Table 15 – O₃ Annual 4th Maximum of Daily Max Eight-Hour Average Trend - Central/Southern Region

	Central/Southern Region							
Trend Length	JLG Supersite	Queen Valley	Tonto NM	Yuma Supersite				
5 Year	2% ↓	5% 个	3% 个	5% ↓				
(2013 - 2017)	Decrease	Increase	Increase	Decrease				
10 Year	1% 个	4% ↓	3% ↓	2% ↓				
(2008 - 2017)	Increase	Decrease	Decrease	Decrease				
Length of Record	1% 个	10% ↓	10% ↓	2% ↓				
	Increase	Decrease	Decrease	Decrease				
	2005-2017	2005-2017	2005-2017	2008-2017				

Table 16 – O₃ Annual 4th Maximum of Daily Max Eight-Hour Average Trend - Northern Region

	Northern Region								
Trend Length	Alamo Lake	Flagstaff Middle School	Prescott College/ Pioneer Park*						
5 Year (2013 - 2017)	4% ↓ Decrease	8% ↓ Decrease	1% 个 Increase						
10 Year (2008 - 2017)	7% ↓ Decrease	5% ↓ Decrease	4% 个 Increase						
Length of Record	5% ↓ Decrease 2005-2017	5% ↓ Decrease 2008-2017	4% 个 Increase 2008-2017						

^{*}Due to site relocation, Prescott College and Prescott Pioneer Park site data were combined to perform the trends calculations.

3.5 Air Quality Index (AQI)

The O_3 daily AQI values for 2017 are categorized into the different levels of health concerns in Table 17. Background levels of O_3 in Arizona are typically elevated in areas without manmade sources due to regional transport. Higher O_3 levels typically occur in the summer months when the sun angle is higher than during the winter months. Graphical representations of the O_3 Daily AQI values for 2017 are shown in Figures 13 and 14.

Table 17 - O₃ Daily AQI Count 2017

		Number of Days								
AQI Values	Levels of Health Concern	Alamo Lake	Flagstaff Middle School	JLG Supersite	Prescott Pioneer Park	Queen Valley	Tonto National Monument	Yuma Supersite		
0 - 50	Good	275	284	237	219	233	261	293		
51 - 100	Moderate	77	78	109	103	109	92	48		
101 – 150	Unhealthy for Sensitive Groups	2	1	13	5	12	8	10		
201 - 300	Unhealthy	0	0	1	0	1	1	0		
301 - 300	Very Unhealthy	0	0	0	0	0	0	0		
301 - 500	Hazardous	0	0	0	0	0	0	0		
	Missing	11	2	5	38	10	3	14		
	Total Days	366	365	365	365	365	365	365		

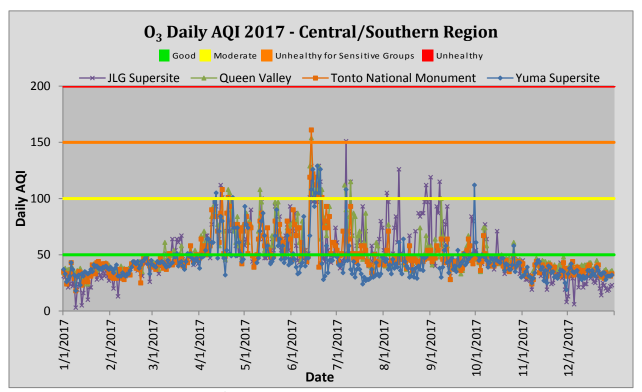


Figure 13 – O₃ Daily AQI 2017 – Central/Southern Region

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data.

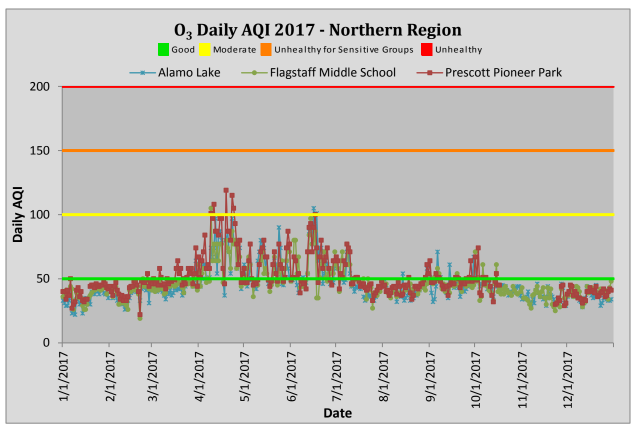


Figure 14 – O₃ Daily AQI 2017 – Northern Region

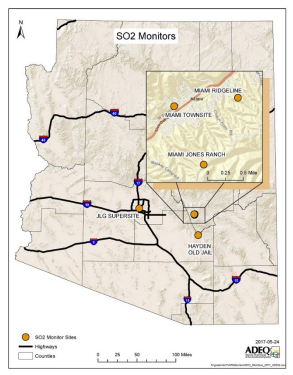
Note: Data are the daily AQI values for 2017 obtained from $\underline{\text{https://www.epa.gov/outdoor-air-quality-data}}$.

4.0 Sulfur Dioxide (SO₂)

Sulfur Dioxide is a colorless gas with a pungent, irritating odor at elevated concentrations. It mainly alters the mechanical function of the upper airway, including increasing the nasal flow resistance and decreasing the nasal mucus flow rate. Current scientific evidence links short-term exposures to SO₂, ranging from five minutes to 24 hours, with an array of adverse respiratory effects, including bronchoconstriction and increased asthma symptoms. These effects are particularly significant for asthmatics at elevated ventilation rates. Studies have shown a connection between short-term exposure and increased visits to emergency departments and hospital admissions for respiratory illnesses, particularly in at-risk populations including children, the elderly, and asthmatics.

4.1 Background

Sulfur Dioxide is removed from the atmosphere through dry deposition on plants and is converted to sulfuric acid and eventually to sulfate. Sulfur Dioxide has extremely low background levels, with elevated concentrations found downwind of large point sources. Concentrations in urban areas are low and are homogeneously distributed. In Arizona, the principal source of SO₂ emissions has been the smelting of sulfide copper ore. Most fuels contain trace quantities of sulfur and their combustion releases both gaseous SO₂ and particulate sulfate. In the 2014 NEI, Arizona showed 53 percent of SO₂ emissions came from industrial point sources including smelting; 38 percent from fuel combustion; 5 percent from fires; 3 percent from mobile sources, such as offroad vehicles and on-road motor vehicles; and 1 percent miscellaneous.



Vehicular emissions of SO_2 and sulfate have been reduced through lowering the sulfur content in diesel fuel and

Figure 15 - Map of ADEQ's SO₂ sites

gasoline. Major controls were installed in Arizona's copper smelters in the 1980s, which reduced SO_2 emissions substantially. Additional improvements to the controls at two of these smelters were started at the end of 2017, with plans of completion by the end of 2018. These are extensive improvements designed to further capture, control, and reduce SO_2 emissions. While these controls have reduced SO_2 emissions throughout the state in recent years, the two significant point sources are affecting nearby air quality. Copper smelting operations from these sources have caused the areas in Miami, AZ and Hayden, AZ to be designated by the EPA as nonattainment areas for the 2010 standard.

Compliance with the new NAAQS standard is being achieved throughout the state besides in these two areas. Historically, there were six copper smelters in operation in AZ. Concentrations and emission of SO₂ have been reduced since four of those major sources were removed.

ADEQ operated four source oriented monitors in 2017. Three are located in and around the Miami, AZ area and one is located in Hayden, AZ. One additional monitor was operated at JLG supersite as part of the NCore station.

4.2 Monitoring Methods

Continuous monitoring for SO₂ is done with pulsed fluorescence instruments. Sulfur Dioxide absorbs a specific wavelength of ultraviolet light. Absorption of light at this wavelength results in the fluorescent emission of light at a different wavelength. This second wavelength of light is detected by a light sensor and is directly proportional to the SO₂ concentration in the sample. Readings are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis and can be retrieved by a data collection system and stored in a database.

4.3 Compliance/Summary of Design Values

On April 30, 1971, the EPA promulgated primary and secondary NAAQS for sulfur oxides, measured as SO₂ under section 109 of the Clean Air Act (36 FR 8186). After periodic reviews of additional scientific information, EPA announced first in 1986 and then in 1996, its decision not to revise the NAAQS for SO₂. In 2010, EPA replaced both the 24-hour and annual standards with a new short-term standard based on the three-year average of the 99th percentile of the yearly distribution of one-hour daily maximum SO₂ concentrations. EPA set the level of this new standard at 75 ppb. The one-hour SO₂ standard added in 2010 is a primary standard. The revision of the SO₂ NAAQS in 2010 did not address the secondary standard, which remained a three-hour standard with a level of 0.5 ppm. This secondary standard was again retained in 2012. Table 18 summarizes the history of the NAAQS for SO₂ during the period of 1971-2012.

Table 18 – History of the National Ambient Air Quality Standards for SO₂ during the period 1971-2012 (Source: USEPA TTN NAAOS)

Source. OSEPA TTN	TVAAQ3)							
Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form			
			24-Hour	0.14 ppm	Not to be exceeded more than			
1971	Primary			0 pp	once per year			
		SO ₂	Annual	0.03 ppm	Annual arithmetic average			
36 FR 8186		302	3-Hour	0.5 ppm	Not to be exceeded more than			
Apr 30, 1971	Secondary		3-11001	0.5 ppm	once per year			
			Annual	0.02 ppm	Annual arithmetic average			
1973								
	Secondary 3-hour SO ₂ standard retained, without revision;							
38 FR 25678	Secondary	,	annual SO_2 standard revoked.					
Sept 14, 1973								

Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form			
1996								
61 FR 25566 May 22, 1996	Primary	Exis	ting primary S	SO ₂ standards	retained, without revision.			
2010	Drimany	SO ₂	1-hour	75 ppb	99 th percentile, averaged over 3 years			
75 FR 35520 Jun 22, 2010	Primary	Р	rimary annua	al and 24-hour	SO ₂ standards revoked.			
2012								
77 FR 20218 Apr 3, 2012	Secondary	Existing s	Existing secondary SO₂ standard (3-hour average) retained, without revision.					

Table 19 summarizes the compliance status for the annual 99^{th} percentile of one-hour daily maximum for SO_2 for the years 2015 to 2017. Hayden Old Jail, Miami Jones Ranch, Miami Ridgeline, and Miami Townsite are sites in violation of the 2010 NAAQS of 75 ppb.

Table 19 – SO₂ One-Hour Compliance Summary

2015 to 2017 Annual 99 th Percentile of One-Hour daily max. SO₂ Compliance (in ppb) (NAAQS primary one-hour 75 ppb)									
Site Name 2015 2016 2017									
Gila County									
Hayden Old Jail	246	359	280	295					
Miami Jones Ranch	242	150	270	221					
Miami Ridgeline ¹	171	120	99#	130*					
Miami Townsite	231	110#	135	159*					
Maricopa County									
JLG Supersite	5	5#	6#	6*					
Number of sites in violation of NAAQS									

¹Monitoring discontinued at Miami Ridgeline in September 2017.

Bold denotes exceedances and sites in violation of the 2010 NAAQS of 75 ppb.

[#]Indicates the data do not satisfy EPA's summary criteria, usually meaning less than 75 percent valid data recovery available in one or more calendar quarters.

^{*}Preliminary Design Value since data completeness was not satisfied.

4.4 Trends

In Arizona, SO_2 monitoring began in the late 1960s and early 1970s. These early sites were predominantly located near facilities where smelting of sulfide copper ore occurred. Monitoring SO_2 at several of these smelting facilities no longer occurs due to the termination of smelting operations in certain areas. However, a long SO_2 monitoring record does exist for the Hayden and Miami areas due to continued smelting operations.

Figure 16 reflects data trends for the length of record for each monitor that was operated by ADEQ in 2017. Hayden Old Jail has the longest length of record, with data collected during the period of 1975-1991, and then again from 1999-2017. The Miami Ridgeline monitor, which was removed in September of 2017, has the second longest length of record with data collected from 1999-2017. Both the Miami Jones Ranch and Miami Townsite monitors have been in operation for a period of five years, from 2013-2017. Monitoring for SO₂ began in the Phoenix area at JLG Supersite in 2005, which later became trace-level monitoring in 2011 as part of EPA's NCore program. The sensitivity of this monitor is far greater than the monitors used earlier, and low concentrations of SO₂ in the urban environments can be assessed with greater confidence. The average trends for a period of 5 years, 10 years, and length of record for each site are provided in Table 20 below.

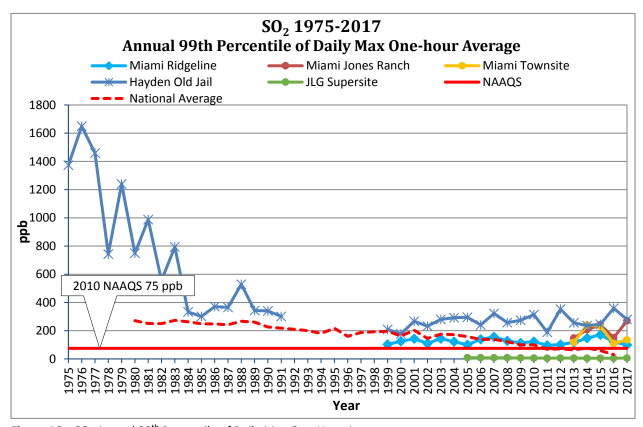


Figure 16 – SO₂ Annual 99th Percentile of Daily Max One-Hour Average Link to National Averages (https://www.epa.gov/air-trends/sulfur-dioxide-trends). 2017 National Averages were not available.

Note: Some years might not satisfy completeness criteria.

Table 20 – SO₂ One-Hour Average Trend

		Site Name									
Trend Length	Hayden Old Jail	JLG Supersite	Miami Jones Ranch	Miami Ridgeline	Miami Townsite						
5-Year (2013 - 2017)	28% 个 Increase	9% 个 Increase	45% 个 Increase	18% ↓ Decrease	20% ↓ Decrease						
10-Year (2008 - 2017)	11% 个 Increase	30% ↓ Decrease	N/A	9% 个 Increase	N/A						
Length of Record	25% ↑ Increase 1999-2017 92% ↓ Decrease 1975-1991	45% ↓ Decrease 2005-2017	45% 个 Increase 2013-2017	1% 个 Increase 1999-2017	20% ↓ Decrease 2013-2017						

4.5 Air Quality Index (AQI)

The SO_2 daily AQI values for 2017 are categorized into the different levels of health concerns in Table 21. Background levels of SO_2 are near zero in areas without manmade sources. Levels of SO_2 are at a maximum near industrial sources where SO_2 is directly related to production activity. Graphical representations of the SO_2 Daily AQI values for 2017 are shown in Figures 17 and 18.

Table 21 - SO₂ Daily AQI Count 2017

401	Levels of Health	Number of Days							
AQI Values	Concern	Hayden Old Jail	JLG Supersite	Miami Jones Ranch	Miami Ridgeline*	Miami Townsite			
0 - 50	Good	133	306	262	188	295			
51 - 100	Moderate	122	0	47	27	34			
101 - 150	Unhealthy for Sensitive Groups	95	0	31	16	18			
151 - 200	Unhealthy	15	0	5	0	1			
201 - 300	Very Unhealthy	0	0	0	0	0			
301 - 500	Hazardous	0	0	0	0	0			
	Missing	0	59	20	37	17			
	Total Days	365	365	365	268	365			

^{*}Miami Ridgeline removed in Sept 2017; 268 possible monitoring days.

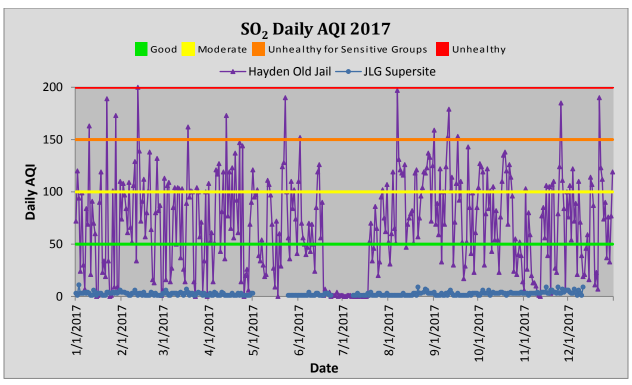


Figure 17 - SO₂ Daily AQI 2017

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data. The 2010 Final Rule for the Primary National Ambient Air Quality Standard for Sulfur Dioxide does not give breakpoints for the higher end of the AQI scale (from 200-500). Thus, an AQI of 200 is reported when concentrations are greater than 305 ppb.

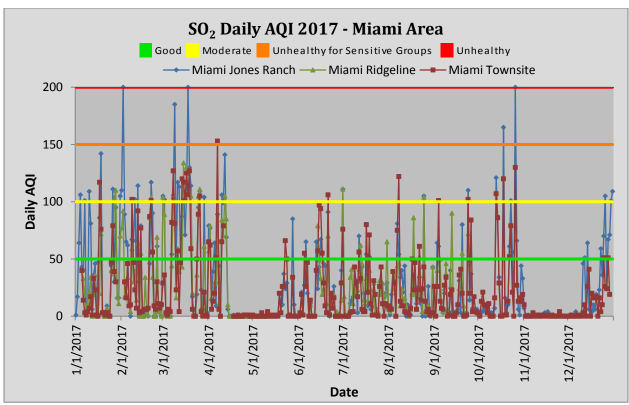


Figure 18 - SO₂ Daily AQI 2017 - Miami Area

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data. The 2010 Final Rule for the Primary National Ambient Air Quality Standard for Sulfur Dioxide does not give breakpoints for the higher end of the AQI scale (from 200-500). Thus, an AQI of 200 is reported when concentrations are greater than 305 ppb.

5.0 Particulate Matter Smaller Than 10 Microns (PM₁₀)

Particulate matter is a collective term describing very small solid or liquid particles that vary considerably in size, geometry, chemical composition, and physical properties. PM_{10} describes particles that are less than 10 microns in diameter. Produced by natural processes (pollen and wind erosion) and by human activity (soot, fly ash, and dust from paved and unpaved roads), particulates contribute to visibility reduction, pose a threat to public health, and cause economic damage through soil disturbances. The size, shape, and chemical composition of particulates determine their health effects. Particles from 2.5 to 10 microns are inhaled and deposited in the upper parts of the respiratory system. Epidemiological studies have shown causal relationships between particulates and excess mortality, aggravation of bronchitis, and small reversible changes in pulmonary function in children.

5.1 Background

PM₁₀ emissions in Arizona are mostly geological in origin and are dominated by dust from three activities: the constant grinding (re-entraining) of dust from paved roads, driving on unpaved roads, and earth moving associated with construction. On days with winds in excess of 15 miles per hour, wind erosion of soil may contribute to this loading. According the 2014 NEI, 55 percent of all PM₁₀ emissions come from dust; 19 percent from industrial processes; 10 percent from wildfires and wood burning; 9 percent from agriculture; 3 percent from mobile sources, including diesel trucks and off highway vehicles; 2 percent miscellaneous; and 2 percent from fuel combustion.

PM₁₀ concentrations are not spatially distributed evenly across Arizona because each monitoring site is strongly influenced by the degree of localized emissions of particulates. PM₁₀ maximum concentrations can occur in any season, provided nearby sources of coarse particulates are present or when strong and gusty winds suspend soil

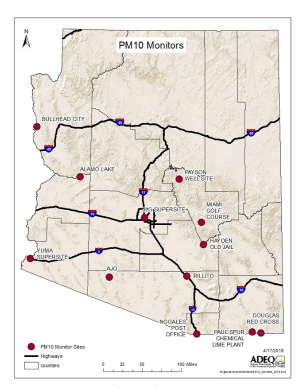


Figure 19 – Map of ADEQ's PM₁₀ sites

disturbed by human activities. Hourly concentrations of particulates tend to peak during those hours of the worst distribution, which is from sunset to midmorning.

Controls to reduce particulates in Arizona have been in place for decades, beginning in the 1960s with a Pima County ordinance that required watering to reduce dust from construction. Maricopa County's umbrella dust abatement rules, Rule 310 and 310.01, have been revised many times through the years and now regulates construction dust, trackout dust (accumulation on tires of vehicles) from construction sites, and dust from unpaved parking and vacant lots. Efforts to reduce dust re-suspended from paved

roads have concentrated on eliminating trackout from construction sites, curbing and stabilizing road shoulders, and investigating more efficient street sweepers. Additionally, techniques have been implemented such as paving of streets, chip sealing, and graveling alleyways. In Maricopa County, the Governor's Agricultural Best Management Practices Committee developed a rule containing best management practices for agricultural activities (AgBMP) to reduce particulate emissions from tilling and harvesting activities of cropland and non-cropland. In the Maricopa and Pinal County 2012 PM₁₀ SIP, the Maricopa Association of Governments (MAG) committed to implement new measures including enhanced enforcement of the county's dust rules and implementation of AgBMP.

Controls on PM_{10} have been successful in limiting anthropogenic emissions in recent years. Due to natural sources of PM_{10} in Arizona, areas of the state as shown in Appendix III- 2017 Area Designations, are in nonattainment for PM_{10} for the 1987 standard. PM_{10} is one of the major pollutants affecting the health of the people of Arizona and as such is a major part of ADEQ's ambient air monitoring. In 2017, ADEQ operated a network of 12 monitors throughout the state.

5.2 Monitoring Methods

Continuous monitoring for PM_{10} is done with Beta Attenuation Mass (BAM) monitor. This method of detection utilizes a radioactive beta source to measure the particles through a paper tape. The rate that a beta source decomposes is a known constant; therefore a sensor similar to a Geiger counter can detect this rate of radiation. Any changes to that rate can be recorded and this change in readings is a direct correlation to the concentration of PM. Particulate matter blocks or interferes with the beta radiation by absorbing or causing the beta radiation to change. Readings on a blank part of the paper tape are taken before sampling, then an airflow rate of 16.7 liters per minute deposits particles on the paper tape for about one hour. Afterward, a reading is taken on the tape where the particles are deposited to determine the concentration of PM. The volume of air sampled is also used to determine the concentration.

Readings from continuous types of instruments (BAM) are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis. Readings are retrieved by a data collection system and stored in a database.

5.3 Compliance/Summary of Design Values

In 2012, EPA revised the suite of standards for PM. EPA retained the 24-hour PM₁₀ standard of 150 μ g/m³ which has been in place since 1987 and revoked the annual PM₁₀ standard. The 24-hour PM₁₀ standard is met when the 24-hour average (rounded to the nearest 10 μ g/m³) does not exceed 150 μ g/m³ more than once per year on average over a three-year period. A history of the NAAQS for PM₁₀ is provided in Table 22 below.

Table 22 – History of the National Ambient Air Quality Standards for PM₁₀ during the period 1971-2012 (Source: USFPA TTN NAAOS)

(Source: USEPA I	Primary/	Indianter	Averaging	Lavial	Farms
Final Rule	Secondary	Indicator	Time	Level	Form
1971	Primary		24-hour	260 μg/m ³	Not to be exceeded more than
13/1	Timary	TSP			once per year
36 FR 8186			Annual	75 μg/m³	Annual Average
Apr 30, 1971	Secondary	TSP	24-hour	150 μg/m³	Not to be exceeded more than
ļ, -	,				once per year
1987			_	. 2	Not to be exceeded more than
2307	Primary and		24-hour	$150 \mu g/m^3$	once per year on average over
52 FR 24634	Secondary	PM ₁₀			a 3-year period
Jul 1, 1987	,		Annual	50 μg/m ³	Annual arithmetic mean,
,					averaged over 3 years
					Initially promulgated 99th
					percentile, averaged over 3
					years; when 1997 standards
1997			_		for PM ₁₀ were vacated, the
	Primary and		24-hour	150 μg/m³	form of 1987 standards
62 FR 38652	Secondary	PM ₁₀			remained in place (not to be
Jul 18, 1997	,				exceeded more than once per
00.1 20, 2007					year on average over a 3-year
					period)
			Annual	50 μg/m³	Annual arithmetic mean,
				1 0,	averaged over 3 years
2006					Not to be exceeded more than
	Primary and	PM ₁₀	24-hour	150 μg/m ³	once per year on average over
71 FR 61144	Secondary			1 3	a 3-year period
Oct 17, 2006					
2212	Primary and			1-0 / 2	Not to be exceeded more than
2012	Secondary	PM ₁₀	24-hour	$150 \mu g/m^3$	once per year on average over
	,				a 3-year period

Table 23 presents the 2014 to 2017 expected exceedance rates for the PM_{10} sites in Arizona, along with the annual maximum 24-hour concentration. The 24-hour primary PM_{10} NAAQS of 150 $\mu g/m^3$ has been exceeded at several sites in the 2014 to 2017 time period. Parts of the following counties are currently designated nonattainment with the 1987 PM_{10} NAAQS of 150 $\mu g/m^3$: Cochise, Pima, Santa Cruz, and Yuma. See Appendix III for the nonattainment area map.

Table 23 – PM₁₀ Compliance Summary

2015 to 2017 Maximum 24-Hour Average PM₁₀ **Compliance (in μg/m³)** Federal Equivalent Methods

(NAAQS 24-hour Average 150 μg/m³)

(NAAQ3 24-Nour Average 130 µg/N		15	20	16	20:	17	Thurs Vacu
Cita Nama	Max		Max		Max		Three-Year
Site Name	24-Hr	Exp.	24-Hr	Exp.	24-Hr	Exp.	Avg Exp. Rate of Exc.
	Avg	Exceed.	Avg	Exceed.	Avg	Exceed	Rate of Exc.
Cochise County							
Douglas Red Cross	89	0	236#	1.1	216	2	1.1*
Paul Spur Chemical Lime Plant	69	0	77	0	133	0	0
Gila County							
Hayden Old Jail	128	0	115	0	126	0	0
Miami Golf Course	46#	0	73	0	121	0	0*
Payson Well Site	62	0	58	0	70	0	0
La Paz County							
Alamo Lake	73	0	90	0	106	0	0
Maricopa County							
JLG Supersite	85	0	223	1	138	0	0.3
Mohave County							
Bullhead City	69	0	119	0	125	0	0
Pima County							
Ajo	67	0	141	0	109	0	0
Rillito	144#	0	192	2	166	2	1.3*
Santa Cruz County							
Nogales Post Office	107#	0	180	1	183	1	0.7*
Yuma County							
Yuma Supersite 400 5 523 9.7 379 4							
Preliminary Number of Sites in V	olation o	f the NAA	\QS				3

^{*}Indicates the data do not satisfy EPA's summary criteria, usually meaning less than 75 percent valid data recovery available in one or more calendar quarters.

Bold denotes exceedances and sites in violation of the 2012 NAAQS of 150 μ g/m³.

5.4 Trends

The analysis of trends in PM_{10} concentrations were divided into two different regions of Arizona: Southern Region, and Northern and Central Region. The division into regions helps to group sites that have similar PM_{10} sources and characteristics.

PM₁₀ Southern Region

Sites evaluated in the southern region of Arizona include Ajo, Douglas Red Cross, Nogales Post Office, Paul Spur Chemical Lime Plant, Rillito, and Yuma Supersite. Figure 20 illustrates the temporal variability of PM_{10} in the southern region over the 1987 to 2017 period in the form of annual means. The average trends for a period of 5 years, 10 years, and length of record for each site are provided in Table 24 below.

^{*}Preliminary Design Value since data completeness was not satisfied.

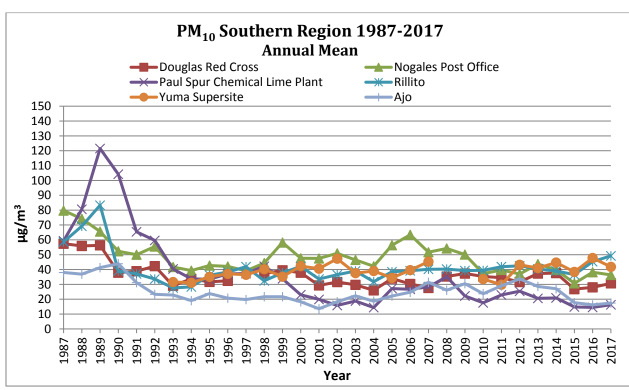


Figure 20 – PM₁₀ Annual Mean Average for the Southern Region **Note:** Some years might not satisfy completeness criteria.

Table 24 - PM10 Annual Mean Trend for the Southern Region

Table 24 – PIVITO Annual Mean Trend for the Southern Region								
	Site Name							
Trend Length Ajo Douglas Re Cross		Douglas Red Nogales Paul Spur Chemical Lime Plant		Chemical	Rillito	Yuma Supersite		
5 Year (2013 - 2017)	47% ↓ Decrease	25% ↓ Decrease	16% ↓ Decrease	29% ↓ Decrease	26% 个 Increase	4% 个 Increase		
10 Year (2008 - 2017)	39% ↓ Decrease	20% ↓ Decrease	31% ↓ Decrease	49% ↓ Decrease	14% 个 Increase	N/A		
Length of Record	31% ↓ Decrease 1987-2017	8% ↓ Decrease 1998-2017 52% ↓ Decrease 1987-1996	37% ↓ Decrease 1987-2017	94% ↓ Decrease 1987-2017	19% ↓ Decrease 1987-2017	33% 个 Increase 2010-2017 26% 个 Increase 1993-2007		

PM₁₀ Northern and Central Region

Sites evaluated in the northern region of Arizona include Bullhead City and Payson Well Site and in Central region include Alamo Lake, Miami Golf Course, Hayden Old Jail and JLG Supersite. Figure 21 illustrates the temporal variability of PM_{10} in the northern and central region over the 1990 to 2017 period in the form of annual means. The trends for a period of 5 years, 10 years, and length of record for each site are provided in Table 25 below.

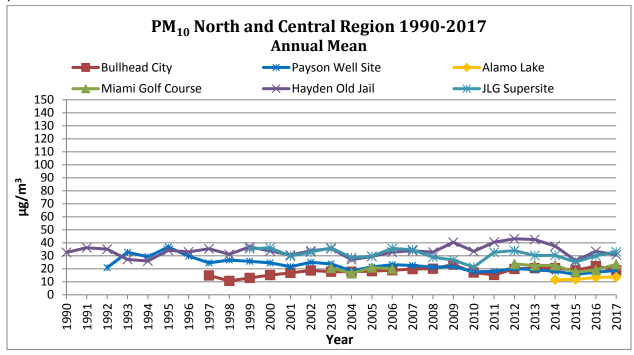


Figure 21 – PM₁₀ Annual Mean Average for the Northern and Central Region *Note:* Some years might not satisfy completeness criteria.

Table 25 – PM₁₀ Annual Mean Trend for the Northern and Central Region

-		ern Sites	Central Sites			
Trend Length	Bullhead City	Payson Well Site	Hayden Old Jail	JLG Supersite	Miami Golf Course	
5 Year	3% ↓	4% ↓	28% ↓	7% 个	3% ↓	
(2013 - 2017)	Decrease	Decrease	Decrease	Increase	Decrease	
10 Year	2% 个	17% ↓	16% ↓	14% 个	N/A	
(2008 - 2017)	Increase	Decrease	Decrease	Increase	N/A	
					11% ↓	
					Decrease	
Length of	45% 个	44% ↓	12% 个	14% ↓	2012-2017	
•	Increase	Decrease	Increase	Decrease		
Record	1997-2017	1992-2017	1990-2017	1999-2017	3% ↑	
					Increase	
					2003-2006	

5.5 Air Quality Index (AQI)

Similar to the analysis of trends in PM₁₀ concentrations, the representation of the AQI daily values for 2017 was divided into three different regions of Arizona: Southern Region (divided into South/Central and Southeastern Regions) and a combined Northern and Central Region.

Background levels of PM_{10} are very low in areas without manmade sources, but can be slightly elevated due to windblown particulates. High PM_{10} levels are influenced by localized emissions (disturbed soil, unpaved roads, and high wind events). High levels can occur during any season when particulates are picked up by strong and gusty winds. The highest PM_{10} AQI values occur when large dust storms form over central and southern Arizona. Wintertime AQI values can be elevated due to woodstove burning and stable atmospheric conditions.

PM₁₀ Southern Region

The PM_{10} daily AQI values for 2017 are categorized into the different levels of health concerns in Table 26. Graphical representations of the PM_{10} Daily AQI values for 2017 are shown in Figures 22 and 23.

Table 26 – PM₁₀ Daily AQI Count 2017 - Southern Region

		Number of Days						
AQI Values	Levels of Health Concern	Ajo	Douglas Red Cross	Nogales Post Office	Paul Spur Chemical Lime Plant	Rillito	Yuma Supersite	
0 - 50	Good	356	329	301	356	239	296	
51 - 100	Moderate	9	27	63	3	124	65	
101 - 150	Unhealthy for Sensitive Groups	0	2	1	0	2	2	
151 - 200	Unhealthy	0	0	0	0	0	1	
201 - 300	Very Unhealthy	0	0	0	0	0	1	
301 - 500	Hazardous	0	0	0	0	0	0	
	Missing	0	7	0	6	0	0	
	Total Days	365	365	365	365	365	365	

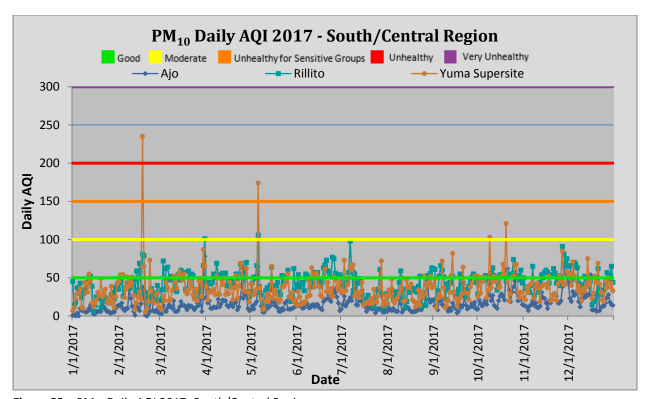
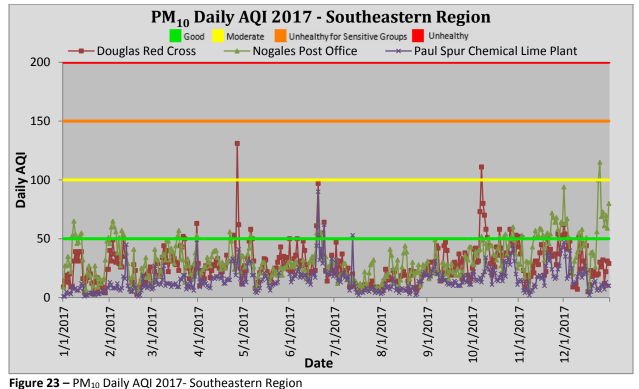


Figure 22 – PM₁₀ Daily AQI 2017- South/Central Region **Note**: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.



Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.

PM₁₀ Northern and Central Region

The PM_{10} daily AQI values for 2017 are categorized into the different levels of health concerns in Table 27. Graphical representations of the PM_{10} Daily AQI values for 2017 are shown in Figures 24 and 25.

Table 27– PM₁₀ Daily AQI Count 2017 - Northern and Central Region

AQI	Levels of Health	Number of Days							
Values	Concern	Alamo Lake	Bullhead City	Hayden Old Jail	JLG Supersite	Miami Golf Course	Payson Well Site		
0 - 50	Good	347	344	341	323	358	358		
51 - 100	Moderate	5	6	23	31	5	4		
101 - 150	Unhealthy for Sensitive Groups	0	0	0	0	0	0		
151 - 200	Unhealthy	0	0	0	0	0	0		
201 - 300	Very Unhealthy	0	0	0	0	0	0		
301 - 500	Hazardous	0	0	0	0	0	0		
	Missing	13	15	1	11	2	3		
	Total Days	365	365	365	365	365	365		

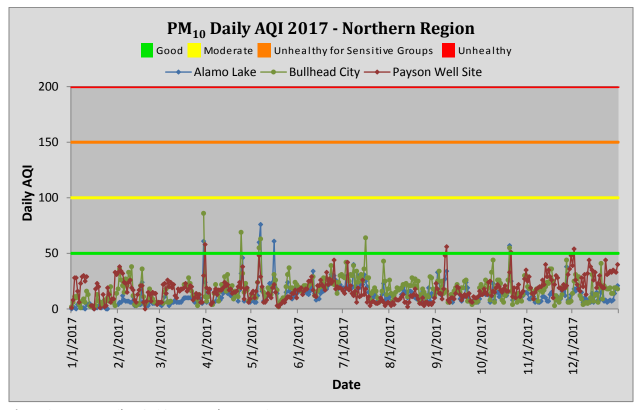


Figure 24 - PM₁₀ Daily AQI 2017 - Northern Region

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.

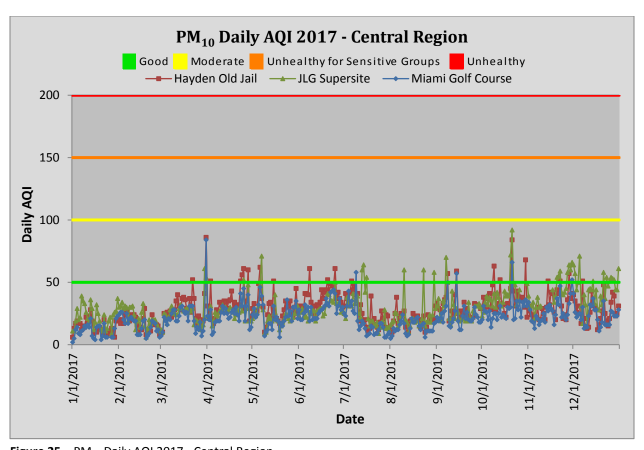


Figure 25 – PM₁₀ Daily AQI 2017 - Central Region **Note**: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.

6.0 Particulate Matter Smaller Than 2.5 Microns (PM_{2.5})

Particulate matter smaller than 2.5 microns (PM_{2.5}) describes particles that are less than 2.5 microns in diameter. These fine particulates are formed by the condensation of vapors or by their subsequent growth through coagulation or agglomeration. Others are emitted directly from sources, either by combustion or from mechanical grinding of soils. Fine particulates are classified as: primary - produced within and emitted from a source with little subsequent change; or secondary - formed in the atmosphere from gaseous emissions. For example, secondary particulate nitrates and sulfates form in the atmosphere from the oxidation of gaseous SO₂ and NO₂. In contrast, most atmospheric particulate carbon is primary, having been emitted directly from combustion sources, although some of the organic carbon in aerosols is secondary, having been formed by the complex photochemistry of gaseous VOCs.

For particulate matter, the primary route of entry into the body is inhalation. Particles smaller than 2.5 microns are respired and enter the pulmonary tissues where they are deposited. Particles in the size range of 0.1 to 2.5 microns are most efficiently deposited in the alveoli, where their effective toxicity is greater than larger particles because of the higher relative content of toxic heavy metals, sulfates, and nitrates. Health studies have shown a significant association between exposure to fine particles and premature death from heart or lung disease. Fine particles can aggravate heart and lung diseases and have been linked to effects such as: cardiovascular symptoms, cardiac arrhythmias, heart attacks, respiratory symptoms, asthma attacks, and bronchitis. Individuals that may be particularly sensitive to fine particle exposure include people with heart or lung disease, older adults, and children.

6.1 Background

With a more diverse chemical composition, fine particulate emissions are distributed among a larger number of sources. According the 2014 NEI, 34 percent of all PM_{2.5} emissions come from wildfires and wood burning; 24 percent from dust; 11 percent from industrial processes; 9 percent from mobile sources including diesel trucks and off highway vehicles; 8 percent from miscellaneous sources; 8 percent from fuel combustion; and 7 percent from agriculture.

Concentrations of PM_{2.5} tend to be at their highest in the central portions of urban areas, diminishing to background levels at the urban fringe. Background concentrations of PM_{2.5} are about 5 μ g/m³, in contrast to the urban maximum of 12 to 15 μ g/m³. Concentrations of fine particulates tend to be higher in



Figure 26 – Map of ADEQ's PM_{2.5} sites

the late fall and winter, when atmospheric dispersion is at a seasonal low, which traps the particulates and allows for a build-up in the ambient air.

Controls to reduce particulates in Arizona have been in place for decades, beginning in the 1960s with a Pima County ordinance that required watering to reduce dust from construction. Secondary fine particulates have been reduced by vehicle emission controls, which reduce precursor gases. For example, reducing gaseous HC emissions has led to reductions in ambient concentrations of secondary organic carbon. Additionally, techniques have been implemented that are similar to PM₁₀ controls for PM_{2.5} reduction such as paving of streets, chip sealing, and graveling alleyways.

Due to the successful nature of the controls on $PM_{2.5}$, most of Arizona is in attainment for both of the 2012 primary $PM_{2.5}$ NAAQS. Areas in nonattainment are shown in Appendix III- 2017 Area Designations. Since fires and blowing dust make up the majority of $PM_{2.5}$ in Arizona, levels can be elevated above the 24-hour standard and are beyond controls on anthropogenic sources.

6.2 Monitoring Methods

ADEQ is utilizing two methods for measuring PM_{2.5}: filter based and continuous instruments. The filter based method is a Federal Reference Method and is used by ADEQ to meet collocation requirements for QA purposes only. Only data monitored with the continuous Federal Equivalent Method are used for compliance purposes in this report.

Particulates can be monitored using a weighing and filter based method. This is done by pulling ambient air



Figure 27 – Nogales Post Office monitoring station

through a filter for 24 hours every sixth day, weighing the filter before and after sampling, and measuring the volume of air sampled. Pollutant concentrations are measured by the weight of pollutant within a standard volume of air, for example micrograms per cubic meter ($\mu g/m^3$). Weight is determined from an automated weighing system by ADEQ's contracted lab. The filter based instruments are fitted with different aerodynamic devices (inlets) to segregate particle size fractions. The particles can be segregated into two size ranges (less than 10 microns and less than 2.5 microns). This method provides one 24-hour concentration.

Another type of instrument called a Beta Attenuation Mass monitor (BAM), utilizes the same inlets as filter instruments to segregate particles. The instrument is the same as the PM_{10} BAM, with the difference being that the $PM_{2.5}$ instrument is fitted with an additional aerodynamic device to segregate particle size fractions greater than 2.5. See section 5.2 of this document.

Readings from continuous instrument, BAM are averaged into hourly, daily, monthly, quarterly, and yearly averages for data analysis. Readings are retrieved by a data collection system and stored in a database. Filter data are stored by air filter lab weighing equipment and uploaded to the database.

6.3 Compliance/Summary of Design Values

Effective December 14, 2012 the EPA changed the annual primary PM_{2.5} NAAQS from 15.0 μ g/m³ (set in 1997) to 12.0 μ g/m³, thereby strengthening this annual health standard. The annual PM_{2.5} standard is met when the three-year average (rounded to the nearest 0.1 μ g/m³) of annual means is less than or equal to 12.0 μ g/m³. This three-year average is determined by calculating the quarterly averages for each year to determine the calendar year average and then averaging the three years together.

The 24-hour primary PM_{2.5} NAAQS of 35 μ g/m³ was retained in December 2012. EPA originally issued this 24-hour standard in 2006 when they changed it from 65 μ g/m³ to 35 μ g/m³. The 24-hour standard is met when the three-year average (rounded to the nearest 1 μ g/m³) of the yearly 98th percentile value is less than or equal to 35 μ g/m³. A history of the NAAQS for PM_{2.5} is provided in Table 28 below.

Table 28 – History of the National Ambient Air Quality Standards for PM_{2.5} during the period 1997-2012

(Source: USEPA TTN NAAQS)

Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form
1997	Primary and	DNA	24-hour	65 μg/m³	98 th percentile, averaged over 3 years
62 FR 38652 Jul 18, 1997	Secondary	PM _{2.5}	Annual	15.0 μg/m³	Annual arithmetic mean, averaged over 3 years
2006	Primary and	DN4	24-hour	35 μg/m ³	98 th percentile, averaged over 3 years
71 FR 61144 Oct 17, 2006	Secondary	PM _{2.5}	Annual	15.0 μg/m ³	Annual arithmetic mean, averaged over 3 years
	Primary		Annual	12.0 μg/m³	Annual arithmetic mean, averaged over 3 years
2012	2012 Secondary		Annual	15.0 μg/m³	Annual arithmetic mean, averaged over 3 years
	Primary and Secondary		24-hour	35 μg/m ³	98 th percentile, averaged over 3 years

The annual primary PM $_{2.5}$ NAAQS of 12.0 $\mu g/m^3$ was met in 2017 by all five sites operated by ADEQ. See Table 29 for a summary of the annual three-year averages at these sites. In order to calculate the three-year average, there must be a designation of a primary monitor at each site. In the 2015 to 2017 time period, all primary monitors were continuous monitors, with samples collected every day (365 sample days in non-leap years).

Table 29 – PM_{2.5} Annual Average Compliance Summary

2015 to 2017 Annual Average PM _{2.5} Compliance (in μg/m³)								
Federal Equivalent Methods								
(NAAQS Annual Average 12 μg/m³)	(NAAQS Annual Average 12 μg/m³)							
Site Name 2015 2016 2017 Three-Year Average								
Cochise County								
Douglas Red Cross	5.2	3.8#	5.9#	5.0*				

2015 to 2017 Annual Average PM _{2.5} Compliance (in μg/m³) Federal Equivalent Methods (NAAQS Annual Average 12 μg/m³)								
Site Name	2015	2016	2017	Three-Year Average				
La Paz County								
Alamo Lake	1.1#	2.4	5.6	3.0*				
Maricopa County								
JLG Supersite	7.7	7.9	5.7	7.1				
Santa Cruz County								
Nogales Post Office	9.0	9.8	8.9	9.2				
Yuma Count								
Yuma Supersite	5.6#	8.3	8.5	7.5*				
Number of Sites in Violation of the NAAQS								

^{*}Indicates the data do not satisfy EPA's summary criteria, usually meaning less than 75 percent valid data recovery available in one or more calendar quarters.

In 2017, all five sites operated by ADEQ were in compliance with the 24-hour primary PM_{2.5} NAAQS of 35 $\mu g/m^3$. See Table 30 for a summary of the 24-hour three-year averages at these sites. Design calculations for this standard were also based on the designated primary monitor, as they were for the annual standard discussed above.

Table 30 – PM_{2.5} 24-Hour Average Compliance Summary

Thuas Voor
Three-Year
Average
11*
9*
21
28
19*
0

^{*}Indicates the data do not satisfy EPA's summary criteria, usually meaning less than 75 percent valid data recovery available in one or more calendar quarters.

^{*}Preliminary Design Value since data completeness was not satisfied.

^{*}Preliminary Design Value since data completeness was not satisfied.

6.4 Trends

Only monitors operated by ADEQ are used for the purpose of assessing PM_{2.5} trends in this report. Figures 28 and 29 illustrate the temporal variability of PM_{2.5} over the 1999 to 2017 period in the form of annual means and the 24-hour averages at the 98^{th} percentile. The average trends for a period of 5 years, 10 years, and length of record for each site are provided in Tables 31 and 32 below.

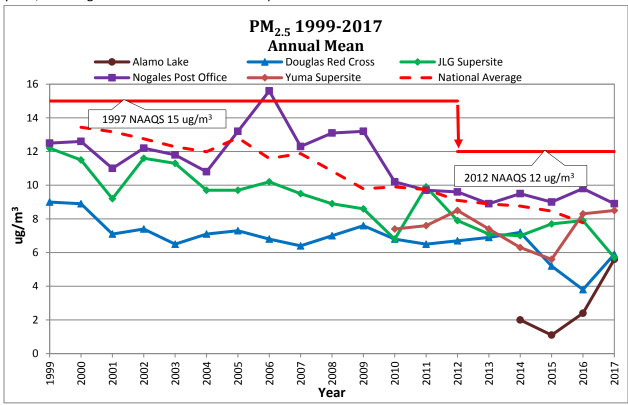


Figure 28 – PM_{2.5} Annual Mean Trend

Link to National Averages (https://www.epa.gov/air-trends/particulate-matter-pm25-trends). 2017 National Averages were not available.

Note: Some years might not satisfy completeness criteria.

Table 31 - PM_{2.5} Annual Mean Trend

	Site Name								
Trend Length	Length Alamo Lake* Douglas Red Cross JLG Supersite		JLG Supersite	Nogales Post Office	Yuma area				
5 Year (2013 - 2017)	N/A	31% ↓ Decrease	10% ↓ Decrease	1% 个 Increase	26% 个 Increase				
10 Year (2008 - 2017)	N/Δ		24% ↓ Decrease	31% ↓ Decrease	N/A				
Length of Record	378% ↑ Increase 2014-2017	32% ↓ Decrease 1999-2017	44% ↓ Decrease 1999-2017	30% ↓ Decrease 1999-2017	2% 个 Increase 2010-2017				

^{*}Background concentrations are monitored at this site. Baseline concentrations are still being established.

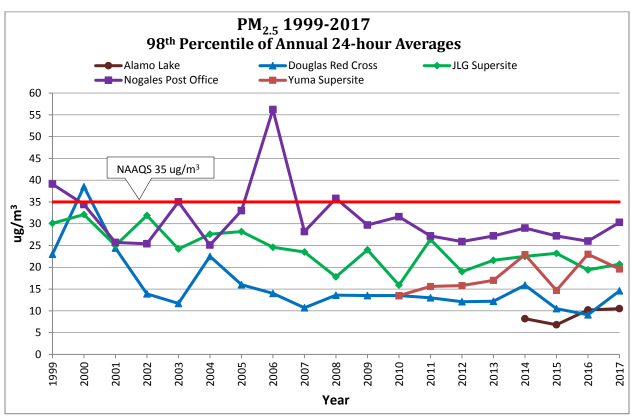


Figure 29 – PM_{2.5} 24-Hour Average Trend

Note: Some years might not satisfy completeness criteria.

Table 32 – PM_{2.5} 98th Percentile of Annual 24-hour Averages Trend

	Site Name								
Trend Length	Alamo Lake*	Douglas Red Cross	JLG Supersite	Nogales Post Office	Yuma area				
5 Year	N/A	6% ↓	9% ↓	5% 个	12% 个				
(2013 - 2017)	IN/A	Decrease	Decrease	Increase	Increase				
10 Year	N/A	11% ↓		16% ↓	N/A				
(2008 - 2017)	IN/A	Decrease	Increase	Decrease	IN/A				
Length of	42% 个	61% ↓	35% ↓	20% ↓	48% 个				
Record	Increase	Decrease	Decrease	Decrease	Increase				
Record	2014-2017	1999-2017	1999-2017	1999-2017	2010-2017				

^{*}Background concentrations are monitored at this site. Baseline concentrations are still being established.

6.5 Air Quality Index (AQI)

The PM_{2.5} daily AQI values for 2017 are categorized into the different levels of health concerns in Table 33. Background levels of PM_{2.5} are very low in areas without manmade sources. Urban background levels are elevated some due to industrial and mobile sources. Wintertime AQI values can be higher due to woodstove burning and stable atmospheric conditions. The highest PM_{2.5} AQI values occur when large dust storms form over central and southern Arizona and also when forest fires occur in the northern areas of Arizona. Graphical representations of the PM_{2.5} Daily AQI values for 2017 are shown in Figures 30 and 31.

Table 33 - PM_{2.5} Daily AQI Count 2017

AQI	Levels of Health	Number of Days						
Values			Douglas Red Cross	JLG Supersite	Nogales Post Office	Yuma Supersite		
0 - 50	Good	345	263	307	276	315		
51 - 100	Moderate	3	10	44	60	48		
101 - 150	Unhealthy for Sensitive Groups	0	0	1	2	2		
151 - 200	Unhealthy	0	0	0	2	0		
201 - 300	Very Unhealthy	0	0	0	0	0		
301 - 500	Hazardous	0	0	0	0	0		
	Missing	17	92	13	25	0		
	Total Days	365	365	365	365	365		

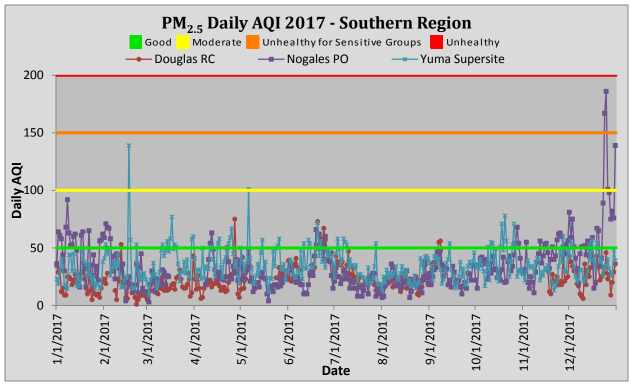
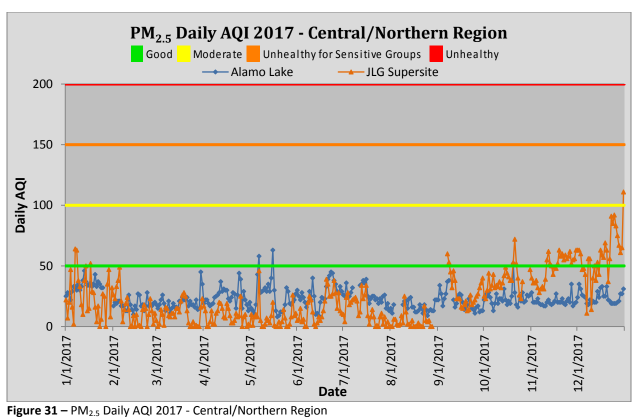


Figure 30 - PM_{2.5} Daily AQI 2017 - Southern Region

Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.



Note: Data are the daily AQI values for 2017 obtained from https://www.epa.gov/outdoor-air-quality-data and include Exceptional Events.

7.0 Lead (Pb)

Lead is a metal found naturally in the earth and a key element in the composition of many rocks and soils. It is a very dense metal that is used in a variety of industrial processes in many forms. It can be used as anodes in car batteries or as ballast in boats or in scuba diving. Lead as defined in terms of air quality, is a component of air particulates that can be inhaled into the lungs. Lead found in car batteries for instance is not considered an air pollutant. Today, the major sources of Pb emissions that pertain to air quality are ore and metals processing and piston-engine aircraft operating on leaded aviation gasoline. The highest ambient air concentrations of Pb in Arizona are usually found near copper smelters.

In addition to exposure to Pb in air, other major exposure pathways include ingestion of Pb in drinking water and Pb-contaminated food as well as incidental ingestion of Pb-contaminated soil and dust. Lead-based paint remains a major exposure pathway in older homes. However, data from these other exposure pathways are not measured by ADEQ Air Quality Division. Once taken into the body in any form, Pb distributes throughout the body in the blood and is accumulated in the bones. Depending on the level of exposure, Pb can adversely affect the nervous system, kidney function, immune system, reproductive and developmental systems, and the cardiovascular system. Lead exposure also affects the oxygen carrying capacity of the blood. The Pb effects most commonly encountered in current populations are neurological effects in children and cardiovascular effects in adults. Infants and young children are especially sensitive to even low levels of Pb, which may contribute to behavioral problems, learning deficits, and lowered IQ.

7.1 Background

Lead emissions in Arizona are made up of two main sources: mobile sources including piston-engine aircraft and other older engines which use leaded gasoline, and copper and metals smelting. According to the 2014 NEI, mobile sources contribute to about 65 percent of Pb emissions in Arizona, smelting and other industrial processes contribute about 27 percent, and all other sources including fuel combustion contribute about 8 percent.

Lead has historically been used in paints, as an additive to fuel, in electronics, and in various other industrial applications. U.S. Regulations have eliminated the use of Pb in almost all of these applications due to its toxicity.

Mobile sources are spread out over the state. There are around 100 small airports that still have some airplanes that require leaded-fuel, but since this is spread out over the whole state, no single airport contributes to a high concentration of Pb at a time. The only other large sources

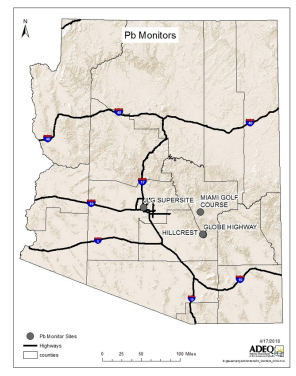


Figure 32 - Map of ADEQ's Pb sites

of Pb are due to metals smelting. Since Arizona has large copper smelting operations, Pb continues to be a concern in the areas surrounding these sources.

Controls to reduce Pb have been in place since its removal from gasoline in the 1970's. The controls that are being implemented to control the major Pb sources at the smelting operations include various fugitive dust control measures and particulate capture systems.

7.2 Monitoring Methods

Lead is monitored using two different sampling techniques. The first uses the same manual filter instruments that are used for PM₁₀ sampling as described in Section II.5. The same PM₁₀ inlet is used, and this method for using the PM₁₀ sampler is approved by the EPA as Pb-PM₁₀. The difference between sampling for Pb and sampling for PM₁₀ is that the filters are not pre-weighed or post weighed; the filters are analyzed by an independent laboratory for Pb content using an EPA approved method involving the digestion or dissolving of



Figure 33 - Hillcrest monitoring station

other contaminants and the isolation of Pb on the filter. Once the Pb is isolated, it can be weighed using a precise scale. Based on the total flow of the sample and the weight, a concentration is calculated.

The second method of lead sampling uses a total suspended particulates (TSP) sampler that does not have an inlet attached. All particles in the air are captured by the sampler on a large filter using a high flow rate. Once the 24-hour sampling period is over, the filter is cut, digested, and Pb is isolated in the same way as with the Pb-PM₁₀ sample filter.

Readings from all the types of instruments are averaged into daily, monthly, quarterly, and yearly averages for data analysis. Readings are retrieved on the samplers manually and by weighing equipment, then stored in a database.

7.3 Compliance/Summary of Design Values

In 2008, the Pb standard was revised from the 1978 rule. This lowered the three-month rolling average from 1.5 μ g/m³ to 0.15 μ g/m³. The rule also set forth a monitoring requirement based on an emissions threshold for each source. This threshold states that sources require monitoring if they are above 0.5 tons per year, based on their most recent emissions inventory. Lead concentrations in Arizona generally do not exceed this standard since leaded fuels were banned in the U.S. starting in 1996. Due to the 2008 rule change, ADEQ operated three source oriented monitors in 2017 around two copper smelters in Miami and Hayden. ADEQ also operated a PM₁₀ Speciated Metals monitor at JLG Supersite as part of the Chemical Speciation Network. Pb is one of the metals monitored for this network. A history of the NAAQS for Pb is provided in Table 34 below:

Table 34 – History of the National Ambient Air Quality Standards for Pb during the period 1978-2016 (Source: USEPA TTN NAAQS)

Final Rule	Primary/ Secondary	Indicator	Averaging Time	Level	Form			
1978 43 FR 46246 Oct 5, 1978	Primary and Secondary	Pb-TSP	Calendar Quarter	1.5 μg/m³	Not to be exceeded			
Feb 21,	Feb 21, 1991 – Agency released multimedia "Strategy for Reducing Lead Exposures"							
2008 73 FR 66964 Nov 12, 2008	Primary and Secondary	Pb-TSP	3-month period	0.15 μg/m³	Not to be exceeded			
2016 81 FR 71906 Oct 18, 2016	Primary and s	econdary st	andards retained	, without revisi	on.			

Table 35 summarizes the maximum three-month rolling average Pb concentrations for the year from 2015 to 2017. Globe Highway did not meet the 2017 NAAQS of 0.15 μ g/m³ with a 2017 Three-Year Design Value of 0.21 μ g/m³. The Hillcrest site began monitoring in 2016. The site was specifically located for Pb maximum concentration determination and chosen in response to elevated readings from an EPA Superfund monitor. There is no Three-Year Design Value for the Hillcrest site due to two years of data availability. However, the maximum three-month rolling averages for 2016 and 2017 exceeded the 0.15 μ g/m³ NAAQS, which will cause violations of the 2018 and 2019 Three-Year Design Values.

Table 35 – Pb Compliance Summary

2015 to 2017 Maximum 3 - Month Average Pb-TSP Compliance (in μg/m³)				
Bold denotes value above the standard.				
(NAAQS 3-Month Rolling Average 0.15 μg/m³)				
Site Name	Max 3-Month Rolling 24-Hr Avg			Three-Year
	2015	2016	2017	Design Value
Gila County				
Globe Highway	0.12	0.14	0.21	0.21
Hillcrest	N/A	0.22	0.28	N/A
Miami Golf Course	0.03	0.04	0.04	0.04
Maricopa County				
JLG Supersite	0.0094	0.0056	0.0059	0.0094
Number of Sites in Violation of the NAAQS				2

7.4 Trends

ADEQ began monitoring for Pb in 2011. These were source oriented monitors near two copper smelters in Miami and Hayden. In 2013 ADEQ began monitoring for Pb at JLG Supersite as part of NCore. Though Pb monitoring at JLG Supersite is no longer an NCore requirement, ADEQ continues to collect Pb data as a part of the National Air Toxics Trends Stations (NATTS) program. Figure 37 reflects data trends for the length or record for each monitor that was operated in 2017.

The average trends for a period of 5 years, 10 years, and length of record for each site are provided in Table 36 below.

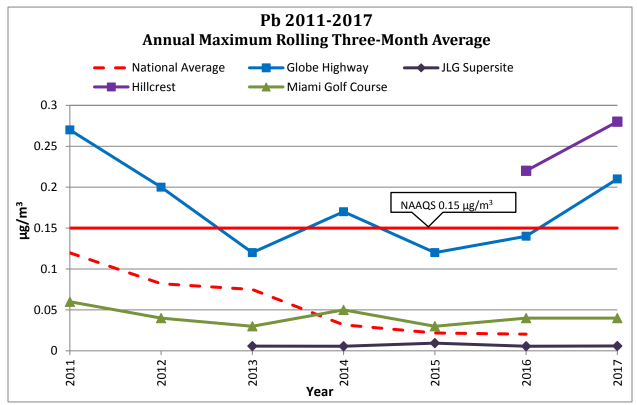


Figure 34 - Pb Three-Month Average

Link to National Averages (https://www.epa.gov/air-trends/lead-trends). 2017 National Averages were not available.

Table 36 – Pb Annual Maximum Rolling Three-Month Average Trend

	Site Name					
Trend Length	Globe Highway	JLG Supersite	Hillcrest	Miami Golf Course		
5 Year (2013 - 2017)	49% 个 Increase	1% 个 Increase	N/A	11% 个 Increase		
10 Year (2008 - 2017)	N/A	N/A	N/A	N/A		
Length of Record	31% ↓ Decrease 2011-2017	1% 个 Increase 2013-2017	27% 个 Increase 2016-2017	27% ↓ Decrease 2011-2017		

7.5 Air Quality Index (AQI)

The AQI scale is not established by EPA for Pb; therefore, there were no AQI values reported for Pb in 2017.

Section III - Non-Criteria Pollutants

This section covers the non–criteria pollutant networks monitored by ADEQ. Non-Criteria pollutants are those that do not have specific standards under the CAA but which can still pose a health threat. There are hundreds of air pollutants classified as non-criteria that are measured under different national programs. These include, but are not limited to, hazardous air pollutants (HAPs) sometimes referred to as air toxics, volatile organic compounds (VOC), heavy metal pollutants, particulate speciation, and many other contaminants and carcinogens. The following sections will discuss the Chemical Speciation Network (CSN), the National Air Toxics Trends Stations (NATTS), and the Photochemical Assessment Monitoring Station (PAMS) programs. Each subsection will provide a background for each program and a synopsis of current monitoring techniques. A trend analysis provides a history of select Non-Criteria pollutants monitored by ADEQ through 2017. The trend analysis includes a summary of the quantitative or compositional increase/decrease of the select pollutants over a period of 5 years, 10 years, and length of record.

1.0 Chemical Speciation Network (CSN)

PM_{2.5}, as described previously, are particles with a diameter less than 2.5 microns and are associated with respiratory and other health-related problems. These negative effects have been correlated to the total mass concentration in ambient air, but whether these negative effects are more correlated to total mass concentration or to the concentration of specific chemical species is not fully understood. To better understand and manage PM_{2.5}, EPA commissioned a companion network in 1997 called the CSN, in addition to the NAAQS PM_{2.5} network, to ascertain the chemical composition of fine particulates. Chemical speciation includes the identification and quantification of individual chemical elements, compounds, or classes of compounds that make up PM_{2.5} aerosols. The PM_{2.5} CSN targets analytes or species that are of particular interest to health officials, epidemiological researchers, and other interested parties.

1.1 Background

ADEQ supports the CSN network by serving as the operator and QA organization for a CSN site in Arizona at JLG Supersite.

The program objectives for the CSN are:

- to characterize annual and seasonal spatial characterization of aerosols
- air quality trends analysis
- tracking progress of control programs
- to compare the chemical speciation data set to the data collected from the IMPROVE network
- development of emission control strategies

The data generated from this network are not used for regulatory decisions concerning the $PM_{2.5}$ NAAQS, but may be used to supplement the $PM_{2.5}$ program.

1.2 Monitoring Methods

CSN instruments operate using the same principles as PM filter-based samplers. A filter is sampled on a specific date for a period of 24-hours, after which the sample is collected and sent to an EPA contracted lab for speciation analysis. EPA sets the schedule so that every CSN site in the U.S. is sampling on the same day. Currently, two different types of speciation samplers produce three 24-hour PM_{2.5} filter samples every three days at JLG Supersite. Samples are collected on three different filter types: Teflon, nylon, and quartz. Polytetrafluoroethylene (PTFE, commonly known as Teflon) filters are used to determine trace elements/crustal constituents that include metals, metalloids, and non-metals. Samples collected on nylon filters are used to determine the cations (sodium, potassium, and ammonium) and the anions (nitrate and sulfate). Samples collected on quartz filters are used to determine elemental and organic carbons.

1.3 Graphs

PM_{2.5} CSN characterization summaries for JLG Supersite are shown in Figures 35 and 36. Major elements show seasonal shifts, but organic carbon emerges as the dominant element throughout the year. Sulfates are more abundant during the spring and summer months as warmer temperatures favor the formation of particulate sulfate by the photochemical oxidation of SO₂. Nitrates are greater during the cooler months due to cooler temperatures favoring the formation of particulate nitrate by the gas-to-particle conversion of ammonium nitrate (NH₄NO₃). The crustal component of PM_{2.5} tends to increase during the months of April, May, and June. This is likely due to these months being the driest months for Arizona, leaving soil very dry and ready to become airborne with winds. Elemental carbon tends to be in higher abundance during the cooler months of year, most likely due to increased burning of biofuels for heat generation. Unidentified constituents are classified as "Other" and tend to increase in cooler months as well. This category exists because the analytical tests do not yield results for all possible species of PM_{2.5} aerosol. Analyzing for all species in PM_{2.5} would be impractical; however, the analytes selected are carefully chosen and often serve as indicators of other species. As more information concerning the chemical composition of fine particulates becomes available to researchers and regulators, the selected analytes may change as they did in early 2009 and 2016 as explained in the following section.

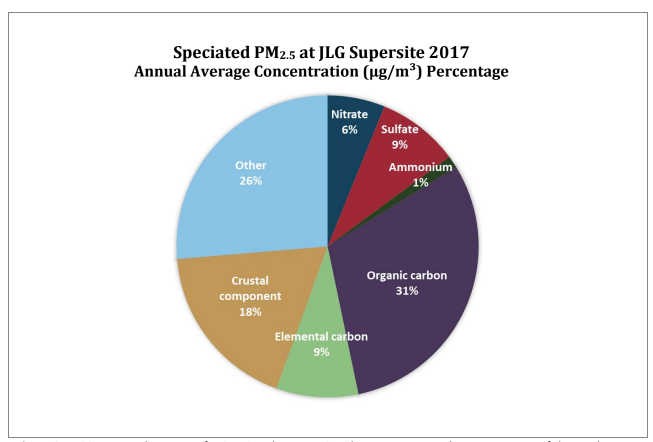


Figure 35 – 2017 Annual Averages for Speciated $PM_{2.5}$ major elements expressed as percentages of the total $PM_{2.5}$ concentration ($\mu g/m^3$) at JLG Supersite

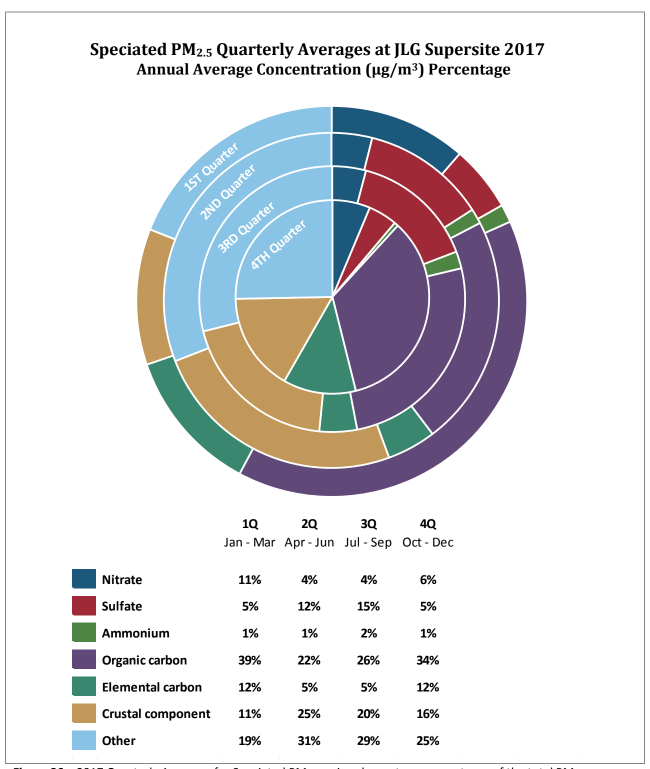


Figure 36 – 2017 Quarterly Averages for Speciated PM_{2.5} major elements as percentages of the total PM_{2.5} concentration ($\mu g/m^3$) at JLG Supersite

1.4 Trends

In 2009 the EPA switched the sampling and analytical methods for organic carbon and elemental carbon nationwide in order to improve data comparability. In 2016 the calculations for organic carbon and elemental carbon were adjusted nationwide in order to improve data comparability between IMPROVE and CSN data. The IMPROVE network tracks visual conditions in and around national parks and monuments, as well as in some urban areas. The organic carbon, elemental carbon, and "Other" concentrations reported from 2010 to 2017 are slightly more accurate and more comparable to national data than those reported prior to 2010, while these concentrations reported in 2016 and 2017 are more comparable to IMPROVE data than those reported prior to 2016. Due to the change in sampling and analytical methods, there was a noticeable decrease in the average organic carbon concentrations beginning in 2010. The percentages of each species compared to the total PM_{2.5} concentration from 2000 to 2017 are shown in Figure 37.

The average change in the percentage of total $PM_{2.5}$ for each major component for a period of 5 years, 10 years, and length of record are provided in Table 37 below.

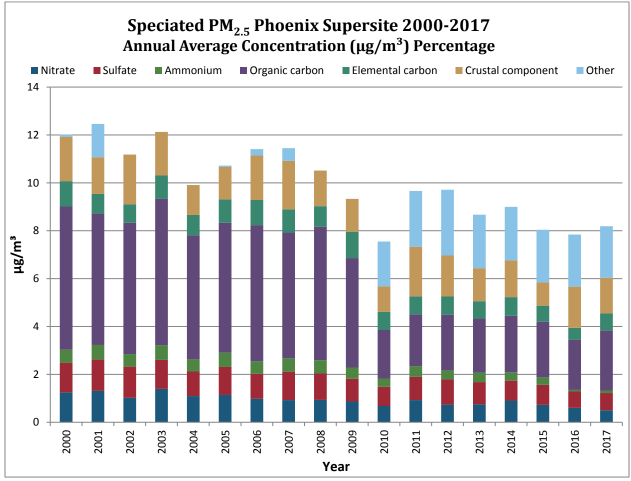


Figure 37 – Speciated PM_{2.5} Phoenix Supersite Annual Average Concentration **Note:** Some years might not satisfy completeness criteria.

Table 37 – Speciated PM_{2.5} Phoenix Supersite Annual Average Percentage of Total PM_{2.5} Trend

		Major Components of PM _{2.5} as Percentage of Total PM _{2.5}					
Trend Length	Nitrate	Sulfate	Ammonium	Organic Carbon	Elemental Carbon	Crustal Component	Other*
5 Year	31% ↓	17% ↓	80% ↓	14% 个	6% ↓	25% 个	6% 个
(2013 - 2017)	Decrease	Decrease	Decrease	Increase	Decrease	Increase	Increase
10 Year	23% ↓	19% ↓	70% ↓	49% ↓	25% ↓	17% 个	274% 个
(2008 - 2017)	Decrease	Decrease	Decrease	Decrease	Decrease	Increase	Increase
Length of Record (2000-2017)	29% ↓ Decrease	13% ↓ Decrease	50% ↓ Decrease	57% ↓ Decrease	0% 个 Increase	23% 个 Increase	830% 个 Increase

^{*}Updates to CSN sampling and analytical methods have improved the accuracy of the Organic Carbon and Elemental Carbon, which has significantly affected the percentage of the 'Other' component.

2.0 National Air Toxics Trends Stations (NATTS)

Air toxics, also known as HAPs, are compounds or elements known or suspected to cause cancer or other serious health effects, such as reproductive, neurological, birth defects, or adverse environmental effects. There are currently 189 HAPs regulated under the CAA. Examples of HAPs include benzene, which is found in gasoline; perchlorethlyene, which is emitted from some dry cleaning facilities; and methylene chloride, which is used as a solvent and paint stripper by a number of industries. Examples of other known air toxics include chloroform, formaldehyde, and metals such as cadmium, chromium, and Pb compounds.

2.1 Background

ADEQ monitors many HAPs including VOCs, carbonyls, and PM_{10} metals at JLG Supersite as part of the NATTS program. NATTS is a national network of 27 monitors that was developed to fulfill the need for long-term, high-quality toxics data to estimate national toxics averages. The principle purpose of the NATTS program is tracking toxics trends to assess progress towards emission and risk reduction. The goal is to reduce air toxics emissions by 75% from 1993 levels to significantly reduce the potential for human health risk. ADEQ entered the NATTS program in 2003 but had been monitoring some of the toxics in prior years. With the minimum pollutants monitored being met, ADEQ monitors a total of 104 HAPs.

The UATMP monitoring program began in 1987 to characterize the magnitude and composition of urban air pollution through monitoring in various cities across the U.S. Urban air pollution consists of many components from a wide range of industrial, motor vehicle, and biogenic and natural sources, some of which are toxics. ADEQ monitored VOCs at the South Phoenix site as part of the EPA's UATMP from 2001-2003 and started monitoring again in 2007. As stated earlier in this report, UATMP data are directly comparable to NATTS data as the monitoring methods and selected analytes are identical, and many of the program objectives overlap.

2.2 Monitoring Methods

To monitor VOCs, ambient air is captured in special evacuated canisters at a constant flow rate for a 24-hour period every sixth day in accordance with the EPA monitoring schedule. The canisters are collected and sent to a laboratory for analysis by a standard procedure (TO-15) appropriate for VOCs, and the results are reported to ADEQ for review before submittal to EPA.

For carbonyls, ambient air is drawn through small cartridges at a constant flow rate for a 24-hour period every sixth day in accordance with the EPA monitoring schedule. The cartridges are made from a high purity silica adsorbent coated with 2,4-dinitrophenylhydrazine (DNPH) which traps carbonyls (e.g. formaldehyde). The cartridges are sent to a laboratory for analysis using a standard procedure (TO-11A) appropriate for carbonyls analysis. The results are reviewed by ADEQ before submittal to EPA.

PM₁₀ metals are collected by drawing ambient air through a standard 47mm filter at a 16.7 LPM for 24-hours, as is done for PM measurements discussed in Section II of this report, every sixth day in accordance

with the EPA monitoring schedule. The filters are sent to a laboratory and processed using a standard procedure (TO-3.5) that is appropriate for metals analysis. The results are reviewed by ADEQ before submittal to EPA.

To sample polycyclic aromatic hydrocarbons (PAH), ambient air is drawn at a high-volume flow rate through a filter and sorbent cartridges containing polyurethane foam (PUF). PUF absorbs PAHs at a high collection efficiency. Approximately 300 m³ of air is drawn through in a 24-hour sample run. Runs occur every sixth day in accordance with the EPA monitoring schedule. The filter and cartridge are sent to a lab for analysis by a standard procedure (TO-13A) appropriate for PAHs. PAHs are analyzed via a Gas Chromatograph and Mass Spectrometer. Using the total air sampled, the concentrations of PAHs in the air sample are found and reported to ADEQ before submittal to EPA.

2.3 Tables

The EPA has not issued monitoring compliance criteria for air toxics (as it has for O₃, for example). Instead it addresses control issues through rules covering emissions from industrial sources such as chemical plants as well as smaller sources, such as dry cleaners and chromium electroplating facilities. As can be seen in Tables 38 and 39, air toxics concentrations at JLG Supersite and South Phoenix are comparable to national averages with some values being higher, and some lower, but none with extreme differences. For the pollutants that are in common between the two sites (i.e., VOCs), the summary data are very similar, which is an indication that the sites do a reasonable job of representing ambient air and are not significantly affected by nearby sources. These data also indicate that in comparison to the 2014 National Averages, manganese may be of local concern as the 2017 annual average exceeds the 2014 national average by a factor of 2.4. Air Toxics is a relatively new field within ambient air monitoring, and these data will continue to be collected to provide more information about urban air pollution.

Table 38 – Air Toxics Data for JLG Supersite

2017 Air Toxics data for JLG Supersite								
			Quar	terly Avera	ige			
Pollutant	Q1	Q2	Q3	Q4	Annual	National (2014)		
VOC (ppb)								
1,3-Butadiene	0.14	0.04	0.03	0.16	0.09	0.05		
Acrolein	0.75	0.57	0.55	0.53	0.61	0.34		
Benzene	0.48	0.19	0.18	0.50	0.33	0.23		
Carbon Tetrachloride	0.10	0.10	0.10	0.10	0.10	0.10		
Chloroform	0.06	0.07	0.05	0.08	0.07	0.04		
Tetrachloroethylene	0.06	0.01	0.02	0.09	0.04	0.02		
Trichloroethylene	0.00	0.00	0.00	0.00	0.00	0.01		
Aldehydes (ppb)								
Acetaldehyde	1.56	1.41	1.46	2.14	1.63	1.00		
Formaldehyde	2.92	3.50	4.59	4.14	3.78	2.30		
PAH(ng/m³)								
Benzo[A]Pyrene (Tsp) STP	0.17	0.01	0.02	0.09	0.06	0.11		
Naphthalene (Tsp) STP	79.64	34.91	32.30	97.19	56.33	66.5		
PM ₁₀ Metals (ng/m³)								
Arsenic	0.96	0.59	0.46	0.97	0.75	0.61		
Beryllium	0.02	0.04	0.03	0.05	0.03	0.01		
Cadmium	0.13	0.07	0.06	0.15	0.10	0.20		
Chromium	2.12	1.72	1.31	3.86	2.27	4.86		
Lead	4.19	2.50	1.98	5.50	3.56	2.93		
Manganese	14.48	17.81	14.72	29.21	19.20	8.02		
Nickel	1.52	1.47	1.14	1.92	1.52	1.11		

Table 39 – Air Toxics Data for South Phoenix

2017 Air Toxics data for South Phoenix site								
	Quarterly Average VOC (ppb)							
Pollutant	Q1	Q2	Q3	Q4	Annual	National (2014)		
1,3-Butadiene	0.13	0.04	0.04	0.16	0.09	0.05		
Acrolein	0.53	0.40	0.53	0.45	0.47	0.34		
Benzene	0.51	0.28	0.76	0.63	0.55	0.23		
Carbon Tetrachloride	0.10	0.10	0.10	0.10	0.10	0.10		
Chloroform	0.03	0.04	0.03	0.06	0.04	0.04		
Tetrachloroethylene	0.03	0.01	0.01	0.06	0.03	0.02		
Trichloroethylene	0.00	0.00	0.00	0.00	0.00	0.01		

2.4 Trends

The VOC trends include data from the JLG Supersite and South Phoenix (SP) sites. Seven VOC species were selected as analytes of interest and are shown in Figures 38 and 39. Two aldehyde species and two PAH species were also selected as analytes of interest and are shown in Figures 40, 41, and 42. These species were chosen due to their high frequency of detections in Arizona and nationwide. South Phoenix VOC concentrations are unavailable from 2005-2006 because ADEQ did not monitor for the UATMP during that time period. ADEQ began monitoring for Acrolein at JLG Supersite and South Phoenix in 2007.

The average trends for a period of 5 years, 10 years, and length of record for the analytes of interest are provided in Tables 40, 41, and 42 below.

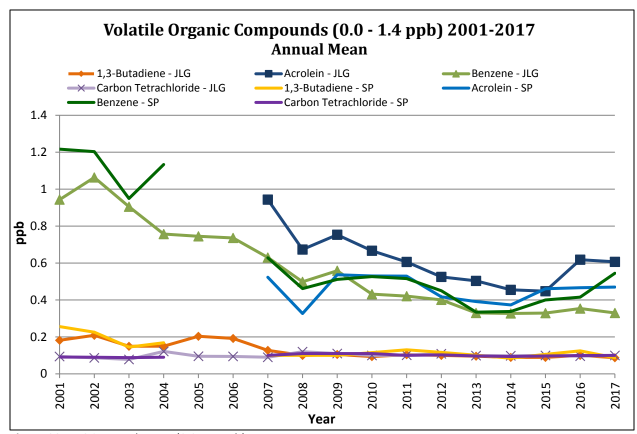


Figure 38 – VOC Annual Mean (0.0-1.4 ppb)

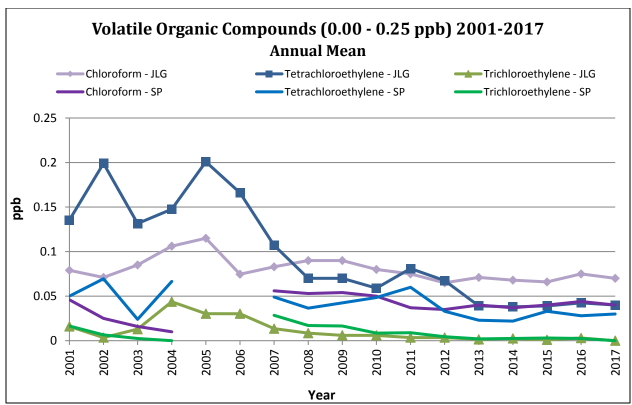


Figure 39 - VOC Annual Mean (0.00-0.25 ppb)

Note: Some years might not satisfy completeness criteria.

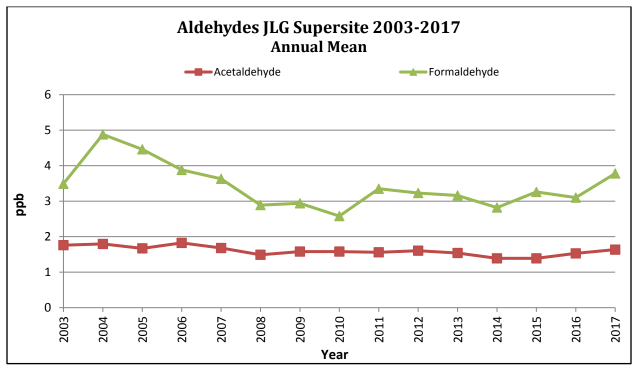


Figure 40 – Aldehydes Annual Mean

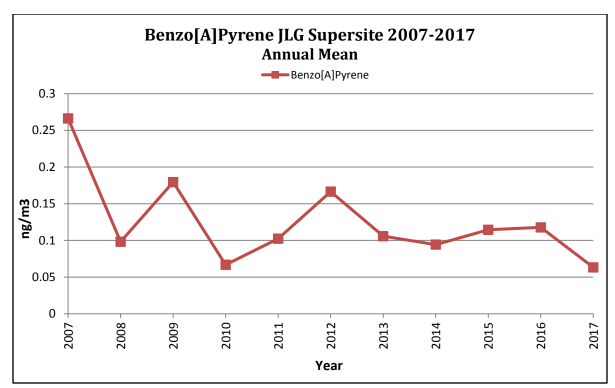


Figure 41 – Benzo[A]Pyrene Annual Mean

Note: Some years might not satisfy completeness criteria.

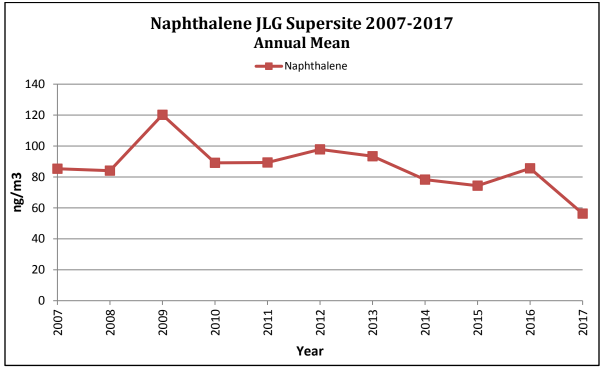


Figure 42 – Naphthalene Annual Mean

Table 40 – JLG VOC Annual Mean Trend

	JLG Supersite VOC Analytes							
Trend Length	1,3- Butadiene	Acrolein	Benzene	Carbon Tetra- chloride	Chloroform	Tetrachloro- ethylene	Trichloro- ethylene	
5 Year	2% ↓	33% 个	3% 个	0% 个	3% 个	7% 个	48% ↓	
(2013 - 2017)	Decrease	Increase	Increase	Increase	Increase	Increase	Decrease	
10 Year	12% ↓	25% ↓	41% ↓	15% ↓	24% ↓	53% ↓	99% ↓	
(2008 - 2017)	Decrease	Decrease	Decrease	Decrease	Decrease	Decrease	Decrease	
Length of	62% ↓	40% ↓	78% ↓	8% 个	25% ↓	89% ↓	106% ↓	
Record	Decrease	Decrease	Decrease	Increase	Decrease	Decrease	Decrease	
Record	2001-2017	2007-2017	2001-2017	2001-2017	2001-2017	2001-2017	2001-2017	

Table 41 – South Phoenix VOC Annual Mean Trend

Table 41 – South Phoenix VOC Annual Mean Trend							
			South P	hoenix VOC A	nalytes		
Trend Length	1,3- Butadiene	Acrolein	Benzene	Carbon Tetra- chloride	Chloroform	Tetrachloro- ethylene	Trichloro- ethylene
5 Year	5% 个	26% 个	65% 个	5% 个	7% 个	34% 个	57% ↓
(2013 - 2017)	Increase	Increase	Increase	Increase	Increase	Increase	Decrease
10 Year	7% ↓	1% ↓	14% ↓	13% ↓	26% ↓	43% ↓	111% ↓
(2008 - 2017)	Decrease	Decrease	Decrease	Decrease	Decrease	Decrease	Decrease
	3% ↓		25% ↓	10% ↓	31% ↓	47% ↓	116% ↓
	Decrease		Decrease	Decrease	Decrease	Decrease	Decrease
Length of	2007-2017	7% ↓	2007-2017	2007-2017	2007-2017	2007-2017	2007-2017
Record		Decrease					
	41% ↓	2007-2017	13% ↓	2% ↓	84% ↓	2% 个	111% ↓
	Decrease		Decrease	Decrease	Decrease	Increase	Decrease
	2001-2004		2001-2004	2001-2004	2001-2004	2001-2004	2001-2004

Table 42 – JLG Carbonyl and PAH Annual Mean Trend

	JLG Supersite Carbonyl and PAH Analytes						
Trend Length	Acetaldehyde	Formaldehyde	Benzo[A]Pyrene	Naphthalene			
5 Year	9% 个	21% 个	22% ↓	29% ↓			
(2013 - 2017)	Increase	Increase	Decrease	Decrease			
10 Year	2% ↓	21% 个	25% ↓	32% ↓			
(2008 - 2017)	Decrease	Increase	Decrease	Decrease			
Length of	17% ↓	9% 个	56% ↓	27% ↓			
Record	Decrease	Increase	Decrease	Decrease			
	2003-2017	2003-2017	2007-2017	2007-2017			

3.0 Photochemical Assessment Monitoring Stations (PAMS)

The PAMS are a collection of nationwide monitoring stations that are used to obtain comprehensive and representative data on O_3 air pollution. Section 182(c)(1) of the 1990 Clean Air Act (CAA) Amendments requires the Administrator to promulgate rules for enhanced monitoring of O3 that includes concurrent monitoring of O_3 , oxides of nitrogen (NO_x), total reactive nitrogen (NO_y), speciated volatile organic compounds (VOC), carbonyls, CO, and meteorology. The principal reasons for requiring the collection of additional ambient air pollutants and meteorological data are the widespread nonattainment of the O_3 NAAQS and the need for a more comprehensive air quality database for O_3 and its precursors. EPA issued a final rule for a reengineering of the PAMS program in October 2015 as part of the 2015 O_3 NAAQS Revision. In 2017, ADEQ collected carbonyl samples during PAMS season (June through August) according to the new reengineered PAMS program specifications at JLG Supersite. 2016 was the final year of operating PAMS at Queen Valley. ADEQ will continue to operate the PAMS program under the new rule at JLG Supersite, which is collocated with the JLG Supersite NCore site as required.

3.1 Background

High O_3 concentrations are caused when sunlight and precursor pollutants react in the lower atmosphere. Higher O_3 levels typically occur in the summer months when the sun angle is higher than during the winter months. This is partially due to the sun being out for a longer period during the summer than the winter. The onset of the ozone season in Arizona is usually observed during early spring. The PAMS monitoring season is during this high concentration time from June-August, which requires many more samples to be taken in order to characterize peak levels of O_3 and O_3 precursor pollutants. The sources for O_3 precursor pollutants include combustion vehicles, biogenics from plants, industrial processes, and electric power plants. Ozone conditions and typical concentrations in Arizona are described in more detail in Section II of this report.

ADEQ maintains the PAMS network in the metropolitan statistical area (MSA) of Phoenix-Mesa-Scottsdale. The first PAMS sites began operation in 1994, and as such, have been in operation for over 15 years. Many changes have occurred during those times that have changed the O₃ problem in the US as well as EPA's understanding of it. As part of the 2015 NAAQS for O₃, the PAMS program was re-engineered with new national objectives, network design, and measurement technologies. ADEQ will continue to operate the PAMS program in the Phoenix-Mesa O₃ nonattainment area. The monitoring includes a required PAMS station at JLG Supersite that will be fully operational by June 2019.

3.2 Monitoring Methods

Methods for monitoring pollutants that are part of PAMS are the same as with other networks. PAMS O_3 is monitored using the same instruments and ultraviolet absorption method as described in Section II of this report.

VOCs and Carbonyls use the same instruments and methods as the air toxics network. During the 2017 PAMS season, three eight-hour Carbonyls samples were collected in addition to the 24-hour NATTS

samples. These samples were collected in order to better characterize precursor pollutant concentrations during the day. Peak levels generally occur during the morning rush hour; therefore, samples of Carbonyls were collected in three eight-hour blocks from 12:00 am to 8:00 am, 8:00 am to 4:00 pm, and 4:00 pm to midnight during PAMS season.

VOCs for PAMS were not measured in 2017 due to PAMS re-engineering changes from the 2015 NAAQS for O₃. ADEQ will measure the VOCs for PAMS in 2018. Starting in 2019, ADEQ will be measuring hourly VOCs, minimally from June – August, using an auto gas chromatograph instrument rather than collecting an air sample in an evacuated cylinder.

Nitrogen oxides and NO_y monitoring use the same chemiluminescence instruments and methods as the NO_2 network. Reactive nitrogen oxides differ from NO_x due to the sampling height. Since O_3 is a scavenger of some oxides of nitrogen, the sample inlet is located above ground level O_3 (~ 10 m). The reactive forms of nitrogen that are normally scavenged by O_3 are measured at this elevated height to give a better representative value of the total nitrogen precursor pollutants.

PAMS CO is monitored using the same nondispersive infrared instruments and methods as described in Section II of this report.

All readings from continuous gas analyzers are averaged into hourly, daily, quarterly, and yearly averages for data analysis. Readings are retrieved by a data collection system and stored in a database. Readings from sample monitors (VOCs and Carbonyls) are sent to ADEQ by the analysis lab and are also averaged for data analysis.

3.3 Trends

Total Non-Methane Organic Compounds (TNMOC) are defined in the PAMS TAD as the unspeciated total concentration of VOC (C2 through C12) in ambient air as determined by "summation of peaks" from GC/FID analysis, expressed in parts per billion carbon (ppbC). Parts per billion carbon is the concentration (in ppb) of the compounds multiplied by the number of carbon molecules in the compound. This unit is useful as it gives the concentration of the individual carbon molecules available to react to form O₃. TNMOC are a precursor to O₃ and emission sources include fossil fuel burning, landfills, and solvents. The annual means used for trend analysis were calculated from 24-hour samples taken during the June 1st through August 31st PAMS sampling season of each year. However, due to PAMS re-engineering, TNMOC samples were not collected in 2017. The TNMOC concentrations from 2007-2016 are shown below in Figure 43.

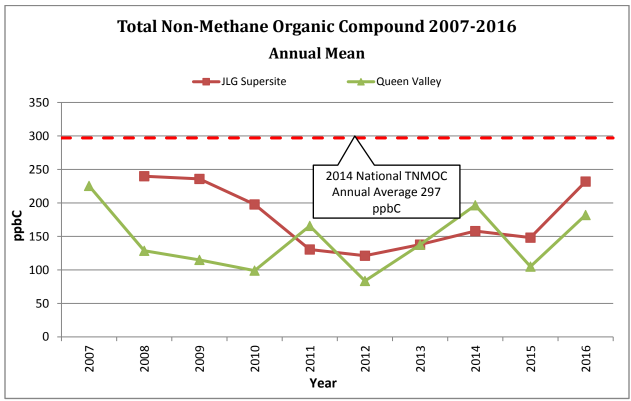


Figure 43 - TNMOC Annual Mean

Section IV – Visibility

Visibility has historically been defined as the maximum distance that a human eye can view a contrasting object. This is a technical definition that also can include other quantitative measurements. Defining visibility does not always lend itself to the quantitative measures that many instruments can produce. Visibility can also be based on the judgments of an observer viewing a beautiful vista. It is the combination of both the technical and the judgmental definition that most closely describes visibility.

Visibility and haze are two terms that are closely related, as haze is the form of air pollution that degrades visibility. Haze is caused when sunlight encounters tiny particles in the air, which reduces the color and clarity of what is seen. Since 1988, the U.S. EPA, States, and Federal land management agencies have conducted monitoring of air pollution and visibility impairment at a number of national parks and wilderness areas



Figure 44 – Pleasant Valley monitoring station.

across the U.S. In 1999, the EPA announced a major effort to improve air quality in national parks and wilderness areas. The Regional Haze Rule (RHR) calls for the improvement of visibility in 156 Class I national parks and wilderness areas throughout the U.S. (formally known as mandatory Federal Class I areas). The RHR outlines the requirements for states to follow in order to address haze impairing pollutants

1.0 Urban Haze

Haze not only affects natural vistas, but can also affect urban skylines. Reducing urban haze mirrors the objectives of the RHR in improving visibility in the urban environment by reducing emissions that directly contribute to haze. House Bill 2538 of the Forty-fifth AZ State Legislature states that ADEQ "shall establish a daily visibility index to be used in evaluating and reporting current visibility conditions and progress toward visibility improvement goals." Pursuant to this bill, ADEQ has established an urban visibility network in the Phoenix area.

1.1 Program Background

ADEQ operates a network of urban visibility instruments that are designed to characterize different optical phenomena in the Phoenix metropolitan area. This network measures the amount of visibility impairing haze using different optical measurements which show the amount of light scattered from one point to another. ADEQ operates several instruments as part of its Urban Haze program including one

transmissometer located in downtown Phoenix, one nephelometer located near central Phoenix, and five high resolution digital cameras in various locations throughout the Phoenix metropolitan area.

1.2 Monitoring Methods

The transmissometer is used to assess visibility impairment by measuring the amount of light lost over a known distance. The instrument consists of a light source (transmitter) and light detector (receiver) which are generally located on an elevated surface, such as a tall building, and are separated by a distance of 1-3 miles. The Phoenix transmitter is located on the roof of the Phoenix Baptist Hospital and the receiver is 2.96 miles away on the roof of the Holiday Inn on the corner of Osborn and 3rd Avenue in downtown Phoenix. The transmitter emits a uniform light beam of constant intensity that is carefully aimed at the receiver. The amount of light transmitted and received is precisely measured. The receiver includes a telescope that gathers the transmitter's light and a computer that compares the measured light intensity with the known transmitter light intensity to calculate the transmission of the intervening atmosphere. The measured transmission can be related to the light lost along the path due to scattering and absorption.

The nephelometer is used to assess visibility impairment by estimating the particle scattering coefficient at a point location. The nephelometer provides a direct measurement of the light scattered by aerosols and gases in a sampled air volume. It employs a light source and a detector set to the side of the source. Light is scattered by particles over a large range of scattering angles, in a defined band of visible wavelengths. The detector picks up the specific wavelengths of scattered light to the side. These wavelengths are particular to a certain type of particle, $PM_{2.5}$. Because the total light scattered out of a path is the same as the reduction of light along a path due to scattering, the integrating nephelometer gives a direct estimate of the particle scattering coefficient and hence impaired visibility.

Photographic documentation is an important aspect of evaluating visibility. Photography is an effective way to document events and trends on a media that is easily interpreted. ADEQ uses high-resolution digital images from locations around Phoenix to document visibility conditions. The digital images are readily available for viewing at www.phoenixvis.net and can be conveniently distributed via the Internet, easily stored, managed, and duplicated without degradation. Each site consists of a high-resolution digital camera housed in a weatherproof, environmental enclosure, and a supporting image capture computer. Digital images are captured every 5 minutes, stored on the system's internal hard drive, and uploaded to the website every 15 minutes.

1.3 Trends

Visibility data from these monitors can be expressed by several different measurement units: deciview, inverse megameters, and visual range. An inverse megameter (Mm⁻¹) (units used by ADEQ) is a representation of the ratio between how much light is not received by a sensor compared to the amount of light that leaves a source. Higher numbers mean worse visibility. As an Mm⁻¹ is an uncommon unit of measure, the visual range will be reported.

Optical measurements of urban visibility have been made continuously in Phoenix since 1994. Light extinction, the degree to which light is reduced by its interaction with particles and gases in the atmosphere, is measured continuously with transmissometers. The units of measurement are Mm⁻¹: the higher the light extinction value in Mm⁻¹, the more visibility is reduced. In Figures 45 and 46, these light extinction data have been plotted as a yearly averages and converted to the preferred units of visual range in miles.

Trend analysis for Phoenix is represented using only the transmissometer data (see Figures XX). The transmissometer measure both the light scattering and light absorption effects of the atmosphere, which is close to how the human eye interprets visibility. The nephelometer instrument measures only the light scattering effects of particulates in the atmosphere and based on the differing wavelengths of scattered light, can measure what types of particulates in the air. While this is important for determining what pollutants are impacting visibility, is not as representative for the human eye. Additionally, the nephelometer instrument was recently relocated to a different location in Phoenix and trend analysis for the new location cannot be determined. Therefore, only data from the transmissometer will be used to show trend analysis for Phoenix visibility.

Figure 45 illustrates the temporal variability of visibility in Phoenix over the 1995 to 2017 period in the form of annual mean, Dirtiest 20%, and Cleanest 20% of days. The average trends for a period of 5 years, 10 years, and length of record are provided in Table 43 below.

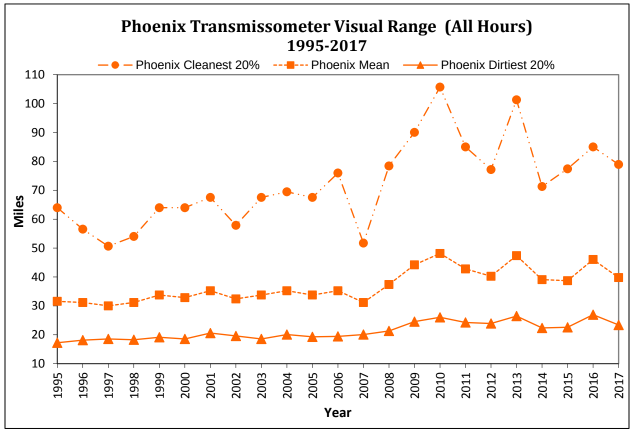


Figure 45 – Phoenix Transmissometer Visual Range (All Hours)

Table 43 – Phoenix Transmissometer Annual Visual Range (All Hours)

	All Hours Transmissometer Visual Range				
Trend Length	Mean of Cleanest 20%	Mean	Mean of Dirtiest 20%		
5 Year 14% ↓		8% ↓	3% ↓		
(2013 - 2017)	Decrease	Decrease	Decrease		
10 Year	11% ↓	2% ↓	3% ↑		
(2008 - 2017)	Decrease	Decrease	Increase		
	59% 个	48% ↑	47% 个		
Length of Record	Increase	Increase	Increase		
	1995-2017	1995-2017	1995-2017		

Figure 46 illustrates the temporal variability of visibility of the morning hours in Phoenix over the 1995 to 2017 period in the form of annual mean, Dirtiest 20%, and Cleanest 20% of days. The average trends for a period of 5 years, 10 years, and length of record are provided in Table 44 below.

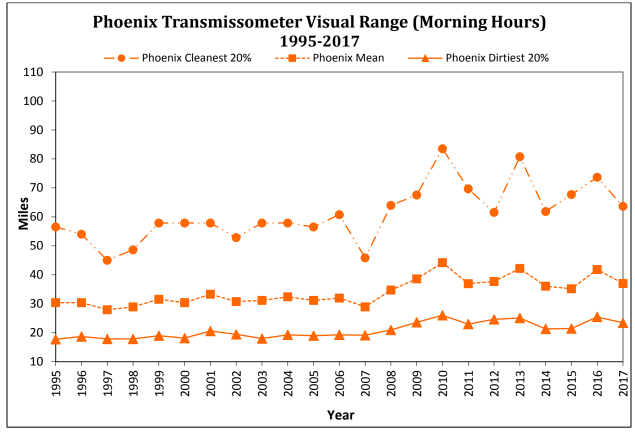


Figure 46 – Phoenix Transmissometer Visual Range (Morning Hours)

Table 44 - Phoenix Transmissometer Annual Visual Range (Morning Hours)

	5am - 11am Hours					
Trend Length	Mean of Cleanest 20% Mean		Mean of Dirtiest 20%			
5 Year	12% ↓	5% ↓	1% 个			
(2013 - 2017)	Decrease	Decrease	Increase			
10 Year	3% ↓	0% 个	2% 个			
(2008 - 2017)	Decrease	Increase	Increase			
Longth of	42% 个	40% 个	42% 个			
Length of Record	Increase	Increase	Increase			
Record	1995-2017	1995-2017	1995-2017			

Transmissometer seasonal variability shows which seasons have higher visual range (Figure 47). The seasons shown are Winter (Jan-Mar), Spring (Apr-June), Summer (July-Sept), and Fall (Oct-Dec). Data for seasonal variability is currently available for the years 2006-2017. The visual range is greater during the spring and summer months than during the fall and winter months. Visual range peaked during the year 2010 in the summer months at around 77 miles. A contributing factor for this was that 2010 was an above normal year for precipitation. Rainfall cleans the air, improving visual range. The average trends for a period of 5 years, 10 years, and length of record are provided in Table 45 below.

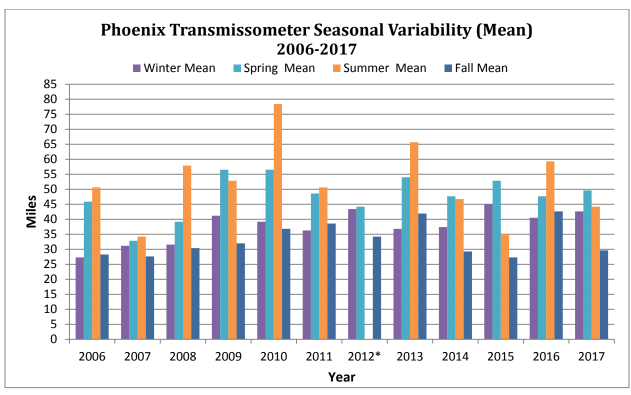


Figure 47 – Transmissometer Seasonal Average Trend *Data did not meet completeness criteria for the summer of 2012.

Table 45 – Phoenix Transmissometer Annual Visual Range (Seasonal)

	Seasonal Mean						
Trend Length	Winter Mean	Spring Mean	Summer Mean	Fall Mean			
5 Year	16% 个	7% ↓	22% ↓	12% ↓			
(2013 - 2017)	Increase	Decrease	Decrease	Decrease			
10 Year	18% 个	2% 个	N/A	0% 个			
(2008 - 2017)	Increase	Increase	IN/A	Increase			
Length of Record	40% 个 Increase 2006-2017	19% 个 Increase 2006-2017	22% ↓ Decrease 2013-2017 40% ↑ Increase 2006-2011	16% 个 Increase 2006-2017			

Examples of photographic visibility conditions are shown in Figure 48. As visibility decreases, the clarity and resolution of the mountains and downtown area decreases and a milky appearance covers the vista. On poor visibility days, the mountain ridgeline and most of the downtown area are no longer visible.

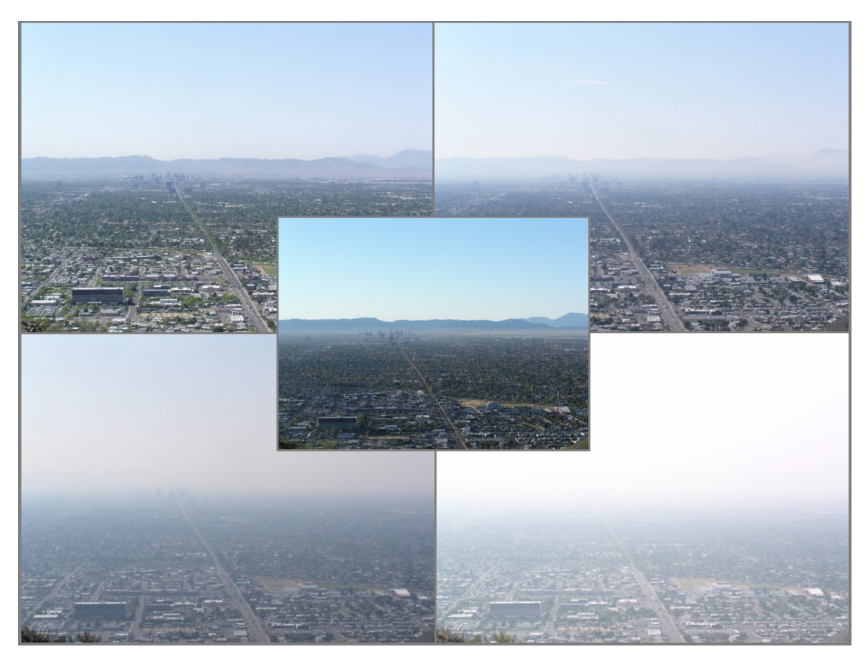


Figure 48 – Examples of Visual Condition in Phoenix. Excellent to Very Poor visibility days taken from the high resolution digital picture. All are images of South Mountain were taken from the camera located at North Mountain. Center-Excellent, Top left-Good, Top right-Fair, Bottom left-Poor, Bottom right-Very Poor.

2.0 IMPROVE

The IMPROVE program is the main supporter of the RHR across the nation. It uses monitors specific to the objectives to increase visibility and decrease haze in the national parks and wilderness areas. The program objectives of IMPROVE are:

- to establish current visibility and aerosol conditions in mandatory Class I areas;
- to identify chemical species and emission sources responsible for existing man-made visibility impairment;
- to document long-term trends for assessing progress towards the national visibility goal; and,
- with the enactment of the RHR, to provide regional haze monitoring representing all visibility-protected Federal Class I areas where practical.

The IMPROVE monitors are closely related to those in the CSN in that they use different types of sample filters to measure the amount of speciated pollutants in the air. Ambient air is drawn into the sampler, where the pollutants are deposited onto the sample filters. There are four different sample filters that are run on the same day, each collecting different types of pollutants. Different aerosols and particulates contribute to haze more than others: therefore, the monitors are designed to capture these specific pollutants.

The IMPROVE network in Arizona has 16 samplers in 12 Class I areas. This program is governed by a steering committee composed of representatives from federal and regional/state organizations. This network was established in 1985 to aid the creation of federal and state implementation plans. ADEQ supports the IMPROVE network by performing filter sample changes at JLG Supersite, Queen Valley, and Douglas Red Cross sites, and by conducting performance audits on protocol samplers in Arizona. ADEQ is one of the many government agencies involved in the IMPROVE network. Users of these data include, but are not limited to, the EPA, visibility researchers, and land management agencies.

More information of the IMPROVE program and data can be accessed at the following website: http://vista.cira.colostate.edu/improve/.

Conclusion

For more information on air quality monitoring and assessments at ADEQ, please go to http://www.azdeq.gov/AQ/monitoring. Monitoring data, previous annual reports and network plans, as well as links to download air quality apps, can be found here.

Appendix I – Abbreviations

Arizona Department of Environmental Quality ADEQ

Best Management Practices for Agricultural Activities **AgBMP**

AQD Air Quality Division AQI Air Quality Index

Air Quality System (EPA database) AQS

Beta Attenuation Monitor BAM Bext **Total Light Extinction Bscat Light Scattering**

Clean Air Act CFR Code of Federal Regulations

CO Carbon Monoxide

CAA

Community Scale Toxics Ambient Monitoring CSATAM

CSN **Chemical Speciation Network**

DM&QA Data Management & Quality Assurance Unit

DNPH 2,4-dinitrophenylhydrazine **Data Quality Indicator** DQI DQO **Data Quality Objective**

EPA Environmental Protection Agency FIP Federal Implementation Plan

Gas Chromatography with Flame Ionization Detector GC/FID

HAP Hazardous Air Pollutant

HC **Hydrocarbons**

IMPROVE Interagency Monitoring of PROtected Visual Environments

IR Infrared

LPM Liters per Minute

MAG Maricopa Association of Governments MQO Measurement Quality Objective **MSA** Metropolitan Statistical Area

 Mm^{-1} **Inverse Megameter**

 mg/m^3 Milligrams per Cubic Meter $\mu g/m^3$ Micrograms per Cubic Meter

National Ambient Air Quality Standard NAAQS **NATTS National Air Toxics Trends Station**

NCore National Core multipollutant monitoring stations

NEI **National Emissions Inventory**

 NH_4NO_3 **Ammonium Nitrate National Monument** NM

NO Nitric Oxide Nitrogen Dioxide NO_2 NOx Nitrogen Oxides

Total Reactive Oxides of Nitrogen NOv

 O_3 Ozone

Polycyclic Aromatic Hydrocarbon PAH

PAMS Photochemical Assessment Monitoring Station Pb Lead

PM Particulate Matter

PM₁₀ Particulate Matter ≤ 10 microns

PM_{10-2.5} Coarse Particulate Matter between 2.5 to 10 micrometers aerodynamic diameter, may

also be denoted as PM_{coarse}

PM_{2.5} Particulate Matter ≤ 2.5 microns

ppb Parts Per Billion ppm Parts Per Million

PQAO Primary Quality Assurance Organization PSD Prevention of Significant Deterioration

PTFE Polytetrafluoroethylene
QA Quality Assurance

QAPP Quality Assurance Program Plan

QC Quality Control

QMP Quality Management Plan

RH Relative Humidity
RHR Regional Haze Rule

SATMI School Air Toxics Monitoring Initiative

SIP State Implementation Plan

SLAMS State and Local Air Monitoring Stations

SO₂ Sulfur Dioxide

SOP Standard Operating Procedure SPM Special Purpose Monitor

STP Standard Temperature Pressure
SVOC Semi-Volatile Organic Compound
TAD Technical Assistance Document

TNMOC Total Non-Methane Organic Compound

TSP Total Suspended Particle

UATMP Urban Air Toxics Monitoring Program

USG Unhealthy for Sensitive Groups

UV Ultraviolet

VOC Volatile Organic Compound

Appendix II – References

http://www.airnow.gov/ - AQI Forecast

http://www.ecy.wa.gov/programs/air/sips/pollutants/naaqs.htm - National and State Ambient Air Quality Standards, Department of Ecology, State of Washington

https://www.epa.gov/criteria-air-pollutants/naaqs-table - National Ambient Air Quality Standards (NAAQS)

http://www.epa.gov/airquality/carbonmonoxide/health.html - Carbon Monoxide Health Effects

https://www.epa.gov/ozone-pollution - Ozone (O₃) Health Effects

https://www.epa.gov/air-trends - National Trends.

http://www.epa.gov/ttn/naaqs/standards/co/s_co_history.html - Carbon Monoxide (CO) Standards - Table of Historical CO NAAQS

 $\frac{\text{http://www.epa.gov/ttn/naaqs/standards/nox/s} \ \ nox \ \ history.html}{\text{historical NO}_2\ NAAQS} - \text{Nitrogen Dioxide (NO}_2)\ Standards - \text{Nitrogen Dioxide (NO}_2) - \text{Nitrogen$

 $\frac{\text{https://www.epa.gov/ozone-pollution/table-historical-ozone-national-ambient-air-quality-standards-naaqs}{\text{naaqs}} \text{ - Ozone } (O_3) \text{ Standards} \text{ - Table of Historical } O_3 \text{ NAAQS}$

http://www.epa.gov/ttn/naaqs/standards/pb/s pb history.html - Lead (Pb) Standards - Table of Historical Pb NAAQS

http://www.epa.gov/ttn/naaqs/standards/pm/s pm history.html - Particulate Matter (PM) Standards - Table of Historical PM NAAQS

http://www.epa.gov/ttn/naaqs/standards/so2/data/so2final.pdf - National Ambient Air Quality Standards for Sulfur Oxides (Sulfur Dioxide)—Final Decision

http://www.epa.gov/ttn/naaqs/standards/so2/s so2 history.html - Sulfur Dioxide (SO₂) Primary Standards - Table of Historical SO₂ NAAQS

http://www.phoenixvis.net/PPMmain.aspx - Non-Regulatory Portable Particulate Monitors

https://www.epa.gov/aqs - EPA's Air Quality System

https://www3.epa.gov/ttnamti1/uatm.html - 2014 National Monitoring Programs Annual Report (UATMP, NATTS, and CSATAM)

Appendix III – 2017 Area Designations Map

