

Northwest Territories Air Quality Report 2007

Department of Environment and Natural Resources



**Northwest
Territories**

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INTRODUCTION

The Environment Division (ED) of the Department of Environment and Natural Resources (ENR) monitors air quality in the Northwest Territories (NWT). ENR maintains and operates the NWT Ambient Air Quality Monitoring Network, consisting of four monitoring stations located in Yellowknife, Fort Liard, Norman Wells and Inuvik. Each station is capable of continuously sampling and analysing a variety of air pollutants and meteorological conditions. The Yellowknife and Inuvik stations are operated in partnership with the National Air Pollution Surveillance (NAPS) program – a joint federal/provincial/territorial monitoring network tracking urban air quality trends throughout Canada.

ENR also monitors acid precipitation at Snare Rapids in cooperation with the Canadian Air and Precipitation Monitoring Network (CAPMoN) and conducts seasonal particulate sampling at Daring Lake.

The 2007 Annual Air Quality Report summarizes the air quality information collected in 2007, along with some discussion of trends. The report also provides information on network operations, the air pollutants monitored and the air quality standards used in assessing the monitoring results. Further information, including ‘almost real time’ air pollutant readings, can be found by visiting the NWT Air Quality Monitoring Network web site at:

<http://isin.rwed-hq.gov.nt.ca/NWTAQ/NetworkSummary.aspx>

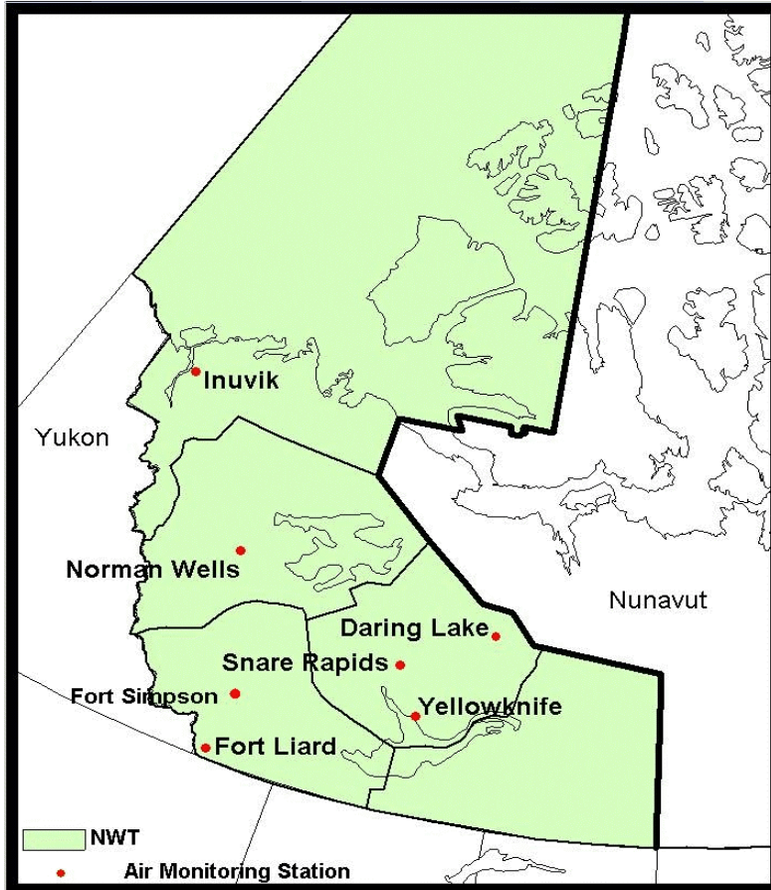
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This report is also available on the Internet at www.enr.gov.nt.ca/eps/environ.htm.

OPERATIONS (NETWORK)



The NWT Air Quality Monitoring Network consists of four permanent monitoring stations located in Yellowknife, Inuvik, Fort Liard and Norman Wells. The stations are climate controlled trailers and include state of the art monitoring equipment capable of continuously sampling and analysing a variety of air pollutants and meteorological conditions. Pollutants monitored vary by station but include sulphur dioxide (SO₂), hydrogen sulphide (H₂S), fine particulate (PM_{2.5}), particulate (PM₁₀), ground level ozone (O₃), carbon monoxide (CO) and nitrogen oxides (NO_x) as well as wind speed, wind direction and temperature. For additional

information on air pollutants see **Appendix B**.

Table 1 shows the breakdown of the NWT air monitoring network by substances and meteorological parameters monitored at each station.

Table 1:

Substances Monitored by Station											
Stations	Particulate Matter			Gaseous				Precipitation	Meteorological Monitoring		
	TSP - Particulate	PM_{2.5} - Fine Particulate	PM₁₀ - Particulate	SO₂ Sulphur Dioxide	H₂S Hydrogen Sulphide	NO_x Nitrogen Oxides	O₃ Ground Level Ozone	CO Carbon Monoxide	Acidic Deposition	Wind Speed & Direction	Air Temperature
Yellowknife	√	√	√	√		√	√	√		√	√
Inuvik		√	√	√	√	√	√			√	√
Norman Wells		√		√	√	√	√			√	√
Fort Liard		√	√	√	√	√	√			√	√
Snare Rapids									√		
Daring Lake		√									√

Using a sophisticated data acquisition system (DAS) and communications software, data from each station is automatically transmitted every hour via telephone line and government intranet to ENR headquarters in Yellowknife, allowing almost real time review of community air quality by ENR staff. The data also undergoes a series of ‘on the fly’ validity checks before being archived by ENR’s data management, analysis and reporting system.

The Yellowknife and Inuvik stations are part of a larger national network that monitors the common or criteria air pollutants in urban centres across Canada. The National Air Pollution Surveillance (NAPS) Network is a joint federal/provincial/territorial program, incorporating approximately 290 stations located in 175 communities, that monitor the same particulate and gaseous substances as those sampled in Yellowknife and Inuvik. Data from both these NWT stations along with data from other cities is summarized and assessed, with results published in the NAPS annual data reports available at:

http://www.etc-cte.ec.gc.ca/publications/napsreports_e.html.

The NAPS Network has a stringent quality assurance/quality control (QA/QC) program that ensures Canada-wide data is comparable. Participation in the NAPS program requires ENR to follow these QA/QC procedures at the Yellowknife and Inuvik sites, and ENR, in turn, applies these procedures at the other stations.

The Fort Liard and Norman Wells stations (and until recently, the Inuvik station) are Territorial stations that were set up in response to increasing resource development activity in the NWT and the potential for the associated emissions to affect air quality. The NAPS Inuvik station still fulfils its original Territorial goals along with its national urban monitoring objective. The primary Territorial objective of these stations is to establish baseline levels of SO₂, H₂S, NO_x, O₃ and PM ahead of development, as well as track the trends and cumulative impacts from source emissions should they occur.

ENR, with assistance from NWT Power Corporation staff, operates an acid precipitation collector at the Snare Rapids hydro-electric site. Daily rain and snow samples are collected and forwarded to the Canadian Air and Precipitation Monitoring Network (CAPMoN) for analysis. CAPMoN is a non-urban monitoring network with 28 measurement sites in Canada and one in the United States. Unlike NAPS, CAPMoN locates sites to limit the effect of anthropogenic sources. Most sites are remote and data is considered representative of background values.

ENR collects fine particulate data at the Daring Lake Tundra Ecosystem Research Station during the summer to establish typical background concentrations in the NWT. This research station is part of a circumpolar initiative called the Polar Continental Shelf Project and is designed as a research facility to conduct long-term research and monitoring of the tundra ecosystem.

Table 2 shows all of the various government affiliations involved with all of the monitoring stations.

Table 2:

NWT Air Quality Network			
	Partnership/Contract	Stations	Network
Environment and Natural Resources – Environment Division  Northwest Territories		Yellowknife and Inuvik	National Air Pollution Surveillance  Environment Canada Environnement Canada
		Fort Liard and Norman Wells	Northwest Territories stations  Northwest Territories
	NWT Power Corporation	Snare Rapids	Canadian Air and Precipitation Monitoring  Environment Canada Environnement Canada
	NWT Tundra Ecological Research Station	Daring Lake	Northwest Territories stations  Northwest Territories

Air quality monitoring in the NWT has evolved over time, beginning with a single TSP monitor in Yellowknife back in 1974, and progressing through various monitoring locations and equipment to reach the current stage of development.

Appendix A traces the history of air quality monitoring in the NWT.

DEVELOPMENTS IN 2007

In 2007, the upgrading of the NWT Air Quality Monitoring Network continued with the addition of monitoring equipment as follows:

- Fort Liard - A BAM Particulate Matter (PM₁₀) monitor and an Ozone (O₃) analyzer were installed and began collecting data in late August.

ENR completed the second phase of the Air Quality Monitoring Network web site, which included database related modifications as well as web design improvements. The ultimate goal for the web site is to provide useful and meaningful information to a broad range of users and be as intuitive as possible. The web site is available at <http://lisin.rwed-hq.gov.nt.ca/NWTAQ/NetworkSummary.aspx> and links with the data management, analysis and reporting system to provide public access to 'almost real-time' air quality data for each monitoring location. Access to archived data is available by querying the database using web-based tools. The web site also provides additional information on the Government of the Northwest Territories (GNWT) Air Quality Monitoring Network, air pollutants, monitoring equipment, as well as links to monitoring information collected in other jurisdictions.

FUTURE PLANS

ENR is exploring options to update the existing air quality data management, analysis and reporting system to the most recent version and continuing modifications will be made to the Air Quality Monitoring Network web site based on user comments and feedback.

A continuous PM₁₀ monitor will be added to the Norman Wells station to complete the particulate monitoring installations throughout the network.

An ultra-low level CO analyzer will be installed at the NAPS designated Inuvik station following additional testing by Environment Canada.

NWT AIR QUALITY STANDARDS

The Government of the NWT has adopted a number of concentration limits for protection of ambient (outdoor) air quality in the NWT. These limits apply to selected pollutants and are contained in the “Guideline for Ambient Air Quality Standards in the Northwest Territories”, established under the *NWT Environmental Protection Act*. They are summarized in **Table 3** below.

The NWT standards are used in the assessment of air quality monitoring data as well as determining the acceptability of emissions from proposed and existing developments. Where NWT standards are not available for a particular pollutant, the Canadian National Ambient Air Quality Objectives (national standards) or limits established in other jurisdictions are used.

Table 3:

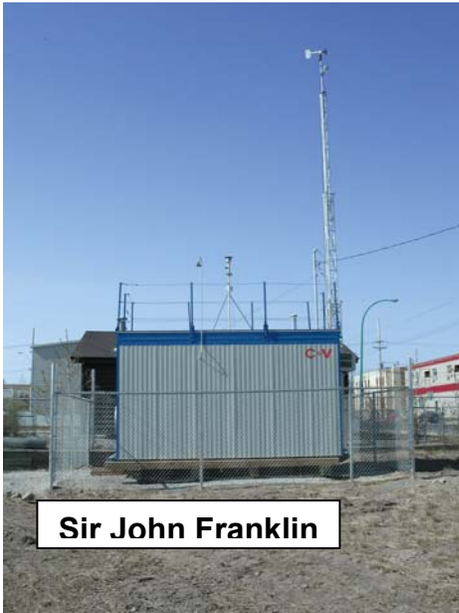
NWT Ambient Air Quality Standards		
Parameter and Standard	Concentration ($\mu\text{g}/\text{m}^3$)*	Concentration (ppbv)**
Sulphur Dioxide (SO_2)		
1-hour average	450	172
24-hour average	150	57
Annual arithmetic mean	30	11
Ground Level Ozone (O_3)		
8-hour running average	127	65
Total Suspended Particulate (TSP)		
24-hour average	120	
Annual geometric mean	60	
Fine Particulate Matter ($\text{PM}_{2.5}$)		
24-hour average	30	

* Micrograms per cubic metre

** Parts per billion by volume

The “Guideline for Ambient Air Quality Standards in the Northwest Territories” provides additional information on the application of the NWT standards and the pollutants of concern. For additional information on air pollutants see **Appendix B**.

YELLOWKNIFE AIR QUALITY



ENR, in partnership with the Canadian NAPS Program, operates the air quality monitoring station in Yellowknife.

This station is located at the Sir John Franklin High School and continuously monitors all criteria air pollutants O₃, SO₂, NO_x, CO, fine particulate (PM_{2.5}), and coarse particulate (PM₁₀). The station also monitors wind speed, wind direction, and temperature, which greatly assist in identifying possible sources of unusual or elevated readings.

Additional non-continuous monitoring at the station consists of a TSP (dust) High Volume sampler (Hi-Vol) and a Partisol Dichotomous fine particulate sampler, formerly located on the roof of the Post Office until 2005.

The air quality monitoring results from the consolidated station at Sir John Franklin are discussed in the following sections, with historical particulate data from the old Post Office site included where applicable.

TOTAL SUSPENDED PARTICULATE (TSP)

Yellowknife's greatest source of TSP is dust from roads, especially in the spring when the snow cover disappears and exposes winter sand and gravel applications on city streets to the effects of wind and vehicle disturbance. Forest fires, mining activities and combustion products from vehicles, heating and electricity generation also raise TSP levels.

Every six days on a predetermined schedule, TSP samples are collected over a 24-hour period (midnight to midnight) using a Hi-Vol air sampler. The sampler draws a measured volume of air through a filter to collect the suspended particulate. The filters are sent to Environment Canada's NAPS laboratory in Ottawa for analysis. TSP is measured as the weight of the particles collected on the filter divided by the volume of air drawn through the filter. Results are expressed in micrograms of particulate per cubic metre of air ($\mu\text{g}/\text{m}^3$). Arsenic levels are also determined in the samples collected.

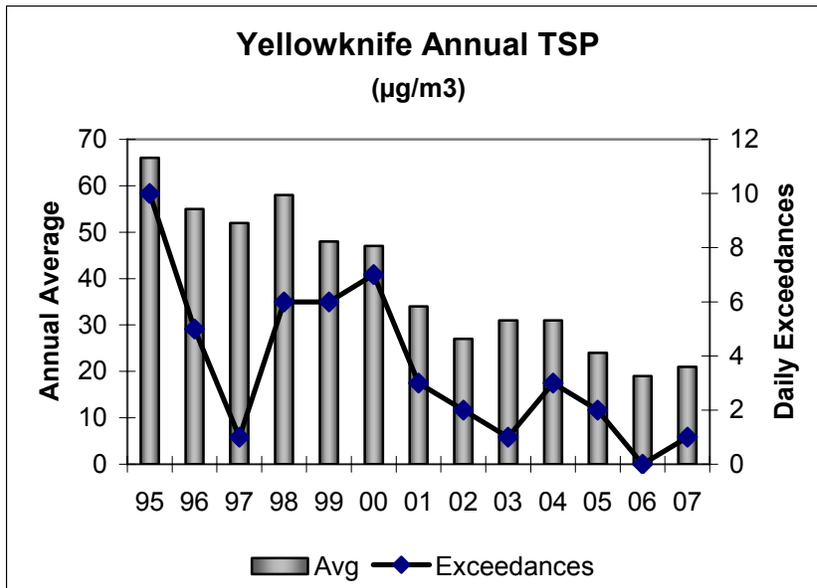
Table 4:

	Sir John Franklin		
	2006	2006	2007
$\mu\text{g}/\text{m}^3$	TSP	Arsenic	TSP
Highest (24-hr.)	90	0.03	121
Lowest (24-hr.)	3	0	2
Annual Average	19	.001	21

Fifty-seven Hi-Vol air samples were collected during 2007. **Table 4** summarizes the 2006 and 2007 TSP results including the annual average (geometric mean) as well as the 2006 arsenic results. The 2007 arsenic data was not available for the release of this report due to analytical delays at the NAPS laboratory.

The bars in **Figure 1** track annual average TSP levels in Yellowknife over the last decade (Post Office station 1995-2005; Sir John Franklin station 2006 and 2007). In 2007, the annual average was $21\mu\text{g}/\text{m}^3$. This is well below the NWT annual standard of $60\mu\text{g}/\text{m}^3$.

Figure 1:



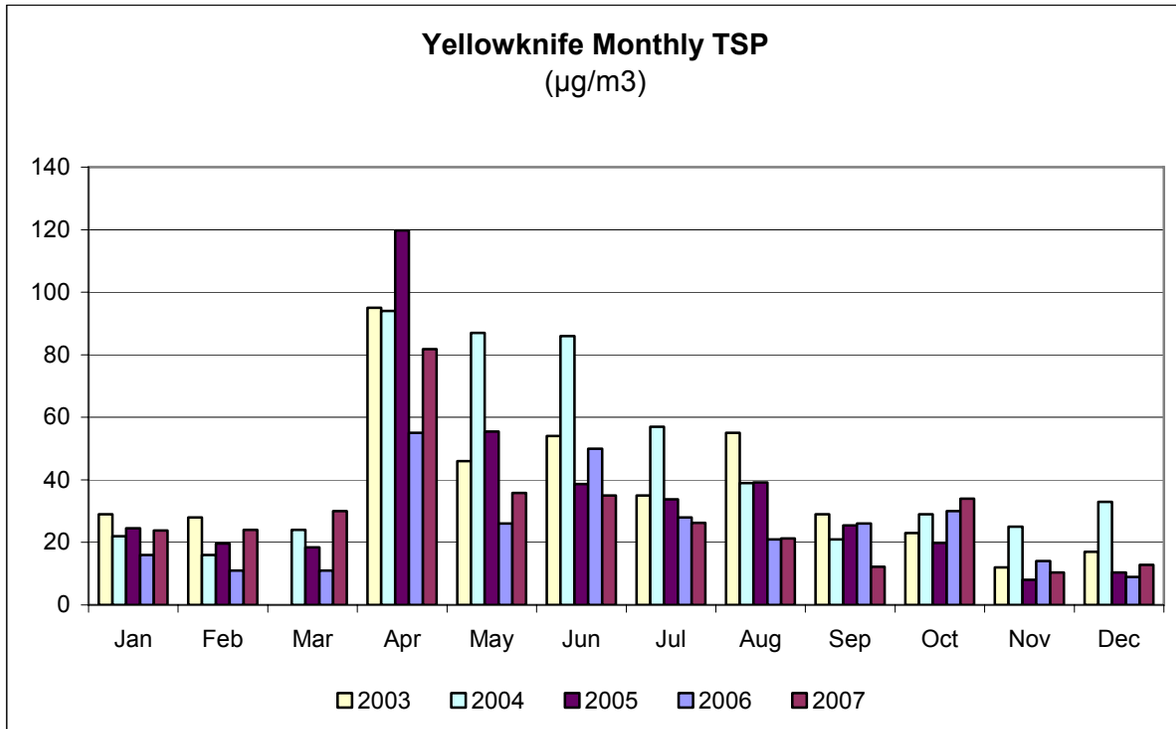
The line in **Figure 1** shows the number of times during the year that the NWT 24-hour standard ($120\mu\text{g}/\text{m}^3$) was exceeded. There was only one exceedance in 2007.

Since the early 1990's, dust conditions in Yellowknife have improved - largely due to the City of Yellowknife's efforts to clean roads throughout

the spring and summer as well as ongoing paving of graveled areas. The overall downward trend in annual TSP levels observed in the late 1990's and early part of the current decade appears to have leveled off in recent years. The recent annual average values may be approaching the lower limits achievable by the City's dust mitigation efforts and represent typical baseline readings. The fluctuations within the recent range of annual readings likely reflect the typical annual variations due to weather conditions and the intermittent sampling schedule.

Figure 2 tracks the average TSP level for each month for the last five years and clearly demonstrates the effects of springtime dust events. 2007 monthly averages followed the usual pattern with higher TSP levels in April through June.

Figure 2:



FINE PARTICULATE (PM_{2.5}) AND COARSE PARTICULATE (PM₁₀)

ENR uses two methods of sampling the smaller size fractions of PM in Yellowknife – a filter-based Partisol Dichotomous sampler and a Beta Attenuation Mass Monitor (BAM). The BAM methodology provides continuous, almost real-time (hourly) analysis of particulate concentrations, while the Partisol samples on a 24-hour basis every six days. The Partisol sampler simultaneously collects both the 2.5µm and less, and the 2.5 to 10µm particulate size fractions on a filter media. The filters require laboratory analysis to determine particulate concentrations and, unlike the BAM, do not provide timely information for real time air quality assessment. However, the filters can also be analysed for a whole suite of additional parameters including metals. The Partisol compliments the BAM in that particle identification can be determined which provides more in depth information about fine particulate. There are two BAMs operating at the Yellowknife station; one measures PM_{2.5} while the other measures PM₁₀.

Only two PM_{2.5} BAM readings at the Sir John Franklin station in 2007 exceeded the NWT 24-hour standard (30µg/m³). These exceedances were influenced by forest fire smoke. The maximum 24-hour PM_{2.5} value occurred on July 6th and coincided with forest fires burning in Lutselk'e, Fort Resolution, Dry Bones

Bay, as well as fires near Wekweètì and Yellowknife. Review and analysis of the results and wind information from the Sir John Franklin station and other meteorological stations clearly indicated the effects of fine particulate transport predominantly from these fires.

Figure 3:

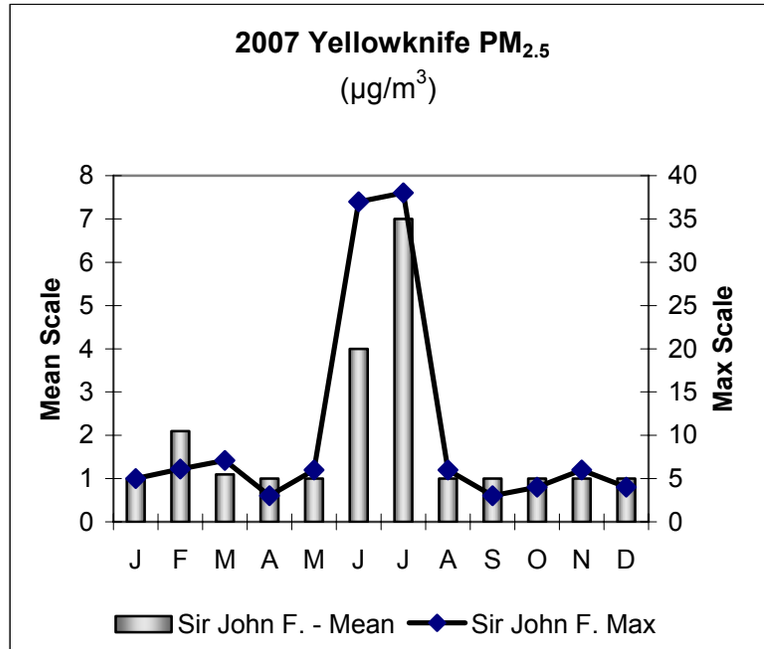


Figure 3 shows the monthly averages and daily maximums measured at the Sir John Franklin station in 2007. The highest daily maximum concentration was 38µg/m³. The highest hourly concentration recorded was 150µg/m³.

Figure 4:

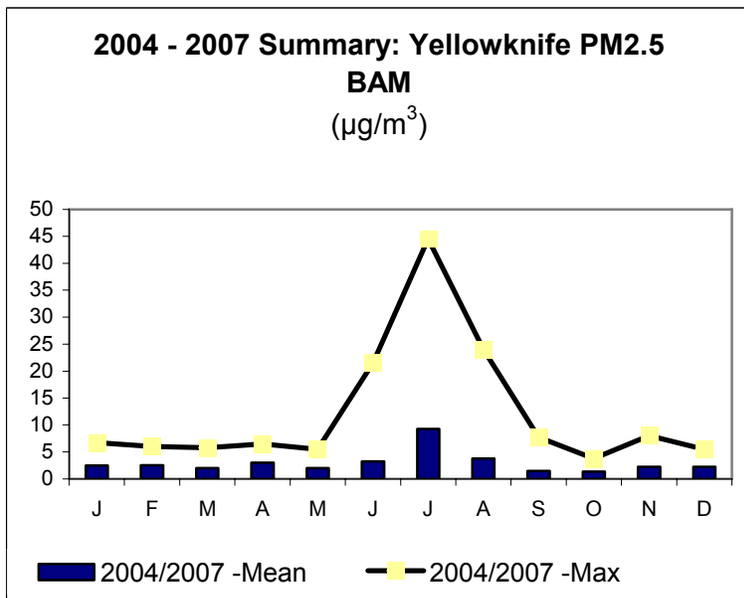


Figure 4 summarizes the PM_{2.5} BAM data over the last four years, showing the overall mean and overall daily maximum for each month. Overall means for each month are low and very consistent ranging from 3-10µg/m³, with an increase in July, which reflects the influence of forest fire events. The July and August maximums were a direct result of smoke from forest fires. If the forest fire events are ignored, then the daily maximums decline considerably, falling within a

range of 5-9µg/m³. Based on the results gathered so far, it appears that PM_{2.5}

concentrations in Yellowknife are consistently low, with the greatest short-term influences being smoke from forest fires.

Because PM₁₀ is a subset of TSP and heavily influenced by the same particulate sources like road dust, monitoring of PM₁₀ levels provides an alternative, more complete and timely method to the Hi-Vol TSP sampling to track the impacts of dust in Yellowknife. Since the Hi-Vol only runs every 6 days, it potentially misses capturing high-level dust events occurring in between the six day cycle. However, the continuous monitoring methodology of the BAM ensures complete coverage throughout the year. In addition, the hourly sampling schedule of the BAM provides almost “real time” data and allows more detailed analysis of the typical 24-hour Hi-Vol sampling period.

The NWT has not adopted a standard for PM₁₀ but several Canadian jurisdictions (e.g. BC, Ontario, Newfoundland and Labrador) have adopted a PM₁₀ concentration of 50µg/m³ (24-hour average) as an acceptable limit. In April 2007, Yellowknife experienced several episodes of very high PM₁₀ levels, resulting in complaints from residents and prompting the City to initiate road sweeping ahead of their normal spring clean up schedule. Overall, there were nine exceedances of the 50µg/m³ limit in 2007 at the Sir John Franklin station, most of which occurred in April.

Figure 5:

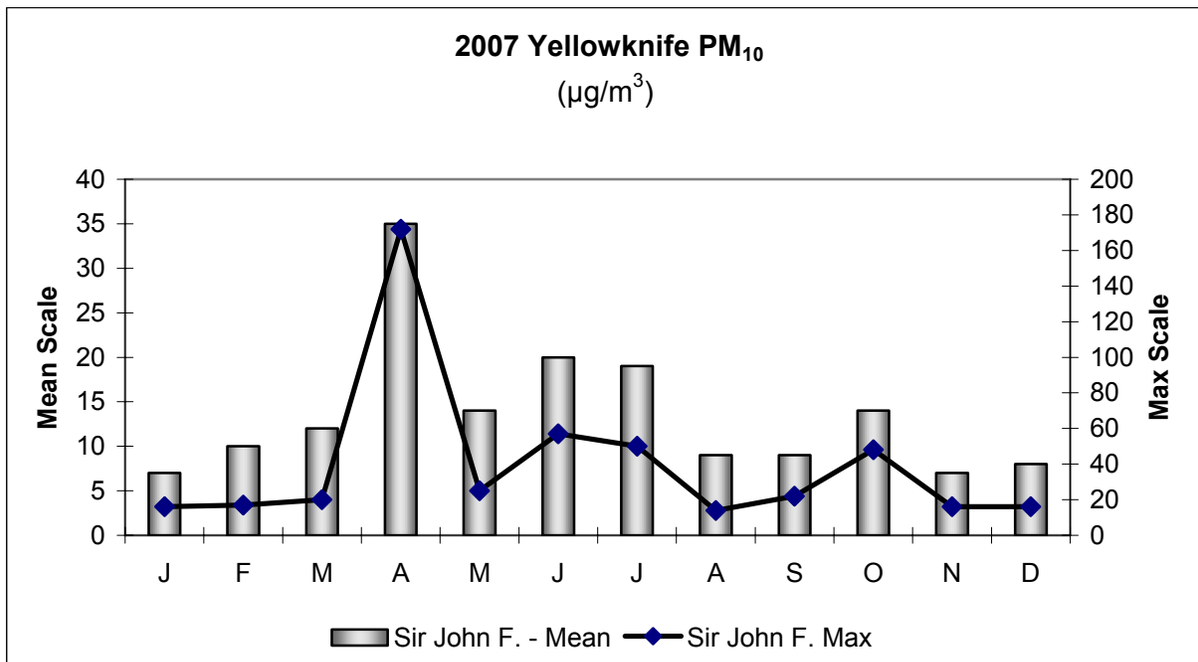


Figure 5 shows the PM₁₀ monthly averages and daily maximums measured at the station in 2007. The highest daily maximum concentration was 172µg/m³ occurring in April, which correlates well with the Hi-Vol data. The highest hourly concentration recorded was 505µg/m³ also in April. The monthly average distribution in **Figure 5** is similar to the distribution for TSP shown in **Figure 2** indicating the close correlation

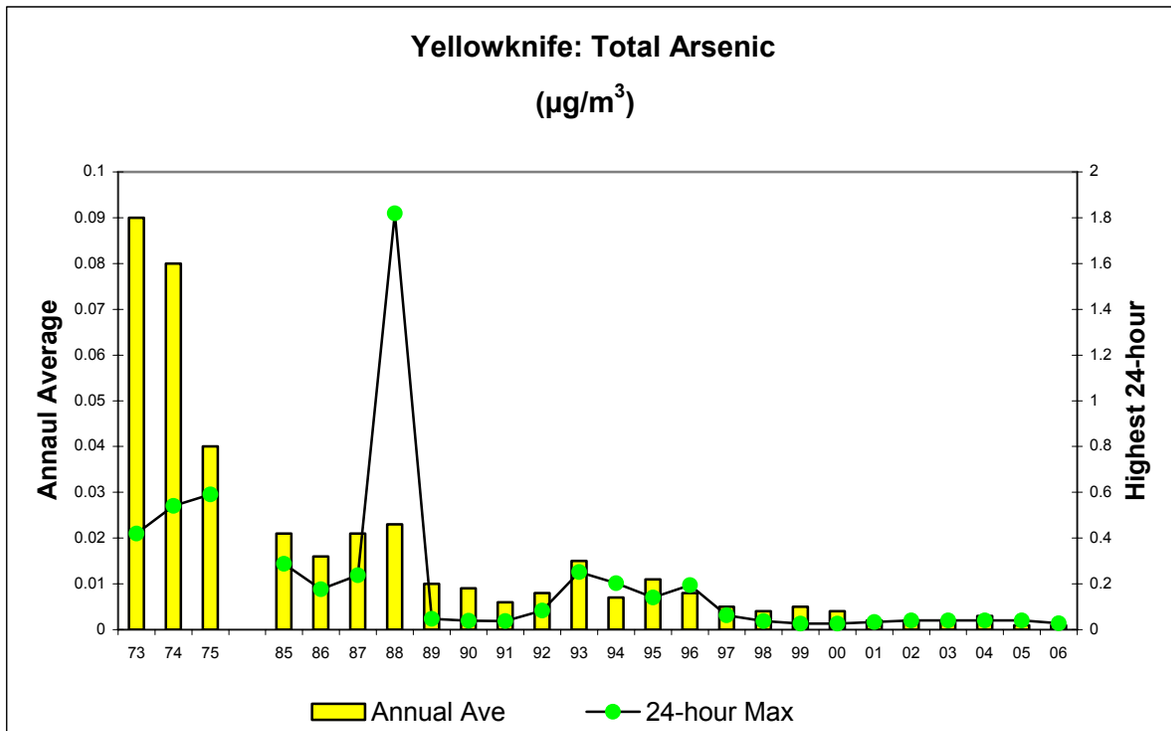
between PM₁₀ and TSP and supporting the use of the BAM PM₁₀ data to track the influence of road dust in Yellowknife.

ARSENIC

TSP filters are analysed to determine arsenic concentrations. Unfortunately, analytical delays at the NAPS laboratory often result in a 1-year time lag between arsenic data and the annual air quality report. Consequently, arsenic data for 2007 is not available for inclusion in this Report, although the 2006 data that was unavailable last year is. Analyses of the 2007 TSP filters will likely be completed by the end of June and results will be included in the 2008 Air Quality Report. ENR is discussing the time lag issue with NAPS in an effort to achieve more timely analysis.

Figure 6 summarizes the historical concentrations of arsenic measured in Yellowknife up to 2006. The bars represent annual averages while the line shows the highest total arsenic level measured over a 24-hour period for a given year. It is apparent that arsenic concentrations in Yellowknife have fallen from the historical elevated concentrations in the 1970s and 1980s. No exceedances of the Ontario Guideline (0.3µg/m³) have occurred since 1988 and the average arsenic concentration over the last decade is 0.006µg/m³ - below all but the most stringent World Health Organisation (WHO) risk estimate concentration (see Appendix B). In recent years, the overall average has decreased further to less than 0.003µg/m³.

Figure 6:



The maximum 24-hour concentration in 2006 was $0.03\mu\text{g}/\text{m}^3$ - almost an order of magnitude lower than the Ontario Guideline. The annual average (geometric mean) concentration in 2006 was $0.001\mu\text{g}/\text{m}^3$, considerably lower than all but the most conservative risk estimate concentration provided by the WHO.

Studies in the United States show an average arsenic concentration in air of $<0.001 - 0.003\mu\text{g}/\text{m}^3$ in remote areas, with concentrations of $0.020 - 0.030\mu\text{g}/\text{m}^3$ in urban areas, while Canadian urban areas range from 0.0005 to $0.017\mu\text{g}/\text{m}^3$ (WHO, 2000). Yellowknife arsenic results over the last decade indicate annual concentrations ranging from 0.002 to $0.015\mu\text{g}/\text{m}^3$ - comparable to concentrations measured in other urban areas. Since 1999, the levels have decreased and the annual average concentrations (0.002 - $0.004\mu\text{g}/\text{m}^3$) are similar to those in remote areas.

Based on the previous discussion, it appears that concentrations of airborne arsenic in Yellowknife are typical of those found in remote areas and that health risks due to inhalation are minimal.

SULPHUR DIOXIDE (SO₂)

Continuous monitoring for SO₂ has been conducted in Yellowknife at several locations since 1992.

In the past, the largest sources of SO₂ in the Yellowknife area were the gold mine ore roasters, the most recent being Giant Mine. The highest levels of SO₂ in the Yellowknife area were measured downwind from the mine. Since the mine was closed in 1999, only background levels have been recorded.

Figure 7:

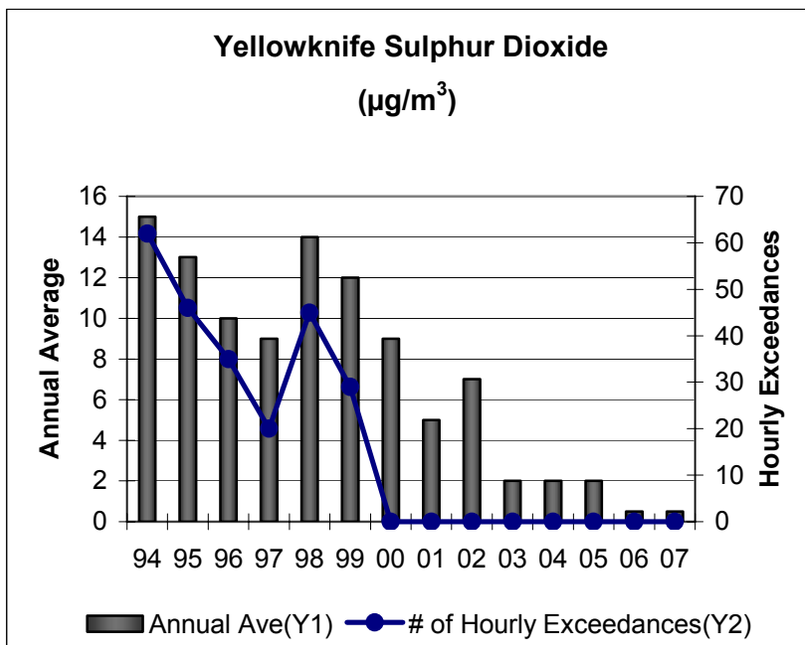


Figure 7 shows the general trends in SO₂ levels measured in Yellowknife air from 1994 to 2007. The bars track the annual average and the line shows the number of times in each year that the NWT 1-hour standard was exceeded. As illustrated by the graph, the number of exceedances has fallen to zero since the closure of Giant Mine in 1999.

The 2007 data continued the trend of recent years, with no exceedances of the NWT hourly ($450\mu\text{g}/\text{m}^3$) and 24-hour ($150\mu\text{g}/\text{m}^3$) standards. The annual average was less than $1\mu\text{g}/\text{m}^3$, a level that is well below the NWT ($30\mu\text{g}/\text{m}^3$) standard.

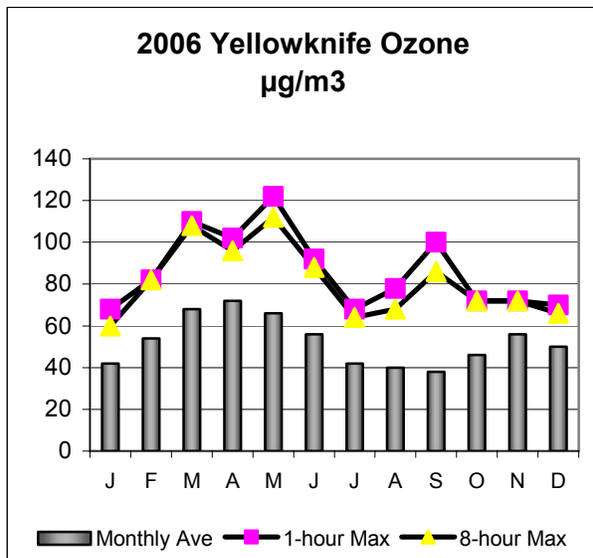
The vast majority of the hourly concentrations recorded in 2007 were only background or slightly greater, with a maximum concentration of $16\mu\text{g}/\text{m}^3$. The concentrations reflect naturally occurring SO_2 , usually in the range of 3 to $4\mu\text{g}/\text{m}^3$, and small amounts from the burning of fossil fuels.

GROUND LEVEL OZONE (O_3)

A continuous O_3 analyzer has been operated in Yellowknife since 1998. The current analyzer has been operating at the Sir John Franklin station since February of 2003.

Detectable concentrations of O_3 exist even in remote areas due to naturally occurring sources of the precursor gases such as forest fires and volatile organic compounds (VOC) emissions from trees and the introduction of stratospheric ozone to lower elevations resulting from atmospheric mixing processes. These background concentrations typically are in the range of 40 to $80\mu\text{g}/\text{m}^3$. In large urban areas (and areas downwind), ozone concentrations can be much higher than typical background due to the additional emissions of precursor gases (see **Appendix B**).

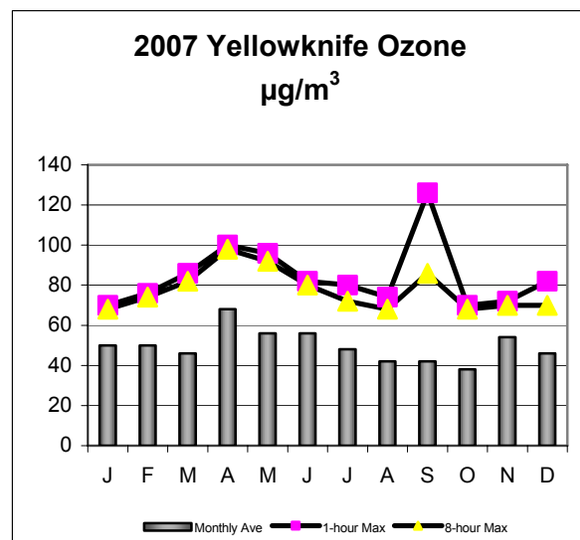
Figure 8:



The **Figures** illustrate the typical spring maximum which commonly occurs at remote monitoring stations located in mid to high latitudes in the Northern hemisphere and the source of which is the subject of considerable scientific

Figures 8 and 9 show the hourly and 8-hour maximums recorded for 2006 and 2007, together with the monthly averages. Neither the 1-hour national standard nor the 8-hour NWT standard was exceeded in Yellowknife during 2006 or 2007.

Figure 9:



debate. Typical monthly ozone concentrations at remote sites in Canada range between 40 and 80µg/m³ and Yellowknife concentrations for both years fell below or within this range, indicating that most of the O₃ detected is likely naturally occurring or background.

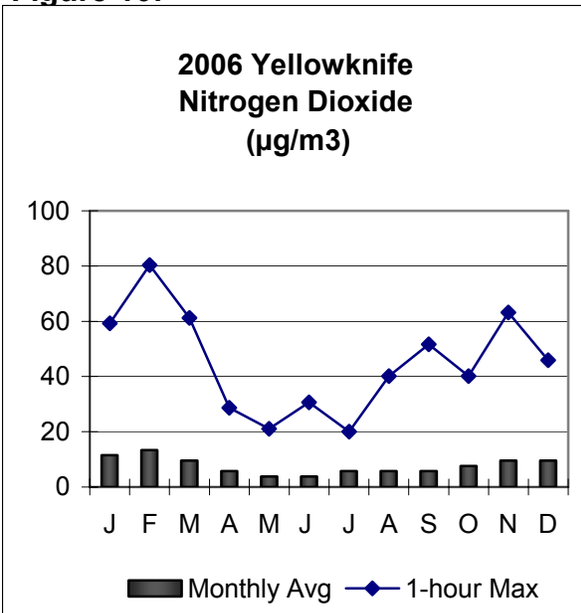
NITROGEN OXIDES (NO_x)

The instrumentation provides continuous information on NO, NO₂ and NO_x. However, the focus is on NO₂ due to the greater health concerns associated with this pollutant and the availability of national air quality standards for comparison (see **Appendix B**).

The 2007 results indicated that there were no exceedances of the 1-hour and 24-hour national standards for NO₂. The maximum 1-hour average was 105µg/m³ and the annual average was 5µg/m³.

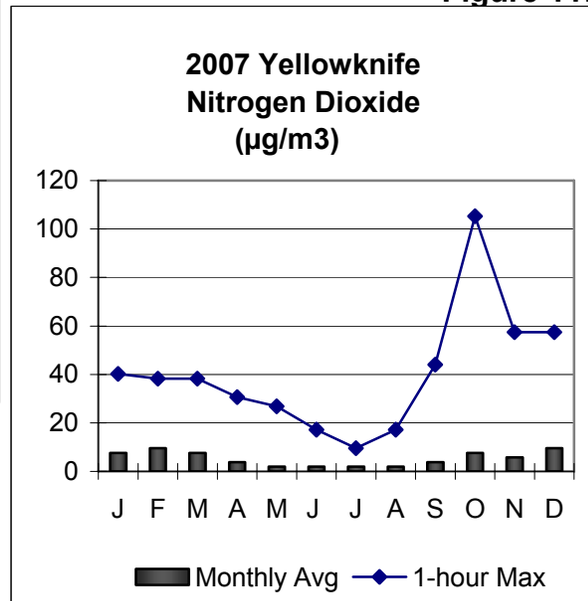
Figures 10 and 11 show the 2006 and 2007 monthly averages and highest hourly concentrations. Both the highest monthly averages and the highest hourly concentrations occurred during the winter months.

Figure 10:



This is likely caused by increased emissions from fuel combustion for residential and commercial heating, and idling vehicles, as well as short-term “rush hour” traffic

Figure 11:

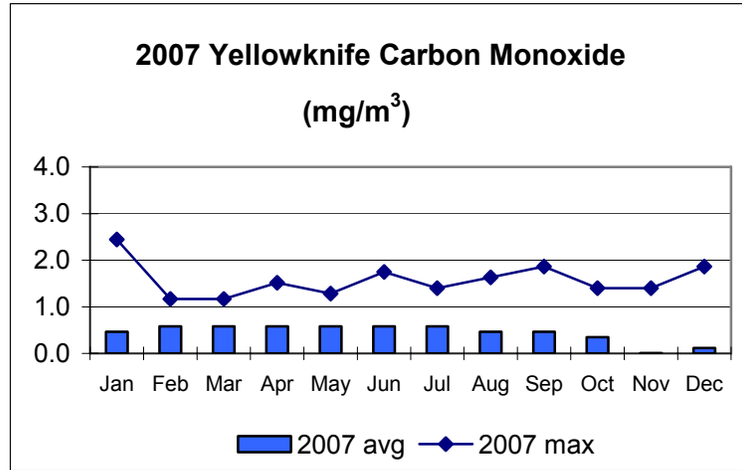


influences. The effects of these emissions on winter-time air quality can be increased when combined with stagnant meteorological conditions. Cold, calm days can result in an atmospheric situation where the normal decrease in air temperature with elevation is reversed and a zone of colder air is present at ground level. This zone of colder air and the lack of wind act to restrict dispersion and trap pollutants close to the ground.

CARBON MONOXIDE (CO)

The 2007 data continued the extremely low CO readings measured in 2006 and were well below the national standards (see **Appendix B**). Given the absence of heavy traffic volumes in Yellowknife, low levels of CO are expected. In 2007 the maximum 1-hour average was 2.4 mg/m³ and the annual average was 0.4 mg/m³. **Figure 12** shows the 2007 monthly averages and highest hourly concentrations.

Figure 12:



INUVIK AIR QUALITY

The focus of the monitoring station in Inuvik is to gather baseline community air quality information and to track trends and cumulative effects of pollutant sources over time. In January 2006, the station was incorporated into the National Air Pollution Surveillance (NAPS) Network and also provides air quality information to this national monitoring network for comparison to other communities in Canada.

The station measures SO₂, H₂S, NO_x, O₃, PM_{2.5}, and PM₁₀. Both the NO_x and the O₃ analyzers were supplied by Environment Canada (Yellowknife office).

HYDROGEN SULPHIDE (H₂S)

The data collected in 2007 continues to indicate very low H₂S concentrations in Inuvik – essentially non-detectable. Most of the readings are less than 1 µg/m³, which is below the detectable limits of the instrumentation and within the ‘noise’ range.

There were no exceedances of the Alberta Guidelines and the maximum recorded 1-hour average was 6 µg/m³.

SULPHUR DIOXIDE (SO₂)

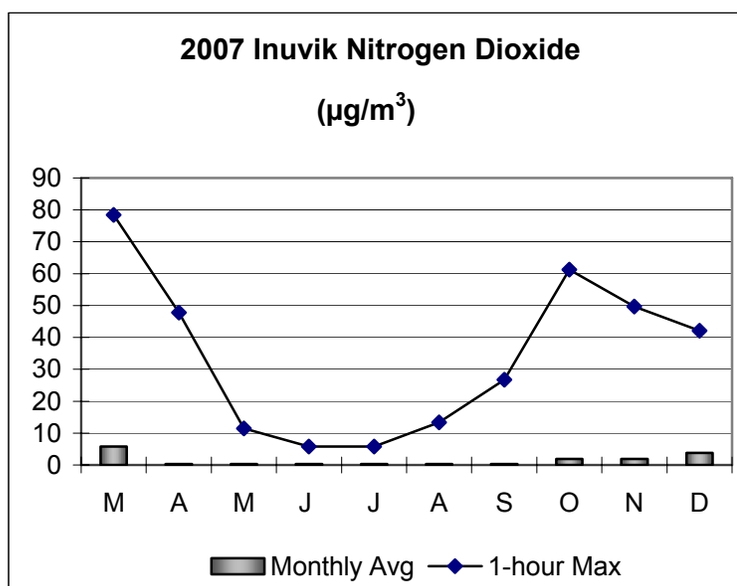
The SO₂ concentrations measured in 2007 were very low and similar to last year’s results, with no exceedances of the NWT standards. The overall average was 1 µg/m³ and the maximum 1-hour average was 11 µg/m³.

NITROGEN OXIDES (NO_x)

As discussed in the Yellowknife section of the report, the focus of NO_x monitoring is on the NO₂ portion in determining air quality, although NO is certainly of interest for other reasons (e.g. assessment of secondary pollutant formation). The data collected in Inuvik in 2007 shows that there were no exceedances of the 1-hour and 24-hour national standards for NO₂ (see **Appendix B**). The maximum 1-hour average was 57 µg/m³ and the annual average was 2 µg/m³.

Figure 13 shows the 2007 monthly averages and highest hourly concentrations. Unfortunately, the analyzer suffered a serious breakdown in Northwest Territories Air Quality Report 2007

Figure 13:



January, which necessitated lengthy repairs and resulted in no data collection for the first two months of 2007.

As with Yellowknife, both the highest monthly averages and the highest hourly concentrations occurred during the winter months. Meteorological data collected over the last three winters has shown that Inuvik is prone to winter inversions and experiences more inversion days than the other three regions. It is not uncommon in the middle of winter to see consecutive days of extremely cold temperatures accompanied with very low wind speeds (calms), reducing dispersal of pollutants.

FINE PARTICULATE ($PM_{2.5}$)

The 2007 BAM readings produced an annual average of $4\mu\text{g}/\text{m}^3$. There were no exceedances of the NWT standard for $PM_{2.5}$, which was expected, as there were no significant impacts from forest fire events during the summer of 2007.

Figure 14:

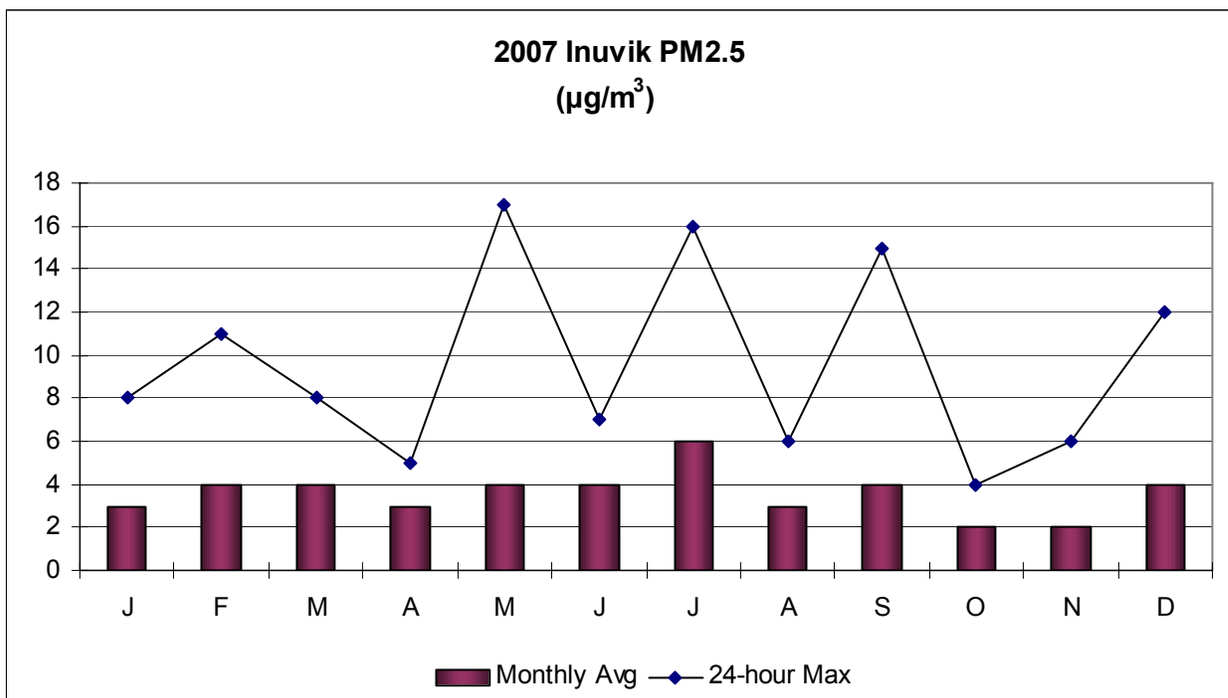


Figure 14 shows the monthly averages and daily maximums measured at the Inuvik station in 2007. The maximum 24-hour $PM_{2.5}$ value of $17\mu\text{g}/\text{m}^3$ occurred during the month of May and was associated with fine dust generated from construction in the area.

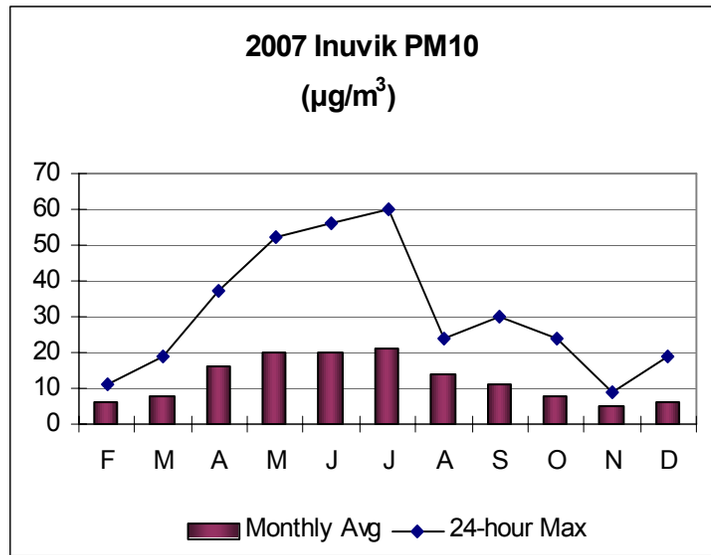
COARSE PARTICULATE (PM_{10})

2007 was the first full year of PM_{10} data collection with the exception of January. This data loss was due to a faulty circuit board in the instrument.

Figure 15:

averages and the daily maximum concentrations. The highest daily maximum was $60\mu\text{g}/\text{m}^3$ recorded in July and there were four exceedances of the $50\mu\text{g}/\text{m}^3$ (24-hour average) acceptable limit used in other jurisdictions. Unlike Yellowknife, the Inuvik readings do not show a 'spring maximum' associated with residual winter gravel, although the highest hourly concentration recorded ($316\mu\text{g}/\text{m}^3$) did occur in April. The summer monthly average values are similar to Yellowknife but the daily maximums in Inuvik are somewhat higher, indicating a longer "dust season" persisting throughout the summer months. This may reflect the greater percentage of unpaved roads and/or exposed areas in Inuvik providing a continuing source of dust throughout the spring and summer months. However, this speculation is based on only one year of complete data and it will be interesting to see if this pattern persists in future years.

Figure 15 shows the monthly



GROUND LEVEL OZONE (O_3)

Neither the 1-hour national standard nor the 8-hour NWT standard was exceeded in 2007. The typical 'spring time maximum' also occurred at this site and is consistent with

the Yellowknife and Norman Wells stations, which also experience a peak in O_3 levels during the spring months. The analyzer had to be taken off line during the months of October and November due to a malfunction.

Figure 16:

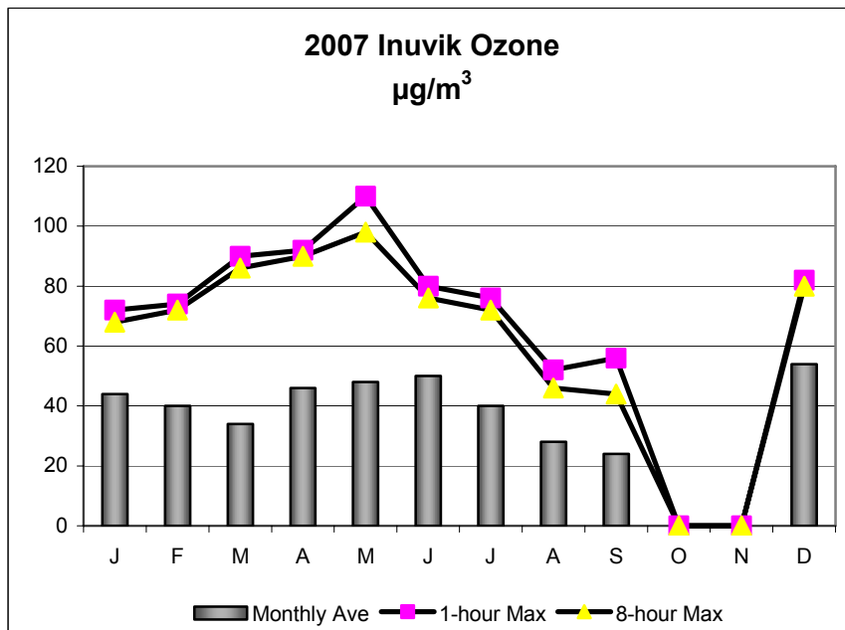


Figure 16 shows the hourly and 8-hour maximums as well as the monthly averages recorded for 2007.

FORT LIARD AIR QUALITY

The focus of the monitoring station in Fort Liard is to gather baseline community air quality information and to track trends and cumulative effects of pollutant sources over time.

HYDROGEN SULPHIDE (H₂S)

The maximum hourly H₂S concentration in 2007 was 3µg/m³ and the vast majority of readings were less than 1µg/m³, essentially within the detection limits or 'noise' range of the analyzer. H₂S in Fort Liard is, therefore, considered largely non-detectable.

SULPHUR DIOXIDE (SO₂)

As in previous years, there were no hourly or 24-hour exceedances of the NWT standards in 2007, with a maximum 1-hour average value of only 5µg/m³. The monthly averages were very low, with values less than 1µg/m³ - consistent with those measured over the last four years.

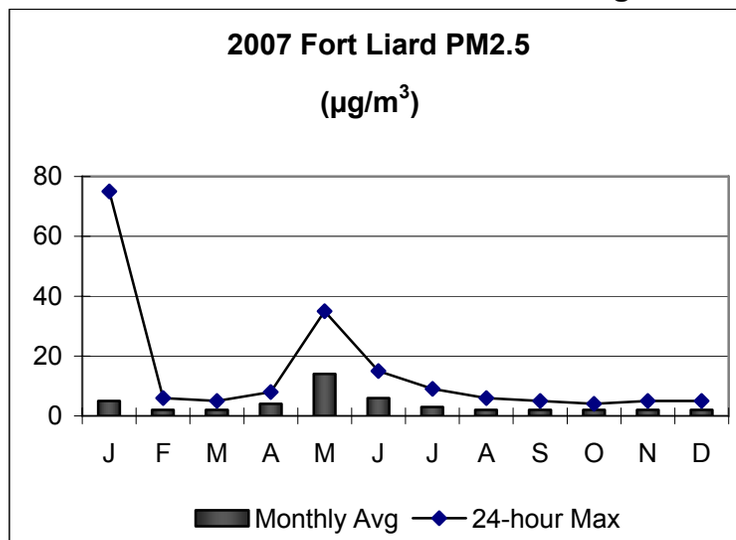
Elevated 1-hour SO₂ values occur occasionally in Fort Liard, but these are rare isolated peaks and still far below the NWT 1-hour standard. Overall, the data indicates that SO₂ concentrations in Fort Liard are within the expected range of background values and there appears to be no impact on community air quality from oil and gas development in the area.

FINE PARTICULATE (PM_{2.5})

Figure 17 shows the monthly averages and daily maximums measured at the Fort Liard station in 2007.

The 2007 annual PM_{2.5} average concentration was 4µg/m³ just 1 µg/m³ higher than the 2006 annual average. The maximum daily average was 75µg/m³ and was attributed directly to localized brush burning on the grounds where the air monitoring trailer is located. There were four exceedances of the NWT 24-hour standard for PM_{2.5}. One occurred in January as a result of brush burning and the other three occurred in May and are associated with transboundary smoke from forest fires burning south of the Northwest Territories.

Figure 17:



COARSE PARTICULATE (PM_{10})

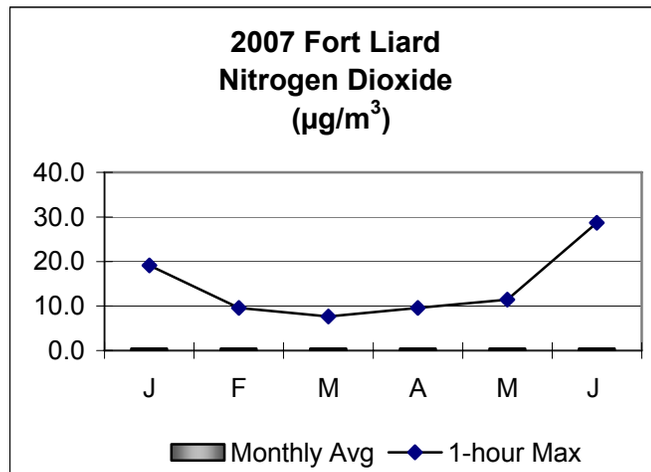
A PM_{10} BAM was added to the station in late August. Based on four months of data collection, the average concentration was only $4 \mu\text{g}/\text{m}^3$ and the highest daily maximum concentration was $17 \mu\text{g}/\text{m}^3$. The highest hourly concentration recorded was $83 \mu\text{g}/\text{m}^3$.

NITROGEN OXIDES (NO_x)

The analyzer was taken off line in July following a major component failure. Attempts at on-site repairs were unsuccessful and it will most likely have to be replaced with a new unit while the original analyzer is shipped back to the manufacturer for repairs. The six months of data collected in 2007 show that NO_2 concentrations continue to remain very low, with no exceedances of the 1-hour and 24-hour national standards. The 'annual' average was less than $1 \mu\text{g}/\text{m}^3$ and the 1-hour maximum concentration was $29 \mu\text{g}/\text{m}^3$.

Figure 18:

Figure 18 shows the 2007 monthly averages and highest hourly concentrations for the six months of instrument operation. The maximum NO_2 concentrations usually occur during winter as was the case in 2006 but limited data in 2007 do not illustrate this in the accompanying chart. The June peak was attributed to local combustion sources near the station. The overall lower levels (compared to the other three stations) are probably reflective of the generally lower emissions due to a smaller community base.



NORMAN WELLS AIR QUALITY

The focus of the monitoring station in Norman Wells is to gather baseline community air quality information and to track trends and cumulative effects of pollutant sources over time.

HYDROGEN SULPHIDE (H_2S)

As in 2006, the 2007 data from the station continued to produce extremely low readings with no exceedances of the Alberta Guidelines. Most hourly averages were essentially zero with a maximum 1-hour concentration of $2\mu\text{g}/\text{m}^3$.

SULPHUR DIOXIDE (SO_2)

The 2007 SO_2 monitoring program produced similar results to the Fort Liard, Yellowknife and Inuvik stations. The 1-hour maximum was $5\mu\text{g}/\text{m}^3$. Overall SO_2 concentrations were generally very low. The annual average was less than $2\mu\text{g}/\text{m}^3$ and no exceedances of the NWT standards occurred.

NITROGEN OXIDES (NO_x)

The 2007 results show that there were no exceedances of the 1-hour and 24-hour national standards for NO_2 . As with the other three network stations, NO_2 levels increase in the winter months. The maximum 1-hour average was $55\mu\text{g}/\text{m}^3$ and the overall average was $3\mu\text{g}/\text{m}^3$.

Figure 19:

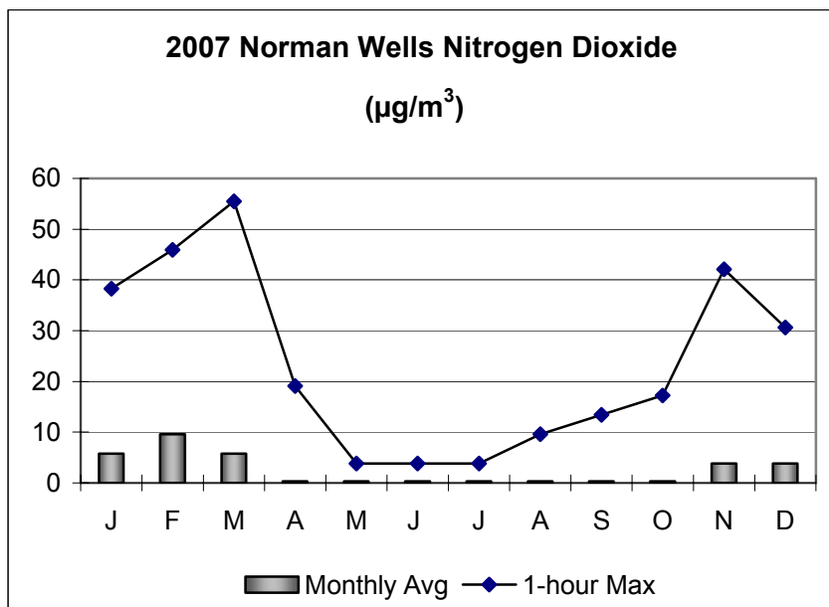
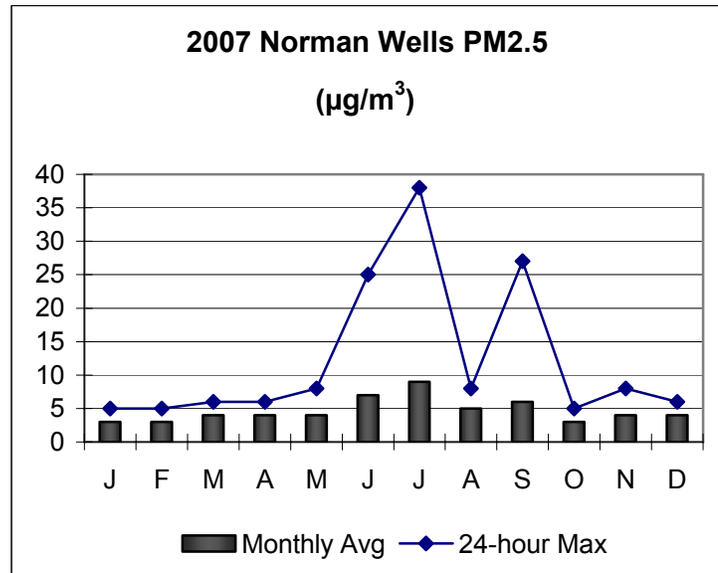


Figure 19 shows the 2007 monthly averages and highest hourly concentrations.

FINE PARTICULATE ($PM_{2.5}$)

Figure 20 shows the monthly averages and daily maximums measured at the Norman Wells station in 2007. The maximum daily concentration was $39\mu\text{g}/\text{m}^3$ and the annual average was $5\mu\text{g}/\text{m}^3$. There was some influence from one specific forest fire located approximately 20 kilometres from the town near the Carcajou River. Smoke from this fire produced the elevated readings during the months of June, July, and September, and resulted in one exceedance of the NWT standard.

Figure 20:



GROUND LEVEL OZONE (O_3)

No exceedances of the 1-hour national standard or the 8-hour NWT standard were observed in 2007. The elevated spring time levels were also evident at this site and are consistent with the Yellowknife and Inuvik stations which also experience a peak in O_3 levels during the spring months.

Figure 21:

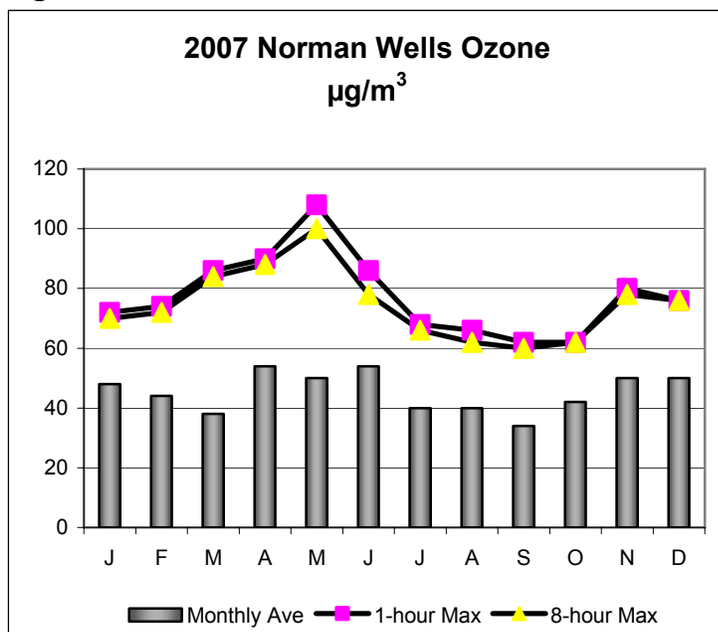


Figure 21 shows the hourly and eight hour maximums as well as the monthly averages recorded for 2007.

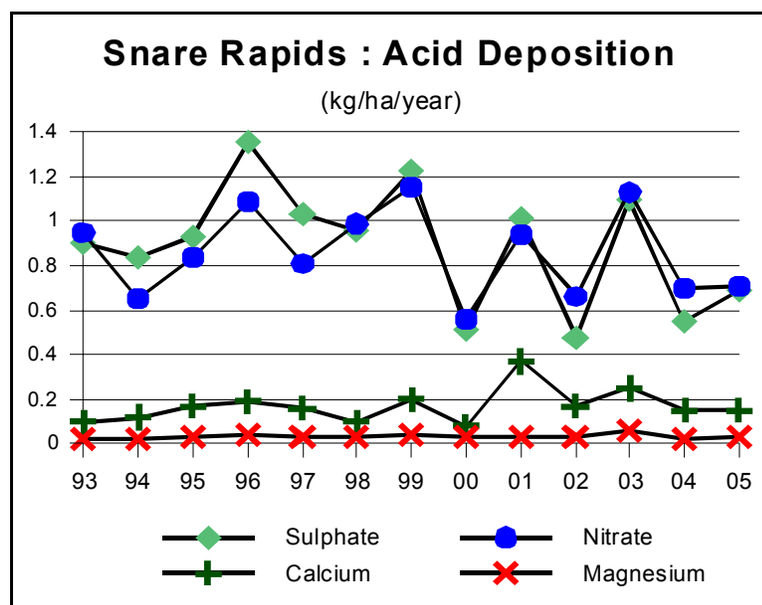
SNARE RAPIDS

Since 1989, ENR has operated a Canadian Air and Precipitation Monitoring (CAPMoN) station at the NWT Power Corporation's Snare Rapids hydro site. This site is located 150 kilometres northwest of Yellowknife. Rain and snow samples are collected on a daily basis and sent to Environment Canada's CAPMoN laboratory in Toronto for analysis. Quality assurance/quality control checks on the 2007 data had not been completed by Environment Canada in time for this report and, unfortunately the 2006 results not available last year are still undergoing validity checks and remain unavailable.

ACID PRECIPITATION

With no updates to report for 2006 and 2007, **Figure 22** shows the deposition rates for sulphate, calcium, nitrate and magnesium up to 2005.

Figure 22:

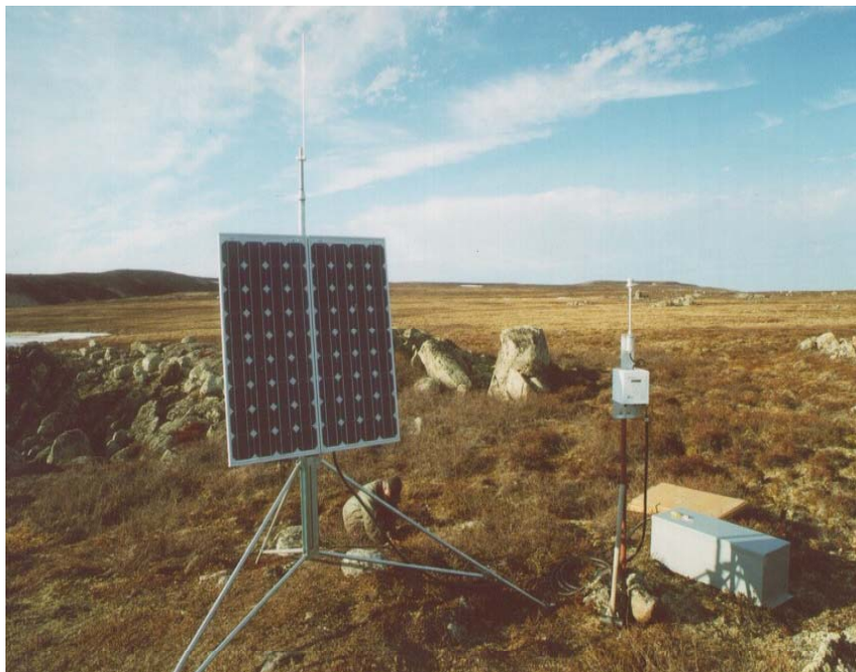


7 kg/ha/yr sulphate is the level considered to protect sensitive ecosystems in the NWT. In areas of eastern Canada, where acid rain is a serious environmental problem, sulphate deposition is in excess of 20 kg/ha/yr. Nitrate deposition at Snare Rapids is similarly low compared to eastern Canada. Sulphate and nitrate deposition rates remain well below levels that could cause an environmental effect in sensitive ecosystems.

DARING LAKE SEASONAL PARTICULATE

In summer of 2007, the short-term seasonal particulate monitoring program continued at the NWT Tundra Ecological Research Station located at Daring Lake. The program was a follow up to the PM₁₀ monitoring program undertaken in summer of 2002 and the PM_{2.5} monitoring program in 2003 to 2006. The monitoring program in 2007 continued the focus on the finer portion of particulate PM_{2.5}. As in the previous five years, a battery operated Mini-Partisol particulate sampler and solar panel were set up to gather information on background concentrations of particulate in the barren lands. The 2007 sampling program began on July 9th and finished on August 21st. The solar-powered system ran without incident in 2007.

A total of 14, 24-hour samples were obtained. Two samples had to be invalidated due to bug fragments on the filters. The overall concentrations were similar to last year, ranging from 0 to 4µg/m³. The 2007 results were typical of background levels and were not influenced by forest fires as in previous years.



APPENDICES

APPENDIX A: MONITORING HISTORY

History of Air Quality Monitoring in the Northwest Territories	
1974	<ul style="list-style-type: none"> Government of the NWT starts monitoring air quality in Yellowknife with the installation of a high-volume air sampler at the Post Office site.
1989	<ul style="list-style-type: none"> Monitoring of acid precipitation at the Snare Rapids hydro-electric site begins.
1992	<ul style="list-style-type: none"> SO₂ analyzer installed at the City Hall site.
1997	<ul style="list-style-type: none"> SO₂ monitoring in N'dilo begins and continues until 2000.
1998	<ul style="list-style-type: none"> O₃ analyzer added in Yellowknife to the City Hall site.
2000	<ul style="list-style-type: none"> A SO₂ analyzer was installed in the ENR building in Fort Liard in March followed by a H₂S analyzer in October.
2002	<ul style="list-style-type: none"> Daring Lake summer sampling of PM₁₀ at Daring Lake begins. City Hall SO₂ analyzer relocated to new air monitoring trailer located at Sir John Franklin High School.
2003	<ul style="list-style-type: none"> Daring Lake summer sampling of PM_{2.5} begins (the same sampler is used for PM₁₀ and PM_{2.5} monitoring) Air monitoring trailers are installed in Inuvik, Norman Wells and Fort Liard. CO and NO_x analyzers added to the Yellowknife station as well as a continuous fine particulate sampler (PM_{2.5}). Norman Wells station monitors SO₂ and H₂S. Inuvik station monitors SO₂, H₂S, NO_x, and PM_{2.5}. Fort Liard station monitors SO₂ and H₂S. A PM_{2.5} sampler is installed late in the year. The O₃ analyzer that was operating at the City Hall location was relocated to the new Sir John Franklin Yellowknife station. ENR initiates the upgrade of the Data Acquisition System moving to a specialized air monitoring system, which will allow more efficient and quality controlled data collection. Continuous PM_{2.5} samplers are installed in Inuvik and Fort Liard. A second high-volume sampler is installed at the Sir John Franklin Yellowknife station.
2004	<ul style="list-style-type: none"> PM_{2.5} sampler is installed in Norman Wells. Data Acquisition System (DAS) is significantly upgraded. New components are installed inside the stations and a new data management, analysis and reporting system is brought on-line.

History of Air Quality Monitoring in the Northwest Territories (cont.)	
2005	<ul style="list-style-type: none"> • NO_x analyzer is installed in March at the Fort Liard station. • O₃ and NO_x analyzers are installed at the Norman Wells station in April. • O₃ analyzer purchased by Environment Canada (Yellowknife office) is installed at the Inuvik station in April. • Due to years of significant data loss caused by extreme cold, the Partisol Dichotomous fine particulate sampler at the Post office station in Yellowknife is relocated indoors at the Sir John Franklin station. • The Yellowknife Post Office station is officially closed after the last TSP sample ran on December 6th, 2005. • Development of an Air Quality web site begins. The web site will link with the data management, analysis and reporting system to provide public access to air quality data for each monitoring location. Access to archived data will also be available by querying the database using web-based tools.
2006	<ul style="list-style-type: none"> • Yellowknife - A BAM Particulate Matter (PM₁₀) monitor was installed and began collecting data in April. • Inuvik - A BAM Particulate Matter (PM₁₀) monitor was installed and began collecting data in October. • The Air Quality Monitoring Network web site was officially released.
2007	<ul style="list-style-type: none"> • Fort Liard - A BAM Particulate Matter (PM₁₀) monitor and an Ozone (O₃) analyzer were installed and began collecting data in late August. • Completed the second phase of the Air Quality Monitoring Network web site, which included database related modifications as well as web design improvements.

APPENDIX B: AIR POLLUTANTS

The NWT air quality monitoring network tracks a number of different air pollutants. With the exception of arsenic and H₂S, these pollutants are known as Criteria Air Contaminants (CAC's). They represent the gases and compounds most often affecting community air quality and targeted by monitoring programs.

Arsenic is monitored in Yellowknife due to its association with metal ore roasting operations in the past and an ongoing concern during remediation work currently being undertaken at the former industrial sites.

H₂S is monitored at the air quality stations in Inuvik, Norman Wells and Fort Liard due to its association with oil and gas development activities.

TOTAL SUSPENDED PARTICULATE (TSP)

Total Suspended Particulate (TSP) is a general term for dust. TSP includes a wide variety of solid and liquid particles found floating in the air, with a size range of approximately 50 micrometers (µm) in diameter and smaller (a human hair is approximately 100 µm in diameter). While TSP can have environmental and aesthetic impacts, it is the smaller particles contained within TSP that are of concern from a human health perspective (see Particulate Matter (PM_{2.5}) and (PM₁₀) later in section). Road dust, forest fires, mining activities and combustion products from vehicles, heating and electricity generation contribute to TSP levels.

The NWT Ambient Air Quality Standard for TSP is 120µg/m³ over a 24-hour period. The standard for the annual average is 60µg/m³ (geometric mean).

ARSENIC

Arsenic is present in the environment in a variety of forms. The most common form in air is the inorganic compound arsenic trioxide. Natural sources of airborne arsenic include volcanoes and windblown dust from arsenic rich soils, while industrial activities such as smelting and burning of coal account for most of the man-made sources. The vast majority of airborne arsenic is associated with dust and therefore analysis of TSP samples provides a good indication of arsenic concentrations.

There are no NWT standards for arsenic compounds. The World Health Organization (WHO) Air Quality Guidelines, (WHO, 2000), state that inhaling inorganic arsenic compounds can contribute to human development of cancer and there is no safe limit. It is, therefore, important to minimize exposure as much as possible, since the cancer risk increases with exposure to higher concentrations.

As the threshold for arsenic compounds has not been set, it is difficult to determine an acceptable level. Ontario continues to use a 24-hour Guideline of 0.3µg/m³ for total arsenic based on general toxicity, but the WHO suggest that a lifetime risk

estimate approach should be used for assessment purposes. The WHO has calculated the following lifetime risk estimates of contracting cancer due to exposure to varying concentrations of arsenic in air:

- 0.066 $\mu\text{g}/\text{m}^3$ results in a theoretical risk of one person in 10,000
- 0.0066 $\mu\text{g}/\text{m}^3$ results in a theoretical risk of one person in 100,000
- 0.00066 $\mu\text{g}/\text{m}^3$ results in a theoretical risk of one person in 1,000,000

In simple terms, the above estimates indicate, for example, that in a population of 10,000 people, a lifetime exposure to an arsenic concentration of 0.066 $\mu\text{g}/\text{m}^3$ could theoretically result in one person contracting cancer. The Ontario Guideline provides a useful comparison for assessment of short-term (24-hour) arsenic measurements but the WHO approach is probably more applicable given the longer-term health risks associated with arsenic exposure.

PARTICULATE MATTER (PM_{2.5}) AND (PM₁₀)

A sub-portion of TSP, these very small particulates are named for the diameter size of the particles contained within each group – PM₁₀ contains particles with a diameter of 10 microns (1 millionth of a metre) or less, while PM_{2.5} (a sub-portion of PM₁₀) contains particles with a diameter of 2.5 microns or less. The significance of these microscopic particles is that they can be inhaled and are associated with health effects including aggravation of existing pulmonary and cardiovascular disease. Generally, the smaller the particle, the greater the penetration into the lung and the greater the associated health risk.

Sources of particulates that can be inhaled include road dust and wind blown soil, which make up the majority of the PM₁₀ particles. Particles in the PM_{2.5} size range primarily result from combustion of fossil fuels for industrial activities, commercial and residential heating, as well as vehicle emissions, forest fire smoke and chemical reactions between other gases emitted to the air.

The national Canada-wide Standards (CWS) process has set an acceptable limit for PM_{2.5} but has not yet established a limit for PM₁₀. The CWS 24-hour average acceptable limit for PM_{2.5} is 30 $\mu\text{g}/\text{m}^3$ and this concentration has been adopted under the *NWT Environmental Protection Act* as the NWT Ambient Air Quality Standard for PM_{2.5}. Several Canadian jurisdictions (e.g. BC, Ontario, Newfoundland and Labrador) have adopted a PM₁₀ concentration of 50 $\mu\text{g}/\text{m}^3$ (24-hour average) as an acceptable limit.

SULPHUR DIOXIDE (SO₂)

SO₂ is a colourless gas, with a pungent odour at elevated concentrations, which can have negative effects on human and environmental health. Certain types of vegetation (especially lichens) are very sensitive to SO₂ impacts. SO₂ also plays a

role in acid deposition and formation of secondary fine particulate through chemical reactions with other pollutants in the air.

There are some natural sources of SO₂ in ambient air (forest fires, volcanoes) but human activity is the major source. Emissions of SO₂ primarily result from the burning of fossil fuels containing sulphur. Sources include natural gas processing plants, gas plant flares and oil refineries, metal ore smelting, power generating plants and commercial or residential heating.

The NWT Ambient Air Quality Standards for SO₂ are 450 µg/m³ (1-hour average), 150 µg/m³ (24-hour average) and 30 µg/m³ (annual average).

HYDROGEN SULPHIDE (H₂S)

Hydrogen sulphide (H₂S) is a colourless gas with a characteristic rotten egg odour. At high concentrations (parts per million range), it can be toxic, but typical ambient (outdoor) concentrations, even in areas impacted by industrial sources, tend to fall in the parts per billion (ppb) range. However, due to its low odour threshold, the presence of H₂S can be offensive and it has been associated with eye irritation and triggering feelings of nausea in sensitive individuals.

Industrial sources include oil and gas extraction, petroleum refining, sewage treatment facilities and pulp and paper mills. Natural sources include sulphur hot springs, swamps and sloughs, which release H₂S as a by-product of organic decomposition.

There are no NWT standards for H₂S. The Alberta Ambient Air Quality Objectives provide an hourly limit of 14µg/m³ (or 10ppb) and a 24-hour limit of 4µg/m³ (or 3ppb), based on avoidance of odour.

NITROGEN OXIDES (NO_x)

Nitrogen oxides (NO_x) consist of a mixture of nitrogen based gases, primarily nitric oxide (NO) and nitrogen dioxide (NO₂). Emissions of both NO and NO₂ results from the high temperature combustion of fossil fuels. The predominant emission is NO, which then rapidly converts to NO₂ through chemical reaction in the atmosphere. NO is a colourless and odourless gas, whereas NO₂ is a reddish-brown colour with a pungent, irritating odour. NO₂ is considered the more toxic and irritating of the two gases and, at elevated concentrations, is associated with both acute and chronic respiratory effects. Both gases play a role in the atmospheric reactions resulting in acid deposition and secondary pollutant formation (i.e. O₃ and fine particulate).

Because of the greater health effects of NO₂, development of air quality standards has focused on this gas, rather than NO or total NO_x. There are no NWT standards for NO₂ but the national standards provide values of 400µg/m³ (1-hour average), 200µg/m³ (24-hour average) and 60µg/m³ (annual average).

GROUND LEVEL OZONE (O₃)

Ground level ozone (O₃) should not be confused with stratospheric O₃, which occurs at much higher elevations and forms a shield that protects life on the planet from the sun's harmful ultraviolet radiation. The gas is the same but at ground level O₃ is regarded as undesirable due to its association with a variety of human health concerns, environmental impacts and property damage. O₃ is a highly reactive gas and is defined as a secondary pollutant. It is not emitted in large quantities from any source but is formed through a series of complex chemical reactions involving other pollutants called precursors (e.g. NO_x and volatile organic compounds or VOCs) in the presence of sunlight.

The national standards provide a Maximum Acceptable Level of 160µg/m³ for O₃ based on a 1-hour average. The Canada-wide Standards (CWS) process has also set an acceptable limit of 65ppb or 127µg/m³ based on an 8-hour average. The CWS 8-hour limit has been adopted under the NWT *Environmental Protection Act* as the NWT Ambient Air Quality Standard for O₃.

CARBON MONOXIDE (CO)

Carbon monoxide (CO) is a colourless, odourless and tasteless gas produced by the incomplete combustion of fuels containing carbon. The primary source is vehicle exhaust especially in cities with heavy traffic congestion. Other sources include industrial processes and fuel combustion for building heating. One natural source is wildfires.

CO affects humans and animals by interfering with the ability of the blood to transport oxygen around the body.

There are no NWT standards for CO but the most stringent national standards provide a value of 15mg/m³ (1-hour average) and 6mg/m³ (8-hour average). CO values are reported in mg/m³ as opposed to other gaseous pollutants, which are reported in µg/m³.

ACID DEPOSITION

Acidity in precipitation is measured in pH units on a scale of 0 to 14. A value of seven indicates neutral, values less than seven indicate acidic conditions and values greater than seven indicate alkaline conditions. Even clean precipitation is slightly acidic—around pH5.6—due to the presence of naturally occurring concentrations of carbon dioxide, and minor amounts of sulphate and nitrate ions. The introduction of sulphur dioxide and nitrogen oxide emissions from combustion of fossil fuels for industrial, commercial and individual activities can result in an increase in acidic compounds in the atmosphere—often in areas far removed from the original emission sources. The removal of these sulphur and nitrogen compounds through

atmospheric washout is reflected in the increased acidity (lower pH values) of precipitation. Calcium and magnesium ions—mostly from natural sources—act to neutralize acidity in precipitation.

Generally, precipitation with a pH value of 5.0 or less is termed 'acidic'. However, assessment of acid precipitation is usually based on deposition to an area over a specified time period (e.g. kilograms per hectare per year, kg/ha/yr) rather than review of specific precipitation event parameters. Also, the degree of impact to a particular environment is influenced by its 'buffering' capacity or ability to tolerate the acidic inputs. Therefore, determination of acceptable limits usually requires a range of values to reflect the differing tolerances of various areas.