# National Air Toxics Trends Study Grand Junction, Colorado

# January through December 2015

Prepared by the Air Pollution Control Division
Technical Services Program
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#### I. EXECUTIVE SUMMARY

The Grand Junction air toxics monitors were originally established as a part of the 2001/2002 Pilot Study for the National Air Toxics Trends Sites (NATTS). The network was created by the Environmental Protection Agency (EPA) in an effort to gather data that were suitable for identifying trends in air toxics concentration levels. Grand Junction was one of the five "rural" sites selected for the study initially. Since that time, and as the population of the Grand Junction area has grown, the EPA has reconsidered, and decided that the site is more indicative of urban concentrations, and has changed the designation of the site from rural to urban.

Most of the compounds detected at Grand Junction in 2015 are found in urban air nationwide. There do not appear to be any compounds of local significance. The majority of compounds can be related to motor vehicle sources. These include formaldehyde, benzene, toluene, ethylbenzene, xylenes, and styrene. Chloroflourocarbons are also present, including chloromethane, dichlorodifluoromethane, trichlorofluoromethane, and trichlorotrifluoroethane. Polycyclic aromatic hydrocarbon compounds naphthalene, phenanthrene and acenaphthene are frequently detected.

This report has two appendices. Appendix A, "Documentation for Grand Junction Urban Air Toxics Trends Monitoring Locations – Site Maps and Photographs" provides information concerning the two air monitoring sites discussed in this report. Appendix B, "Air Toxics Summary: Compounds Contributing to Cancer and Non-cancer Risks – Overview of Sources and Health Effects," provides a brief summary of many of the compounds monitored. That document discusses the chemical formula, sources, and uses of each compound. It also profiles potential health effects, such as carcinogenicity, the compound's potential to cause birth defects, and whether it damages target organs in the body.

## II. INTRODUCTION

# Background

The NATTS Network collects ambient air toxics monitoring data as a part of the Urban Air Toxic Strategy (UATS). Under Section 112 of the Clean Air Act (CAA), the EPA established a list of 187 toxic air pollutants, also known as hazardous air pollutants (HAPs). These are pollutants that are known, or suspected, to cause cancer, or other major health issues. People who are exposed to these HAPs at sufficient concentration levels may have an increased chance of getting cancer, having their immune systems damaged, etc. Most air toxics originate from mobile sources (such as cars, trucks, and buses) or stationary sources (such as factories, refineries, and power plants). Some air toxics can come from indoor sources as well (for example: cleaning solvents or building materials).

Since it is not practical, or possible, to monitor for each of the 187 compounds, the EPA developed a subset of HAPs that have the greatest impact on the public and the environment. For the purposes of the NATTS Study, 62 HAPs are monitored. Thirty three of these are on the "Urban HAP List." The remaining 29 compounds were chosen based on risk factors that were developed by the EPA. From the list of 62 compounds, a "core" list of 19 toxic air pollutants that must be monitored at all times was created. These compounds are considered to be "priority compounds" because they are major health risk drivers, based on a relative ranking performed by the EPA. They are referred to as the "Method Quality Objective (MQO) Core Analytes." These compounds can be seen in Table 1.

<sup>&</sup>lt;sup>1</sup> Technical Assistance Document for the National Air Toxics Trends Stations Program." US Environmental Protection Agency. Revision 3, October 2016.

https://www3.epa.gov/ttnamti1/files/ambient/airtox/NATTS%20TAD%20Revision%203\_FINAL%20October%202016.pdf

<sup>&</sup>lt;sup>2</sup> Ibid.

<sup>&</sup>lt;sup>3</sup> *Ibid*.

Table 1. NATTS HAPs with Mandatory Monitoring Requirements

VOCs	Carbonyls	PAHs	PM <sub>10</sub> Metals	TSP Metals
1,3-Butadiene	Acetaldehyde	Benzo(a)pyrene	Arsenic	Hexavalent Chromium
Acrolein	Formaldehyde	Naphthalene	Beryllium	
Benzene			Cadmium	
Carbon Tetrachloride			Lead	
Chloroform			Manganese	
Tetrachloroethylene			Nickel	
Trichloroethylene				
Vinyl Chloride				

The Grand Junction air toxics monitoring site was established in 2004. This site measures air toxics to determine the success of the National Air Toxics Strategy in reducing the U.S. population exposure to cancercausing substances in the air. The primary test is a comparison of mean concentrations of compounds for the first three years (2004-2006), versus the mean concentrations for successive three-year periods (2007-2009, 2010-2012, etc.), starting from 2004 and continuing to the present. Data collected beyond the initial six year study scope will be used for trending analyses.

This report presents data from January 2015 through December 2015. It is separated into sections covering the various compounds of interest. Sections 3, 4, 5, and 6 discuss the compounds monitored as a part of this study. Sections 7, 8 and 9 compare the PM<sub>10</sub>, PM<sub>2.5</sub>, and meteorological data collected as a part of the regular monitoring conducted in Grand Junction by the Colorado Department of Public Health and Environment (CDPHE) to show compliance with the national ambient air quality standards (NAAQS). Each section begins with summary statistics for the compounds analyzed followed by the percentage of samples in which each chemical was detected. Summary graphs of certain compounds are presented. It is important to note here that sampling for hexavalent chromium was discontinued in June 2013 due to an extremely low detection rate in addition to the lack of any apparent sources in the vicinity. Historical data for this compound can be found in prior years' NATTS reports.

#### Site Information

The NATTS Study at Grand Junction collects samples at two separate locations. These two sites (Powell and Pitkin sites) are in close proximity to one another. The Powell site is located on top of the Powell Building (approximately three stories in height) at 650 South Avenue, and the Pitkin site is located approximately 50 meters to the NNW of the Powell Building, on the roof of a small shelter, near ground level, at 645-1/4 Pitkin Avenue. The particulate/metals samplers are located on the Powell Building, and the carbon monoxide analyzer, air toxics samplers (VOC/carbonyl/PAH), and meteorological tower are located at the Pitkin site. Due to the different sampling heights, staff at Region VIII of the EPA suggested the sites be separately catalogued in the national air monitoring database [AQS IDs: 080770017 (Powell), and 080770018 (Pitkin)]. Documentation regarding these sites, including maps, photographs, and aerial views, is available as Appendix B in this document. The sites are located on the southern end of the downtown area, in an area of commercial/light industrial land use.

# III. CARBONYLS

#### Summary Statistics

The carbonyls discussed in this section are the group of organic chemicals that contain a functional group composed of a carbon atom double bonded to an oxygen atom. The generalized symbol for the carbonyl group is R-C=O, where the "R" is some other carbon compound. Thirteen compounds were measured for this study in 2015. A listing of these compounds, as well as a summary of the collected data, is shown in Table 2. Of the thirteen carbonyl compounds analyzed for, two are included on the mandatory monitoring list of 19 core HAPs. They are shown in bold type in Table 2. In 2012, 2-butanone was added to the carbonyl

analyses. It has previously been analyzed for via EPA's Method TO-15. It was moved to the Method TO-11A list because the TO-11A methodology provides better results at lower levels for this compound. The previous years' values from Method TO-15 are indicated with an asterisk.

Carbonyl sampling was attempted on an every-sixth-day basis in 2015, in accordance with the EPA's national sampling schedule. This should have yielded 60 samples during 2015. Thirteen samples were either missed or had to be voided due to site equipment problems or operator errors. Seven of those samples were made up for on alternate dates, for a total of 55 samples recovered during the year. This data recovery rate of 91.6% exceeds the EPA goal of over 85% sample recovery. However, a large number of samples from February and March were voided, compromising the completeness requirements for the 1<sup>st</sup> quarter. APCD was unable to perform make up samples during the same time period, as the issues were not discovered until performing quality assurance reviews of the data several months later.

The annual mean concentrations for each carbonyl compound, from 2004 through 2015, are listed in Table 2. The annual means were calculated by replacing all "non-detect" values with one-half of the sample method detection limit (MDL). This is an accepted conservative technique for calculating annual values when some of the samples were less than the laboratory's ability to detect. The most prevalent carbonyls in the ambient air in Grand Junction are formaldehyde, acetone, and acetaldehyde. The other ten compounds measured in this study occurred at concentration levels significantly below those of the top three compounds. Since 2004, the annual average concentrations for many of the carbonyl compounds have dropped. The 2015 averages are similar to the 2014 averages.

All of the carbonyls, except for isovaleraldehyde and 2,5-dimethylbenzaldehyde, were present in greater than 90% of the samples. During the pilot phase of this study in 2001-2002, 2,5-dimethylbenzaldehyde was detected 34 percent of the time. That number dropped to 4.8 percent in 2005, and the compound has not been detected since 2006.

Isovaleraldehyde has not been detected since 2010. Note that the true annual means of 2,5-dimethylbenzaldehyde and isovaleraldehyde may be well below the numbers reported in the table due to the fact that these compounds were not detected, and one-half of the detection limit was used for the estimated concentration of the non-detects. Actual concentrations could have been at lower levels than these estimates.

It should be noted here that there was a shift in the data processing methods for this report. In previous years, data was presented for compounds that were present in at least 90% of the samples taken. For this (and future) reports the data presented will be from compounds that were present in at least 10% of the samples taken. While this move did not have a noticeable affect on the carbonyl data (as all but 3 compounds were detected in all samples taken), it did allow for inclusion of more of the VOC compounds discussed in later sections. For purposes of consistency, this new 10% detection criterion was applied to all carbonyl, VOC, metal, and SVOC data sets.

Table 2. 2015 Sampling Statistics and Annual Averages

	GA G	2015	2015		Annual Averages (μg/m³)										
Compound	CAS Number	# of ND's	% ND	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014	2015
2,5-Dimethylbenzaldehyde	5779-94-2	55	100%	0.08	0.06	0.02	0.03	0.03	0.00	0.01	0.01	0.01	0.01	0.01	0.01
2-Butanone	78-93-3	0	0%	2.56*	0.43*	1.23*	0.99*	0.98*	1.03*	1.46*	1.08*	0.54	1.35	0.46	0.48
Acetaldehyde	75-07-0	0	0%	10.53	5.39	4.25	5.03	4.48	2.89	1.95	2.43	2.85	3.76	2.76	1.58
Acetone	67-64-1	0	0%	18.39	11.08	9.69	12.45	12.35	5.57	5.13	4.92	5.46	6.38	4.63	4.50
Benzaldehyde	100-52-7	0	0%	1.11	0.95	1.45	1.41	1.3	0.34	0.31	0.41	0.39	1.41	0.16	0.28
Butyraldehyde	123-72-8	0	0%	0.91	1.18	1	1.06	0.92	0.35	0.34	0.39	0.33	0.66	0.3	0.26
Crotonaldehyde	123-73-9	0	0%	0.67	0.62	0.5	0.57	0.55	0.22	0.2	0.16	0.16	0.24	0.16	0.19
Formaldehyde	50-00-0	0	0%	3.45	3.83	4.94	4.94	5.04	4.01	2.74	2.74	2.98	6.41	3.86	3.08
Hexaldehyde	66-25-1	0	0%	0.56	0.43	0.46	0.43	0.52	0.12	0.13	0.1	0.11	0.52	0.1	0.15
Isovaleraldehyde	590-86-3	55	100%	0.04	0.07	0.15	0.08	0.08	0.01	0.01	0.00	0.01	0.01	0.00	0.01
Propionaldehyde	123-38-6	0	0%	0.39	0.75	0.74	0.73	0.91	0.39	0.35	0.35	0.34	0.43	0.32	0.29
Tolualdehydes	NA	2	4%	0.61	0.63	1.11	0.98	0.77	0.18	0.19	0.19	0.18	0.4	0.11	0.15
Valeraldehyde	110-62-3	0	0%	0.18	0.71	0.59	0.06	0.52	0.15	0.11	0.08	0.09	0.28	0.09	0.09

ND = Not Detected

**Bold = MQO Core Analyte**Italic = Less than 90% detection rate (2004-2014), less than 10% detection rate (2015)

<sup>\* =</sup> Results obtained by different analytical method

# **Graphs**

The summary data for carbonyl compounds measured during 2015 are graphed in Figure 1. The compounds in these graphs are ordered by ranking their average concentrations from highest to lowest. The graphs show that acetone, formaldehyde, and acetaldehyde had the highest annual averages with values of  $4.50 \pm 2.08$ ,  $3.08 \pm 1.89$ , and  $1.58 \pm 0.65$  micrograms per cubic meter, respectively. The maxima for acetone and formaldehyde observed in 2015 were larger than their respective maxima in 2014. The maximum value for acetaldehyde in 2015 was slightly lower than the 2014 maximum.

In comparison, the national average concentrations in 2013 for acetone, formaldehyde, and acetaldehyde for all air toxics monitoring sites participating in the EPA's National Monitoring Program (NMP), were 1.12  $\pm$  0.88, 2.30  $\pm$  1.90, and 1.00  $\pm$  0.70 micrograms per meter cubed, respectively. The national data reports for 2014 and 2015 had not yet been released at the time this report was written.

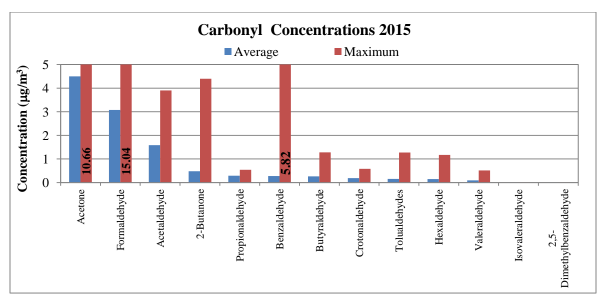


Figure 1. Annual Mean and Maximum Carbonyl Concentrations for 2015

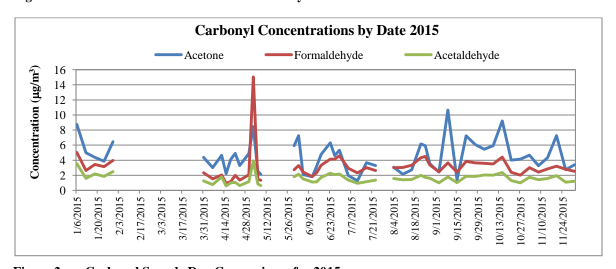


Figure 2. Carbonyl Sample Day Comparisons for 2015

[5]

<sup>&</sup>lt;sup>4</sup> "2013 National Monitoring Programs Annual Report (UATMP, NATTS, CSATAM). US EPA. October 2015.

Figure 2 shows the concentrations for the acetone, formaldehyde, and acetaldehyde compounds in chronological order for 2015. It is difficult to make conclusions regarding seasonal variation at this site during the year given the large data gaps. Generally, more formaldehyde is formed photochemically during the summer period of higher solar radiation. Formaldehyde plays a role in the formation of ozone, a chemical that usually peaks during the summer. The data does show a large spike in formaldehyde concentrations on May 3, 2015.

Figure 3 is a graph of the weekday versus weekend average carbonyl concentrations in 2015. Each of the carbonyl compounds exhibited very similar weekday and weekend averages. 2,5-dimethylbenzaldehyde, and isovaleraldehyde, have weekday and weekend average concentrations that are equal, because their concentrations are merely half the value of their respective MDLs for the entire year, since they were non-detectable in all samples. Formaldehyde, acetaldehyde, 2-butanone, propionaldehyde, butyraldehyde, benzaldehyde, the tolualdehydes, and hexaldehyde all have weekend averages that are slightly higher than their weekday counterparts.

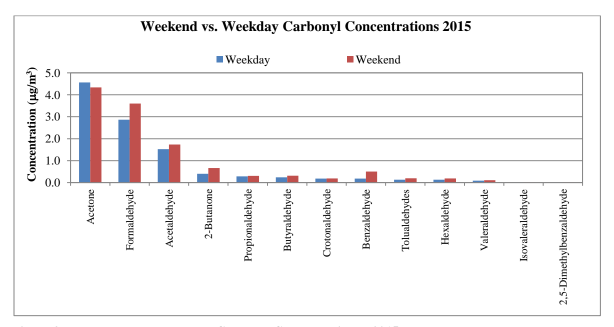


Figure 3. Weekday vs. Weekend Carbonyl Concentrations - 2015

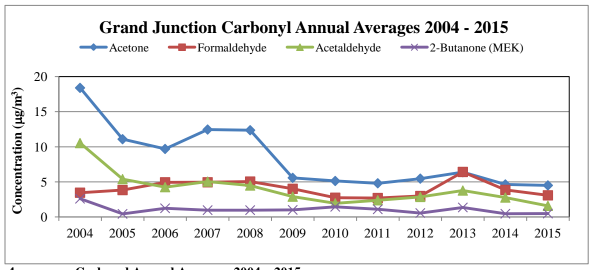


Figure 4. Carbonyl Annual Averages 2004 – 2015

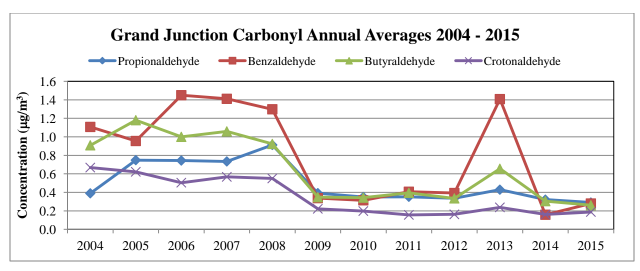


Figure 5. Carbonyl Annual Averages 2004 – 2015, ctd.

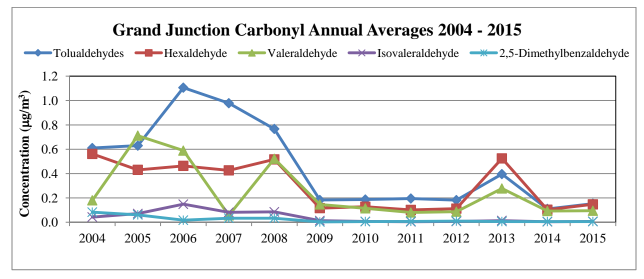


Figure 6. Carbonyl Annual Averages 2004 – 2015, ctd.

Figure 4 through Figure 6 are graphs of the annual average carbonyl concentrations at the Grand Junction site, for 2004 through 2015. The overall trend from 2004 through 2011 appears to be that the carbonyl concentrations are decreasing for most compounds. During 2011 and 2012 the annual average concentrations were very similar, with slight increases or decreases for some compounds. The averages in 2013 are higher than previous years, and do not follow the overall decreasing trend of the data. This is due to a large wildfire in the Grand Junction area during the summer of 2013. The 2014 and 2015 averages are lower than the 2013 averages. The 2015 averages, however, are slightly higher than their 2014 counterparts for several compounds.

The NATTS program was initially established to monitor the 3-year average concentrations of air toxics compounds, with the thought that successive 3-year averages would show at least a 15% drop in concentration values. Figure 7 through Figure 9 below show the 3 year average concentrations for eleven of the thirteen carbonyl compounds. 2,5-Dimethylbenzaldehye and isovaleraldehyde 3-year averages were not calculated as the compounds have not been detected for many years. The 3-year averages are taken from 2004 through 2006, 2007 through 2009, 2010 through 2012, and 2013 through 2015. Generally, the 3-year average concentrations decreased from 2004 to 2012. The 2013 through 2015 averages decreased for only one compound, 2-butanone. The remaining averages increased, or stayed the same largely as a result of the contribution from the 2013 data which was heavily influenced by the wild fires that year.

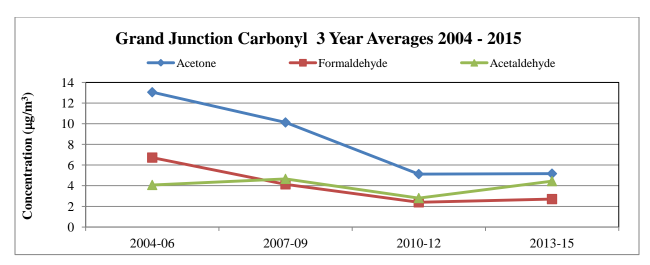


Figure 7. Carbonyl 3-Year Averages 2004 – 2015

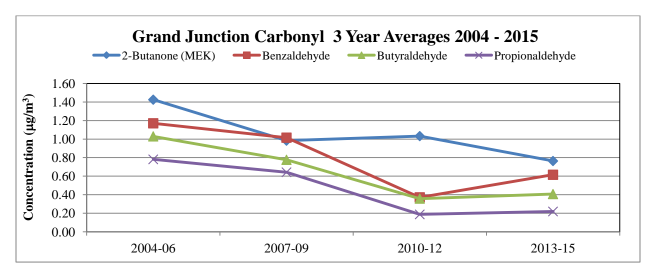


Figure 8. Carbonyl 3-Year Averages 2004-2015, ctd.

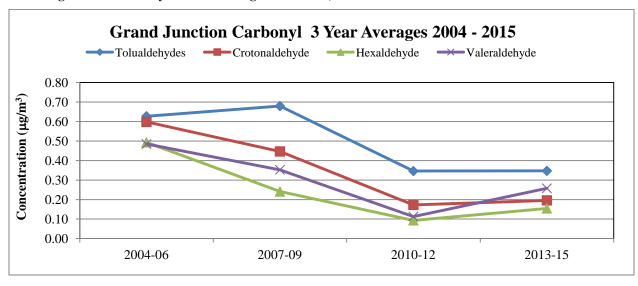


Figure 9. Carbonyl 3-Year Averages 2004-2015, ctd.

# Quality Assurance/Quality Control

#### **Blanks**

Field blanks were collected twelve times per year by attaching a blank sample cartridge to the sampler briefly, and then removing it. The purpose of these blanks was to assess contamination that might exist in the cartridge media, sample installation, or shipping. Most cartridges had very small amounts of formaldehyde, acetaldehyde, acetone, and propionaldehyde. Detailed information regarding field blank results is available upon request.

## **Precision of Sample Results**

This project collected precision data in order to assess both sampling and analytical procedures. Six times during the year, a second carbonyl cartridge was sampled simultaneously with the primary sample. These additional samples, or duplicates, were collected to assess the precision (repeatability) of the sampling method. In general, agreement between the two samples was excellent. Detailed information regarding precision results is available upon request.

# IV. VOLATILE ORGANIC COMPOUNDS

# **Summary Statistics**

Volatile organic compound (VOC) data collected at the Grand Junction – Powell station from January through December 2015 are presented in this section. There were 58 VOCs analyzed for this study. The list of these VOCs and the number of times each was detected in samples during the study is found in Table 3. The compounds in bold type are MQO Core Analytes. These are the same VOCs collected by all of the sites participating in the national air toxics study. VOCs are typically sampled on an every-sixth-day basis, for a total of 60 possible sample days in 2015. In all, 60 samples were attempted, with 12 samples that were missed or voided for various reasons, and 11 samples that were made up, for a 98% sample recovery rate.

Table 3. VOC List with 2015 Detection Rates

	CAS		
Compound	Number	# of ND's	% ND
1,2,4-Trimethylbenzene	95-63-6	0	0%
1,3-Butadiene	106-99-0	0	0%
Acetonitrile	75-05-8	0	0%
Acetylene	74-86-2	0	0%
Benzene	71-43-2	0	0%
Bromomethane	74-83-9	0	0%
Carbon Disulfide	75-15-0	0	0%
Carbon Tetrachloride	56-23-5	0	0%
Chloroform	67-66-3	0	0%
Chloromethane	74-87-3	0	0%
Dichlorodifluoromethane	75-71-8	0	0%
Dichloromethane	75-09-2	0	0%
Dichlorotetrafluoroethane	76-14-2	0	0%
Ethylbenzene	100-41-4	0	0%
m,p-Xylene	100-01-6	0	0%
Methyl Isobutyl Ketone	108-10-1	0	0%
n-Octane	111-65-9	0	0%
o-Xylene	95-47-6	0	0%

	CAS		
Compound	Number	# of ND's	% ND
Propylene	115-07-1	0	0%
Toluene	108-88-3	0	0%
Trichlorofluoromethane	75-69-4	0	0%
Trichlorotrifluoroethane	76-13-1	0	0%
Styrene	100-42-5	1	2%
1,3,5-Trimethylbenzene	108-67-8	2	3%
Tetrachloroethylene	127-18-4	2	3%
1,2-Dichloroethane	107-06-2	3	5%
Acrolein	107-02-8	8	14%
Chloroethane	75-00-3	11	19%
Hexachloro-1,3-butadiene	87-68-3	11	19%
1,1,1-Trichloroethane	71-55-6	12	20%
1,2,4-Trichlorobenzene	120-82-1	16	27%
Ethyl tert-Butyl Ether	637-92-3	22	37%
p-Dichlorobenzene	106-46-7	30	51%
Acrylonitrile	107-13-1	39	66%
Dibromochloromethane	124-48-1	45	76%
Trichloroethylene	79-01-6	52	88%
Methyl Methacrylate	80-62-6	53	90%
Vinyl chloride	75-01-4	58	98%
o-Dichlorobenzene	95-50-1	55	93%
1,1,2,2-Tetrachloroethane	79-34-5	56	95%
1,1-Dichloroethene	75-35-4	56	95%
Bromodichloromethane	75-27-4	56	95%
m-Dichlorobenzene	541-73-1	57	97%
trans-1,2-Dichloroethylene	156-60-5	57	97%
1,2-Dibromoethane	106-93-4	58	98%
1,2-Dichloropropane	78-87-5	58	98%
Bromoform	75-25-2	58	98%
cis-1,3-Dichloropropene	10061-01-5	58	98%
1,1,2-Trichloroethane	79-00-5	59	100%
1,1-Dichloroethane	75-34-3	59	100%
Bromochloromethane	74-97-5	59	100%
Chlorobenzene	108-90-7	59	100%
Chloroprene	126-99-8	59	100%
cis-1,2-Dichloroethylene	156-59-4	59	100%
Ethyl Acrylate	140-88-5	59	100%
Methyl tert-Butyl Ether	1634-04-4	59	100%
tert-Amyl Methyl Ether	994-05-8	59	100%
trans-1,3-Dichloropropene	10061-02-6	59	100%

ND = Not Detected, **Bold = MQO Core Analyte** *Italic = Less than 10% detection rate (2004-2014)*In 2015, there were 38 compounds detected in at least 10% of the samples taken, while in 2014, there were

39 that met the same criterion. Eight of those 38 VOC compounds are on the core list of 19 HAPs to be monitored by NATTS stations. Seven of those eight core compounds were detected in greater than 90% of the samples taken in 2015. Vinyl chloride is the only core compound that was detected in less than 10% of the samples taken in 2015. Table 4 is an alphabetical listing of the 38 compounds most frequently detected in 2015. Bolded compounds are on the list of 19 core HAPs. The italicized compound was detected in less than 10% of the samples.

Table 4. VOCs Detected in Greater Than 10% of 2015 Samples

10% Detection Rate									
1,1,1-Trichloroethane	Carbon Tetrachloride	Methyl Methacrylate							
1,2,4-Trichlorobenzene	Chloroethane	n-Octane							
1,2,4-Trimethylbenzene	Chloroform	o-Xylene							
1,2-Dichloroethane	Chloromethane	p-Dichlorobenzene							
1,3,5-Trimethylbenzene	Dibromochloromethane	Propylene							
1,3-Butadiene	Dichlorodifluoromethane	Styrene							
Acetonitrile	Dichloromethane	Tetrachloroethylene							
Acetylene	Dichlorotetrafluoroethane	Toluene							
Acrolein	Ethyl tert-Butyl Ether	Trichloroethylene							
Acrylonitrile	Ethylbenzene	Trichlorofluoromethane							
Benzene	Hexachloro-1,3-butadiene	Trichlorotrifluoroethane							
Bromomethane	m,p-Xylene	Vinyl chloride							
Carbon Disulfide	Methyl Isobutyl Ketone								

Bolded compounds are on the list of 19 core HAPs

Italicized compounds were detected in less than 10% of samples taken.

There were ten compounds that were not detected at all during 2015, which is an increase from the nine non-detect compounds in 2014. There were 19 compounds that were detected in five percent, or less, of the samples in 2015. This is an increase from 2014, where only seven compounds were detected in five percent, or less, of the samples. This list of 20 compounds includes many compounds that are chiefly emitted by stationary sources. It appears that these source types are not present in the immediate vicinity of the Grand Junction station.

Table 5 summarizes the annual mean concentrations for each of the 58 VOCs measured during the study, from 2004 through 2015. Compounds that have bolded values are MQO Core Analytes. Compounds with italicized values were detected in less than 10% of the samples for the year. It should be noted here that there was a shift in the data processing methods for this report. In previous years, data was presented for compounds that were present in at least 90% of the samples taken. For this (and future) reports the data presented will be from compounds that were present in at least 10% of the samples taken. This move had a noticeable effect on the VOC data by including more compounds into the discussion. For purposes of consistency, this new 10% detection criterion was applied to all carbonyl, VOC, metal, and SVOC data sets.

The annual means were calculated by replacing all "non-detect" values with one-half of the sample MDL. This is an accepted conservative technique for calculating annual values when some of the samples were less than the laboratory's ability to measure. As a result of this technique, the average and maximum concentrations are the same if the compound was never detected. A year to year variation of the average concentrations for these non-detect compounds will be seen due to the reevaluation of the MDLs annually. The compounds are listed in alphabetical order. There are several things to note about this table. First, the acetonitrile values for all of 2004, and the first three-and-a-half months of 2005 were voided due to a contamination problem in the sampling equipment. Acrolein was not analyzed until 2005, and carbon disulfide was added to the list of analytes in 2006. Removed from this list for 2012 were the compounds of chloromethylbenzene, and methyl ethyl ketone (MEK). MEK was added to the carbonyl analysis. Chloromethylbenzene has never been detected in greater than 10% of samples.

Table 5.VOC Data Summary 2015

		Annual Average VOC Concentrations (μg/m³)											
Analyte	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014	2015	
1,1,1-Trichloroethane	0.14	0.15	0.12	0.10	0.09	0.09	0.09	0.07	0.05	0.05	0.05	0.04	
1,1,2,2-Tetrachloroethane	0.17	0.16	0.05	0.06	0.03	0.01	0.03	0.08	0.06	0.09	0.05	0.06	
1,1,2-Trichloroethane	0.22	0.15	0.02	0.05	0.02	0.01	0.04	0.07	0.06	0.05	0.06	0.05	
1,1-Dichloroethane	0.10	0.07	0.01	0.03	0.01	0.01	0.03	0.02	0.03	0.03	0.04	0.03	
1,1-Dichloroethene	0.10	0.09	0.03	0.05	0.01	0.01	0.02	0.02	0.03	0.02	0.03	0.02	
1,2,4-Trichlorobenzene	0.67	0.58	0.06	0.15	0.11	0.02	0.07	0.13	0.08	0.09	0.20	0.19	
1,2,4-Trimethylbenzene	1.21	1.01	0.81	0.64	0.50	0.47	0.52	0.70	0.59	0.44	0.43	0.33	
1,2-Dibromoethane	0.19	0.16	0.07	0.05	0.03	0.01	0.04	0.07	0.07	0.06	0.06	0.07	
1,2-Dichloroethane	0.12	0.10	0.03	0.04	0.02	0.02	0.03	0.04	0.08	0.07	0.08	0.07	
1,2-Dichloropropane	0.16	0.12	0.08	0.05	0.02	0.01	0.05	0.05	0.04	0.04	0.04	0.04	
1,3,5-Trimethylbenzene	0.41	0.33	0.25	0.21	0.16	0.15	0.19	0.23	0.23	0.16	0.15	0.10	
1,3-Butadiene	0.21	0.20	0.20	0.16	0.15	0.17	0.14	0.14	0.18	0.15	0.17	0.13	
Acetonitrile	VOID	17.182*	0.59	1.70	6.61	1.24	20.33	0.54	6.03	1.58	5.65	28.47	
Acetylene	2.26	2.05	1.80	1.46	2.02	2.05	1.55	1.38	1.44	1.26	1.31	1.26	
Acrolein		0.81	0.62	0.63	0.68	1.02	1.37	0.74	1.09	0.82	1.13	1.06	
Acrylonitrile	0.11	0.07	0.09	0.04	0.13	0.14	0.04	0.07	0.03	0.17	0.51	0.06	
Benzene	2.25	1.95	1.85	1.46	1.62	1.93	1.41	1.33	1.28	0.99	0.99	0.88	
Bromochloromethane	0.24	0.16	0.05	0.05	0.02	0.01	0.04	0.02	0.04	0.04	0.04	0.04	
Bromodichloromethane	0.13	0.12	0.02	0.06	0.02	0.01	0.06	0.08	0.07	0.06	0.07	0.06	
Bromoform	0.31	0.22	0.09	0.08	0.03	0.01	0.05	0.13	0.10	0.11	0.08	0.09	
Bromomethane	0.11	0.08	0.04	0.05	0.06	0.06	0.08	0.04	0.11	0.08	0.10	0.25	
Carbon Disulfide			8.51	8.71	10.94	13.61	1.19	1.50	1.42	3.02	7.59	0.51	
Carbon Tetrachloride	0.52	0.49	0.59	0.53	0.68	0.66	0.53	0.54	0.67	0.58	0.58	0.60	
Chlorobenzene	0.09	0.07	0.01	0.03	0.02	0.02	0.03	0.05	0.06	0.04	0.04	0.04	
Chloroethane	0.14	0.09	0.03	0.03	0.03	0.04	0.02	0.02	0.03	0.02	0.03	0.08	
Chloroform	0.10	0.11	0.08	0.09	0.11	0.12	0.09	0.09	0.09	0.11	0.11	0.10	
Chloromethane	1.27	1.32	1.21	1.22	1.42	1.47	1.34	1.27	1.24	1.15	1.22	1.35	
Chloroprene	0.09	0.07	0.04	0.03	0.02	0.01	0.02	0.01	0.02	0.02	0.03	0.02	
cis-1,2-Dichloroethylene	0.12	0.09	0.03	0.04	0.01	0.01	0.07	0.02	0.04	0.03	0.03	0.03	
cis-1,3-Dichloropropene	0.11	0.08	0.03	0.04	0.02	0.01	0.03	0.05	0.03	0.03	0.05	0.04	
Dibromochloromethane	0.30	0.20	0.04	0.06	0.02	0.02	0.04	0.09	0.07	0.07	0.06	0.06	

	Annual Average VOC Concentrations (μg/m³)											
Analyte	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013	2014	2015
Dichlorodifluoromethane	3.07	3.18	2.78	2.70	2.79	3.22	2.90	2.76	2.57	2.54	2.54	2.57
Dichloromethane	0.49	0.43	0.41	0.38	3.43	1.96	91.65	1.31	40.12	15.89	40.28	83.49
Dichlorotetrafluoroethane	0.11	0.12	0.12	0.12	0.11	0.14	0.13	0.13	0.12	0.12	0.12	0.12
Ethyl Acrylate	0.12	0.11	0.02	0.03	0.04	0.01	0.02	0.04	0.03	0.03	0.04	0.02
Ethyl tert-Butyl Ether	0.10	0.10	0.02	0.02	0.01	0.04	0.02	0.02	0.02	0.06	0.16	0.05
Ethylbenzene	1.20	1.36	0.66	0.61	0.47	0.53	0.51	0.62	0.69	0.48	0.45	0.39
Hexachloro-1,3-butadiene	0.85	0.99	0.07	0.19	0.09	0.01	0.06	0.19	0.12	0.14	0.15	0.18
m,p-Xylene	3.73	4.62	2.29	2.05	1.53	1.70	1.55	1.97	2.10	1.45	1.35	1.03
m-Dichlorobenzene	0.21	0.17	0.01	0.05	0.05	0.02	0.03	0.10	0.07	0.08	0.05	0.07
Methyl Isobutyl Ketone	0.27	0.18	0.21	0.17	0.17	0.15	0.17	0.16	0.17	0.15	0.17	0.15
Methyl Methacrylate	1.29	0.79	0.26	1.34	0.49	0.05	0.05	0.05	0.06	0.07	0.06	0.06
Methyl tert-Butyl Ether	0.13	0.12	0.01	0.02	0.01	0.01	0.02	0.02	0.02	0.03	0.05	0.03
n-Octane	0.33	0.34	0.24	0.24	0.20	0.23	0.30	0.37	0.42	0.29	0.28	0.21
o-Dichlorobenzene	0.12	0.15	0.02	0.05	0.05	0.02	0.03	0.10	0.06	0.07	0.05	0.07
o-Xylene	1.55	1.97	0.83	0.73	0.56	0.60	0.55	0.71	0.79	0.53	0.53	0.49
p-Dichlorobenzene	0.18	0.14	0.09	0.07	0.04	0.07	0.04	0.09	0.08	0.06	0.05	0.08
Propylene	1.41	1.32	1.11	0.91	0.88	1.01	0.88	0.86	0.95	0.79	0.84	0.72
Styrene	2.19	1.05	0.37	0.58	1.26	0.63	2.57	1.45	2.96	1.91	3.47	16.47
tert-Amyl Methyl Ether	0.15	0.13	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.03	0.02
Tetrachloroethylene	0.31	0.27	0.34	0.32	0.33	0.43	0.40	0.26	0.32	0.27	0.23	0.15
Toluene	5.58	5.53	4.06	4.22	2.91	3.82	3.23	4.01	3.66	2.96	2.91	3.18
trans-1,2-Dichloroethylene	0.10	0.09	0.04	0.03	0.01	0.01	0.03	0.02	0.03	0.02	0.03	0.03
trans-1,3-Dichloropropene	0.12	0.10	0.02	0.04	0.02	0.01	0.03	0.06	0.04	0.04	0.05	0.05
Trichloroethylene	0.13	0.12	0.05	0.06	0.03	0.11	0.06	0.09	0.14	0.05	0.05	0.05
Trichlorofluoromethane	2.17	1.63	1.52	1.46	1.51	1.71	1.60	1.52	1.59	1.45	1.33	1.39
Trichlorotrifluoroethane	0.78	0.81	0.76	0.83	0.68	0.85	0.72	0.75	0.66	0.63	0.61	0.63
Vinyl chloride	0.05	0.05	0.01	0.03	0.01	0.01	0.02	0.01	0.01	0.01	0.02	0.01

2004 NOTE: Acetonitrile VOID due to contamination in sampler.

2005 NOTE: Acetonitrile VOID thru 4/10/2005 due to contamination in sampler.

**Bold = MQO Core Analyte** 

Italic = Less than 90% detection rate (2004-2014), less than 10% detection rate (2015)

In general, the concentrations from 2015 compared well with the 2014 data. However, some compounds did show average concentrations that were significantly different than their 2014 values. For instance, dichloromethane and acetonitrile showed much larger annual average concentrations in 2015 than in 2014. In 2014, their respective annual average concentrations were  $40.28 \pm 82.51$  and  $5.65 \pm 8.77 \ \mu g/m^3$ , while in 2015 they were  $83.49 \pm 214.8$ , and  $28.47 \pm 54.95 \ \mu g/m^3$ . The 2013 National Monitoring Programs Annual report, written by Eastern Research Group for the EPA, shows the averages for these compounds to be  $8.16 \pm 141.4$  and  $16.96 \pm 84.95 \ \mu g/m^3$ , respectively. The large change in concentrations arises from significantly elevated concentrations of these compounds on several sample days in 2015. Elevated dichloromethane concentrations were also seen in 2010 and 2012. At this point in time it has not been determined what is causing these highly variable concentrations.

# Graphs

Figure 10 through Figure 13 are graphs showing the 24 hour maximum, and annual mean concentrations for each of the 37 compounds that were detected in greater than 10% of the samples in 2015 and vinyl chloride, which was not detected in any samples but is included because it is on the core HAPS list. These graphs are ordered from highest to lowest annual mean concentration. Note that the graphs' scales vary from a full-scale level at 20 micrograms per meter cubed to a full-scale value of 0.25 micrograms per meter cubed. The compounds with the five largest annual average concentrations are dichloromethane, acetonitrile, styrene, toluene, and dichlorodifluoromethane. Their values are  $83.49 \pm 214.8$ ,  $28.47 \pm 54.95$ ,  $16.47 \pm 18.73$ ,  $3.18 \pm 2.32$ , and  $2.57 \pm 0.44$  micrograms per meter cubed, respectively. In comparison, the 2013 national averages for the same compounds are  $8.16 \pm 141.4$ ,  $16.96 \pm 84.95$ ,  $0.089 \pm 3.30$ ,  $1.13 \pm 2.46$ , and  $2.52 \pm 0.36$  micrograms per meter cubed, respectively.

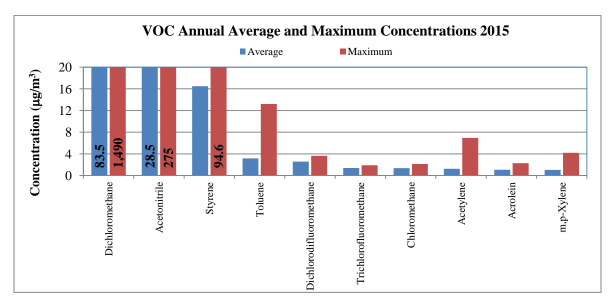


Figure 10. VOC Annual and Maximum Concentrations 2015

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<sup>&</sup>lt;sup>5</sup> "2013 National Monitoring Programs Annual Report (UATMP, NATTS, CSATAM). US EPA. October 2015.

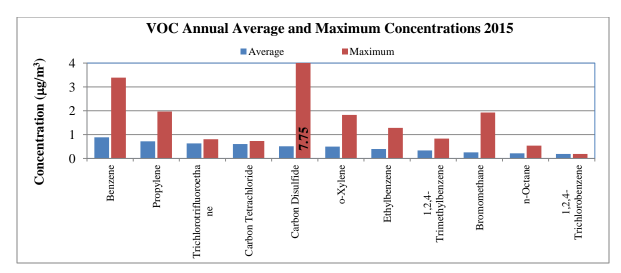


Figure 11. VOC Annual and Maximum Concentrations 2015, ctd.

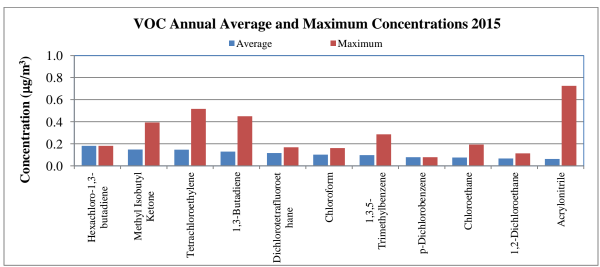


Figure 12. VOC Annual and Maximum Concentrations 2015, ctd.

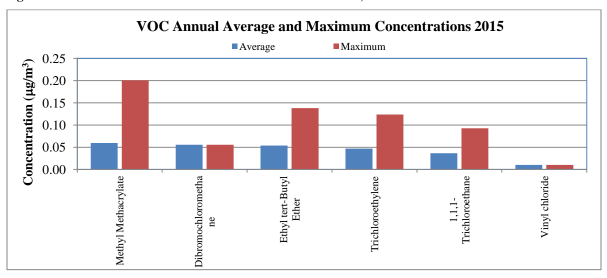


Figure 13. VOC Annual and Maximum Concentrations 2015, ctd.

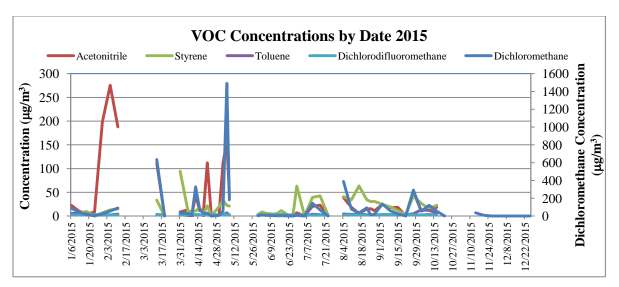


Figure 14. VOC Concentrations by Date 2015

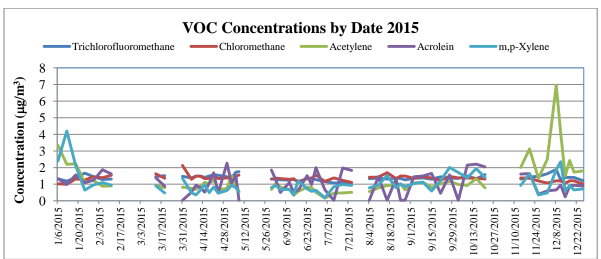


Figure 15. VOC Concentrations by Date 2015, ctd.

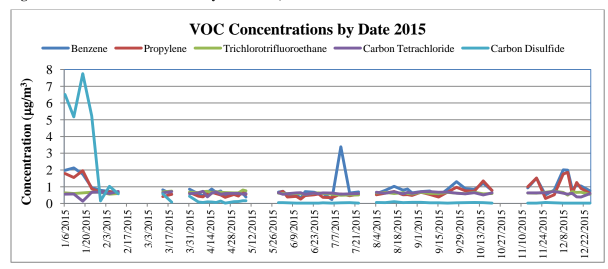


Figure 16. VOC Concentrations by Date 2015, ctd.

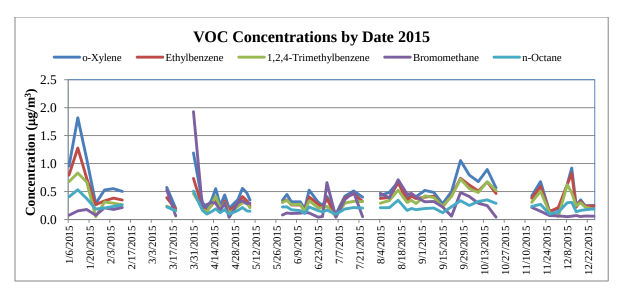


Figure 17. VOC Concentrations by Date 2015, ctd.

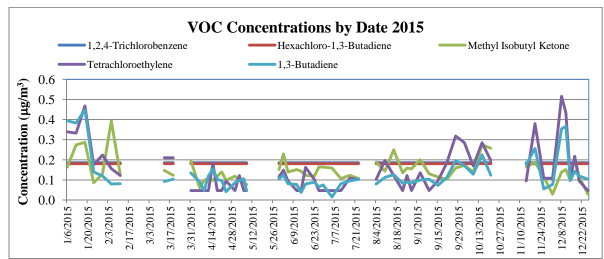


Figure 18. VOC Concentrations by Date 2015, ctd.

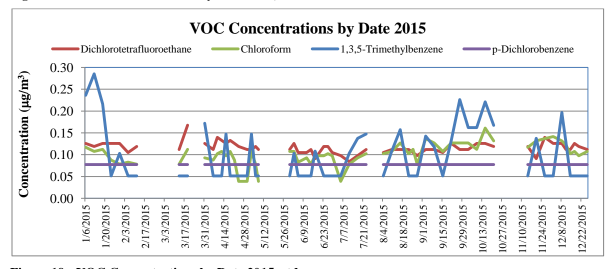


Figure 19. VOC Concentrations by Date 2015, ctd.

Figure 14 through Figure 19 show the concentrations of the 37 most detected VOCs, as well as vinyl chloride, in chronological order. There are several gaps in the 2015 data. This was due to having to void several canister samples as a result of operator errors or equipment malfunction. The remaining sample concentrations tended to trend well with each other. It should be noted, in Figure 14, the dichloromethane values had to be plotted on a separate scale from the other compounds. The scale on the right hand side of the graph, with a range of zero to 1,600 micrograms per cubic meter, applies only to the dichloromethane concentrations. The other compound concentrations are scaled against the left hand side of the graph. Some of the compounds do show a seasonal variation in their concentrations, though it is difficult to discern the variation in this year's data set given the large gaps of missing data. There are several compounds that show this seasonal variability in Figure 18. VOC concentrations are typically higher in the summer due to the higher temperatures, and longer availability of ultraviolet rays for the photolytic process to occur.

Figure 14 also shows large concentrations of dichloromethane for several months during 2015, exhibiting a maximum value of 1,490 micrograms per cubic meter. This value is more than double the next closest maximum seen in 2012, with a value of over 700 micrograms per cubic meter. No direct link has yet been found to explain why these values are increasing.

Figure 20 through Figure 23 graphically illustrate the weekday versus weekend VOC concentrations in 2015 for the 38 compounds. The compounds are separated into four groups: alkanes, alkenes, alkynes, and aromatics. The alkane compounds have carbon atoms with only single bonds. The alkenes have carbon atoms with double bonds, and the alkynes have triple bonds. The aromatics are ring structures (e.g., benzene) with other substituents bonded to the ring.

In 2015, weekday concentrations for 23 of the 38 compounds were larger than concentrations captured during the weekend sampling. This is expected, as many of the compounds emitted are associated with automobile emissions, and traffic in the area usually decreases on the weekends. There were several compounds with larger weekend averages, however. Six of the 38 compounds had higher weekend concentrations than weekday concentrations. The remaining compounds had weekday and weekend averages that were equal. This is most likely attributable to the practice of substituting of one-half the value of the MDL for the non-detect and less than MDL sample values.

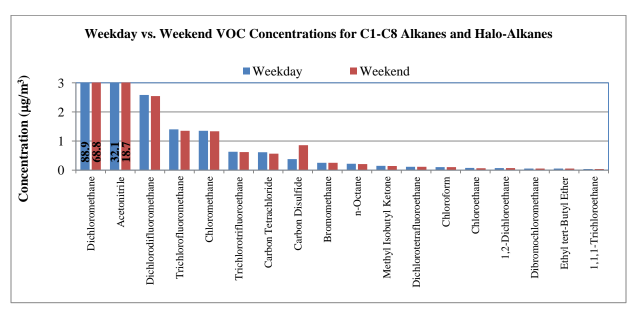


Figure 20. VOC Weekday vs. Weekend Comparison for C1-C8 Halo-Alkanes

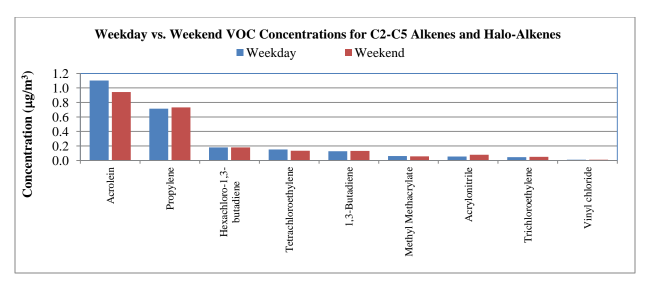


Figure 21. VOC Weekend vs. Weekday Concentrations for C2-C4 Halo-Alkenes

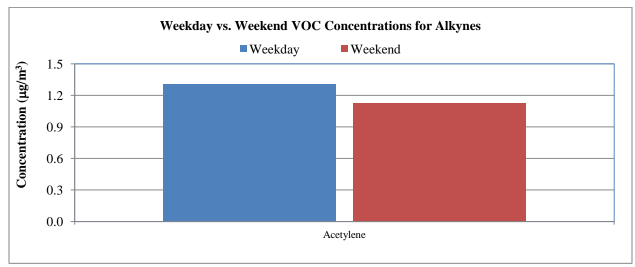


Figure 22. VOC Weekend vs. Weekday Concentrations for Alkynes

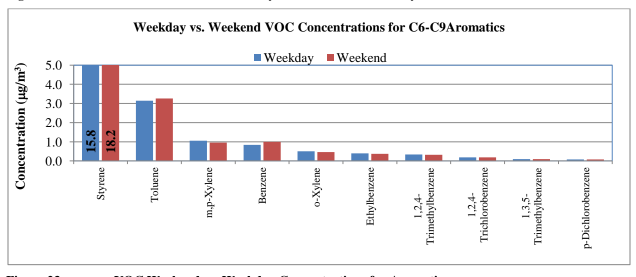


Figure 23. VOC Weekend vs. Weekday Concentrations for Aromatics

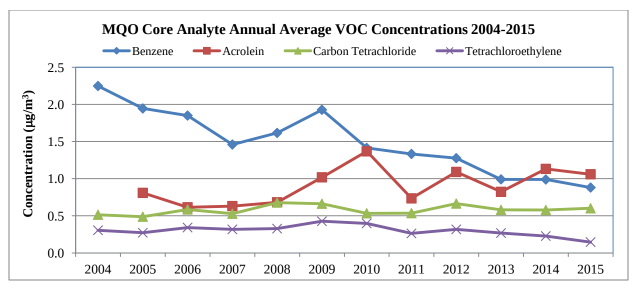


Figure 24. MQO Core Analyte VOC Concentrations 2004 – 2015

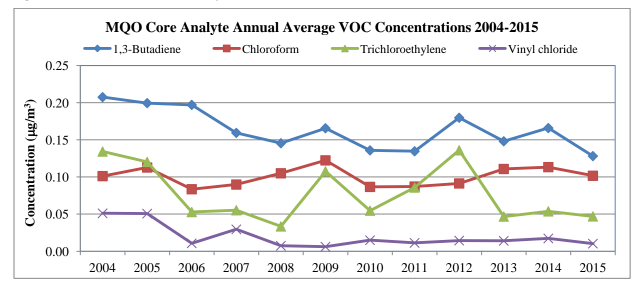


Figure 25. MQO Core Analyte VOC Concentrations 2004 – 2015, ctd.

Figure 24 and Figure 25 graph the annual average concentrations of the eight VOCs that are a part of the mandatory monitoring subset of 19 HAPs. The benzene and 1,3-butadiene graphs appear to show a downward trend since 2004. The remaining graphs do not appear to indicate a general trend in concentration values since 2004. Annual average benzene concentrations have trended downward since 2004. Figure 26 and Figure 27graphically illustrate how the 3-year average concentrations of the eight MQO Core Analyte VOCs have trended since the Pilot Study began in 2004.

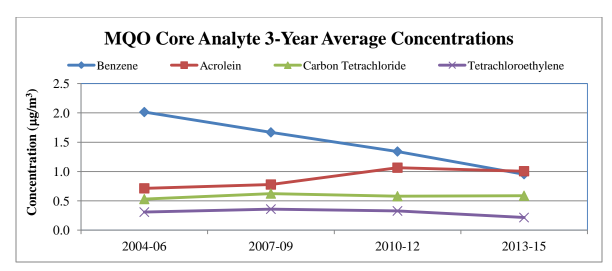


Figure 26. MQO Core Analyte 3 year Average VOC Concentrations 2004 – 2015

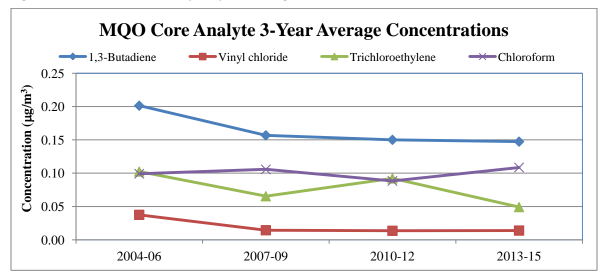


Figure 27. MQO Core Analyte 3 Year Average VOC Concentrations 2004 – 2015

## Quality Assurance/Quality Control

#### **Blanks**

The volatile organic compound sampling method involves sampling in stainless steel canisters with specially treated interior surfaces. The canisters are re-used. After a full canister is analyzed, it is pumped out repeatedly to a high vacuum. This procedure cleans it for the next use. Periodically, one canister from each cleaning batch is tested to make sure the method is performing adequately. The test canister is filled with ultra-pure air, and then analyzed. If it shows no contamination, the batch is released for use. If contamination is found, the entire batch is sent through the cleaning process for a second time. The canisters arrive in the field closed, and under a vacuum of 20 to 30 inches of mercury. Therefore, field blanks are not used in this method. The method blank canister test is performed at the laboratory prior to shipping to the field.

## **Precision of Sample Results**

On six random sampling dates per year, a second canister was sampled simultaneously with the primary

sample. These additional samples, known as duplicates, were collected in order to assess the precision (repeatability) of the canister sampling method. In general, repeatability for the two collocated samples was acceptable. Information regarding precision and accuracy results is available upon request to the Air Pollution Control Division.

# V. POLYCYCLIC AROMATIC HYDROCARBONS

# **Summary Statistics**

In April 2008, the Grand Junction National Air Toxics Trends Site added a sampler for polycyclic aromatic hydrocarbon (PAH) compounds. A good definition of these chemicals is:

Polycyclic aromatic hydrocarbons (also known as polynuclear aromatic hydrocarbons) are composed of two or more aromatic (benzene) rings which are fused together when a pair of carbon atoms is shared between them. The resulting structure is a molecule where all carbon and hydrogen atoms lie in one plane. Naphthalene ( $C_{10}H_8$ , MW = 128.16 g), formed from two benzene rings fused together, has the lowest molecular weight of all PAHs. The environmentally significant PAHs are those molecules which contain two (e.g., naphthalene) to seven benzene rings (e.g., coronene with a chemical formula  $C_{24}H_{12}$ ; MW = 300.36 g). In this range, there are a large number of PAHs which differ in number of aromatic rings, position at which aromatic rings are fused to one another, and number, chemistry, and position of substituents on the basic ring system. (Source: Ambient Water Quality Criteria for Polycyclic Aromatic Hydrocarbons (PAHs) Ministry of Environment, Lands and Parks, Province of British Columbia. By N. K. Nagpal, Ph.D., Water Quality Branch, Water Management Division, British Columbia, Canada, Ministry of Environment, February, 1993).

In 2015, 60 PAH samples were possible. Of those 60, 10 were either missed or invalidated by the lab for errors during the analytical process. Five were made up, for a total of 55 out of 60 samples (91.7% sample recovery rate). Twenty-two compounds were measured for this study. The list of these compounds and the summary of the collected data are shown in Table 6 and Table 7. All of the 22 compounds analyzed for were detected in more than 10% of the samples taken. Perylene was found to have the lowest detection rate, with only 38% of the samples taken showing levels above the compound's MDL. Thirteen compounds were detected in every sample taken. This is an increase from the 2014, where only eleven compounds were present in all samples taken.

Table 6. PAH Sample Summary Data 2015

	CAS		
Compound	Number	# of ND's	% ND
9-Fluorenone	486-25-9	0	0%
Acenaphthene	83-32-9	0	0%
Acenaphthylene	208-96-8	15	27%
Anthracene	120-12-7	0	0%
Benzo (a) anthracene	56-55-3	0	0%
Benzo (a) pyrene	50-32-8	5	9%
Benzo (b) fluoranthene	205-99-2	0	0%
Benzo (e) pyrene	192-97-2	1	2%
Benzo (g,h,i) perylene	191-24-2	0	0%
Benzo (k) fluoranthene	207-08-9	30	55%
Chrysene	218-01-9	0	0%
Coronene	191-07-1	1	2%
Cyclopenta[cd]pyrene	27208-37-3	22	40%
Dibenz (a,h) anthracene	53-70-3	16	29%

Compound	CAS Number	# of ND's	% ND
Fluoranthene	206-44-0	0	0%
Fluorene	86-73-7	5	9%
Indeno(1,2,3-cd)pyrene	193-39-5	0	0%
Naphthalene	91-20-3	0	0%
Perylene	198-55-0	34	62%
Phenanthrene	85-01-8	0	0%
Pyrene	129-00-0	0	0%
Retene	483-65-8	0	0%

ND = Not Detected

**Bold = MQO Core Analyte** 

Table 7 summarizes the annual mean concentrations for each PAH measured during the study, from 2008 through 2015. The compounds that were detected in less than 10% of the samples taken in 2015 are italicized to show that their averages are dependent on their respective MDL values. From 2008 through 2014 the italicized results indicate compounds that were detected in less than 90% of the samples taken. It should be noted here that there was a shift in the data processing methods for this report. In previous years, data was presented for compounds that were present in at least 90% of the samples taken. For this (and future) reports the data presented will be from compounds that were present in at least 10% of the samples taken. While this move did not have a noticeable effect on the carbonyl or PAH data, it did allow for inclusion of more of the VOC compounds discussed in earlier sections. For purposes of consistency, this new 10% detection criterion was applied to all carbonyl, VOC, metal, and SVOC data sets.

Bolded compounds are listed among those on the list of 19 core HAPs to be monitored. The annual means were calculated by replacing all "non-detect" values with one-half of the sample MDL. This is an accepted conservative technique for calculating annual values when some of the samples were less than the laboratory's ability to detect. Naphthalene had the largest annual average of the PAH compounds with a value of just over 90 nanograms per meter cubed in 2015. This is more than four times greater than the next closest average concentration, which is phenanthrene, with 22.1 nanograms per meter cubed.

Table 7. PAH Annual Average Values 2008 - 2015

	2008	2009	2010	2011	2012	2013	2014	2015
Analyte	Average (ng/m³)							
9-Fluorenone	1.53	2.67	2.34	2.13	2.74	2.42	2.09	2.26
Acenaphthene	8.41	11.34	7.30	10.54	20.5	8.07	7.17	8.29
Acenaphthylene	2.12	3.68	2.50	2.22	2.28	2.51	1.33	1.17
Anthracene	0.63	1.65	0.89	0.77	0.83	0.82	0.66	1.28
Benzo (a) anthracene	0.20	0.39	0.25	0.26	0.22	0.31	0.22	0.26
Benzo (a) pyrene	0.18	0.33	0.20	0.22	0.18	0.24	0.19	0.24
Benzo (b) fluoranthene	0.36	0.72	0.50	0.48	0.41	0.56	0.44	0.31
Benzo (e) pyrene	0.19	0.39	0.24	0.23	0.19	0.25	0.22	0.22
Benzo (g,h,i) perylene	0.26	0.43	0.28	0.25	0.21	0.26	0.25	0.23
Benzo (k) fluoranthene	0.10	0.21	0.14	0.14	0.12	0.14	0.12	0.09
Chrysene	0.35	0.68	0.49	0.48	0.42	0.54	0.41	0.48
Coronene	0.15	0.23	0.13	0.11	0.09	0.10	0.12	0.12
Cyclopenta[cd]pyrene	0.16	0.19	0.10	0.13	0.12	0.11	0.07	0.18
Dibenz (a,h) anthracene	0.06	0.06	0.03	0.05	0.04	0.03	0.02	0.05
Fluoranthene	2.52	3.79	3.30	3.35	3.55	3.36	2.60	5.48

Analyte	2008 Average (ng/m³)	2009 Average (ng/m³)	2010 Average (ng/m³)	2011 Average (ng/m³)	2012 Average (ng/m³)	2013 Average (ng/m³)	2014 Average (ng/m³)	2015 Average (ng/m³)
Fluorene	5.15	9.20	6.44	7.67	12.6	6.89	5.75	7.34
Indeno(1,2,3-cd)pyrene	0.21	0.37	0.24	0.23	0.19	0.25	0.25	0.20
Naphthalene	112	189	147	158	204	137	100	90.3
Perylene	0.07	0.08	0.09	0.07	0.06	0.04	0.03	0.04
Phenanthrene	12.0	17.9	13.9	14.0	18.7	13.3	11.1	22.1
Pyrene	1.81	2.87	2.28	2.19	2.20	2.30	1.79	3.01
Retene	0.67	1.37	1.04	0.85	0.77	1.06	0.74	0.71

**Bold = MQO Core Analyte** 

Italic = less than 90% detection rate 2008 – 2014

# **Graphs**

Chronological graphs of the concentration data for the PAH compounds are shown in Figure 28 through Figure 31. Naphthalene is the most variable, with concentrations ranging from 11.1 to 320 nanograms per meter cubed. Naphthalene had the largest annual average concentration, followed by phenanthrene, with values of 90.3, and 22.1 nanograms per meter cubed, respectively. In comparison, the National Monitoring Program (NMP) national averages for these compounds in 2013 were  $75.3 \pm 73.0$ , and  $9.86 \pm 17.9$  nanograms per meter cubed, respectively.<sup>6</sup>

Overall, the PAH compound concentrations tended to follow the same general trend. Many of the compounds showed larger concentrations during the summer months, as is expected. This is likely due to smoke in the area caused by regional forest fires during that time period. Several other compounds exhibited a seasonal variation, with larger concentrations in the winter months, and lower concentrations in the summer months. This also makes sense, since the primary source of many PAHs in air is the incomplete combustion of wood and fuel.<sup>7</sup> PAHs are a product of combustion from common sources like automobiles, wood-burning stoves and furnaces, cigarette smoke, etc. The natural sources of PAHs include volcanoes, forest fires, crude oil, and shale oil.<sup>8</sup>

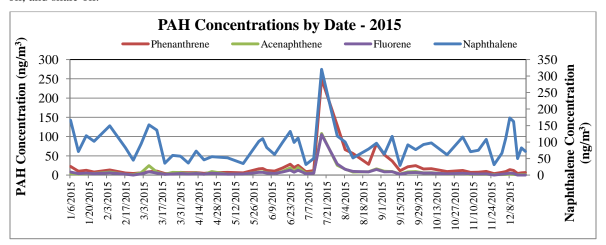


Figure 28. PAH Concentrations by Date 2015

[24]

<sup>6 &</sup>quot;2013 National Monitoring Programs Annual Report (UATMP, NATTS, CSATAM). US EPA. October 2015.

<sup>&</sup>lt;sup>7</sup> "Toxicological Profile for Polycyclic Aromatic Hydrocabons." US Department of Health and Human Services, Agency for Toxic Substances and Disease Resigtry. August 1995. http://www.atsdr.cdc.gov/ToxProfiles/tp69.pdf

<sup>&</sup>lt;sup>8</sup> Ibid.

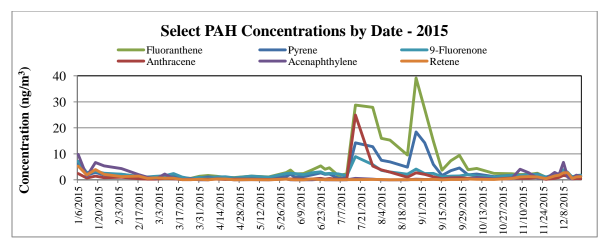


Figure 29. PAH Concentrations by Date 2015, ctd.

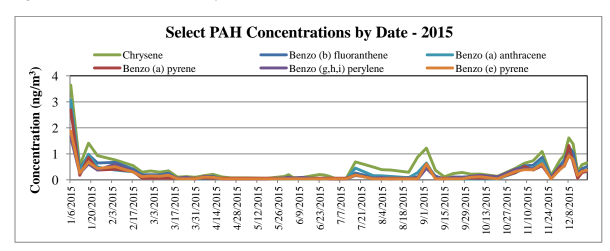


Figure 30. PAH Concentrations by Date 2015, ctd.

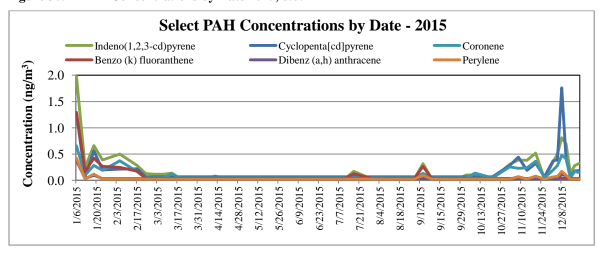


Figure 31. Select PAH Concentrations by Date 2015, ctd.

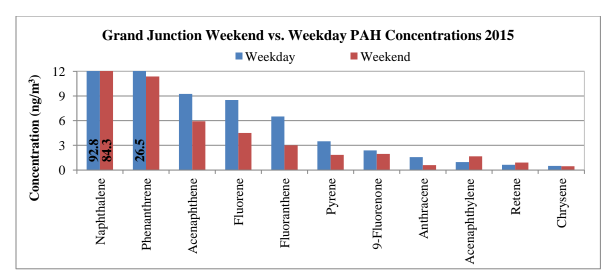


Figure 32. PAH Weekend vs. Weekday Concentrations 2015

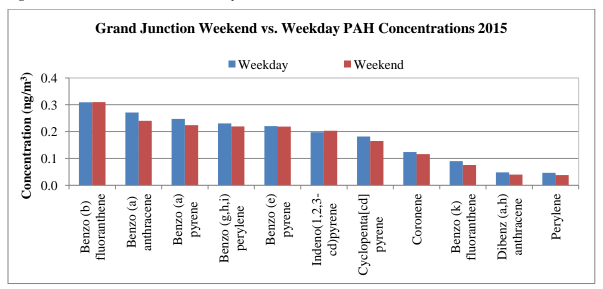


Figure 33. PAH Weekend vs. Weekday Concentrations 2015, ctd.

Figure 32 and Figure 33 are graphs of the weekend versus weekday concentrations for all the PAH compounds in 2015. The weekday averages were larger than the weekend values for all but 4 compounds: acenaphthylene, retene, benzo(b)fluoranthene, and indeno(1,2,3-cd)pyrene. Acenaphthylene was detected in only 40 of the 55 samples taken, while the other three compounds were detected in all samples. The average for benzo(b)fluoranthene is equal for the weekdays and weekends. This would seem to indicate that ambient concentrations are at background levels. The values for naphthalene are off the chart with a weekday average of 92.8 nanograms per meter cubed, and a weekend average of 84.3 nanograms per meter cubed, which are both lower than their respective 2014 values. Figure 34 through Figure 37 are graphs of the annual average concentrations for all the PAH compounds. The graphs show that several of the annual average compound concentrations have increased since 2012, while others have decreased or remained fairly constant.

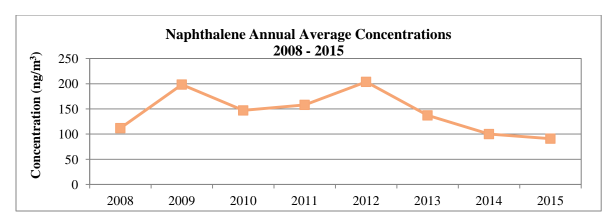


Figure 34. Naphthalene Annual Average Concentrations 2008 – 2015

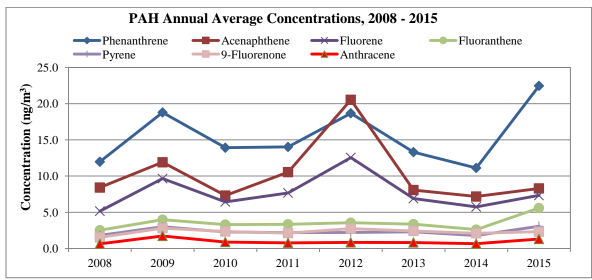


Figure 35. PAH Annual Average Concentrations 2008 – 2015

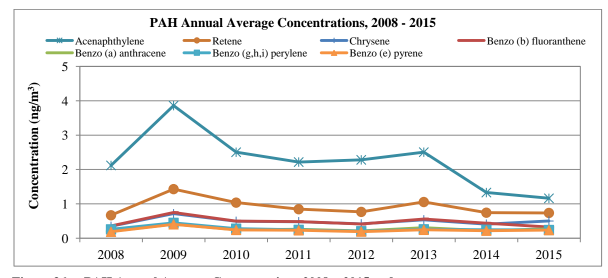


Figure 36. PAH Annual Average Concentrations 2008 – 2015, ctd.

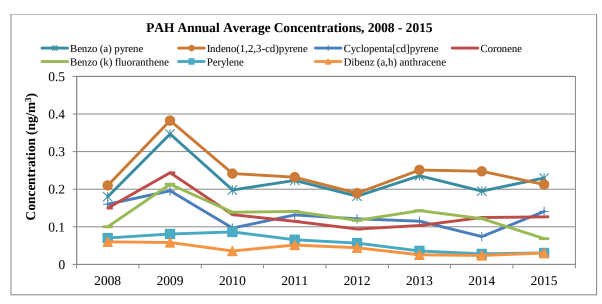


Figure 37. PAH Annual Average Concentrations 2008 - 2015, ctd.

# Quality Assurance/Quality Control

#### **Blanks**

Periodically, the laboratory analyzes a "blank," or unused, filter for PAH compounds. The purpose of this extra analysis is to determine if there was any contamination of the filter during manufacturing, or during laboratory processing. In 2015, the laboratory analyzed 11 "filter blanks," filters which never left the lab. Several compounds were detected at very low levels in many of the filter blanks. The blank concentrations were averaged, and subtracted from the sample concentrations before annual averages were calculated. Results of the blank filters are available upon request from CDPHE.

#### **Precision of Sample Results**

Precision air samples were not run at this site in 2015. Assessing precision requires a collocated sampler at the site, and the NATTS group chose to take precision samples at other locations in the nationwide network.

# VI. PM<sub>10</sub> METALS

In 2014, CDPHE switched to a different contract lab for metals analyses, as previous years' data were found to have various errors due to the contracted laboratory not following correct procedures for establishing the method detection limits (MDLs). The reported concentrations for some of the metals rely heavily on the MDL values, as one-half the value of the MDL is substituted for the concentration in instances where the metal is not detected during the analysis, or is detected but it's value is less than the MDL. Because it was impossible to go back and calculate the MDLs being used for the previous data, a new MDL study was performed by the lab in 2014. The values obtained as a result of that study are used for the analysis of 2010-2013 metals data, in an effort to keep from losing several years' worth of valuable data.

In previous years, antimony and total chromium were also a part of the suite of compounds CDPHE had the lab analyze for. These two compounds are not required as a part of the NATTS program. As such, when the new MDL study was performed they were dropped from the list of compounds to be recalculated. Any data associated with those two compounds for the years 2010 through 2013 has consequently been removed.

## **Summary Statistics**

In 2015, as in past years, metals were sampled on the every sixth day schedule, for a total of 60 samples attempted. Of those 60 samples, three were missed or voided, leaving a total of 57 samples collected (95% sample recovery). Table 8 shows the percentage of the samples in which each metal was either not detected, or had a concentration value that was below the MDL value for that compound. Some compounds were detected on many of the sample dates, but at concentrations that were less than their respective MDL. For instance, chromium was detected in all of the samples taken, having an average of 4.26 ng/m³ (using the raw data). However, the MDL for chromium is 11.92 ng/m³. Therefore, a value equal to one-half the MDL (5.96 ng/m³) was substituted in place of the listed raw value as a conservative, reliable estimate of the compound's concentration. Since the actual raw value is under the MDL, it is difficult to use it with any certainty. Although there is a peak on the graph at the proper time indicating that it is chromium, it could also be attributed to instrument noise as it is less than the instrument's ability to accurately determine the compound. The fact that the chromium average in the blank samples taken throughout the year was near the raw average of the samples (3.57 ng/m³ for blanks versus 4.26 ng/m³ for samples) supports this.

There were a total of only nine actual non-detect values for the entire metals data set in 2015. These non-detect values were spread between four compounds: beryllium, cadmium, lead, and selenium. Beryllium was listed as non-detect for 5 of those nine samples, selenium for two, and cadmium and lead for one each. The non-detect percentages seen for mercury and chromium in Table 8 are due to the compounds having detected values that were less than their respective MDLs.

Table 8. Metals Detection Rates, 2015

	CAS			
Compound	Number	# of ND's	% ND	
Antimony	7440-36-0	0	0%	
Arsenic	7440-38-2	0	0%	
Beryllium	7440-41-7	1	2%	
Cadmium	7440-43-9	0	0%	
Chromium	7440-47-3	57	100%	
Cobalt	7440-48-4	0	0%	
Lead	7439-92-1	0	0%	
Manganese	7439-96-5	0	0%	
Mercury	7439-97-6	53	93%	
Nickel	7440-02-0	8	14%	
Selenium	7782-49-2	0	0%	

**Bold = MQO Core Analyte** 

Table 9 summarizes the annual mean concentrations for each of the metals measured during the study, from 2004 through 2015. The compounds that are listed in bold type are on the list of 19 core HAPs. The italicized compounds are those that were detected in less than 10% of the samples taken. Annual means were calculated by using one-half of the detection limits in place of the non-detect and less than MDL samples. In instances where subtracting the blank value made the resulting concentration less than zero the data were treated as non-detect samples. Non-detect sample values are those that were listed as "ND" by the lab, or those mentioned above. Results show that manganese was the compound with the highest annual average. The other metals were present at lower concentrations. The data for antimony and chromium from 2010 through 2013 has been removed due to lab issues discussed previously. CDPHE's new contract lab analyzes for more compounds than are required by the NATTS technical assistance document. As such, there are new compounds that have been added to the analytical suite. CDPHE has no data for these compounds prior to 2014.

Table 9. **Metals Data Summary 2015** 

Year	Antimony (ng/m³)	Arsenic (ng/m³)	Beryllium (ng/m³)	Cadmium (ng/m³)	Chromium (ng/m³)	Cobalt (ng/m³)	Lead (ng/m <sup>3</sup> )	Manganese (ng/m³)	Mercury (ng/m³)	Nickel (ng/m³)	Selenium (ng/m³)
2004	0.36	0.27	0.08	0.05	1.86		4.91	13.0		0.63	
2005	1.34	2.13	0.91	0.35	31.7		4.01	12.0		0.91	
2006	1.47	2.88	0.59	0.26	13.1		4.33	15.0		1.19	
2007	0.99	4.22	0.68	0.24	16.8		4.26	15.2		1.44	
2008	1.08	2.43	0.19	0.14	8.75		2.48	14.7		1.43	
2009	0.54	0.87	0.13	0.23	8.83		2.09	8.70		0.88	
2010		1.32	0.14	0.20			2.05	8.34		1.80	
2011		0.67	0.14	0.21			2.79	8.82		2.11	
2012		0.57	0.13	0.13			3.03	10.8		2.26	
2013		0.37	0.08	0.11			2.75	7.89		2.64	
2014	0.96	0.30	0.02	0.08	7.86	0.17	2.64	9.39	0.02	0.57	0.71
2015	0.73	0.31	0.01	0.06	5.96	0.12	2.09	7.09	0.01	0.55	0.68

**Bold = MQO Core Analyte**Italic = less than 90% detection rate 2004-2014, less than 10% detection rate 2015.

# Graph

The annual average metals concentrations are graphed in Figure 38. The figure shows that manganese and lead were the metals with the largest average concentrations, having values of 7.00 and 2.08 nanograms per meter cubed, respectively. While the concentration for chromium is shown for comparative purposes, it is important to note that the value is dependent only on it's MDL, as it was never detected at levels higher than that. In comparison, the NMP national average concentrations for these compounds in 2013 were  $7.87 \pm 9.43$  and  $3.57 \pm 4.28$  nanograms per meter cubed, respectively. Figure 39 and Figure 40indicate that most of the metals were at low concentration levels throughout the year. There does not appear to be any seasonal trending in the metals values based on the 2015 data. Manganese has the largest amount of variability in the concentration values recorded, with values ranging from 0.68 to over 21 nanograms per meter cubed. The data for mercury and chromium are not graphed due to their MDL dependence.

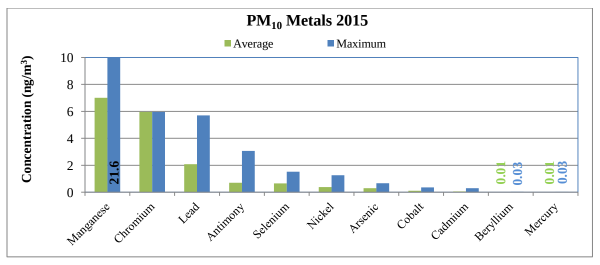


Figure 38. PM<sub>10</sub> Metals Average and Maximum Concentrations 2015

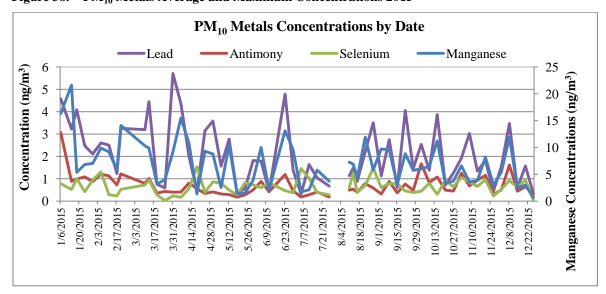


Figure 39. PM<sub>10</sub> Metals Concentrations by Date 2015

[31]

 $<sup>^9</sup>$  "2013 National Monitoring Programs Annual Report (UATMP, NATTS, CSATAM). US EPA. October 2015.

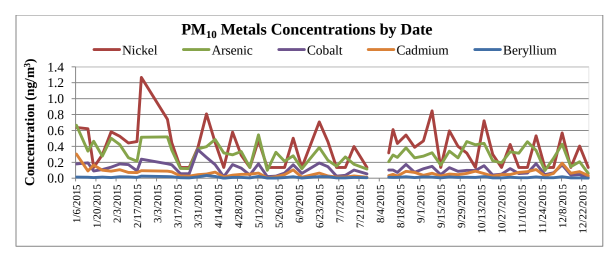


Figure 40. PM<sub>10</sub> Metals Concentrations by Date 2015, ctd.

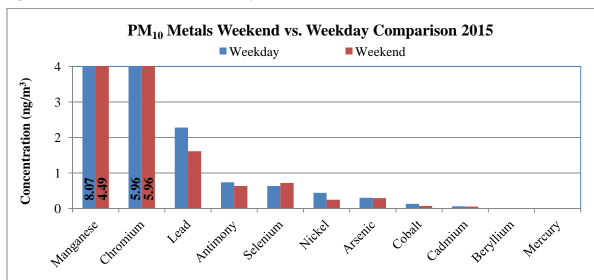


Figure 41. PM<sub>10</sub> Metals Weekend versus Weekday Comparison 2015

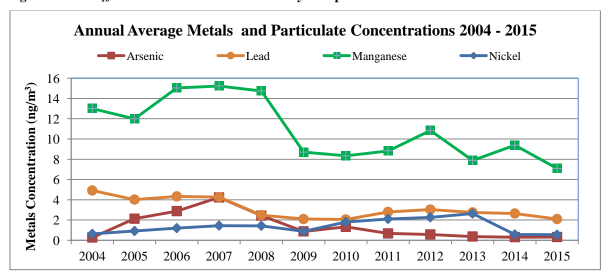


Figure 42. PM<sub>10</sub> Metals Annual Average Concentrations 2004 – 2015

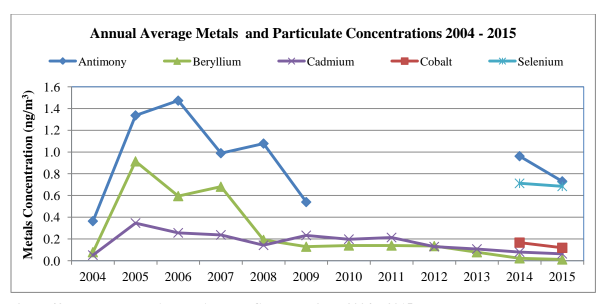


Figure 43. PM<sub>10</sub> Metals Annual Average Concentrations, 2004 – 2015, ctd.

Figure 41 is a chart of the weekend versus weekday concentrations for the  $PM_{10}$  metals. All of the compounds except selenium had weekend averages that were equal to or less than the weekday averages. Chromium and mercury were not detected often, meaning the concentration values are heavily dependent on their MDL values, resulting in equal weekend and weekday concentrations. Figure 42 and Figure 43 are graphs of the annual average concentrations (2004 – 2015) for each of the  $PM_{10}$  metals, except chromium and mercury, as they were detected at levels lower than their respective MDLs. The graphs appear to show a general downward trend in the concentration values for many of the metals. A calculation of the 3-year averages from 2004 to 2006, 2007 to 2009, 2010 to 2012, and 2013 to 2015 shows a decrease in concentrations (going from the first 3-year average to the last 3-year average) for most compounds. These can be seen in Figure 44 and Figure 45.

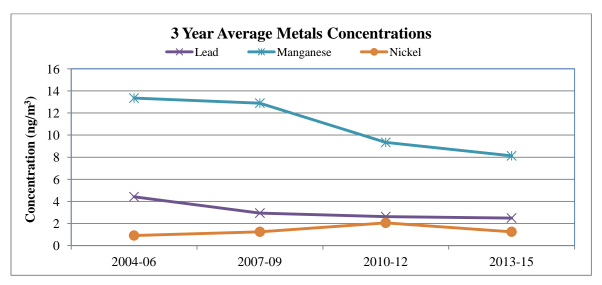


Figure 44. 3-Year Average Metals Concentrations

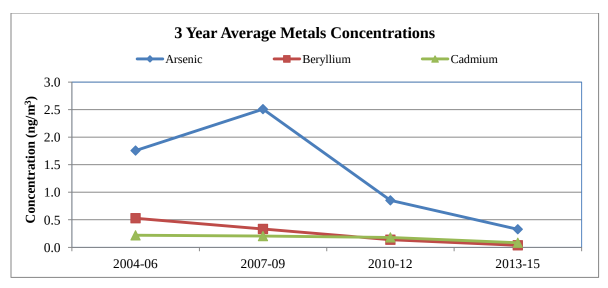


Figure 45. 3-Year Average Metals Concentrations (ctd.)

## Quality Assurance/Quality Control

#### **Blanks**

Periodically, the laboratory analyzes a "blank" or unused, filter for metals. The purpose of this extra analysis is to determine if there was any contamination of the filter during manufacturing, the sample setup/tear down or transport processes, or during laboratory processing. In 2015, there were six blank samples taken and analyzed by the lab. The concentration values seen in the blank samples were found to be less than their respective MDL value, with the exceptions of antimony and cobalt. These two compounds had blank filter average concentration values that were slightly larger than their MDLs. Those two averages were subtracted from each sample concentration found before taking the annual average of the data.

#### **Precision of Sample Results**

Once per month a duplicate sampler was run simultaneously with the primary sampler. These additional samples, known as duplicates, were collected in order to assess the precision (repeatability) of the metals sampling method. In general, repeatability for the two collocated samples was acceptable for the compounds that were detected in greater than 10% of the samples taken. Information regarding precision and accuracy results is available upon request to the Air Pollution Control Division.

# VII. PM<sub>10</sub>

#### Sample Statistics Summary

The Colorado Department of Public Health and Environment operates samplers for particulate matter 10 microns or less in diameter ( $PM_{10}$ ) at the Grand Junction – Powell station. These samplers serve to indicate the status of Grand Junction regarding the National Ambient Air Quality Standards (NAAQS) for  $PM_{10}$ . Results of the statewide particulate matter monitoring network are discussed in "Colorado: 2015 Air Quality Data Report" by the Air Pollution Control Division. In 2015, 121 samples were attempted, and 110 were collected on the primary sampler, bringing the final data recovery rate to 91%. The duplicate sampler was in operation for four of the missed samples. Therefore, the value from the duplicate sampler was used to replace the missing value, bringing the data completion rate to 94%.

Table 10.  $PM_{10}$  Average and Maximum Concentrations 2004 - 2015

	PM <sub>10</sub> Concentrations			
	3rd Day		6th day	
Year	Average (μg/m³)	Maximum (μg/m³)	Average (μg/m³)	Maximum (μg/m³)
2004	29	102	29	102
2005	26	198	25	62
2006	30	98	30	66
2007	30	85	29	69
2008	30	116	30	84
2009	25	68	26	68
2010	22	155	19	57
2011	18	41	18	39
2012	22	77	20	44
2013	19	55	20	48
2014	18	46	19	46
2015	15	37	14	34

Table 100 lists the average and maximum concentrations observed at the Grand Junction site from 2004 through 2015, for both the primary (every third) and collocated (every sixth day) samplers. The averages are similar for the third and sixth day sampling, and are far lower than the 24-hour standard level of 150 micrograms per meter cubed. The maximum value observed in 2015 was 37 micrograms per meter cubed. The maximums are similar for some years, but quite different for others. To date, the highest concentration observed was 198 micrograms per meter cubed in 2005.

## Graphs

The graphs in this section will cover data for the NATTS sampling calendar (every sixth day). Any data discussed will be derived from the every sixth day (or primary sampler) values. Figure 46 is a graph of the  $PM_{10}$  concentration data recorded every sixth sampling day. The graph does not appear to indicate any seasonal variation in 2015.

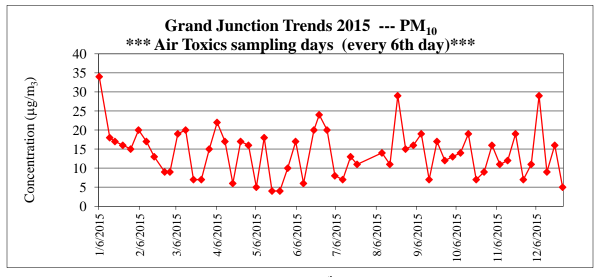


Figure 46. PM<sub>10</sub> Concentrations by Date, 2015 (every 6<sup>th</sup> Day)

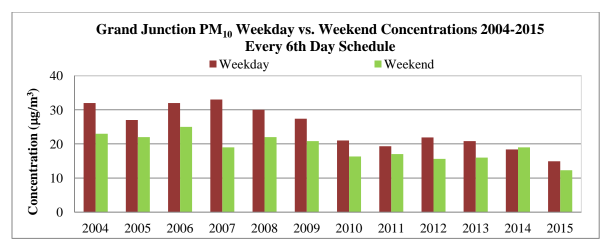


Figure 47. PM<sub>10</sub> Weekend vs. Weekday Comparison, 2004-2015 (every 6<sup>th</sup> day)

Figure 47 is a graph of the weekend versus weekday concentrations for  $PM_{10}$  on the every sixth day sampling schedule. The weekday average is larger than the weekend average for all years except 2014.  $PM_{10}$  is dominated by surface disturbance of earth materials (e.g., street sand, windblown dust, etc.). The  $PM_{10}$  levels are subject to change due to daily weather conditions. Figure 48 is a graph of the annual average  $PM_{10}$  concentrations from 2004 through 2015, for the every sixth day sampling period. Figure 49 is a graph of the 24-hour maximum  $PM_{10}$  concentrations from 2004 through 2015, for the every sixth day sampling period. The maximum  $PM_{10}$  concentrations observed in each year are used for comparison to the NAAQS in this graph.

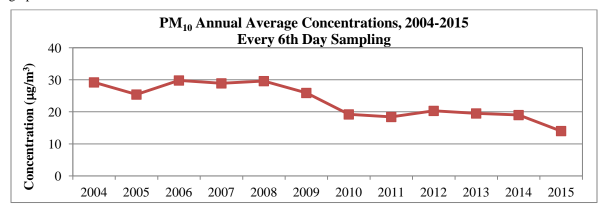


Figure 48. PM<sub>10</sub> Annual Average Concentrations 2004 – 2015

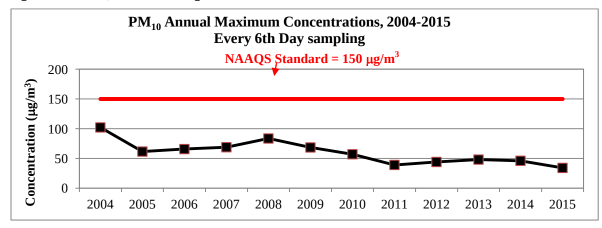


Figure 49. PM<sub>10</sub> Annual Maximum Concentrations 2004 – 2015

### Quality Assurance/Quality Control

#### **Blanks**

Blank filters are collected for the purposes of determining any bias in the sampling setup, removal, transport, or laboratory weighing processes. Data from these field blanks is available on request.

#### **Precision of Sample Results**

Collocated samples were run once every sixth day, half as frequently as the primary samples were run. This is done in an effort to validate the collected data. There is good agreement between the primary and collocated sampler concentrations. Information regarding precision and accuracy results is available upon request to the Air Pollution Control Division.

### VIII.PM<sub>2.5</sub>

### Sample Statistics Summary

The Colorado Department of Public Health and Environment operates samplers for particulate matter 2.5 microns or less in diameter ( $PM_{2.5}$ ) at the Grand Junction – Powell station. This sampler serves to indicate the status of Grand Junction regarding the National Ambient Air Quality Standards (NAAQS) for  $PM_{2.5}$ . Results of the statewide particulate matter monitoring network are discussed in "Colorado: 2015 Air Quality Data Report" by the Air Pollution Control Division. The National Air Toxics Trends Study chose to monitor air toxics in Grand Junction due to the availability of a  $PM_{2.5}$  speciation sampler, which gives insight into air toxics in particulate matter. It should be noted here, however, that the speciation sampler previously located in Grand Junction was removed, and relocated to the state's NCore site in Denver at the end of 2009. The  $PM_{2.5}$  data discussed here is the gravimetric filter data only, and does not include any speciated results. In 2015, 123 samples were attempted, and 119 were collected on the 3 day sampling schedule, giving a 97% sampling rate.

Table 11. PM<sub>2.5</sub> Average Concentrations 2004-2015

	3rd day		6th day		98 <sup>th</sup> %ile
Year	Average (μg/m³)	1 <sup>st</sup> Max (µg/m³)	Average (μg/m³)	1 <sup>st</sup> Max (µg/m³)	Max (μg/m³)
2004	10.4	36.3	10.4	31.6	31.6
2005	8.4	19.0	7.9	18.2	18.0
2006	9.7	28.5	9.8	28.5	24.0
2007	9.5	30.7	9.0	30.7	26.0
2008	9.1	27.8	8.9	27.8	25.0
2009	9.8	59.1	10.5	59.1	41.0
2010	9.0	43.3	8.4	43.3	37.0
2011	7.1	23.9	6.8	23.9	22.0
2012	7.3	28.3	7.2	28.3	24.0
2013	8.8	42.2	8.6	40.3	40.0
2014	6.3	21.7	6.3	20.9	16.0
2015	5.4	22.2	5.5	22.2	19.0

Table 11 lists the annual average and first maximum  $PM_{2.5}$  concentrations at the Grand Junction sites for 2004 through 2015, for both every third day and every sixth day sampling. There is very little difference between the averages for the two sampling schedules, and they are less than the current annual standard level

of 12 micrograms per cubic meter.  $PM_{2.5}$  emissions are generated by agriculture, and the combustion of automobile fuels, coal, wood, etc., as well as by secondary formation from other available atmospheric compounds. Table 11 also lists the  $98^{th}$  percentile maximum values for 2004 through 2015. The maxima for the third and sixth day sampling in 2015 are the same at 22.2 micrograms per cubic meter.

To meet the primary  $PM_{2.5}$  national standard, the three year average of the annual mean concentrations must be less than 12 micrograms per cubic meter. The design value for this site is 5.7 micrograms per meter cubed, thus the primary standard is met. To meet the secondary  $PM_{2.5}$  national standards, the three year average of the annual mean must be less than 15 micrograms per cubic meter, and the 3 year average of the  $98^{th}$  percentile value must be less than 35 micrograms per meter cubed. Those values for this site are 5.7 and 25 micrograms per meter cubed, respectively. Therefore, the secondary standards are met. The standard values were calculated using the data from the every third day sampling set.

### **Graphs**

A graph of the daily concentration values for the every sixth day sampling subset is shown in Figure 50. They show that the PM<sub>2.5</sub> concentrations are generally consistent throughout the year, but tend to vary more during the winter months, when there is more smoke in the air from agricultural and household wood burning.

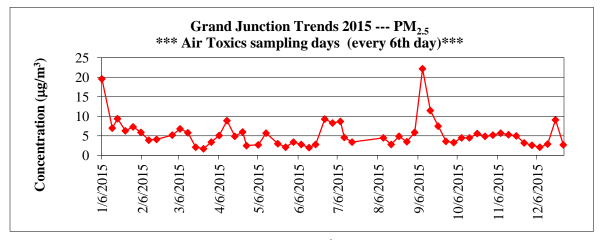


Figure 50. PM<sub>2.5</sub> Concentration by Date, 2015 (every 6<sup>th</sup> day)

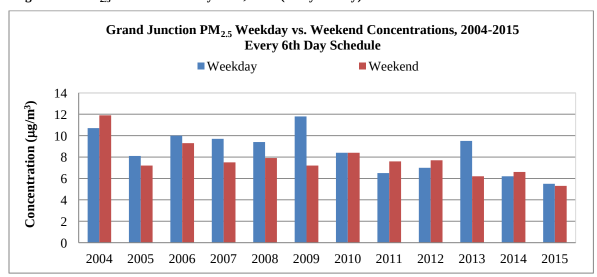


Figure 51. PM<sub>2.5</sub> Weekend vs. Weekday Comparison, 2004-2015 (every 6<sup>th</sup> day)

Figure 51 shows how the weekend versus weekday average concentrations compare for 2004 through 2015, for the every 6<sup>th</sup> day sampling schedule. In 2004, 2011, 2012, and 2014, weekday averages were less than the weekend averages. Since beginning sampling in 2004, the average fine particulate matter concentrations have varied from five to twelve micrograms per meter cubed. Figure 52 shows the annual average concentrations for PM<sub>2.5</sub> for 2004 through 2015. Overall, the average trend appears to be decreasing.

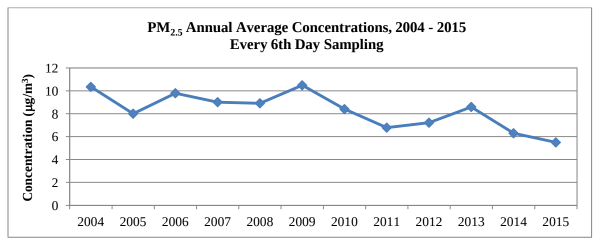


Figure 52. PM<sub>2.5</sub> Annual Average Concentrations 2004 – 2015

# Quality Assurance/Quality Control

### **Precision of Sample Results**

No collocated samples were run for  $PM_{2.5}$ , as there is currently only one  $PM_{2.5}$  instrument available to place at the Grand Junction site.

#### IX. METEOROLOGY

A meteorological tower at the Pitkin shelter site measures wind speed, wind direction, relative humidity, and temperature. The 2015 wind rose is shown below in Figure 53. The "arms" of this diagram show the percentage of the time that the wind blew from each direction. The shading on each arm indicates the wind speeds associated with each direction. Each of the concentric rings, moving outward, signifies an additional 1.6 percent of the time. For example, about 7 to 8% of the winds are from the east. Wind speeds in the ranges of 0.5 to 2.1 meters per second (m/s) are the most frequent.

The wind rose shows that winds follow a daily pattern typical of river valleys. At night, the winds come from the southeast quarter, flowing down river. During the day, heating of the air causes flow reversals, and flow comes from the northwest.

A look at the highest concentrations days for each pollutant indicated that some days showed maxima for more than one air pollutant. Many of these dates are in the fall or winter period, which indicates possible local temperature inversions and limited air mixing, thus allowing pollutants of all types to build up in the area.

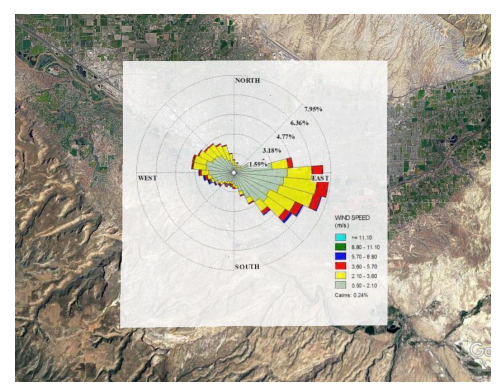


Figure 53. Wind Rose for Grand Junction 2015

# X. DATA CORRELATIONS AND DISCUSSION

The data presented below are the results of several correlation comparisons between the particulate concentrations and various other air toxics compound concentrations.

# Carbonyl Correlations and Sample Composition

Carbonyl compounds are known to have adverse effects on human health. They can be emitted directly from primary sources (motor vehicle emissions, and incomplete combustion), or can be formed secondarily via atmospheric photo-oxidation reactions. They play an important role in the formation of ozone in the atmosphere, and are of great interest to atmospheric researchers, as is particulate matter. Particulates are a mixture of solid particles and liquid droplets found in the air. Of interest to researchers are two different classes of particulates: coarse (having a diameter of 10 micrometers or less), and fine (having a diameter of 2.5 micrometers or less). Fine particles are small enough to be inhaled deep into the lungs, and cause serious health problems. Fine particulates are the major cause of visibility issues in many parts of the U.S. A correlation of the annual average carbonyl concentration data was performed with both the  $PM_{10}$ , and  $PM_{2.5}$  annual average data sets. The results of the correlation are presented in Table 12.

Wang et al., "Seasonal Variation and Source Apportionment of Atmospheric Carbonyl Compounds in Urban Kaohsiung, Taiwan." Aerosol and Air Quality Research, 10: 559–570, 2010. http://aaqr.org/VOL10\_No6\_December2010/5\_AAQR-10-07-OA-0059\_559-570.pdf

Table 12. Correlation Coefficient Values for Carbonyls-Particulates

Analyte	r - PM <sub>10</sub>	r-PM <sub>2.5</sub>
2-Butanone (MEK)	0.40	0.62
Acetaldehyde	0.68	0.59
Acetone	0.80	0.59
Benzaldehyde	0.71	0.53
Butyraldehyde	0.80	0.49
Crotonaldehyde	0.84	0.55
Formaldehyde	0.43	0.43
Hexaldehyde	0.69	0.53
Propionaldehyde	0.76	0.38
Tolualdehydes	0.83	0.52

**Bold** = **MQO** Core Analyte

Several of the carbonyl compounds tended to correlate well with the  $PM_{10}$  data, having "r" values of 0.7 or larger. It should be noted here that the correlation was performed only for the carbonyl compounds that were detected in 10% or more of the samples taken. Crotonaldehyde shows the strongest correlation with an "r" value of 0.84. One of the two MQO Core Analyte carbonyls, acetaldehyde, did show some correlation with the course particulate concentrations. There was little correlation between any of the carbonyls and the fine particulate concentrations. 2-Butanone had the highest "r" value of the group at 0.62. A graph of the four carbonyls with "r" values greater than or equal to 0.80 for the  $PM_{10}$  correlation is shown in Figure 54.

The final graph presented in this section is a snapshot of the chemical make-up of the carbonyls group from 2004 through 2015. Figure 55 shows the percentage each carbonyl compound contributed to the overall total carbonyl concentration from year to year. Acetone, acetaldehyde, and formaldehyde clearly dominate the carbonyl concentrations yearly.

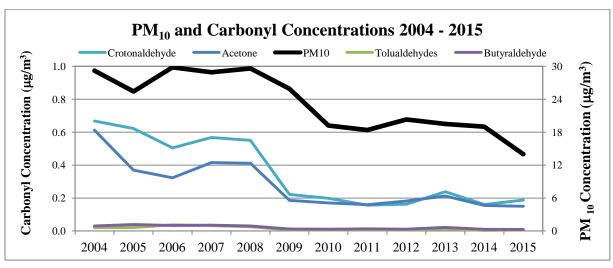


Figure 54. PM<sub>10</sub> – Carbonyl Concentration Comparison

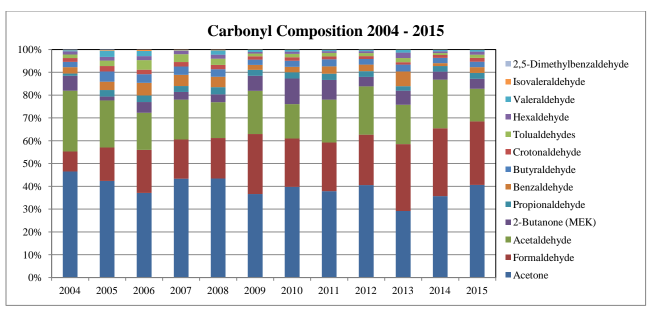


Figure 55. Annual Carbonyl Composition

# **VOC Correlations and Sample Composition**

VOCs are organic compounds which have a high vapor pressure at room temperature. Because of this high vapor pressure, which is the result of a low boiling point, large numbers of VOC molecules can evaporate or sublimate from a liquid or solid form and enter the ambient air. The NATTS program monitors for 60 of these compounds, many of which are never detected in any samples. The VOC correlation data used and discussed in this section is based upon the subset of 35 compounds that were detected in greater than 10% of the samples taken since 2004. The MQO Core Analytes are bolded in the table below.

Table 13 is a listing of the correlation coefficients (r) for each of the 35 VOC compound data sets, with both  $PM_{2.5}$ , and  $PM_{10}$  data sets. For the VOC -  $PM_{10}$  correlation, acetylene, benzene, and carbon disulfide correlated fairly well with the course particulate concentrations with correlation coefficient values of 0.77, 0.80, and 0.80, respectively. Figure 56 is a graph of the carbon disulfide, benzene, acetylene, and  $PM_{10}$  annual average concentrations from 2004 through 2015.

Table 13. VOC – Particulate Correlation Coefficient Values

Compound	$PM_{10}$	PM <sub>2.5</sub>
1,1,1-Trichloroethane	0.79	0.68
1,2,4-Trimethylbenzene	0.57	0.45
1,2-Dichloroethane	-0.14	-0.19
1,3,5-Trimethylbenzene	0.51	0.43
1,3-Butadiene	0.62	0.49
Acetonitrile	-0.58	-0.53
Acetylene	0.77	0.73
Acrolein	-0.65	-0.35
Acrylonitrile	-0.16	-0.22
Benzene	0.80	0.79
Bromomethane	-0.60	-0.59
Carbon Disulfide	0.80	0.70
Carbon Tetrachloride	-0.01	0.02
Chloroethane	0.25	0.20

Compound	$PM_{10}$	PM <sub>2.5</sub>
Chloroform	-0.01	0.08
Chloromethane	0.13	0.17
Dichlorodifluoromethane	0.50	0.62
Dichloromethane	-0.73	-0.57
Dichlorotetrafluoroethane	-0.31	-0.02
Ethyl tert-Butyl Ether	-0.16	-0.23
Ethylbenzene	0.42	0.30
m,p-Xylene	0.47	0.33
Methyl Isobutyl Ketone	0.58	0.51
Methyl Methacrylate	0.68	0.45
n-Octane	-0.26	-0.22
o-Xylene	0.40	0.26
p-Dichlorobenzene	0.32	0.26
Propylene	0.64	0.57
Styrene	-0.65	-0.65
Tetrachloroethylene	0.54	0.77
Toluene	0.49	0.40
Trichloroethylene	0.14	0.21
Trichlorofluoromethane	0.45	0.62
Trichlorotrifluoroethane	0.63	0.66
Vinyl chloride	0.35	0.22

**Bold = MQO Core Analyte** 

*Italic = Did not meet detection requirements* 

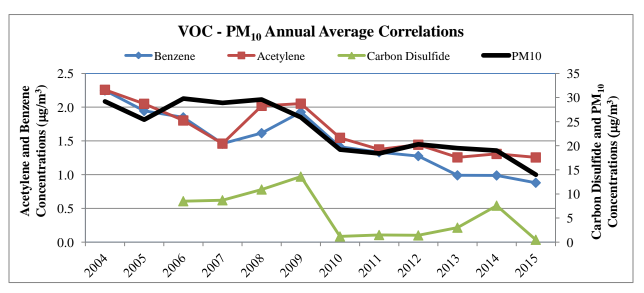


Figure 56. VOC – PM<sub>10</sub> Concentration Comparison

The VOC –  $PM_{2.5}$  correlation showed only one compound with a relatively strong correlation. Benzene correlated well with the fine particulate matter concentrations, showing a positive r-value of 0.79. Acetylene and carbon disulfide also showed some correlation with the fine particulate matter, with r-values of 0.73, and 0.70, respectively. Figure 57 shows the annual average concentrations for benzene, acetylene, carbon disulfide, and  $PM_{2.5}$ .

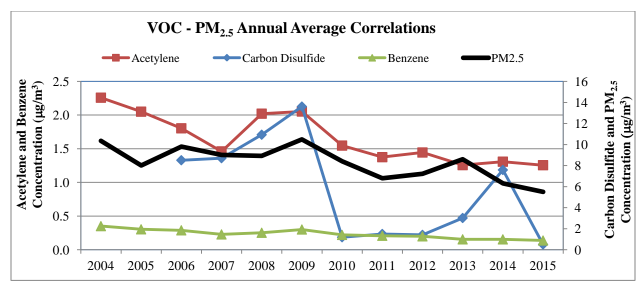


Figure 57. VOC – PM<sub>2.5</sub> Concentration Comparison

The chemical make-up of the VOC compounds tends to be much more variable from year to year than the carbonyl compounds. This is especially true for the C1 through C2 carbon chains of the VOCs, which can be seen in Figure 58. The graph shows data from 2004 through 2015. The year to year variability is easily seen. Carbon disulfide was not sampled for during the 2004 and 2005 campaigns, but was added in 2006. It was a major component of the VOCs for 2006 through 2009, but has not been a major contributor since 2009. The data from 2010, 2012, 2013, 2014, and 2015 show a shift to dichloromethane as a large component of this VOC group, comprising 50 to 75 percent of the C1through C2 VOC composition.

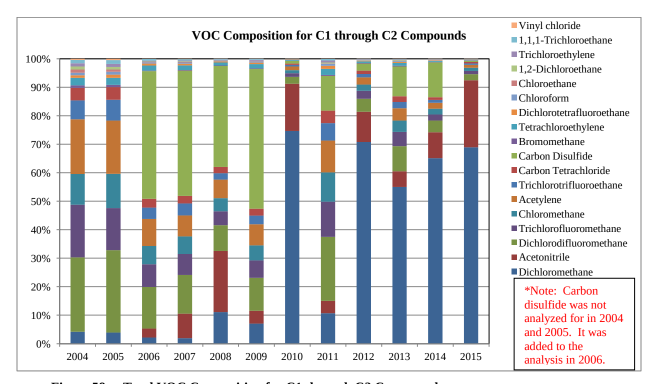


Figure 58. Total VOC Composition for C1 through C2 Compounds

Figure 59 shows the chemical composition of the C3 though C9 carbon chain compounds. These compounds tend to show a more consistent make-up from year to year, as opposed to the lighter end alkanes of the C1 and C2 chains. It should be noted that this grouping contains straight chain alkanes, as well as aromatic compounds. It seems likely that the major source for these C3 through C9 compounds is from motor vehicle traffic, due to the consistent nature of the chemical makeup, and the site's nearness to a major road. In 2015, styrene concentrations dominated the mix, comprising more than 60% of the matrix. This is a large jump from previous years.

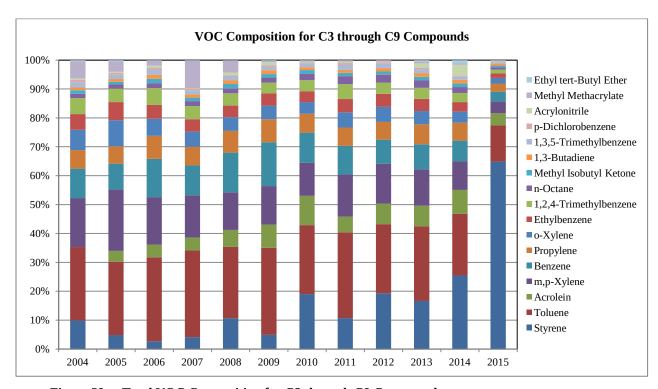


Figure 59. Total VOC Composition for C3 through C9 Compounds

#### PAH Correlations and Sample Composition

Polycyclic Aromatic Hydrocarbons are often found naturally in the environment, but are also man-made. They can enter the air through the incomplete combustion of fuels and garbage. They are a concern because of their persistence in the environment. Table 14 lists the correlation coefficient values for each of the PAH compounds. All but one of the compounds show little to no correlation with the  $PM_{10}$  values. This is reasonable, since  $PM_{10}$  is largely from geologic sources.

This particular set of compounds did tend to trend better with the fine particulate matter concentrations. All but two compounds showed positive correlations with the  $PM_{2.5}$  concentrations. The strongest correlation between the PAH and  $PM_{2.5}$  concentrations was seen with acenapthylene. A correlation coefficient of 0.90 was obtained for this compound. PAHs can exist in liquid or solid phases, so their positive relationship with the smallest diameter particles, which develop from gaseous condensation, is easily explained. PAHs and  $PM_{2.5}$  are also both direct combustion products. The compounds with correlation coefficient values that are greater than 0.7 are graphed in Figure 60.

Table 14. PAH – Particulate Correlation Coefficient Values

PAH correlations	r - PM <sub>10</sub>	r - PM <sub>2.5</sub>
9-Fluorenone	-0.28	0.18
Acenaphthene	0.09	0.03
Acenaphthylene	0.57	0.90
Anthracene	0.00	0.34
Benzo (a) anthracene	0.09	0.54
Benzo (a) pyrene	0.17	0.54
Benzo (b) fluoranthene	0.31	0.75
Benzo (e) pyrene	0.22	0.61
Benzo (g,h,i) perylene	0.46	0.76
Benzo (k) fluoranthene	0.39	0.76
Chrysene	0.00	0.51
Coronene	0.55	0.68
Cyclopenta[cd]pyrene	0.56	0.56
Dibenz (a,h) anthracene	0.74	0.55
Fluoranthene	-0.54	-0.31
Fluorene	-0.07	0.07
Indeno(1,2,3-cd)pyrene	0.33	0.68
Naphthalene	0.27	0.51
Perylene	0.56	0.68
Phenanthrene	-0.34	-0.17
Pyrene	-0.28	0.10
Retene	0.23	0.74

**Bold = MQO Core Analyte** 

Italic = Detected in less than 10% of samples taken

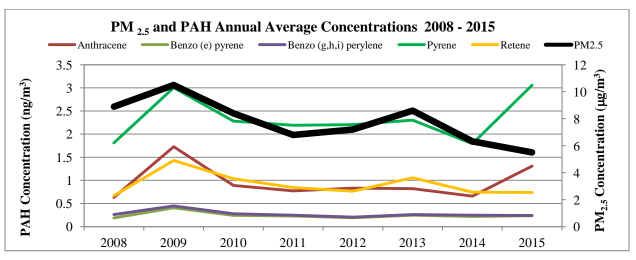


Figure 60. PAH – PM<sub>2.5</sub> Concentration Comparison

Figure 61 is a graph showing the percentage contribution each of the PAH compounds to the total PAH concentration. Clearly, naphthalene is the dominant compound of the group, consistently making up more than 60% of the PAH composition. The composition of the PAH group does not appear to vary much from year to year. This may imply that PAH sources are consistent over time.

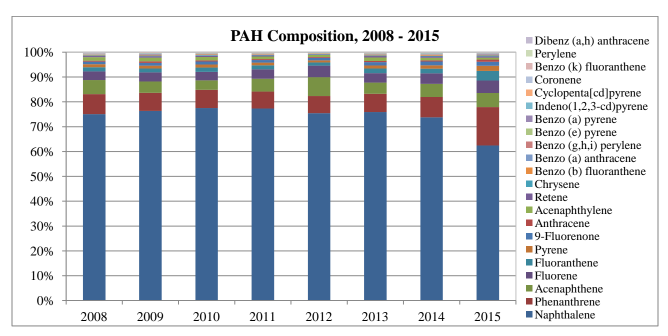


Figure 61. PAH Chemical Composition 2008 – 2015

# Metals Correlations and Sample Composition

The metals in this group are sampled via a  $PM_{10}$  filter based monitor. Six of the metals sampled for are listed as MQO Core Analytes. The correlation coefficients for the metals compounds (having enough data to correlate) with the two different particulate classes are shown in Table 15. Manganese concentrations correlated well with the  $PM_{10}$  concentrations, having an r-value of 0.88. There were no significant correlations between any of the metals compounds and the  $PM_{2.5}$  concentrations. This suggests that select metals may be coming from geologic crustal, rather than combustion or secondary formation, sources. Nickel sources include various metal alloys, electroplating, motor vehicle exhaust, and geologic crustal material. A graph of the  $PM_{10}$  and manganese concentrations is seen in Figure 62.

Table 15. Metals – Particulates Correlation Coefficients

Analyte	r-PM <sub>10</sub>	r-PM <sub>2.5</sub>
Antimony	0.19	-0.14
Arsenic	0.66	0.36
Beryllium	0.54	0.26
Cadmium	0.38	0.28
Chromium	0.17	-0.04
Cobalt		
Lead	0.66	0.44
Manganese	0.88	0.51
Mercury		
Nickel	-0.20	-0.03
Selenium		

**Bold = MQO Core Analyte** 

<sup>&</sup>lt;sup>11</sup> http://scorecard.goodguide.com/chemical-profiles/html/nickel.html

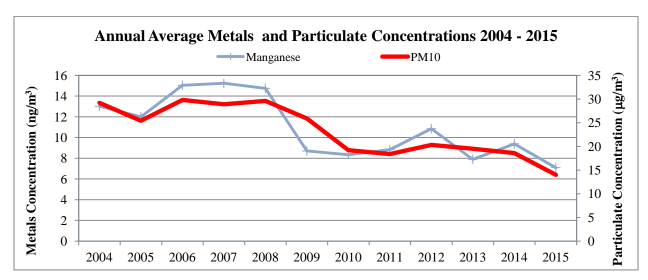


Figure 62. Metals – PM<sub>10</sub> Concentration Comparison

Figure 63 is a graph showing the percentage contribution of each of the individual metals compounds to the overall total. The concentrations vary somewhat from year to year, but not as much as the C1 through C4 compounds of the VOC section.

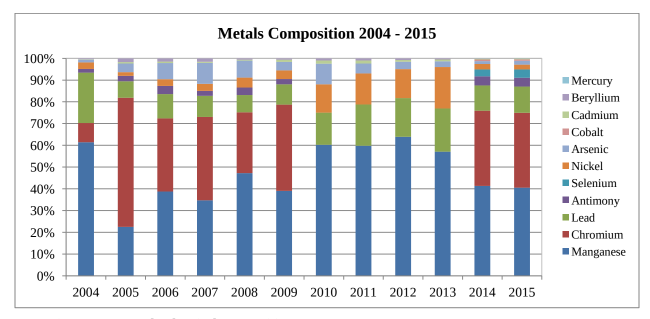


Figure 63. Metals Chemical Composition 2004 – 2015

# XI. SUMMARY AND CONCLUSIONS

The National Air Toxics Trends Study in Grand Junction for 2015 showed similar results to prior years. The highest carbonyls in air were formaldehyde, acetaldehyde, and acetone. A correlation analysis was run between the particulate concentrations and the carbonyl concentrations.  $PM_{10}$  concentrations tended to correlate with many of the carbonyl compounds. A correlation value (r) of 0.84 was obtained when

comparing annual average  $PM_{10}$  to crotonaldehyde concentrations. This value was the highest obtained for the  $PM_{10}$ -carbonyl correlation. A comparison of the  $PM_{2.5}$  concentrations with the carbonyls showed that 2-butanone correlated the best, but had a moderate correlation coefficient value of 0.62. Many of the carbonyls showed no correlation at all with the  $PM_{2.5}$  values.

Thirty-five volatile organic compounds are ubiquitous, having been detected in at least 10% of the air samples taken in 2015. From 2004 to 2015, the makeup of the C1 to C2 group was highly variable, with large concentrations of carbon disulfide from 2006 through 2009, and moderate concentrations in 2011, 2013, and 2014. There were also large concentrations of dichloromethane in 2010, 2012, 2013, 2014 and 2015. The C3 through C8 group showed more consistency in the constituent concentrations from 2004 to 2015, with the exception of styrene, which showed a marked increase in concentration from 2014 to 2015. Correlations with particulate data showed that carbon disulfide, benzene, and acetylene tracked most closely with the  $PM_{10}$  concentrations, with r-values of 0.0.80, 0.80, and 0.77, respectively. Benzene correlated best with the fine particulate concentrations, having an r-value of 0.79.

The highest polycyclic aromatic hydrocarbons concentrations were naphthalene, acenaphthene, and phenanthrene, none of which correlated well with  $PM_{10}$  concentrations. The compounds that did correlate well with the fine particulate matter were acenaphthylene, benzo(b)fluoranthene, benzo(g,h,i) perylene, benzo(k)fluoranthene, and retene, with r-values of 0.90, 0.75, 0.76, 0.76, and 0.74, respectively. Several of the other PAH compounds also correlated somewhat with  $PM_{2.5}$  values, but their r-values were under the 0.70 cutoff.

For the metals, lead and manganese showed the highest average concentrations. Manganese had the highest correlation value with the coarse particulate matter at 0.88. It is important to note here that many of the metals concentrations rely heavily on their respective MDLs. None of the metal compounds showed any correlation with fine particulate matter.

In general, it appears that the concentrations of many of the compounds of interest are dropping since the inception of the NATTS program in Grand Junction. The study will continue in 2016, as one of the major goals is to run the site long term, for comparison of the mean concentrations for each pollutant during the first three years to the means for successive three year intervals. Calculation of the three year average concentrations to date has shown a decrease in the majority of the concentrations of the compounds of interest.

# Appendix A: Compounds Contributing to Cancer and Non-cancer Risks - Overview of Sources and Health Effects

Chemicals can be released to the environment as a result of their use and manufacture. Some chemicals may also form, as other chemicals react with sunlight and one another in outdoor air. A brief summary of the potential sources and health effects of some prevalent chemicals in the ambient air is provided below. This information is adopted from the following main sources: EPA Air Toxic Website, EPA Office of Pollution Prevention and Toxics (OPPT), EPA Integrated Risk Information System (IRIS), Agency for Toxic Substances and Disease Registry (ATSDR), New Jersey Department of Health and Senior Services, Occupational Safety and Health Administration (OSHA), National Institute of Occupational Safety and Health (NIOSH), and the California Air Resources Board (CARB).

#### **CARBONYLS**

Three of the twelve carbonyl compounds sampled are discussed below. These three are believed to be significant health risk drivers, at the national level.

#### **ACETALDEHYDE**

Acetaldehyde is a hydrocarbon with the formula  $CH_3CHO$ . It is thus closely related to formaldehyde, HCHO. Like formaldehyde, it exists in the atmosphere as a gas with a pungent odor. Acetaldehyde is ubiquitous in the ambient environment. It is mainly used as an intermediate in the synthesis of other chemicals, such as acetic acid, acetic anhydride, chloral, and glyoxal. It is employed in the food processing industry as a food and fish preservative, a flavoring agent, and in gelatin fibers. The tanning and paper industries use acetaldehyde, as do the perfume and dye manufacturers (CARB Acetaldehyde Fact Sheet).

Acetaldehyde can be released to the environment as a product of incomplete combustion in fireplaces and wood stoves, forest and wild fires, pulp and paper production, stationary internal combustion engines and turbines, vehicle exhaust, and petroleum refineries. Waste water processing is also a source. It is important to note that residential fireplaces and woodstoves are the two highest sources of emissions, followed by various industrial emissions.

Although it is used in industry, the California Air Resource Board believes that the largest sources of acetaldehyde in outdoor air are combustion and production from photochemical reactions (CARB Acetaldehyde Fact Sheet). Acetaldehyde itself can break down in these complex reactions between air pollutants and sunlight, forming formaldehyde.

The health effects of acetaldehyde are very similar to those of its chemical relative formaldehyde. It irritates the eyes and mucous membranes. It can paralyze the respiratory muscles, act as a narcotic to prevent coughing, and speed up pumping of the heart. Exposure can lead to headaches and sore throat. (Kirk Othmer, Vol 1, page 107). It should be noted that most of these health effects have been observed in factory workers, who are exposed to acetaldehyde concentrations thousands of times greater than those occurring in outdoor air. Acetaldehyde is believed to be a probable human carcinogen, leading to cancer of the nose and throat. Acetaldehyde has been shown to cause birth defects in animals, but no human research is available. (CARB Acetaldehyde Fact Sheet).

EPA's Technology Transfer Network Air Toxic Website provides information on the potential health effects of acetaldehyde. According to this source, the primary acute effects of acetaldehyde are irritation of the eyes, skin, and respiratory tract in humans. At higher exposure levels erythema, coughing, pulmonary edema, and necrosis may happen. Chronic toxicity symptoms in humans resemble those of alcoholism.

The EPA has established a Reference Concentration (RfC) for inhalation exposure to acetaldehyde based on degeneration of the olfactory epithelium in rats. No information is available on the reproductive and developmental effects of acetaldehyde in humans. Animal studies data indicate that acetaldehyde may be a potential developmental toxin. EPA has classified acetaldehyde as a Group B2, probable human carcinogen, based on increased incidence of nasal tumors in male and female rats and laryngeal tumors in male and female hamsters after inhalation exposure.

The California Air Resources Board observed an annual mean of 1.33 ppb acetaldehyde in its state-wide network during 1996 (CARB Acetaldehyde Fact Sheet). The mean observed in this Grand Junction study, 3.2 ppb, is a bit above the California data, but acetaldehyde in Grand Junction occurs at levels typical of large urban areas. Acetaldehyde levels are a national problem related primarily to the use of motor vehicles.

#### CROTONALDEHYDE

Crotonaldehyde with the chemical formula of C<sub>4</sub>H<sub>6</sub>O is also known as propylene aldehyde, betamethylacrolein, crotinin aldehyde and butenal. Crotonaldehyde is a colorless liquid with a pungent, suffocating odor.

Crotonaldehyde can be emitted to the environment from the combustion of gasoline, the burning of wood, paper, cotton, plastic, and tobacco. It can also be released through industrial use. It is found naturally in emissions of some vegetables and volcanoes.

According to the ATSDR Medical Management Guidelines, inhaled crotonaldehyde is highly toxic. It is irritating to the upper respiratory tract even at low concentrations. Crotonaldehyde vapor is heavier than air. Therefore, higher levels of crotonaldehyde vapors would be found nearer to the ground. The mechanism of toxicity of crotonaldehyde is not known, but it is highly reactive. Crotonaldehyde is also a skin irritant and can cause eye irritation and damage to the cornea. After an acute, relatively high concentration exposure, people may become sensitized to crotonaldehyde. Except for rare cases of sensitization, no health effects have been reported in humans exposed to relatively low concentrations of crotonaldehyde. No studies have been found that address reproductive or developmental effects of crotonaldehyde in humans. The compound has been shown to cause degeneration of spermatocytes in mice. No teratogenic effects from acute exposures have been reported.

The Department of Health and Human Services has determined that crotonaldehyde may be a possible carcinogen. The EPA IRIS has classified crotonaldehyde as a possible carcinogen based on the fact that there is no human data, but an increased incidence of hepatic tumors in male rats. The possible carcinogenicity of crotonaldehyde is supported by genotoxic activity and the expected reactivity of croton oil and aldehyde. The EPA IRIS, however, has not derived a cancer toxicity value for the compound. The EPA HEAST (Health Effects Summary Tables) has established an oral cancer toxicity value for crotonaldehyde. The Agency for Research on Cancer has determined that crotonaldehyde is not classifiable as to its carcinogenicity to humans.

Information concerning typical concentrations of crotonaldehydes in air could not be located.

#### **FORMALDEHYDE**

Formaldehyde is a hydrocarbon compound with the formula HCHO. It exists in the atmosphere as a colorless gas with a pungent odor. It is used in the manufacture of urea-formaldehyde resins which are used in particleboard and plywood products. Therefore, high levels of airborne formaldehyde can also be found in indoor air as a result of release from various consumer products such as building materials and home furnishings. Another source of formaldehyde in indoor air is smoking. It is also employed in chemical manufacturing of pharmaceuticals, herbicides, and sealants. Textile finishes, such as used for "permanent press" clothes, contain formaldehyde (Kirk-Othmer, Vol 11, pages 245 - 246).

EPA's Technology Transfer Network Air Toxic Website provides information on the potential sources and health effects of formaldehyde. According to this source, the major sources of formaldehyde emissions to the

ambient air include power plants, manufacturing facilities, incinerators, forest and wild fires, stationary internal combustion engines and turbines, pulp and paper plants, petroleum refineries, and automobile traffic. In urban areas, combustion of automotive fuel is the dominant source for much of the year. However, formaldehyde can also form photochemically in the air, as other hydrocarbons and oxides of nitrogen from automobile traffic break down to form ozone. Complicating the situation is the fact that the complex ozone-producing atmospheric reactions may both create and destroy formaldehyde, as the chains of chemical reactions proceed along various pathways.

The Agency for Toxic Substances and Disease Registry (ATSDR), lists a number of possible health effects that may occur from inhalation of formaldehyde. Formaldehyde is an irritant. The major acute toxic effects via inhalation exposure are eye, nose, and throat irritation and effects on the nasal cavity. At 0.4 - 3 ppm, it may cause the eyes to tear. Other effects observed in humans from exposure to high levels of formaldehyde are coughing, wheezing, chest pain, and bronchitis (EPA's Technology Transfer Network Air Toxic Website). Formaldehyde is believed to be carcinogenic to humans. However, the body can quickly break down formaldehyde, so it does not accumulate in fatty tissue. Currently, ATSDR believes that formaldehyde does not cause birth defects in humans (ATSDR Toxicological Profile for Formaldehyde). Thus, the main concerns with this compound are its irritant properties and its potential ability to cause cancer of the nose and throat.

Chronic inhalation exposure to formaldehyde in humans has been associated with respiratory symptoms and eye, nose, and throat irritation. EPA has not established an inhalation Reference Concentration (RfC) for formaldehyde. However, the ATSDR has established an inhalation reference concentration called a Minimal Risk Level (MRL) for formaldehyde based on respiratory effects in humans. Developmental effects, such as birth defects, have not been observed in animal studies. EPA has classified formaldehyde as a Group B1, probable human carcinogen, based on limited evidence in humans and sufficient evidence in animals. Occupational studies have shown statistically significant increases in incidence of lung and nasopharyngeal cancer. This evidence is considered limited because of possible exposure to other agents. Animal studies have reported an increased incidence of nasal squamous cell carcinoma by inhalation exposure. Please see EPA IRIS for a detailed discussion on the carcinogenicity of formaldehyde.

ATSDR states that typical levels of formaldehyde in urban air are 10-20 ppb. ATSDR cites concentrations of 0.2 ppb for rural areas, and 2-6 ppb for suburban areas (ATSDR Toxicological Profile for Formaldehyde). The mean level observed in Grand Junction during this study, 2.3 ppb, is within the "suburban" range.

#### **VOLATILE ORGANIC COMPOUNDS**

Volatile organic compounds commonly present included 1,3 – butadiene, benzene, carbon tetrachloride, tetrachloroethylene, 1,3,5 – trimethylbenzene and 1,2,4 - trimethylbenzene. Some health summary and source information regarding these compounds is given below.

#### BENZENE

Benzene is a hydrocarbon compound with the formula  $C_6H_6$ . It exists in the atmosphere as a colorless gas with a sweet odor. It is used in chemical manufacturing of medicines, detergents, explosives, shoes, dyes, leather, resins, paints, plastics and inks (CARB Fact Sheet on Benzene). It is also present in gasoline.

The largest sources of benzene in ambient air are automobiles, gasoline service stations, refineries, and chemical plants. Burning of vegetative matter in forest fires and woodstoves is also a source. In ambient air, benzene reacts with hydroxyl (OH) radicals within a few hours. Since hydroxyl radicals are common in outdoor air, this chemical transformation prevents the build-up of large concentrations of benzene.

Benzene is a serious concern from a toxicological standpoint. Unlike many of the compounds discussed here, benzene is a proven human carcinogen. It damages the blood-forming capacity of the body, leading to anemia or leukemia. Like the other volatile organic compounds, breathing large amounts can cause lightheadedness, headache, vomiting, convulsions, coma and death. It also irritates the skin and eyes, exerting a drying effect.

However, these health effects are usually seen in workplaces, where levels are thousands of times higher than those in outdoor air. Experiments with laboratory animals suggest that benzene exposure may be associated with numerous cancers. It may cause bone marrow damage and bone formation problems for a developing fetus (ATSDR Toxicological Profile for Benzene). Thus, EPA has had concern about whether levels of benzene in outdoor air are associated with cancer and leukemia. While no link with outdoor air concentrations has been unequivocally proven, EPA has acted to reduce air concentrations of this pollutant.

The EPA has established a Reference Concentration for inhalation exposure to benzene based on decreased lymphocyte count in an occupational epidemiologic study. Benzene is classified as a "known" human carcinogen for all routes of exposure by the EPA IRIS based on the increased incidence of leukemia in epidemiologic and case studies.

The Agency for Toxic Substances and Disease Registry (ATSDR) cites national 1984 to 1986 data from 300 cities, which indicate an average benzene level of 1.8 ppb for urban and suburban areas (ATSDR Toxicological Profile for Benzene). The Grand Junction – Powell site mean of 0.7 ppb observed in this study is somewhat lower.

### 1,3-BUTADIENE

1,3-Butadiene is a hydrocarbon compound with the formula  $C_4H_6$ . It exists in the atmosphere as a colorless gas with an odor similar to gasoline. It is used in making rubber and plastics. The most important use is in tire production. It is also used in the production of chemicals such as 1,4-hexadiene (NIOSH Current Intelligence Bulletin 41).

According to the California Air Resources Board, most emissions of 1,3-butadiene come from combustion of fuels in diesel and gas-powered motor vehicles. Other sources that they list include petroleum refining, tire wear, residential wood heating, and forest fires. Rubber and chemical production plants also have emissions. Breathing of cigarette smoke is another source of 1,3-butadiene exposure (ATSDR Fact Sheet)

1,3-Butadiene is of concern toxicologically because it is characterized as carcinogenic to humans based on the new EPA guidelines for cancer risk assessment and it also has adverse effects on reproduction and fetal development. Exposure to high concentrations can cause irritation and central nervous system effects such as eye irritation, cough, sore throat, headache, drowsiness, nausea, unconsciousness, and death. Rats and mice exposed to this compound in laboratory tests developed multiple cancers within single individuals. The animals had damaged testes and ovaries, and offspring of the animals had skeletal problems. Other effects seen in animals at low levels of inhalation exposure for one year include kidney and liver disease, and damaged lungs (ATSDR Fact Sheet). Generally, the acute health effects have not been seen at concentrations existing in outdoor air. However, EPA considers that the levels of 1,3-butadiene in air may represent a significant portion of the cancer risk related to ambient airborne chemicals.

The EPA has established a Reference Concentration for inhalation exposure to 1,3-butadiene based on ovarian atrophy in mice. The EPA has characterized 1,3-butadiene as carcinogenic to humans by inhalation based on the following total evidence: sufficient evidence from epidemiologic studies showing increased lymphohematopoietic cancers and leukemia; tumors at multiple sites in animal studies, and strong evidence suggesting that the carcinogenic effects are mediated by genotoxic metabolites of 1,3-butadiene.

ATSDR estimates that urban and suburban areas have an average concentration of 0.3 ppb 1,3-butadiene, while rural areas have 0.1 ppb (ATSDR Toxicological Profile for 1,3-Butadiene). The annual average at Grand Junction - Powell is 0.09 ppb.

#### **CARBON TETRACHLORIDE**

Carbon tetrachloride, also known as tetrachloromethane or methane tetrachloride, is a chlorinated hydrocarbon with the formula CCl<sub>4</sub>. It exists in the atmosphere as a gas. It has a sweet odor. The primary uses of

carbon tetrachloride were as a dry cleaning solvent, a grain fumigant, as a refrigerant, and as an aerosol propellant. Carbon tetrachloride has a long atmospheric half-life, so it can travel to the higher reaches of the atmosphere and damage the earth's ozone layer. Due to its toxicity and ozone-damaging qualities, most uses of carbon tetrachloride have been banned. It is still in use in industrial settings for producing refrigerants.

Carbon tetrachloride is emitted to the air from industrial sources and from petroleum refineries (California Air Resources Board Toxic Air Contaminant Identification List Summary for Carbon Tetrachloride). Carbon tetrachloride is also a common indoor air contaminant due to releases from building materials and products, such as cleaning agents used in homes (Air Toxic Website). There are no natural sources of carbon tetrachloride; it is produced by man (ATSDR Toxicological Profile for Carbon Tetrachloride).

As is true for many of the chlorinated hydrocarbons, breathing large concentrations of carbon tetrachloride has central nervous system effects including lightheadedness, coma, convulsions, double vision, intoxication, and death. It can also cause vomiting. In animal studies, it had effects on the liver and kidney. Male rats exposed to carbon tetrachloride had lower sperm production. Female rats exposed to it had stunted offspring with birth defects. These health effects are generally observed in occupational settings, where people had exposure to very high levels over a number of years.

EPA has not established a Reference Concentration for carbon tetrachloride. The CalEPA has established a Reference Exposure Level for carbon tetrachloride based on liver effects in guinea pigs. Carbon tetrachloride has been associated with liver and kidney cancer in animals. EPA considers it a Class B2 Carcinogen (probable human carcinogen) based on liver tumors in animals.

The California Air Resources Board has monitored carbon tetrachloride at a number of locations, and found a mean value of 0.078 ppb (California Air Resources Board Toxic Air Contaminant Identification List Summary for Carbon Tetrachloride). The 0.08 ppb annual mean observed at Grand Junction – Powell is at the same level.

#### **TETRACHLOROETHYLENE**

Tetrachloroethylene, also known as perchloroethylene, is a chlorinated hydrocarbon with the formula  $C_2Cl_4$ . It exists in the atmosphere as a gas. It has a "chloroform-like" odor (NIOSH Pocket Guide to Chemical Hazards, Tetrachloroethylene). The primary uses of tetrachloroethylene are as a dry cleaning solvent, metal cleaning solvent, or for chemical production. Tetrachloroethylene is used in paints, inks, aerosols, glues, polishes, silicones and rubber products (CARB Fact Sheet on Tetrachloroethylene and OPPT Chemical Fact Sheet on Tetrachloroethylene).

Most emissions of tetrachloroethylene come from degreasing, dry cleaning, or chemical production facilities. There are microorganisms that can produce tetrachloroethylene (ATSDR Toxicological Profile For Tetrachloroethylene).

As is true for many of the chlorinated hydrocarbons, breathing large concentrations of tetrachloroethylene has central nervous system effects including lightheadedness, coma, convulsions, double vision, intoxication, and death. It also can cause vomiting. In animal studies, it had effects on the liver and kidney. It also is an irritant to eyes, lungs, and skin. However, many of these health effects were observed in occupational settings, where exposure is much higher than in outdoor air. Some animal studies suggest that tetrachloroethylene exposure may lead to leukemia (NIOSH Registry of Toxic Effects of Chemical Substances Information for Tetrachloroethylene). Tetrachloroethylene has been associated with liver and kidney cancer in animals.

The ATSDR has established a Minimal Risk Level (MRL) based on nervous system effects in humans. It is important to note that EPA is currently re-evaluating the toxic potential of tetrachloroethylene, including its carcinogenicity, and therefore no relevant information is available in IRIS. In the interim, EPA recommends the use of CalEPA toxicity values as provisional values. The CalEPA cancer toxicity value is derived by considering data on liver tumors in male and female mice and mononuclear cell leukemia in male and female rats. EPA is currently working to revise the toxicity assessment for tetrachloroethylene.

The California Air Resources Board has monitored tetrachloroethylene at a number of locations within their state, and found a mean value of 0.019 ppb during 1996 (California Air Resources Board Toxic Air Contaminant Identification List Summary for Tetrachloroethylene). The annual mean at Grand Junction - Powell was 0.05 ppb. These levels are greater than the network-wide mean value for California. However, this compound was detected less than half the time.

# 1,3,5-TRIMETHYBENZENE AND 1,2,4-TRIMETHYLBENZENE

1,3,5-trimethylbenzene and 1,2,4-trimethylbenzene are isomers of the hydrocarbon formula  $C_9H_{12}$ . In pure form they are colorless liquids. They are used in chemical manufacturing of medicines, detergents, dyes, paints and inks. Trimethylbenzenes are a large component of distilled petroleum. They are also used as gasoline additives.

The largest sources of trimethylbenzenes in ambient air are likely to be automobiles, gasoline service stations, refineries, and chemical plants. In ambient air, trimethylbenzenes have a half-life of less than a day (EPA OPPT Chemical Summary For 1,2,4-Trimethylbenzene).

Health effects of trimethylbenzenes are similar to those of benzene. It damages the blood-clotting capacity of the body. Like the other volatile organic compounds, breathing large amounts can cause lightheadedness, headache, vomiting, convulsions, coma and death. It also irritates the skin and eyes, exerting a drying effect. Long-term exposure can lead to cough, reduced lung capacity, and bronchitis. However, these health effects are usually seen in workplaces, where levels are thousands of times higher than those in outdoor air. It is not known whether these compounds are carcinogenic. Some animal experiments suggest that they may cause bone formation problems for a developing fetus (EPA OPPT Chemical Summary For 1,2,4-Trimethylbenzene).

The Environmental Protection Agency cites national data indicating that average atmospheric concentrations of 1,2,4-trimethylbenzene are 0.58 ppb in rural areas, and 1.20 ppb in cities (EPA OPPT Chemical Summary For 1,2,4-Trimethylbenzene). The Grand Junction - Powell site had a mean value of 0.09 ppb. As the EPA citation is for 1988, it is likely that concentrations have gone down in recent years.

#### **METALS**

Arsenic and manganese are discussed below. Levels of lead observed in Grand Junction were below the Colorado state standard of 1.5 ug/m3 for a monthly average.

#### ARSENIC

Arsenic is a metal-like element that occurs naturally in the earth's crust. Its chemical symbol is "As". It exists in the atmosphere as particulate matter, in compounds formed from combination with other atoms such as oxygen, chlorine, and sulfur (ATSDR Public Health Statement for Arsenic). In the past, arsenic was used as a pesticide for orchard crops. Today, the chief use is in chromated copper arsenate (CCA) used to "pressure-treat" wood, to preserve it from decay in marine or in-ground usage. It is also used in metal alloy, glass-making, and electrical semi-conductors.

Emission sources of arsenic include smelters, coal-fired power plants, wood-burning, metals operations, mining operations, and incinerators. Arsenic occurs naturally in many soils, so wind-blown dusts from exposed land can contain it. Mine tailings piles generally contain enriched levels of arsenic, resulting in emissions of arsenic in the particulate emissions that occur under windy conditions. Soils contaminated by smelter fall-out can also be a source of emissions during high winds. Burning wood treated with CCA also leads to arsenic emissions.

Arsenic's toxicity has led to its use as a poison. Orally ingesting large amounts can be fatal. The effects of inhalation are similar to the oral effects. Arsenic disturbs the gastro-intestinal system, leading to abdominal pain, vomiting, and diarrhea. It affects the central nervous system, leading to nerve damage in the legs and arms. It can damage the liver and kidney. Arsenic also has effects on the skin, causing dark patches (hyperpigmentation), and skin cancer. Arsenic also irritates the eyes, lungs, and skin. These effects have been observed in situations of

occupational exposure that are significantly higher than concentrations seen in outdoor air. Exposure can lead to effects in the blood, such as anemia.

EPA has not established a Reference Concentration for arsenic. The Cal EPA has established a chronic reference level based on the developmental effects in mice; and other target organs included the cardiovascular system and nervous system. Arsenic exposure is known to cause lung cancer. EPA classifies arsenic in Group A, the known human carcinogens, based on an increased lung cancer mortality in multiple human populations exposed primarily through inhalation.

The Agency for Toxic Substances and Disease Registry (ATSDR) states that remote areas have concentrations of 0.001 to 0.003  $\mu$ g/m3 arsenic in air, while urban locations range from 0.020 to 0.100  $\mu$ g/m3 (ATSDR Toxicological Profile on Arsenic). The mean level of 0.0003  $\mu$ g/m3 at Grand Junction – Powell site falls below the cited rural range. It is likely that national levels of arsenic have decreased in recent years.

### MANGANESE

Manganese is a metal that occurs naturally in the earth's crust. Its chemical symbol is "Mn". It exists in the atmosphere as particulate matter, in compounds formed from combination with other atoms. Manganese is used as an additive in metal processing and steel production. It is also used in ceramics, matches, glass, dyes, batteries, and as a pigment in paints (California Air Resources Board Fact Sheet on Manganese). It is also employed in wood preservatives. Organic forms of manganese are used as pesticides and for disease prevention in crops such as fruits, vegetables, and cotton.

Emission sources of manganese include petroleum refineries, steel producers, cement producers, coal-fired power plants, wood-burning, metals operations, mining operations, and incinerators. Manganese occurs naturally in some soils, so wind-blown dusts from exposed land can contain it. Soils contaminated by smelter fall-out can also be a source of emissions during high winds.

Manganese is considered an essential micronutrient in the human body. The body tends to regulate manganese concentrations, so oral exposure to small amounts naturally present in food is rarely a problem. Exposure of manganese by inhalation can lead to health effects. Manganese health effects on the respiratory system include lung irritation, chemical pneumonia, cough, and bronchitis. Manganese may damage the central nervous system. The disease known as "manganism", which results from manganese poisoning, includes psychological and nervous system damage. Individuals with manganism have a mask-like face, depression, uncontrollable laughter, and lethargy. The central nervous system effects include trouble with tremors, balance and walking that is similar to that of Parkinson's disease. Central nervous system damage can occur at exposure levels below those that lead to manganism. Examples are decreases in visual reaction time, hand steadiness, and eye-hand coordination. Manganese also affects the gastro-intestinal tract and the kidneys. However, it should be noted that these health effects have been observed in workers with long-term exposure to manganese fumes and dusts in industrial settings. These exposures were at levels hundreds or thousands of times higher than manganese levels in outdoor air.

EPA classifies manganese as Group D, unclassifiable as to carcinogenic potential. This is because there is little evidence to link it to cancer health effects. EPA has established a Reference Concentration for manganese based on an impairment of neurobehavioral function in humans in occupational exposure studies.

The California Air Resources Board monitored manganese in 1996. They report a network-wide average of  $0.0212~\mu g/m3$  total manganese (CARB Fact Sheet on Manganese). The 0.0130 annual mean measured at Grand Junction is below the California average.

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# Appendix B: Documentation for Grand Junction Urban Air Toxics Trends Monitoring Locations

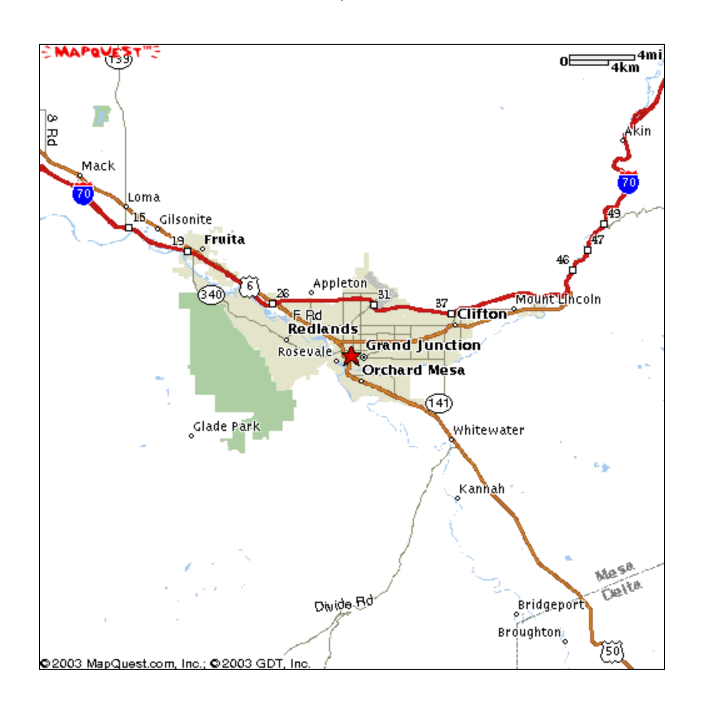
# **REGIONAL MAP (5 - 30 miles)**

AQS ID: <u>08-077-0017</u> Site Name: <u>Grand Junction – Powell Building</u>

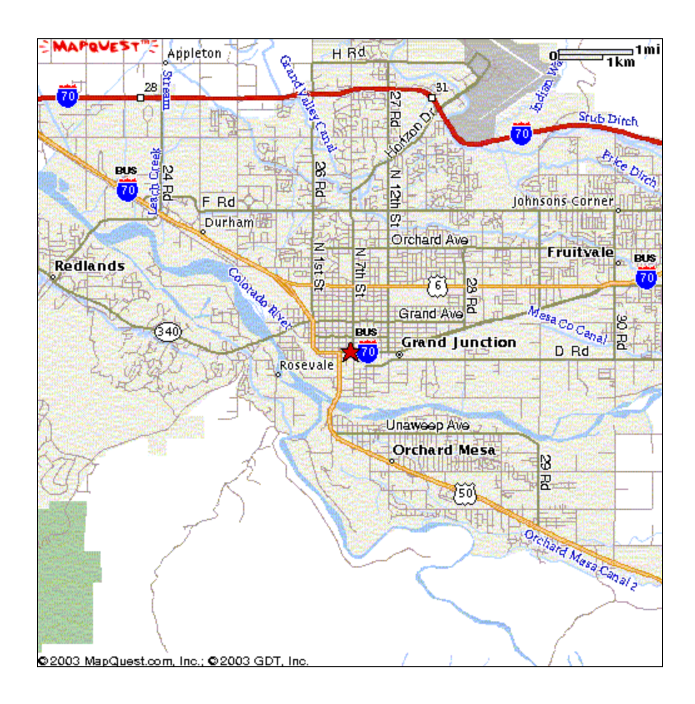
650 South Avenue, Grand Junction, CO 81501 GPS: Zone 12, 710962 E, 4326741 N, elev. 1396m 39° 03' 51" N, 108° 33' 42" W

AQS ID: <u>08-077-0018</u> Site Name: <u>Grand Junction – Pitkin Shelter</u>

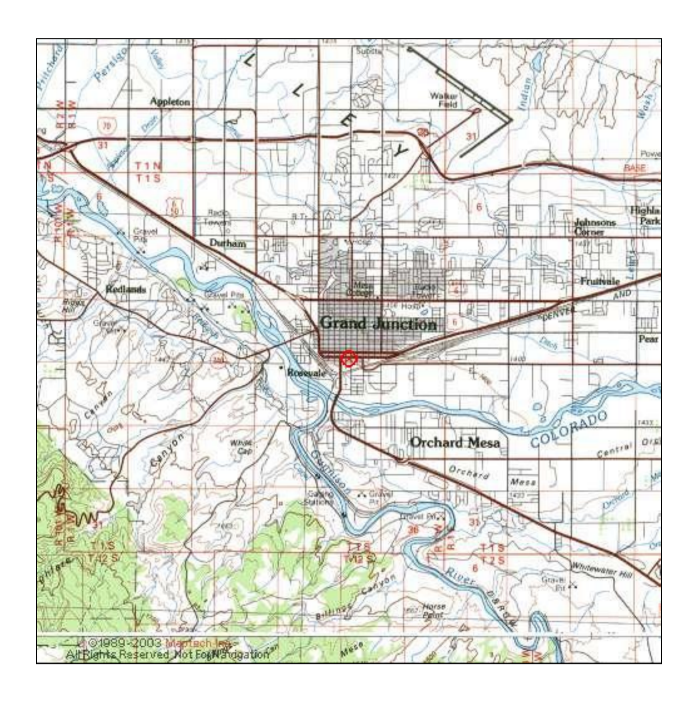
645 <sup>1</sup>/<sub>4</sub> Pitkin Avenue, Grand Junction, CO 81501 GPS: Zone 12, 710962 E, 4326741 N, elev. 1396m 39° 03' 51" N, 108° 33' 42" W



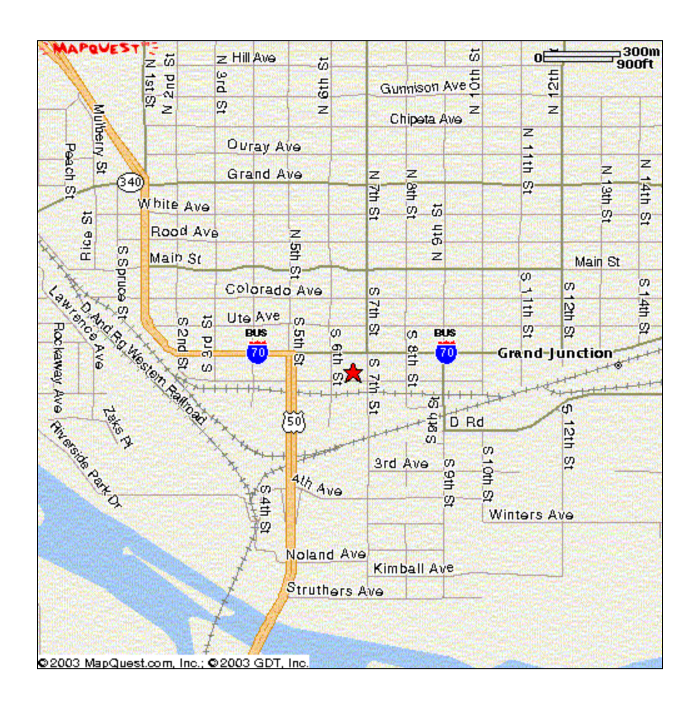
# **REGIONAL MAP (5 - 30 miles)**



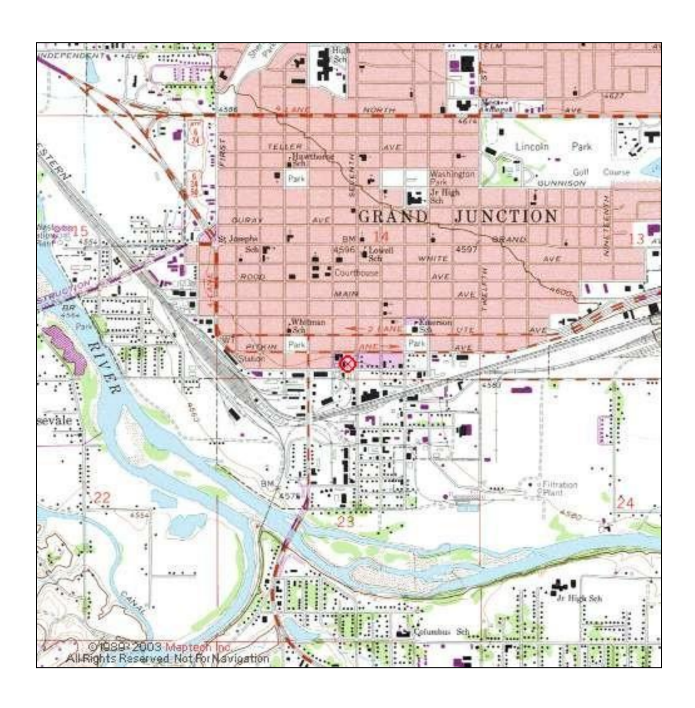
# **REGIONAL MAP (5 - 30 miles)**



# **SITE MAP (1/4 - 1 mile)**



# **SITE MAP (1/4 - 1 mile)**



# **SITE MAP (1/4 - 1 mile)**



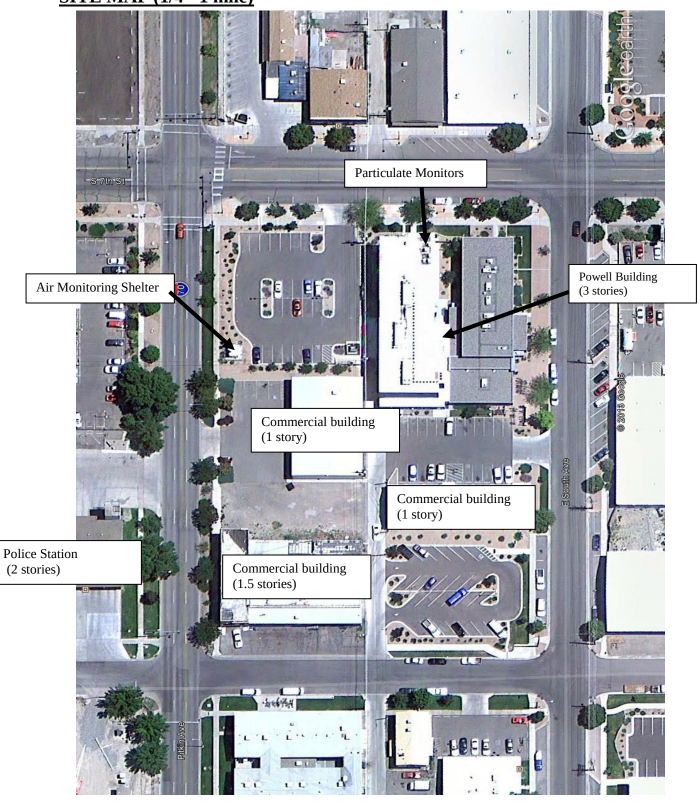


## **SITE MAP (1/4 - 1 mile)**



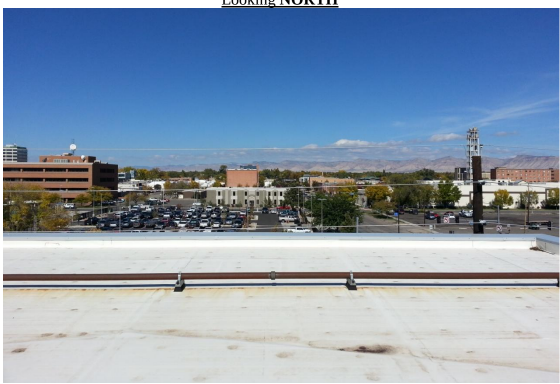


## **SITE MAP (1/4 - 1 mile)**



AQS ID: <u>08-077-0017</u> Site Name: Grand Junction – Powell Building
Photo Date: 10/16/2013





Looking NORTHEAST

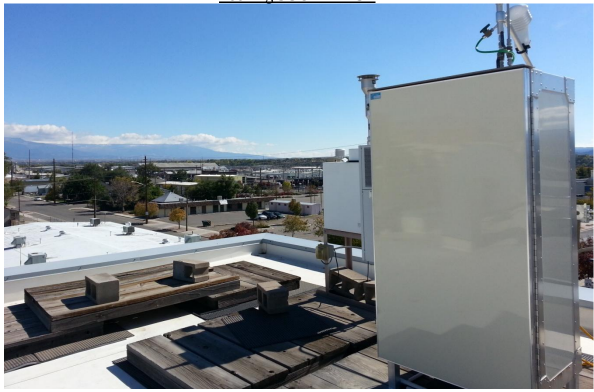


AQS ID: <u>08-077-0017</u> Site Name: <u>Grand Junction – Powell Building</u>
Photo Date: <u>10/16/2013</u>







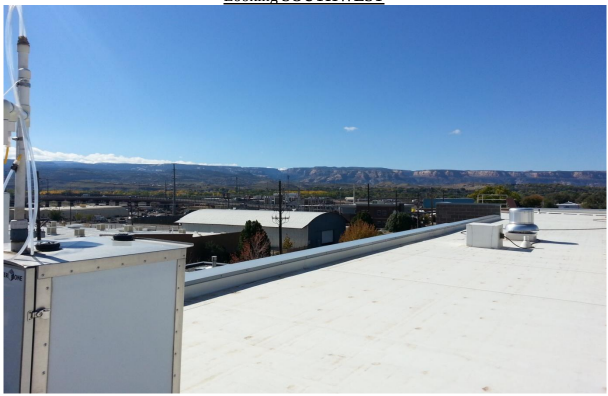


AQS ID: <u>08-077-0017</u> Site Name: <u>Grand Junction – Powell Building</u>
Photo Date: <u>10/16/2013</u>





Looking SOUTHWEST



AQS ID: <u>08-077-0017</u> Site Name: Grand Junction – Powell Building
Photo Date: 10/16/2013





Looking NORTHWEST



AQS ID: <u>08-077-0017</u> Site Name: Grand Junction – Powell Building
Photo Date: 10/16/2013





AQS ID: <u>08-077-0018</u> Site Name: Grand Junction – Pitkin Shelter
Photo Date: 04/24/2017





Looking NORTHEAST



AQS ID: <u>08-077-0018</u> Site Name: Grand Junction – Pitkin Shelter
Photo Date: 04/24/2017





Looking SOUTHEAST



AQS ID: <u>08-077-0018</u> Site Name: <u>Grand Junction – Pitkin Shelter</u>

Photo Date: <u>04/24/2017</u>

Looking **SOUTH** 



Looking **SOUTHWEST** 



AQS ID: <u>08-077-0018</u> Site Name: Grand Junction – Pitkin Shelter
Photo Date: 04/24/2017

Looking WEST



Looking NORTHWEST



Site Name: <u>Grand Junction – Pitkin Shelter</u> Photo Date: <u>04/24/2017</u> AQS ID: <u>08-077-0018</u>



