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**2017 Results Of Environmental  
Monitoring Programs**

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Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>2 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table of Contents**

	<b>Page</b>
List of Tables and Figures.....	5
Revision Summary.....	9
Executive Summary.....	10
<b>1.0 INTRODUCTION.....</b>	<b>12</b>
1.1 Program Objectives.....	12
1.2 Overview of Pickering and Darlington Nuclear Sites.....	13
1.2.1 Site Description.....	13
1.2.2 Nuclear Generation Performance.....	14
<b>2.0 EFFLUENT MONITORING PROGRAM.....</b>	<b>15</b>
2.1 Radiological Emissions.....	15
2.1.1 Radiological Emissions Graphs.....	16
2.1.2 OPG Nuclear Carbon-14 Inventory Data.....	20
2.2 Conventional Emissions.....	20
<b>3.0 ENVIRONMENTAL MONITORING PROGRAM.....</b>	<b>22</b>
3.1 Design of EMPs.....	22
3.1.1 Environmental Risk Assessments.....	22
3.2 EMP Sampling Plan.....	23
3.2.1 Radiological Contaminants.....	23
3.2.2 Conventional Contaminants.....	24
3.3 Environmental Monitoring Program Results.....	25
3.3.1 Protocol for Reporting Data and Uncertainties.....	26
3.3.2 Atmospheric Sampling.....	27
3.3.2.1 Tritium Oxide.....	27
3.3.2.2 Carbon-14.....	28
3.3.2.3 Noble Gas Detectors.....	29
3.3.3 Terrestrial Sampling.....	31
3.3.3.1 Fruits and Vegetables.....	31
3.3.3.2 Milk and Animal Feed.....	33
3.3.3.3 Eggs and Poultry.....	35
3.3.3.4 Soil Sampling.....	36
3.3.4 Aquatic Sampling.....	37
3.3.4.1 Water Supply Plants.....	37
3.3.4.2 Well Water.....	40
3.3.4.3 Lake Water.....	41
3.3.4.4 Fish.....	42
3.3.4.5 Beach Sand.....	45
3.3.4.6 Sediment.....	46

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>3 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

3.4	Supplementary Studies .....	46
3.4.1	EMP Supplementary Study - Air Kerma Rate from the PWF at the PN Site .....	46
3.4.1.1	Method .....	47
3.4.1.2	Results .....	47
3.4.1.3	Conclusions and Recommendations .....	47
3.4.2	EMP Supplementary Study – Contaminants of Potential Concern in Soil at PN .....	48
3.4.2.1	Method .....	48
3.4.2.2	Results .....	48
3.4.2.3	Conclusions and Recommendations .....	48
3.4.3	EMP Supplementary Study – Sediment and Water Sampling of Non-radiological Contaminants at PN .....	49
3.4.3.1	Method .....	49
3.4.3.2	Results .....	49
3.4.3.3	Conclusions and Recommendations .....	49
3.4.4	EMP Supplementary Study – 2015/16 Entrainment Study for DN .....	50
3.4.4.1	Method .....	50
3.4.4.2	Results .....	50
3.4.4.3	Conclusions and Recommendations .....	50
3.4.5	EMP Supplementary Study – Effluent Characterization Study at DN .....	51
3.4.5.1	Method .....	51
3.4.5.2	Results .....	51
3.4.5.3	Conclusions and Recommendations .....	51
3.4.6	EMP Supplementary Study – 2016 Benthic Study for DN .....	51
3.4.6.1	Method .....	51
3.4.6.2	Results .....	52
3.4.6.3	Conclusions and Recommendations .....	52
3.5	Other Studies .....	52
3.5.1	Potassium in Lake Water .....	52
3.6	Areas of Regulatory Interest and Other Monitoring Programs .....	52
3.6.1	Thermal Monitoring Program .....	53
3.6.2	Impingement and Entrainment Monitoring Program .....	55
3.6.3	Groundwater Monitoring Program .....	55
<b>4.0</b>	<b>ASSESSMENT OF RADIOLOGICAL DOSE TO THE PUBLIC .....</b>	<b>56</b>
4.1	Modelling .....	57
4.1.1	Integrated Model for Probabilistic Assessment of Contaminant Transport (IMPACT) ..	57
4.1.2	Calculated Atmospheric Dispersion Factors .....	57
4.1.3	Meteorological Data .....	58
4.2	Critical Group Dose .....	59
4.2.1	Exposure Pathways .....	60
4.2.2	Age Classes .....	61
4.2.3	Basis of Dose Calculation .....	61
4.2.4	Uncertainty in Dose Calculation .....	61
4.3	Darlington Nuclear Public Dose .....	61
4.3.1	Darlington Nuclear Potential Critical Groups .....	61
4.3.2	Dose Calculation Results .....	62
4.3.3	Discussion of Results .....	63

**Report**

<b>Public Information</b>		
Document Number:	<b>N-REP-03443-10017</b>	Usage Classification:
Sheet Number:	<b>N/A</b>	Revision Number:
		Page:
	<b>R000</b>	<b>4 of 132</b>

**Title:**  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

4.4	Pickering Nuclear Public Dose .....	64
4.4.1	Pickering Nuclear Potential Critical Groups .....	64
4.4.2	Dose Calculation Results .....	64
4.4.3	Discussion of Results .....	66
4.5	Natural and Anthropogenic Data .....	67
<b>5.0</b>	<b>QUALITY ASSURANCE AND PERFORMANCE .....</b>	<b>67</b>
5.1	Laboratory Quality Assurance and Quality Control .....	68
5.1.1	Laboratory Quality Control .....	68
5.1.2	Laboratory Performance Testing .....	68
5.2	Equipment Calibrations/Maintenance .....	69
5.3	Program Quality Assurance .....	69
5.3.1	Audits .....	69
5.3.2	Self-Assessments .....	70
5.4	Third-Party Verification of Annual EMP Report .....	70
5.5	Program Performance .....	70
5.5.1	Sample Unavailability .....	70
5.6	Annual Assessment of the EMPs .....	73
5.6.1	Summary of Darlington Results .....	73
5.6.2	Summary of Pickering Results .....	74
<b>6.0</b>	<b>OUTLOOK FOR 2018 .....</b>	<b>75</b>
<b>7.0</b>	<b>REFERENCES .....</b>	<b>76</b>
	Appendix A: Radiological Units and Conversions .....	81
	Appendix B: Glossary of Acronyms and Symbols .....	82
	Appendix C: Maps of Environmental Monitoring and Critical Group Locations .....	85
	Appendix D: Environmental Monitoring Data .....	89
	Appendix E: Potential Critical Group Descriptions .....	102
	Appendix F: Dose Calculation Procedure and Concentrations .....	106
	Appendix G: Tables of Public Doses by Radionuclide, Pathway and Age Group for Darlington Nuclear and Pickering Nuclear Potential Critical Groups .....	113
	Appendix H: Supplementary Study Data .....	118
	Appendix I: Compliance with Regulatory Document REGDOC-3.1.1 .....	132



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>5 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**List of Tables and Figures**

	<b>Page</b>
Figure 2-1: Darlington Nuclear Airborne Elemental Tritium Emissions .....	16
Figure 2-2: Darlington Nuclear Tritium Oxide Air Emissions .....	17
Figure 2-3: Pickering Nuclear Tritium Oxide Air Emissions .....	17
Figure 2-4: Darlington Nuclear C-14 Air Emissions .....	18
Figure 2-5: Pickering Nuclear C-14 Air Emissions .....	18
Figure 2-6: Darlington Nuclear Tritium Oxide Water Emissions .....	19
Figure 2-7: Pickering Nuclear Tritium Oxide Water Emissions .....	19
Figure 2-8: Darlington Nuclear Gross Beta-Gamma Water Emissions .....	20
Figure 2-9: Pickering Nuclear Gross Beta-Gamma Water Emissions .....	20
Figure 3-1: DN Annual Average HTO in Air .....	28
Figure 3-2: PN Annual Average HTO in Air .....	28
Figure 3-3: DN Annual Average C-14 in Air .....	29
Figure 3-4: PN Annual Average C-14 in Air .....	29
Figure 3-5: PN Annual Average Ar-41 Dose Rate in Air .....	30
Figure 3-6: DN Annual Average HTO in Vegetation .....	32
Figure 3-7: PN Annual Average HTO in Vegetation .....	32
Figure 3-8: DN Annual Average C-14 in Vegetation .....	33
Figure 3-9: PN Annual Average C-14 in Vegetation .....	33
Figure 3-10: DN Annual Average HTO in Milk .....	34
Figure 3-11: PN Annual Average HTO in Milk .....	34
Figure 3-12: DN Annual Average C-14 in Milk .....	35
Figure 3-13: PN Annual Average C-14 in Milk .....	35
Figure 3-14: DN Annual Average HTO in Eggs .....	36
Figure 3-15: DN Annual Average C-14 in Eggs .....	36
Figure 3-16: DN Annual Average HTO in Poultry .....	36
Figure 3-17: DN Annual Average C-14 in Poultry .....	36
Figure 3-18: Bowmanville WSP – Annual Average HTO in Water .....	39
Figure 3-19: Newcastle WSP – Annual Average HTO in Water .....	39
Figure 3-20: Oshawa WSP – Annual Average HTO in Water .....	39
Figure 3-21: Ajax WSP – Annual Average HTO in Water .....	39
Figure 3-22: Scarborough Horgan WSP – Annual Average HTO in Water .....	39
Figure 3-23: Toronto Harris WSP – Annual Average HTO in Water .....	39
Figure 3-24: Whitby WSP – Annual Average HTO in Water .....	40
Figure 3-25: DN Annual Average HTO in Well Water .....	41
Figure 3-26: PN Annual Average HTO in Well Water .....	41
Figure 3-27: DN Annual Average HTO in Lake Water .....	42
Figure 3-28: PN Annual Average HTO in Lake Water .....	42
Figure 3-29: DN Annual Average HTO in Fish .....	43
Figure 3-30: PN Annual Average HTO in Fish .....	43
Figure 3-31: DN Annual Average C-14 in Fish .....	44
Figure 3-32: PN Annual Average C-14 in Fish .....	44
Figure 3-33: DN Annual Average Cs-137 in Fish .....	45
Figure 3-34: PN Annual Average Cs-137 in Fish .....	45
Figure 3-35: Long term trends in Lake Ontario winter temperatures (Dec 1 <sup>st</sup> to March 31 <sup>st</sup> ) .....	54
Figure 3-36: Long term trends in Lake Ontario winter temperatures (Dec 15 <sup>th</sup> to January 15 <sup>th</sup> ) .....	54

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>6 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Figure 4-1: Model of Exposure Pathways from Site Emissions .....	56
Figure 4-2: Darlington Nuclear Annual Public Dose Trend .....	63
Figure 4-3: Comparison of Darlington Nuclear Public Dose to Background Dose.....	64
Figure 4-4: Pickering Nuclear Annual Public Dose Trend.....	66
Figure 4-5: Comparison of Pickering Nuclear Public Dose to Background Dose .....	66
Table 1-1: OPG Public Dose Estimates - 2017.....	11
Table 2-1: DN and PN Annual Site Radiological Emissions 2017 .....	15
Table 2-2: DN and PN Annual Total Site Emissions of Conventional Hazardous Substances – 2016 .....	21
Table 3-1: Routine Environmental Samples Used for the DN and PN EMPs.....	25
Table 3-2: Water Supply Plants Monitored and Distance from Stations .....	37
Table 4-1: Darlington Nuclear Annual Boundary Dispersion Factors – 2017 .....	58
Table 4-2: Pickering Nuclear Annual Boundary Dispersion Factors – 2017.....	58
Table 4-3: Darlington and Pickering Nuclear – 2017 Annual Average Wind Frequency by Direction (at 10 m height) .....	59
Table 4-4: 2017 Annual Darlington Nuclear Critical Group Doses .....	62
Table 4-5: 2017 Darlington Nuclear Public Dose .....	62
Table 4-6: 2017 Annual Pickering Nuclear Critical Group Doses .....	65
Table 4-7: 2017 Pickering Nuclear Public Dose .....	65
Table 4-8: Typical Doses from Exposure to Natural and Anthropogenic Sources .....	67
Table 4-9: Naturally Occurring Annual Public Effective Doses .....	67
Table 5-1: Summary of Analytics Performance Test Results – 2017 .....	69
Table 5-2: Unavailability of EMP Sample Data Used for Dose Calculation Purposes .....	72
Table D-1: Annual Average Concentrations of Tritium-in-Air – 2017 .....	89
Table D-2: Annual Average Concentrations of Carbon-14 in Air – 2017 .....	90
Table D-3: Annual Average Dose Rates of Noble Gases and Ir-192 Skyshine in Air – 2017 ....	91
Table D-4: Fruits and Vegetables – 2017 .....	92
Table D-4: Fruits and Vegetables – 2017 (Continued) .....	93
Table D-4: Fruits and Vegetables – 2017 (Continued) .....	94
Table D-5: Animal Feed – 2017 .....	95
Table D-6: Annual Average Concentrations in Milk – 2017 .....	96
Table D-7: Annual Average Concentrations in Eggs and Poultry – 2017 .....	97
Table D-8: Annual Average Drinking Water and Lake Water Concentrations – 2017 .....	98
Table D-9: Lake Fish – 2017.....	99
Table D-10: Beach Sand – 2017.....	100
Table D-11: Soil – 2017 .....	101
Table F-1: Radionuclides and Pathways Measured and Modeled in the Dose Calculation.....	108
Table G-1: Darlington Nuclear – Farm Doses – 2017.....	113
Table G-2: Darlington Nuclear – Dairy Farm Doses – 2017 .....	114
Table G-3: Darlington Nuclear – Rural Resident Doses – 2017 .....	115
Table G-4: Pickering Nuclear – Dairy Farm Doses – 2017.....	116
Table G-5: Pickering Nuclear – Industrial/Commercial Doses – 2017 .....	116
Table G-6: Pickering Nuclear – Correctional Institute (C2) Doses – 2017.....	117
Table G-7: Pickering Nuclear – Urban Resident Doses – 2017.....	117
Table H-2: Screening of Soil COPCs for Ecological Risk Assessment.....	119
Table H-2: Screening of Soil COPCs for Ecological Risk Assessment (Continued).....	120
Table H-2: Screening of Soil COPCs for Ecological Risk Assessment (Continued).....	121

**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>7 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table I-1: OPG EMP Report Compliance with Regulatory Document-3.1.1, Reporting Requirements for Nuclear Power Plants ..... 132

**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>8 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Acknowledgement**

Ontario Power Generation would like to thank the residents of the local communities in the vicinity of Pickering Nuclear and Darlington Nuclear stations and throughout the province of Ontario, who voluntarily participate in our environmental monitoring programs. Their support in allowing OPG to maintain air monitoring equipment on their properties and in supplying samples of vegetables, fruits, soil, animal feed, milk, eggs, poultry, and water helps to ensure that the annual public dose estimates are realistic.

**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>9 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Revision Summary**

<b>Revision Number</b>	<b>Date</b>	<b>Comments</b>
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**Report**

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>10 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Executive Summary**

Ontario Power Generation (OPG) maintains Environmental Monitoring Programs (EMPs) in the vicinity of Darlington Nuclear (DN) and Pickering Nuclear (PN) stations in accordance with operating licence requirements. The EMPs comply with the Canadian Standards Association (CSA) N288.4-10 standard for Environmental Monitoring Programs at Class I Nuclear Facilities and Uranium Mines and Mills. The program scope encompasses protection of both the public and the environment from nuclear substances, hazardous substances, and physical stressors resulting from the operation of DN and PN sites, including the on-site waste management facilities.

The EMPs are designed to satisfy the following four primary objectives of CSA N288.4-10:

1. Assess the impact on human health and the environment of contaminants and physical stressors of concern resulting from operation of OPG nuclear facilities.
2. Demonstrate compliance with limits on the concentration and/or intensity of contaminants and physical stressors in the environment or assess their effect on the environment.
3. Demonstrate the effectiveness of containment and effluent control, and provide public assurance of the effectiveness of containment and effluent control, independent of effluent monitoring.
4. Verify the predictions made by the Environmental Risk Assessments (ERAs), refine the models used, and reduce the uncertainty in the predictions made by these assessments and models.

Additionally, environmental sampling and analyses for the EMPs support the calculation of annual public dose resulting from operation of OPG nuclear facilities, as required by Canadian Nuclear Safety Commission (CNSC) REGDOC-3.1.1, Reporting Requirements for Nuclear Power Plants.

The 2017 program results contained in this report include concentrations of radionuclides in the air, water, milk, vegetation, animal feed, eggs, poultry, beach sand, soil and fish samples taken in the vicinity of DN and PN, and the associated public radiation dose assessments. Samples from provincial-background locations were used to determine background radiation levels in areas considered to be outside the influence of the nuclear stations.

In addition, a supplementary study was conducted in 2017 on the air kerma (kinetic energy released in matter) rate from the Pickering Waste Management Facility (PWMF) at the PN Site. The results from this study demonstrate that the air kerma rate due to the waste storage facilities cannot be detected at distances greater than 400 m from the facilities and that the skyshine dose from this source is, therefore, not significant for potential critical groups outside the 1 km boundary.

The EMP designs address the monitoring of non-radiological substances through scheduled supplementary studies. Supplementary studies were conducted on contaminants of potential

**Report**

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>11 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

concern in soil, and sediment and water sampling of non-radiological contaminants at PN, as well as entrainment, effluent characterization and the benthic community at DN.

In 2017, OPG operated nine of ten nuclear reactors, producing 40.7 terawatt hours (TWh) of electricity. Unit 2 at DN was under refurbishment in 2017. Site radiological emissions remained at a very small fraction of their licensed Derived Release Limits (DRLs).

A total of 979 laboratory analyses were performed on a variety of environmental media used for the annual public dose calculation. The availabilities of PN and DN samples analyzed for the dose calculation met the annual performance requirements.

IMPACT 5.4.0 software was used for the dose calculations and is consistent with the method of dose calculation described in the CSA N288.1-08 standard, Guidelines for Calculating Derived Release Limits for Radioactive Material in Airborne and Liquid Effluents for Normal Operation of Nuclear Facilities.

The 2017 critical group doses resulting from the operation of the PN and DN sites continue to be a very small fraction of both the annual legal limit of 1,000 microsieverts ( $\mu\text{Sv}$ ) and the estimated annual average background radiation dose around DN and PN of 1,400  $\mu\text{Sv}$ . The 2017 public doses for the DN and PN sites are similar to those observed in 2016 and are summarized in Table 1-1:

**Table 1-1: OPG Public Dose Estimates - 2017**

Site	Critical Group (Receptor)	Effective Dose ( $\mu\text{Sv}$ )	Percentage of Legal Limit (%)	Percentage of Background Radiation around DN and PN (%)
Darlington Nuclear	Dairy Farm (Infant)	0.7	0.1	< 0.1
Pickering Nuclear	Urban Resident (Adult)	1.8	0.2	0.1



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>12 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

## 1.0 INTRODUCTION

Ontario Power Generation (OPG) owns and operates the Pickering Nuclear (PN) and Darlington Nuclear (DN) Generating Stations. To ensure nuclear activities at these sites are conducted in a manner that minimizes any adverse impact on the public and the natural environment, OPG has established an Environmental Management Program that is consistent with the Canadian Nuclear Safety Commission (CNSC) Regulatory Document REGDOC-2.9.1 [R-1] at both stations. Additionally, this program is registered to the International Organization for Standardization (ISO) 14001 Environmental Management Systems standard.

As part of this program, each site has an Environmental Monitoring Program (EMP), which identifies the contaminants and physical stressors to be monitored and conducts monitoring in the environment surrounding the site. The EMP designs use a risk-based approach and rely on the results of site Environmental Risk Assessments (ERAs), as described in Section 3.1.1. Locations considered to be outside the influence of PN and DN site operations are also monitored to allow for a baseline comparison with background values.

The EMPs are maintained in accordance with the operating licences issued to PN and DN and are required to comply with the Canadian Standards Association (CSA) N288.4-10 standard, Environmental Monitoring Programs at Class I Nuclear Facilities and Uranium Mines and Mills [R-2]. This report is prepared and submitted to the CNSC in accordance with their Regulatory Document REGDOC-3.1.1, Reporting Requirements for Nuclear Power Plants [R-3]. It is also made available to the public.

The emissions and environmental data collected for each site during the 2017 sampling year, their interpretations, and the estimates of radiation doses to the public resulting from the operation of PN and DN sites are provided in this report.

Emissions and environmental data are summarized in Sections 2.0 and 3.0, respectively. Assessment of the doses to the public is provided in Section 4.0.

## 1.1 Program Objectives

The PN and DN EMPs are designed to satisfy the following primary objectives:

- (a) To assess the impact on human health and the environment of contaminants and physical stressors of concern resulting from operation of OPG nuclear facilities.
- (b) To demonstrate compliance with limits on the concentration and/or intensity of contaminants and physical stressors in the environment or assess their effect on the environment.
- (c) To demonstrate the effectiveness of containment and effluent control, and provide public assurance of the effectiveness of containment and effluent control, independent of effluent monitoring.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>13 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

- (d) To verify predictions made by ERAs, refine the models used in ERAs, or reduce uncertainty in the predictions made by ERAs.

The EMPs are also designed to facilitate realistic estimates of radiation doses to the public resulting from the operation of PN and DN sites, and to demonstrate that these doses remain below the regulatory limit specified in the current Radiation Protection Regulations under the Nuclear Safety and Control Act [R-4].

## 1.2 Overview of Pickering and Darlington Nuclear Sites

### 1.2.1 Site Description

DN and PN Generating Stations have a combined generating capacity of about 6,600 megawatts (MW). A brief description of the two stations is as follows:

#### Darlington Nuclear

The DN Generating Station is an OPG CANDU (CANadian Deuterium Uranium) nuclear generating station. It is a four-unit station with a total output of 3,500 MW and is located on the shores of Lake Ontario in the Municipality of Clarington in Durham Region. It provides about 20% of Ontario’s electricity needs.



The DN site also contains the Tritium Removal Facility (TRF), where tritium is extracted from tritiated heavy water, and the Darlington Waste Management Facility (DWMF) for used fuel dry storage and processing. The EMP encompasses all the facilities on the DN site.

The immediate area around the Darlington station is mostly rural and farm lands with some industrial/commercial areas. The urban residential locations of Oshawa, Bowmanville and West/East Beach are more than 3 km from the site.

Based on the results of site-specific surveys, the residents around DN are grouped into categories which best represent their locations and/or lifestyle characteristics. The categories are known as potential critical groups and are further described in Appendix

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>14 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

E, Section E.1.0. The DN EMP design focuses primarily on the farm, dairy farm, and rural resident potential critical groups, as described in Section 4.0.

### Pickering Nuclear

The PN site is located on the shores of Lake Ontario, in the city of Pickering. The site contains the PN Generating Stations and the Pickering Waste Management Facility (PWMF) which consists of sites located inside and outside of the station protected area. The EMP encompasses all the facilities on the PN site.



PN has six operating CANDU reactors. This station has a total output of 3,100 MW. PN Units 2 and 3 are in a safe storage state.

Unlike DN, the area around PN is mainly urban residential and industrial/commercial. The closest farm lands are more than 6 km from the station.

Based on the results of site-specific surveys, the residents around PN are grouped into categories which best represent their locations and/or lifestyle characteristics. The categories are known as potential critical groups and are further described in Appendix E, Section E.2.0. The PN EMP design focuses primarily on the urban resident, dairy farm, industrial/commercial worker, and correctional institute occupant potential critical groups, as described in Section 4.0.

#### 1.2.2 Nuclear Generation Performance

In 2017, OPG operated nine of ten nuclear reactors that produced 40.7 terawatt hours (TWh) of electricity. This production is broken down as follows:

**Darlington Nuclear:** Net electrical output in 2017 was 19.3 TWh.

**Pickering Nuclear:** Net electrical output in 2017 was 21.4 TWh.

**Report**

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>15 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**2.0 EFFLUENT MONITORING PROGRAM**

**2.1 Radiological Emissions**

The radiological emissions from DN and PN sites in 2017 remain at a very small fraction of the site Derived Release Limits (DRLs). These licensing limits represent radionuclide release rates that correspond to an exposure at the legal public dose limit of 1,000 microsieverts per year ( $\mu\text{Sv}/\text{year}$ ) for the most affected critical group. See Section 4.0 for the description of a critical group.

Table 2-1 shows the 2017 total airborne and waterborne emissions for radionuclides measured at the DN and PN sites, including the waste management facilities, and the percentage of their respective DRLs.

**Table 2-1: DN and PN Annual Site Radiological Emissions 2017**

Site Emissions <sup>(d)</sup>	DN		PNA (Units 1-4)		PNB (Units 5-8)	
	Bq	% DRL	Bq	% DRL	Bq	% DRL
<b>AIR</b>						
Tritium Oxide	2.4E+14	0.41	3.1E+14	0.26	3.8E+14	0.20
Elemental Tritium <sup>(a)</sup>	1.4E+14	1.6E-02	NA	NA	NA	NA
Noble Gas <sup>(b)</sup>	1.5E+13	0.03	1.5E+14	0.47	3.5E+12	<0.01
I-131 <sup>(c)</sup>	1.5E+08	<0.01	9.6E+06	<0.01	4.3E+06	<0.01
Particulate	2.6E+07	<0.01	6.9E+06	<0.01	2.0E+08	2.8E-02
C-14	1.4E+12	0.40	1.3E+12	0.06	1.3E+12	0.07
<b>WATER</b>						
Tritium Oxide	5.6E+14	1.1E-02	1.1E+14	0.03	2.7E+14	0.04
Gross Beta/Gamma	2.6E+10	3.7E-02	6.5E+09	0.38	2.0E+10	0.63
C-14 <sup>(e)</sup>	1.7E+09	<0.01	NA	NA	1.9E+09	<0.01

NOTES: NA = Not Applicable, Bq = Bequerels

(a) Emissions from Darlington Tritium Removal Facility

(b) Units for noble gas emissions are Bq-MeV

(c) Weekly samples are usually < Method Detection Limit (MDL)

(d) Annual air emissions are the sum of continuous samples analysed weekly.

Note that if interim Noble Gas sampling is in place, samples may not be continuous.

Annual water emissions are the sum of monthly composite samples for C-14, and weekly composite samples for tritium oxide and gross beta/gamma.

(e) While reported under PNB emissions in this table, the 2017 C-14 waterborne emission value is the total for all Pickering units.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>16 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

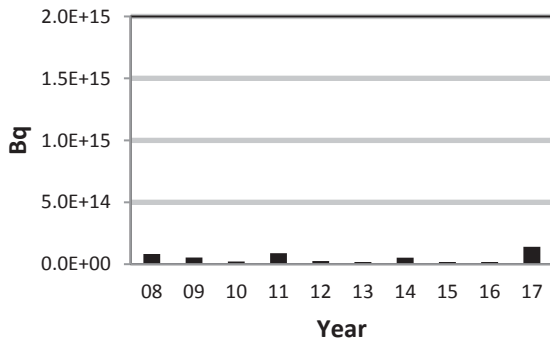
### 2.1.1 Radiological Emissions Graphs

Graphs displaying the past ten years of tritium and C-14 emissions to air and tritium emissions to water from DN and PN are provided in Figures 2-1 to 2-7. DN and PN gross beta-gamma emissions to water are provided in Figures 2-8 and 2-9. Given that the reported noble gas stack emissions are often below the instrument detection limits, the results of environmental noble gas monitors are used to trend the station noble gas emissions as described in Section 3.3.2.3. Iodine and particulate in airborne emissions and C-14 waterborne emissions are not graphed because their contribution to the overall public dose is minimal.

#### Elemental Tritium Airborne Emissions

##### DN – Figure 2-1

As indicated in Figure 2-1, the elemental tritium (HT) emissions from DN have remained at low levels. In 2017, the HT emissions were  $1.4 \times 10^{14}$  becquerels (Bq). As of 2017, these emissions include HT emissions from the powerhouse. However, the increase in elemental tritium emissions observed in 2017 is primarily attributed to a valve that was inadvertently opened and vented to the TRF stack on May 8, 2017. Corrective actions were immediately taken to redirect residual elemental tritium to the Air Clean up System and a procedure update was subsequently initiated to rectify the deficiencies.



**Figure 2-1: Darlington Nuclear Airborne Elemental Tritium Emissions**

##### PN

PN does not experience substantial HT emissions as it does not have a TRF.

#### Tritium Oxide Airborne Emissions

##### DN – Figure 2-2

In 2014, a small increase was observed in DN tritium oxide (HTO) airborne emissions, which was attributed to both dryer performance and TRF restart activities. During 2015



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>17 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

and 2016, work plans were executed to refurbish dryers throughout the station resulting in a decrease in emissions. Airborne HTO emissions in 2017 were  $2.4 \times 10^{14}$  Bq.

PN – Figure 2-3

PN HTO airborne emissions decreased from 2008 to 2010 and again in 2013 as a result of improvements in emissions management, reliability and operation of vapour recovery dryers, and reduction of HTO source terms. The increase in 2016 was primarily attributed to the presence of tritiated water in Fuel Transfer Conveyor Tunnel, and the resulting airborne HTO emissions being vented to a monitored stack. Mitigating actions were taken to reduce HTO airborne emissions from this source. Airborne HTO emissions in 2017 were  $6.9 \times 10^{14}$  Bq. The slight increase observed in 2017 was primarily attributed to dryer performance issues and a rupture disk failure on Unit 1, which has since been corrected.

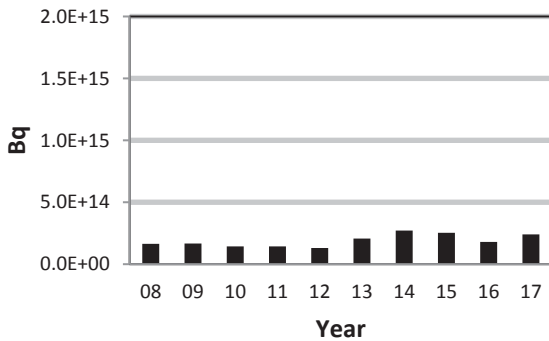


Figure 2-2: Darlington Nuclear Tritium Oxide Air Emissions

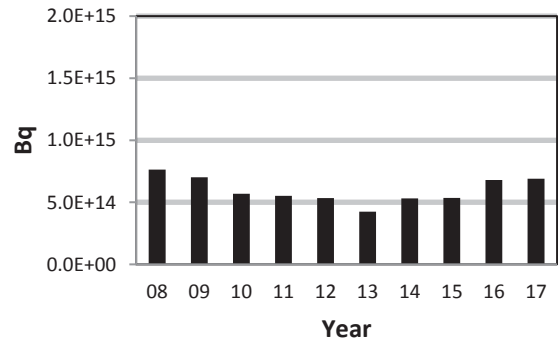


Figure 2-3: Pickering Nuclear Tritium Oxide Air Emissions

**Carbon-14 Airborne Emissions**

DN – Figure 2-4

DN C-14 airborne emissions remain stable. The 2017 C-14 airborne emissions were  $1.4 \times 10^{12}$  Bq.

PN – Figure 2-5

Considerably lower PN C-14 airborne emissions have been observed since 2009. The 2017 C-14 airborne emissions were  $2.6 \times 10^{12}$  Bq, which are similar to emissions in 2016.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>18 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

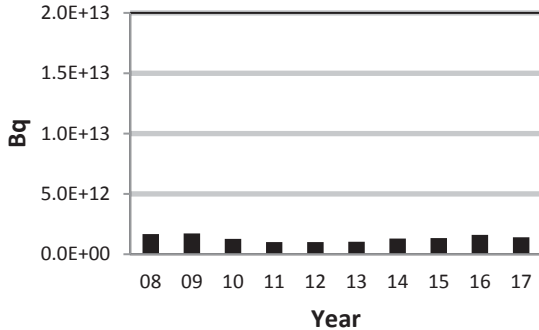


Figure 2-4: Darlington Nuclear C-14 Air Emissions

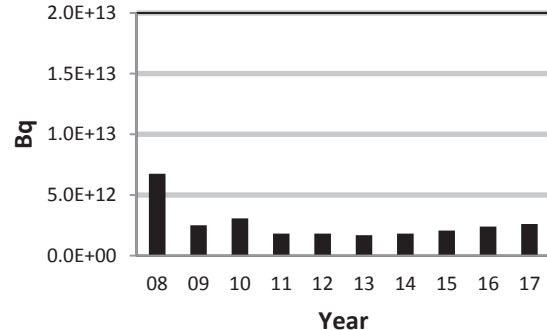


Figure 2-5: Pickering Nuclear C-14 Air Emissions

**Tritium Oxide Waterborne Emissions**

DN – Figure 2-6

The 2009 DN vacuum building outage (VBO) required system drainage in 2007 and 2008, which resulted in slightly elevated DN HTO to water emissions during these years. Similarly, drainage and discharge activities associated with the 2015 VBO took place in 2014 and 2015. The 2017 DN tritium to water emission was  $5.6 \times 10^{14}$  Bq. The increase in emissions observed in 2016 and 2017 are primarily attributed to the processing and discharge of condensate from reactor building air conditioning units (ACUs) through the active liquid waste system. A program to replace ACU coils is currently underway and will continue through 2018.

PN – Figure 2-7

The PN waterborne HTO emissions remain stable. The slightly elevated emissions in 2008 and 2009 were due to a minor heavy water leak from a Unit 1 shutdown cooling heat exchanger and a small Unit 1 boiler tube leak, respectively. The PN tritium to water emission in 2017 was  $3.8 \times 10^{14}$  Bq. The slight increase observed in 2017 is attributed to a leak in the Unit 5 moderator pit. Tritiated water from the moderator room was processed and discharged through the active liquid waste system. Sealing and repair work to the moderator pit was completed in April 2017.



Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>19 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

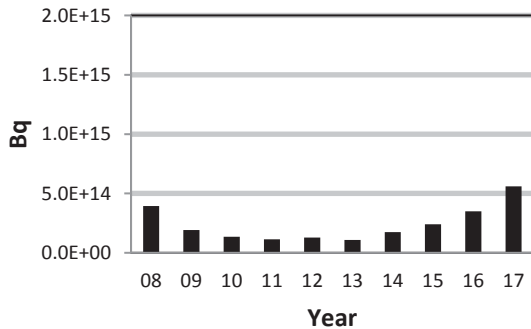


Figure 2-6: Darlington Nuclear Tritium Oxide Water Emissions

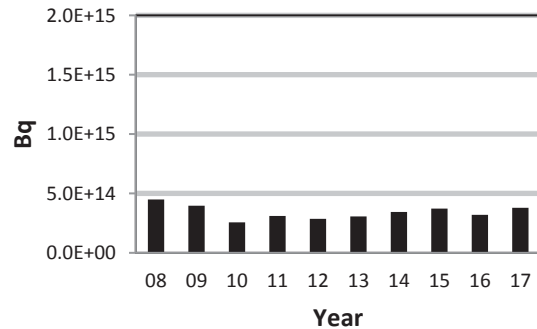


Figure 2-7: Pickering Nuclear Tritium Oxide Water Emissions

**Gross Beta-Gamma Waterborne Emissions**

DN – Figure 2-8

The DN gross beta-gamma emissions to water remain low. The slightly elevated emission values in 2015 and 2016 do not reflect a true increase in emissions, but rather the use of an alternate counter with a higher detection limit than the main counter. The 2017 gross beta-gamma water emission was  $2.6 \times 10^{10}$  Bq.

PN – Figure 2-9

The PN gross beta-gamma emissions to water remain low. The increase in 2009 and 2010 was due to anomalously high activity of several samples. Mitigating actions from OPG’s investigation and third-party review of this matter have been implemented. Since 2011, the emissions have generally returned to pre-2009 levels, as shown in Figure 2-9. The increase in gross beta-gamma waterborne emissions seen in 2016 were primarily attributed to spontaneous release of concentrated, entrained active lake sediment materials from the Reactor Building Service Water system, and not a station generated source of activity. In 2017, gross beta-gamma water emission returned to low levels of  $2.7 \times 10^{10}$  Bq.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>20 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

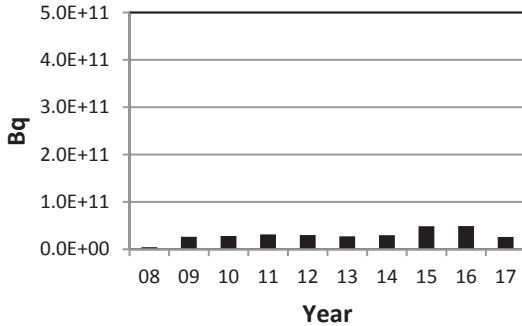


Figure 2-8: Darlington Nuclear Gross Beta-Gamma Water Emissions

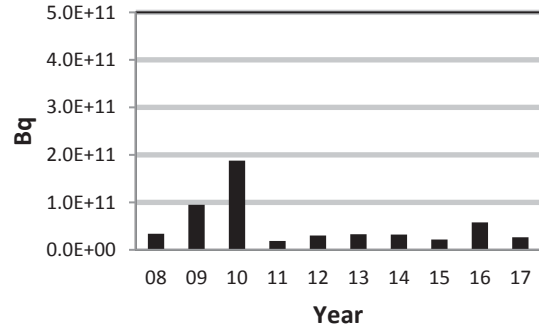


Figure 2-9: Pickering Nuclear Gross Beta-Gamma Water Emissions

### 2.1.2 OPG Nuclear Carbon-14 Inventory Data

The C-14 inventories within the DN and PN stations are included in this report to fulfill a regulatory commitment to the CNSC [R-5]. The 2017 estimates of C-14 inventory within the DN and PN stations are  $7.0 \times 10^{14}$  Bq and  $8.7 \times 10^{14}$  Bq, respectively [R-6].

## 2.2 Conventional Emissions

OPG monitors conventional substances emitted to air and water as a result of DN and PN site operations. Reports on emissions of both conventional hazardous and non-hazardous substances are prepared in accordance with regulatory requirements and submitted to provincial and federal agencies throughout the year. As the submission of 2017 reports continues through 2018, the complete set of conventional hazardous substances released from DN and PN sites in 2016 is provided in Table 2-2. 2017 emissions will be summarized in the 2018 EMP report.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>21 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table 2-2: DN and PN Annual Total Site Emissions of Conventional Hazardous Substances – 2016**

Hazardous Material	DN	PN
	Mg	Mg
<b>AIR</b>		
SO <sub>2</sub> to Air <sup>(a)(b)</sup>	1.0E+00	2.1E+00
NO <sub>2</sub> to Air <sup>(b)</sup>	2.2E+01	4.5E+01
CO <sub>2</sub> to Air <sup>(a)(b)</sup>	8.4E-02	4.0E-01
Ammonia to Air	4.6E+00	4.5E+00
Hydrazine to Air <sup>(c)</sup>	2.1E-02	5.1E-03
Ozone Depleting Substances (ODS) Releases <sup>(d)</sup>	2.5E-02	4.0E-02
<b>WATER</b>		
Ammonia to Water	2.0E+00	5.9E-01
Hydrazine to Water <sup>(c)</sup>	4.1E-01	2.4E-01

NOTES:

Mg = Megagrams

- (a) Reported in OPG Sustainable Development Report as an OPGN aggregate value.
- (b) Based on annual fuel consumption.
- (c) Based on annual consumption.
- (d) Based on estimated quantity when a release occurs.

**Sulphur Dioxide, Nitrogen Oxides and Carbon Dioxide Emissions**

DN and PN have standby diesel generators to provide back-up electrical power to the station if required, which account for sulphur dioxide, nitrogen oxides and carbon dioxide emissions. These generators are routinely tested to ensure availability. There were no regulatory non-compliances associated with the air emissions from these generators in 2016 from DN or PN.

**Hydrazine and Ammonia**

Hydrazine and ammonia are used in station water systems to prevent corrosion. These chemicals are released when steam is vented to the atmosphere and when water is drained to Lake Ontario. There were no regulatory non-compliances associated with hydrazine and ammonia emissions in 2016 for DN or PN.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>22 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****Ozone Depleting Substances**

Ozone-depleting substances (ODS) are used in refrigeration systems. Refrigerant leaks to air are minimized through routine inspections and maintenance of equipment. There were no releases of ODS that were reportable as spills in 2016 for DN or PN. ODS releases between 10 kg and 100 kg are reported in semi-annual halocarbon release reports.

**3.0 ENVIRONMENTAL MONITORING PROGRAM****3.1 Design of EMPs**

The EMP designs were developed using the guidance in CSA N288.4-10 to address site specific objectives covering the aspects of regulatory requirements, ERA results, confirmation of effluent control, areas of regulatory interest, and stakeholder commitments.

**3.1.1 Environmental Risk Assessments**

The PN and DN site ERAs assess potential human health and ecological risks from exposure to radiological contaminants, conventional contaminants, and physical stressors present in the environment as a result of site operations. The ERAs help to identify which monitoring to include in the EMPs.

The most recent DN ERA was completed at the end of 2016 in accordance with the requirements of CSA N288.6-12, Environmental Risk Assessments at Class I Nuclear Facilities and Uranium Mines and Mills [R-9]. Changes to the EMP as a result of the latest ERA will be identified and captured in the next EMP Design Review, which will be undertaken in 2018.

The PN ERA was updated in 2017 in accordance with the requirements of CSA N288.6-12 [R-9]. Changes to the EMP as a result of the latest ERA will be identified and captured in the next EMP Design Review in 2018.

The 2013 PN ERA identified a recommendation to sample water at Hydro Marsh to confirm the assumption that the effects from airborne tritium deposition in the marsh are minor [R-10]. The result of a supplementary study conducted in 2016 confirmed that there is only a minor difference in dispersion factors between Hydro Marsh and Frenchman's Bay. Therefore, it is not necessary to consider Hydro Marsh as a separate potential assessment location in future ERAs [R-60].

Beyond obtaining data to clarify potential risks identified by the ERAs, the EMPs also fulfill the CSA N288.4-10 and regulatory requirements of estimating public dose from radiological emissions, confirming effluent control, clarifying risks and refining ERA models and predictions.

## Public Information

Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>23 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****3.2 EMP Sampling Plan**

The EMP sampling plan outlines the contaminants monitored, the sampling locations, the sample types, and the frequency of collection. Samples collected, analyses performed, and data interpreted aim to support the EMP objectives as follows:

## 1) Public Dose Calculation

To ensure that the public dose estimation from radiological emissions is as realistic as possible, various exposure pathways, such as food ingestion, inhalation, and water ingestion are assessed for radionuclide concentrations resulting from site operations. Samples are collected at site boundary locations as well as at potential critical group locations. A description of critical groups is provided in Section 4.0, Assessment of Radiological Dose to the Public. For sample types that are not available at potential critical group locations, contaminant concentrations are estimated from concentrations measured at the boundary locations using ratios of modeled atmospheric dispersion factors.

## 2) Demonstration of Emissions Control

To meet this objective, environmental measurements at the site boundary are used to confirm that concentrations are as expected based on effluent monitoring. Similarly, lake water/drinking water monitoring demonstrates waterborne emissions are properly controlled. Environmental monitoring provides an independent ongoing check on the effectiveness of containment and effluent control.

## 3) Refining ERA Models and Predictions

Sampling to verify ERA predictions and to assist in refining models used in the ERAs is included in the EMP designs and handled through supplementary studies, which are documented in the annual EMP report. Refer to Section 3.4. for a discussion of the supplementary studies that took place in 2017.

**3.2.1 Radiological Contaminants**

Radionuclides that are emitted as a result of PN and DN site operations and monitored in the EMPs are listed below. They are identified through the pathway analyses as discussed in Section 4.2 of this document. The routine sample analyses used in the public dose calculation are indicated in Table 3-1.

Carbon-14 (C-14) – is produced from the operation of nuclear stations. It is also a naturally occurring radionuclide and a by-product of past nuclear weapons testing with average background concentrations between 220 becquerels per kilogram carbon (Bq/kg-C) and 250 Bq/kg-C for air. C-14 values detected above background are included in the dose calculations.

Tritiated Water (HTO) – is a normal station emission of CANDU plants. Concentrations measured in plants and animals refer to the HTO concentration in the free water portion of the sample.

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>24 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

Tritiated Hydrogen Gas (HT) – is emitted to air primarily as a result of the operation of the TRF at DN. HT concentration in air is modeled from emissions and not monitored in the environment. However, much of the HT is converted to HTO in the environment, and this HTO is monitored.

Organically Bound Tritium (OBT) – is tritium that is bound to the organic molecules in organisms and is not readily exchanged with other hydrogen atoms. In accordance with CSA N288.1-08, OBT concentrations used in the dose calculation are modeled from HTO concentrations measured in sample media at each potential critical group location and in fish. OPG dose calculations incorporate dose from OBT via intake of terrestrial plants and animal products, and from fish. OBT is measured in a few environmental samples for informational purposes and these results are presented in Appendix D.

Noble Gases – Radioactive noble gases released from the DN and PN plants are mostly Argon-41 (Ar-41), Xenon-133 (Xe-133) and Xenon-135 (Xe-135). The environmental detectors that measure noble gas doses may also detect Iridium-192 (Ir-192) skyshine from industrial radiography carried out in the stations.

Iodine-131 – The dose from radioiodine emissions is modeled from I-131 emissions, with the assumption that I-131 emissions are accompanied by an equilibrium mixture of other short lived iodine fission products (i.e., I-132, I-133, I-134 and I-135) or mixed fission products [I(mfp)].

Particulates and gross beta-gamma – Atmospheric particulate emissions are represented by Cobalt-60 (Co-60) and liquid effluent beta-gamma emissions are represented by Cesium-137 (Cs-137) as this provides the most conservative assignment of dose based on the pathway analyses in the program design reviews [R-53][R-54]. Cs-137 is also present in the environment as a result of historic weapons testing. Co-60 and Cesium-134 (Cs-134) are representative of station emissions and are analyzed together with Cs-137, which helps distinguish between the Cs-137 resulting from station operations with that from past weapons testing.

### 3.2.2 Conventional Contaminants

Conventional contaminants emitted as a result of PN and DN operations may be monitored in the environment as part of the EMPs for ERA confirmation and/or demonstration that concentrations fall below benchmark values. The monitoring of these contaminants is achieved through supplementary studies.

Supplementary studies were conducted on contaminants of potential concern in soil, and sediment and water sampling of non-radiological contaminants at PN. An effluent characterization was undertaken at DN. Results of these supplementary studies are summarized in Section 3.4.

# Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>25 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table 3-1: Routine Environmental Samples Used for the DN and PN EMPs**

Environmental Medium of Interest	Monitored For	Sampling Frequency	Analyses Frequency
<b>SAMPLES USED FOR PUBLIC DOSE CALCULATIONS</b>			
<b>Atmospheric Sampling</b>			
Air	HTO (active monitor)	Continuous	Monthly
Air	C-14 (passive monitor)	Continuous	Quarterly
Air	Noble gases (Ar-41, Xe-133, Xe-135), Ir-192 <sup>(a)</sup>	Continuous	Reported monthly
<b>Terrestrial Sampling</b>			
Fruits and Vegetables <sup>(c)</sup>	HTO and C-14	3 grab samples/year	3 times/year
Animal Feed	HTO and C-14	Bi-annual grab samples	Bi-annual
Eggs	HTO and C-14	Quarterly grab samples	Quarterly
Poultry	HTO and C-14	Annual grab samples	Annual
Milk <sup>(b)</sup>	HTO and C-14	Monthly grab samples	Monthly
<b>Aquatic Sampling</b>			
Municipal Drinking Water	HTO	2-3 grab samples/day	Weekly composite
Well Water	HTO	Monthly grab samples	Monthly
Lake Water	HTO	Monthly grab samples	Monthly
Fish	HTO, C-14, Cs-137, Cs-134, Co-60	Annual grab samples	Annual
Beach Sand	Cs-137, Cs-134, Co-60	Annual grab samples	Annual
<b>SAMPLES FOR OTHER EMP OBJECTIVES</b>			
Vegetables	OBT	Annual grab samples	Annual
Soil	Cs-137, Cs-134, Co-60	Grab samples every five years	Every five years
Milk	OBT	Monthly grab samples	Monthly
Municipal Drinking Water	Gross beta	2-3 grab samples/day	Monthly composite
Fish	OBT	Annual grab samples (composite)	Annual
Sediment	C-14, Cs-137, Cs-134, Co-60	Grab samples every five years (composite)	Every five years
Lake water	Potassium	Grab samples every three years (composite)	Every three years

(a) Air kerma is measured and converted to external air immersion dose.

(b) Sampling frequency is quarterly for provincial-background locations.

(c) Sampling frequency is annual for provincial-background locations.

### 3.3 Environmental Monitoring Program Results

This section contains the results of the EMPs for the DN and PN sites and those of the provincial-background locations. All sampling locations are shown in Appendix C, Figures C1 to C3, and are selected based on the pathway analyses and site specific survey reviews as described in Section 4.2 of this report.



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>26 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

### 3.3.1 Protocol for Reporting Data and Uncertainties

Statistical analyses typically performed on datasets include determination of the mean and standard deviation, trend analysis, demonstration that the concentrations of contaminants are below the benchmark value, and dataset comparison.

Trend analysis is performed on most EMP data, however, it is more meaningful when sampling locations and frequencies remain consistent throughout the trending period. As the air monitors around the site boundary are sensitive to changes in location, only locations that were active for the entire trending period are used in the trend analysis of boundary air data. For other sample media, all locations that are currently active are included in the trend analysis. Fruits and vegetables are the exception in that all sample locations, both current and historical, are included in the trend analysis since these sample locations change frequently. Therefore, for the trend analysis of EMP environmental sample media other than air, there is a degree of inaccuracy when comparing year to year averages since the same set of locations may not have been used for the entire trending period.

Radionuclide concentrations in the environment are low and at times below levels which can be detected by routine analytical techniques. In these situations the analytical result is reported as being below the detection limit (Ld) or critical level (Lc).

**Lc:** The critical level is the level (relative to background) below which a quantity cannot reliably be measured. More specifically, the critical level is the largest value of the quantity for which the probability of a wrong conclusion that a quantity is present exceeds a specified probability [R-2]. The EMPs use a probability of 5%. For the EMPs, Lc is approximately equal to half of the Ld.

**Ld:** The detection limit is the level (relative to background) above which a quantity can confidently be measured. More specifically, the detection limit is the smallest value of the quantity for which the probability of a wrong conclusion that the quantity is not present does not exceed a specified probability [R-2]. The EMPs use a probability of 5%.

When reporting the analytical data in Appendix D tables, the following conventions are used:

- Where a measured value is below the analytical Ld but above the Lc, the measured value is reported in bold type.
- Where a measured value is below the Lc, then “< Lc” is reported without an uncertainty measure.
- Where a measured value is censored at the Ld, it is reported as “< Ld”. This is the case for gamma spectrometer results, noble gas data, and conventional contaminants.
- For a dataset comprised of a single measured value, the associated uncertainty is the laboratory analytical uncertainty for that particular sample.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>27 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

- For a dataset without any data censored at the Ld, the arithmetic mean is reported and associated uncertainty is two times the standard deviation of the dataset.
- For a dataset containing some data censored at the Ld, the Kaplan-Meier (KM) estimation method is used. The KM mean is reported and associated uncertainty is two times the KM standard deviation of the dataset. An asterisk “\*” is used to identify these datasets.
- For a dataset that consists entirely of data censored at the Ld, the average is reported as “<Ld” without an uncertainty measure.
- For a dataset that consists entirely of data below the Lc (with no censored data), the average is reported as “< Lc” without an uncertainty measure.

See Appendix F.2.0 for treatment of background data for dose calculation purposes.

### 3.3.2 Atmospheric Sampling

Samples of air are collected to monitor the environment around the DN and PN sites. Background samples are also collected to allow determination of net values for dose calculations. The radionuclide analyses performed and the sample collection frequency are detailed in Table 3-1 and results are summarized in Sections 3.3.2.1 to 3.3.2.3. Detailed data are given in Appendix D, Environmental Monitoring Data, Tables D1 to D3.

#### 3.3.2.1 Tritium Oxide

The active tritium-in-air sampler collects water vapour by passing air continuously at a steady rate through two molecular sieve canisters in series. The active samplers are located at six site boundary EMP monitoring locations around DN (D1, D2, D5, D9, D10 and D11) and six around PN (P2, P3, P4, P6, P10, and P11), as identified in Figures C1 and C2 in Appendix C. These samples are collected and analyzed monthly.

The background concentration of HTO in air is measured at Nanticoke, which is considered to be far from the influence of nuclear stations. The annual average HTO in air measured at the background location is consistently at or below the active sampler detection limit of 0.2 Bq/m<sup>3</sup>.

The 2017 annual average results of airborne HTO at the DN, PN, and background monitoring locations are summarized in Appendix D, Table D1. The levels of HTO observed in the environment depend on station emissions, wind direction, wind speed, ambient humidity, and seasonal variations. As such, fluctuations from year to year are expected even if site HTO emissions remain constant.

For the purpose of statistical trend analyses, Figures 3-1 and 3-2 utilize only locations which were active for all of the last 10 years in order to provide a representative year

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>28 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

to year comparison. For DN this includes locations D1, D2, and D5. For PN this includes locations P2, P3, P4, P6, P10 and P11.

DN – Figure 3-1

The 2017 HTO in air annual average concentrations measured at DN boundary locations ranged from 0.2 to 1.4 Bq/m<sup>3</sup>, with an average concentration of 0.7 Bq/m<sup>3</sup>. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend over the past 10 years.

PN – Figure 3-2

The 2017 HTO in air annual average concentrations measured at PN boundary locations ranged from 1.2 to 12.6 Bq/m<sup>3</sup>, with an average concentration of 5.8 Bq/m<sup>3</sup>. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend over the past 10 years.

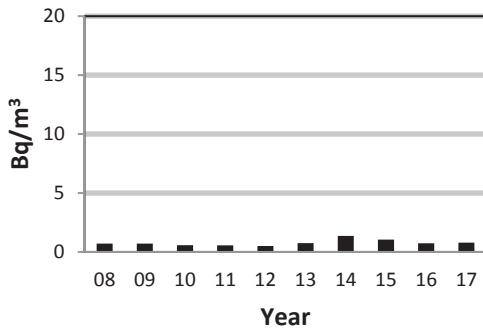


Figure 3-1: DN Annual Average HTO in Air

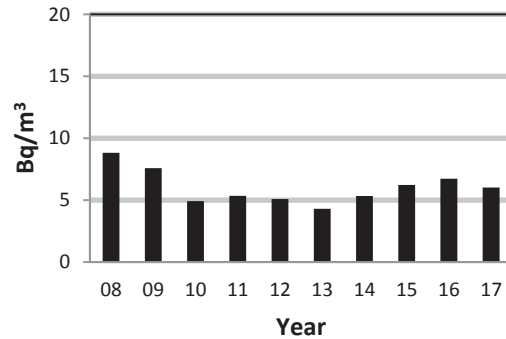


Figure 3-2: PN Annual Average HTO in Air

**3.3.2.2 Carbon-14**

C-14 in air is sampled using passive sampling technology. The passive C-14 sampler works by absorption of CO<sub>2</sub> in air into soda lime pellets exposed for a period of an annual quarter. Samples are analyzed after each quarter.

C-14 is naturally occurring in the environment but is also a by-product of past nuclear weapons testing from the early 1960s. C-14 background concentrations around the world are decreasing as weapons test C-14 levels naturally decay over time. Pre-atmospheric weapons test levels were measured at 226 Bq/kg-C [R-11]. The annual average C-14 in air concentration observed at the Nanticoke EMP background location in 2017 was 238 Bq/kg-C.

In the EMP designs, C-14 in air is monitored at four boundary locations for DN (D1, D2, D5, and D10) and four boundary locations for PN (P3, P4, P6, and P10). Appendix D, Table D2, provides the 2017 annual averages of airborne C-14 measured at the DN, PN, and background sampling locations.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>29 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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For the purpose of statistical trend analyses, Figures 3-3 and 3-4 utilize only locations which were active for all of the last 10 years in order to provide a representative year to year comparison. For DN this includes locations D1, D2, and D10. For PN this includes locations P6 and P10.

DN – Figure 3-3

The 2017 annual average C-14 in air concentrations measured at DN boundary locations ranged from 234 to 274 Bq/kg-C, with an average concentration of 259 Bq/kg-C. A Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend over the past 10 years.

PN – Figure 3-4

The 2017 annual average C-14 in air concentrations measured at PN boundary locations ranged from 289 to 425 Bq/kg-C, with an average concentration of 339 Bq/kg-C. A Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend over the past 10 years. The higher level observed in 2008 is in line with the emissions patterns, as discussed in Section 2.1.1.

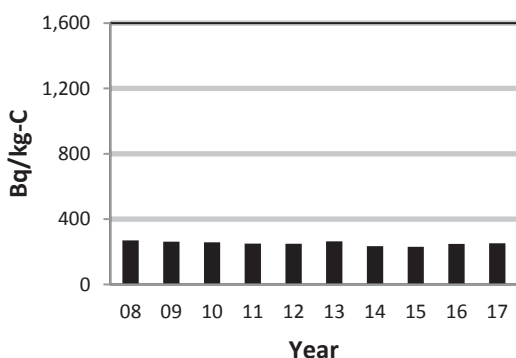


Figure 3-3: DN Annual Average C-14 in Air

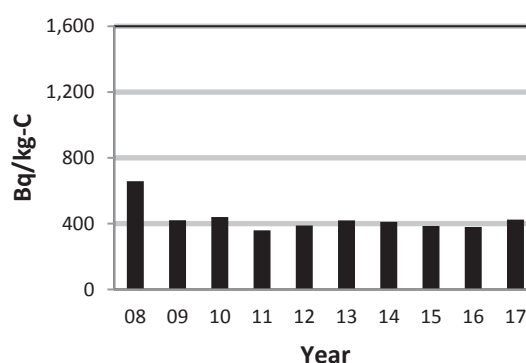


Figure 3-4: PN Annual Average C-14 in Air

**3.3.2.3 Noble Gas Detectors**

Under a Memorandum of Understanding (MOU) between OPG and Health Canada (HC), established in 2009, HC operates and maintains OPG’s network of noble gas detectors. In exchange, OPG allows HC to release the detector results on their public website as part of their Fixed Point Surveillance (FPS) network [R-12].

In years past, OPG and HC would each calculate noble gas dose from raw data using different analysis and processing software, yielding comparable results. Starting in 2014, OPG began using the noble gas dose results generated by HC for calculation of the annual public dose. Noble gas data generated by HC is reviewed by OPG on a quarterly basis.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>30 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

External gamma radiation doses from noble gases and Ir-192 are measured using sodium iodide (NaI) spectrometers set up around the DN and PN sites. There are a total of eight detectors around the DN site and eight detectors around the PN site that monitor the dose rate continuously. Natural background dose has been subtracted from noble gas detector results.

The annual boundary average noble gas dose rate is estimated from the monthly data from each detector. Results obtained in 2017 from the noble gas detectors are summarized in Appendix D, Table D3 and discussed below.

DN

Due to a different station design, DN does not experience the same level of noble gas emissions as PN. The DN boundary average dose rates for Ar-41, Xe-133, Xe-135, and Ir-192 are typically below the detection limits. Therefore, no trend graph is presented for DN.

PN – Figure 3-5

Ar-41 is the predominant radionuclide measured in noble gas around PN followed by Xe-133 and Xe-135. The PN boundary average Ar-41 dose in air was 285 nanogray (nGy)/month in 2017.

Figure 3-5 illustrates the boundary average Ar-41 dose rate for PN from 2008 to 2017, which represents the period of time when all six PN units were operational, in units of nGy/month. A Mann-Kendall trend analysis at the 95% confidence level indicates an increasing trend over the past 10 years for Ar-41. Ar-41 emissions are largely related to the number of operating days of PN Units 1 and 4, therefore higher Ar-41 in the environment is typically attributed to a higher number of operating days from these two units.

Xe-133 and Xe-135 were also, at times, measured above the detection limit at PN. Measured boundary average values in 2017 were 9 nGy/month for Xe-133 and <3 nGy/month for Xe-135. No Ir-192 was detected at PN in 2017.

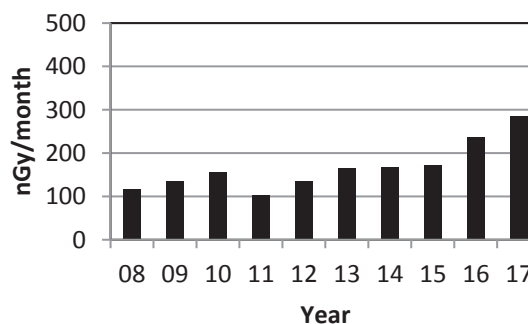


Figure 3-5: PN Annual Average Ar-41 Dose Rate in Air

Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>31 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****3.3.3 Terrestrial Sampling**

Terrestrial biota receive exposure from both airborne and waterborne emissions as indicated in Figure 4-1. Cow's milk, for example, is affected by the air, plants, and water sources that the cow consumes. It is therefore important to consider the combined effect of all these pathways when assessing the station impact on terrestrial samples.

Samples of soil, fruits, vegetables, animal feed, milk, eggs, and poultry are collected to support the public dose calculation for DN and PN sites. Background samples are also collected for calculating net concentrations for dose calculations. The radionuclides monitored and the sample collection frequencies are summarized in Table 3-1 and the 2017 results are discussed in the following sections. Detailed data are given in Appendix D, Tables D4 to D7.

**3.3.3.1 Fruits and Vegetables**

In the EMP designs, fruits and vegetables are sampled three times from each location for a representation of the entire growing season. Each sample is analysed for C-14 and HTO. Sampling locations for 2017 are shown in Appendix C.

A total of 10 fruit and vegetable locations were sampled around DN and 10 were sampled around PN. Fruits and vegetables were sampled from four background locations.

The results for vegetation are provided in Appendix D, Table D4.

**Tritium Oxide**

HTO concentrations in vegetation around the nuclear sites tend to vary from year to year due to prevailing winds, HTO emissions, humidity, etc. Furthermore, the number of samples and their locations change over the years. These variations should be considered when reviewing the following graphs and trend analysis results.

The average HTO concentrations measured in fruits and vegetables from the background locations in 2017 were < 2.2 Bq/L.

**DN – Figure 3-6**

The 2017 average concentration for HTO was 19.2 Bq/L in fruits and 19.2 Bq/L in vegetables. Figure 3-6 illustrates the combined DN fruit and vegetable annual average HTO results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.

**PN – Figure 3-7**

The 2017 average concentration for HTO was 45.9 Bq/L in fruits and 78.6 Bq/L in vegetables. Figure 3-7 illustrates the combined PN fruit and vegetable annual average

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>32 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

HTO results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.

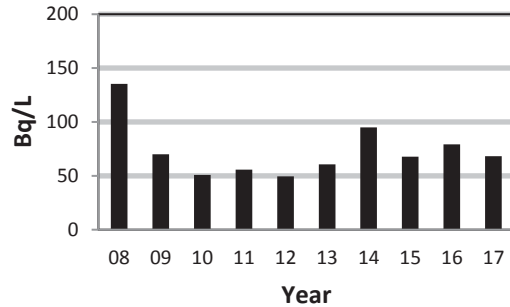
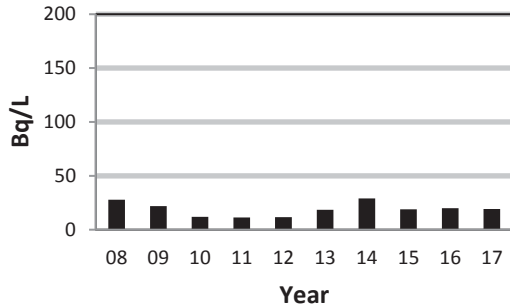


Figure 3-6: DN Annual Average HTO in Vegetation

Figure 3-7: PN Annual Average HTO in Vegetation

**Carbon-14**

The number of fruit and vegetable samples, their locations, and sampling frequencies have changed over the years, which should be considered when reviewing the following graphs and trend analysis results.

The average C-14 concentrations measured in fruits and vegetables from the background locations in 2017 were 235Bq/kg-C and 220 Bq/kg-C respectively.

**DN – Figure 3-8**

The 2017 average concentration of C-14 was 247 Bq/kg-C in fruits and 246 Bq/kg-C in vegetables. Figure 3-8 illustrates the combined DN fruit and vegetable annual average C-14 results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.

**PN – Figure 3-9**

The 2017 average concentration of C-14 at PN locations was 257 Bq/kg-C in fruits and 262 Bq/kg-C in vegetables. Figure 3-9 illustrates the combined PN fruit and vegetable annual average C-14 results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>33 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

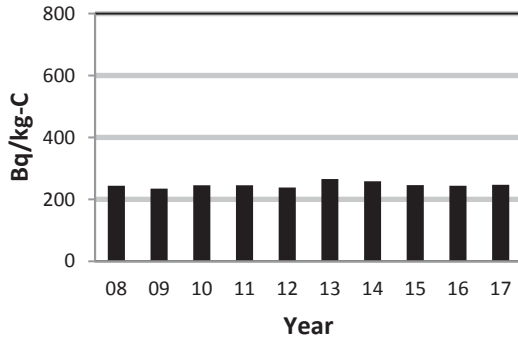


Figure 3-8: DN Annual Average C-14 in Vegetation

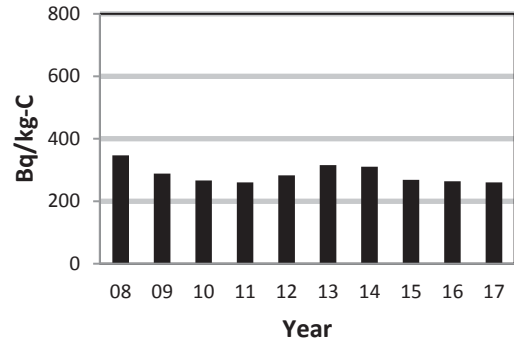


Figure 3-9: PN Annual Average C-14 in Vegetation

**3.3.3.2 Milk and Animal Feed**

Milk sampling is used to estimate the portion of dose received from milk ingestion for the Dairy Farm potential critical group. Milk consumed by other members of the public comes from commercial dairies whose products consist of composites from many dairy farms across Ontario. Values in this report are only applicable to residents of the surrounding dairy farms who consume raw milk and are not representative of milk that is sold at a grocery store.

Milk samples are collected on a monthly basis from dairy farms around DN and PN and analysed for HTO and C-14. Samples are collected from three dairy farms around DN and two dairy farms located around PN. Quarterly milk samples are collected from a background location with three replicates collected per quarter.

Locally grown animal feed is collected from four dairy farms around DN, twice a year, with two replicates collected per visit. Animal feed is collected from one dairy farm around PN and one dairy farm from a background location twice a year, with four replicates collected per visit. Since 2013, dry feed (grains, hay, etc.) and wet feed (forage) are collected separately. Animal feed is analysed for HTO and C-14.

Annual average values of HTO and C-14 in animal feed and milk are provided in Appendix D, Table D5 and D6, respectively.

The annual average HTO and C-14 in milk measurements around the nuclear sites vary from year to year due to changes in prevailing winds, emissions, humidity, cow's diet, feed sources, and water sources. These variations should be considered when reviewing the following graphs.

**Tritium Oxide**

The background average HTO in milk concentration was < 2.1 Bq/L and HTO in animal feed was 5.7 Bq/L for dry feed, and 6.6 Bq/L for wet feed (forage).

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>34 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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DN – Figure 3-10

The 2017 average concentration of HTO in milk was 5.6 Bq/L based on three dairy farms around DN. Figure 3-10 illustrates DN HTO in milk results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.

The average HTO concentration in animal feed was 16.4 Bq/L for dry feed (grains, hay, etc.) and 9.8 Bq/L for wet feed (forage). No trend analysis was performed on animal feed since, beginning in 2013, wet feed and dry feed have been sampled separately, resulting in changes to sampling frequency and replicates.

PN – Figure 3-11

The 2017 average concentration of HTO in milk was 14.6 Bq/L based on two dairy farms located within 12 km of PN. Figure 3-11 illustrates PN HTO in milk results over the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend for PN HTO in milk.

The average HTO concentration in animal feed was 20.7 Bq/L for dry feed (grains, hay, etc.) and 33.7 Bq/L for wet feed (forage). No trend analysis was performed on animal feed since, beginning in 2013, wet feed and dry feed have been sampled separately, resulting in changes to sampling frequency and replicates.

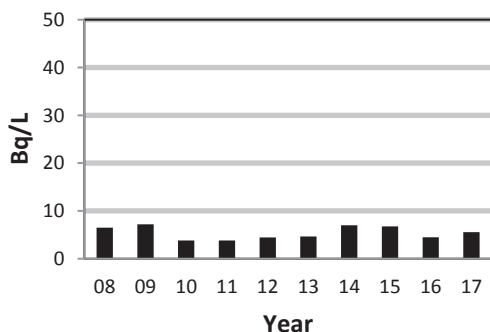


Figure 3-10: DN Annual Average HTO in Milk

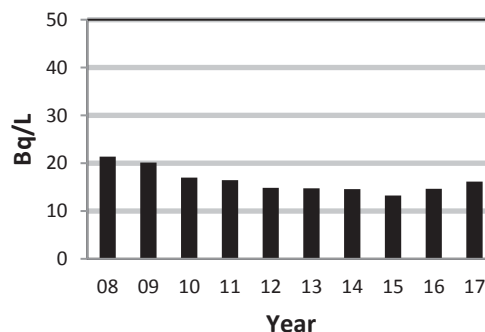


Figure 3-11: PN Annual Average HTO in Milk

Carbon-14

The background average C-14 in milk was 243 Bq/kg-C. C-14 in dry feed was 236 Bq/kg-C and 233 Bq/kg-C in wet feed (forage).

The C-14 level in animal feed consumed by the cows is the main contributing factor to the C-14 levels in milk. Animal feed contains C-14 from the previous year when it was grown, therefore emissions from the previous year would affect the C-14 values measured in milk in the current year for cows consuming local feed.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>35 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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DN – Figure 3-12

The 2017 average concentration of C-14 in milk from dairy farm locations in the vicinity of DN was 233 Bq/kg-C. Figure 3-12 illustrates that C-14 levels in milk around DN have been stable and near background levels for the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend.

The average C-14 concentration in animal feed was 239 Bq/kg-C for dry feed (grains, hay, etc.) and 247 Bq/kg-C for wet feed (forage). No trend analysis was performed on animal feed since, beginning in 2013, wet feed and dry feed have been sampled separately, resulting in changes to sampling frequency and replicates.

PN – Figure 3-13

The 2017 average concentration of C-14 in milk from dairy farm locations in the vicinity of PN was 233 Bq/kg-C. Figure 3-13 illustrates that C-14 levels in milk around PN have been stable and near background levels for the past 10 years. A Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend for PN C-14 in milk.

The average C-14 concentration in animal feed was 252 Bq/kg-C for dry feed (grains, hay, etc.) and 256 Bq/kg-C for wet feed (forage). No trend analysis was performed on animal feed since, beginning in 2013, wet feed and dry feed have been sampled separately, resulting in changes to sampling frequency and replicates.

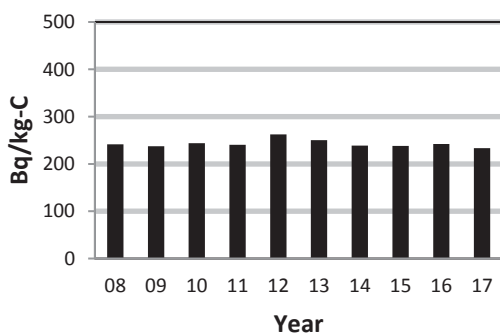


Figure 3-12: DN Annual Average C-14 in Milk

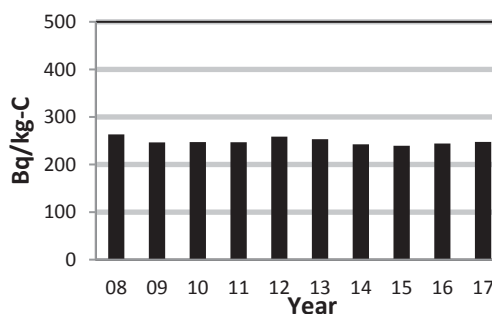


Figure 3-13: PN Annual Average C-14 in Milk

**3.3.3.3 Eggs and Poultry**

Eggs and poultry were added to the EMP sampling program in 2013. Eggs are sampled on a quarterly basis and three sample replicates are collected per visit. Poultry is collected annually with eight sample replicates collected per visit. Both eggs and poultry are analysed for HTO and C-14.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>36 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

One farm location around DN is sampled for eggs (D10) and one farm location is sampled for poultry (F16). No farm location selling fresh eggs and poultry could be found in the PN vicinity, and therefore these pathways are modeled for PN. One background location is sampled for both eggs and poultry.

The background concentration of HTO was 2.6 Bq/L for eggs and 2.9 Bq/L for poultry. The background concentration of C-14 was 233 Bq/kg-C for eggs and 236 Bq/kg-C for poultry.

The concentrations of HTO in DN eggs was 4.0 Bq/L and HTO in poultry was 8.9 Bq/L. C-14 in DN eggs was 233 Bq/kg-C and C-14 in poultry was 235 Bq/kg-C. Refer to Table D7 in Appendix D for detailed data. No trend analysis was performed as only four years of data have been collected from these locations thus far.

**3.3.3.4 Soil Sampling**

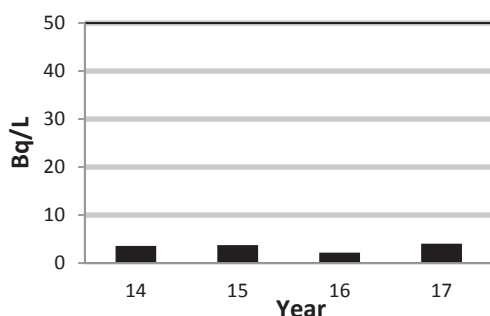


Figure 3-14: DN Annual Average HTO in Eggs

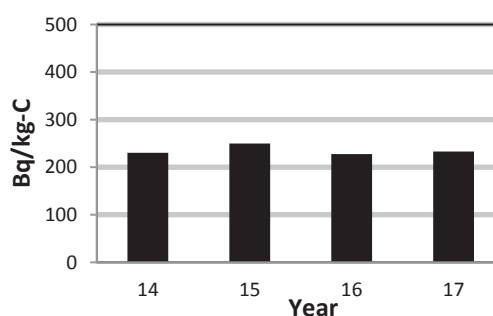


Figure 3-15: DN Annual Average C-14 in Eggs

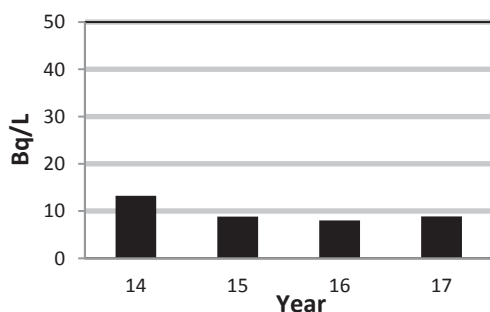


Figure 3-16: DN Annual Average HTO in Poultry

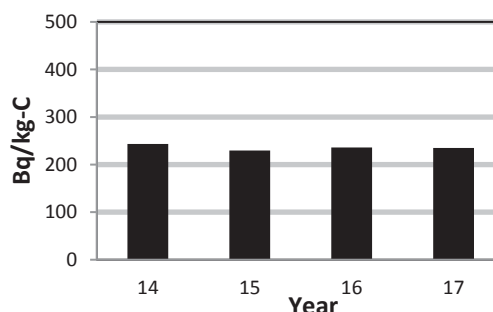


Figure 3-17: DN Annual Average C-14 in Poultry

Soil is sampled every five years to identify possible radionuclide accumulation over time. Sampling previously took place in 2012 [R-13] and most recently in 2017. The 2017 results for soil sampling are provided in Appendix D, Table D11.

## Public Information

Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>37 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****Gamma Spectrometry**

Background values of Cs-137 are present in the soil as a result of historic weapons testing fallout. Co-60 and Cs-134, if detected, would be a result of emissions from DN or PN.

In 2017, Cs-137 concentrations in background soil samples ranged from 1.7 to 9.0 Bq/kg. All measured Cs-137 concentrations around the sites are within the range of values seen at the background locations, ranging from 3.0 to 7.2 Bq/kg. At the DN locations, Cs-137 concentrations in soil ranged from 5.1 to 7.2 Bq/kg, and at the PN locations Cs-137 concentration in soil ranged from 3.0 to 4.5 Bq/kg. There is no indication of a buildup of activity in soil. Neither Cs-134 nor Co-60 were detected in any PN or DN soil samples in 2017. Therefore, the Cs-137 measured in these soil samples is from historic weapons testing fallout and not from OPG operations.

**3.3.4 Aquatic Sampling**

Samples of drinking water sources (municipal and well water), lake water, lake sediment, beach sand and fish are collected to monitor the aquatic environment around the DN and PN sites. Background samples are also collected to provide a comparison benchmark and to allow determination of net values for dose calculations. The radionuclides monitored and the sample collection frequencies are detailed in Table 3-1. Detailed data for the results of aquatic sampling are given in Appendix D, Tables D8 to D10, and discussed in the following sections.

**3.3.4.1 Water Supply Plants**

Samples of drinking water are taken during each 8-12 hour shift at water supply plants (WSPs) that supply water to Durham Region and the City of Toronto. Weekly composites of these samples are analyzed for HTO and monthly composites are analyzed for gross beta activity.

The locations of the WSPs sampled relative to the nearest nuclear station discharge are indicated in Table 3-2. The results of water sampled are provided in Appendix D, Table D8.

**Table 3-2: Water Supply Plants Monitored and Distance from Stations**

	Distance from Site
<b>DN AREA WSPs</b>	
Bowmanville WSP	7 km ENE of DN
Newcastle WSP	13 km E of DN
Oshawa WSP	8 km W of DN
<b>PN AREA WSPs</b>	
R.C. Harris WSP	22 km WSW of PN
Horgan WSP	11 km SW of PN
Ajax WSP	7 km ENE of PN
Whitby WSP	12 km ENE of PN

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>38 of 132</b>

Title:

### **2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

The impact of HTO emissions from OPG stations on the nearby WSPs varies depending upon distance from the station, lake current direction, location and depth of the WSP intake pipe and general dispersion conditions. Annual average HTO levels at all WSPs are well below the Ontario Drinking Water Quality Standard of 7,000 Bq/L [R-14].

A single sample hypothesis test was performed to demonstrate that the annual average at each WSP is below OPG's commitment to maintain HTO in drinking water below 100 Bq/L. Results from Ajax, Bowmanville, Whitby, Oshawa, Harris, Horgan, and Newcastle WSPs all showed annual averages < 100 Bq/L.

### **Tritium Oxide**

HTO in Lake Ontario, along with all the Great Lakes, originates from several sources: natural cosmogenic tritium, residual tritium fallout from atmospheric weapons testing, current emissions from nuclear plants, and residual HTO from past emissions of nuclear plants. For the purpose of calculating public dose resulting from OPG operations, the sum of contributions from current emissions and residual HTO from past emissions was used. The background HTO value, subtracted from HTO measurements, includes only natural cosmogenic tritium and residual weapons fallout tritium. This produces a conservative estimate of dose from tritium in lake water. This Lake Ontario background component for 2017 was conservatively estimated to be 1.4 Bq/L, using the Great Lakes Time-Concentration Tritium Model [R-15].

The WSPs annual average concentrations of tritium in drinking water are shown in Figures 3-18 through 3-24. A statistical trend analysis was performed for each WSP over a 10 year period.

#### **DN – Figures 3-18 to 3-20**

Annual average HTO concentrations measured at the Bowmanville, Newcastle, and Oshawa WSPs ranged from 6.4 to 10.8 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for HTO at any DN WSP location.

#### **PN – Figure 3-21 to 3-24**

Annual average HTO concentrations measured at the Ajax, Horgan, Harris, and Whitby WSPs ranged from 4.6 to 8.3 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for HTO at any PN WSP location.

Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>39 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

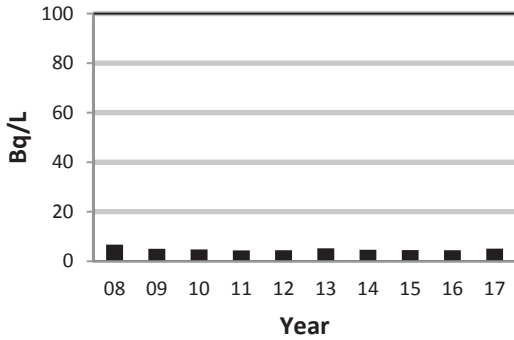


Figure 3-18: Bowmanville WSP – Annual Average HTO in Water

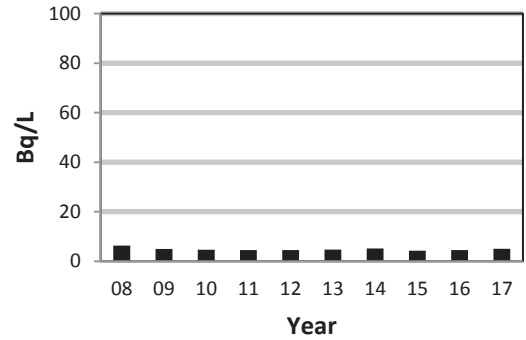


Figure 3-19: Newcastle WSP – Annual Average HTO in Water

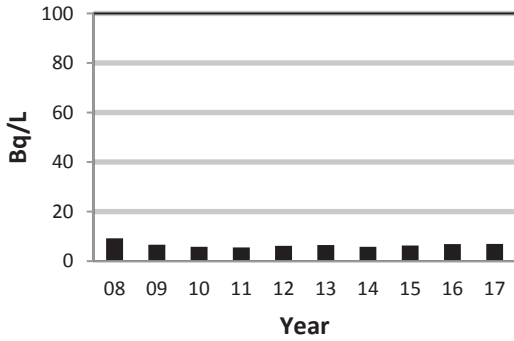


Figure 3-20: Oshawa WSP – Annual Average HTO in Water

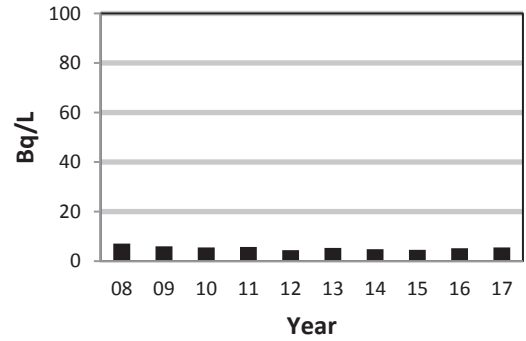


Figure 3-21: Ajax WSP – Annual Average HTO in Water

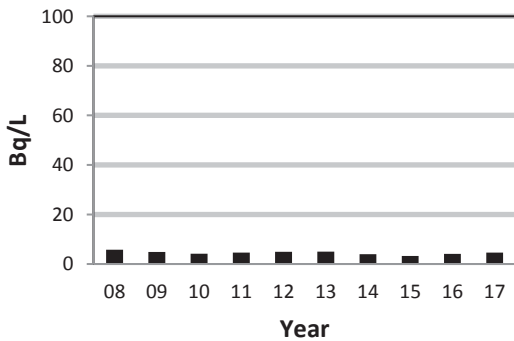


Figure 3-22: Scarborough Horgan WSP – Annual Average HTO in Water

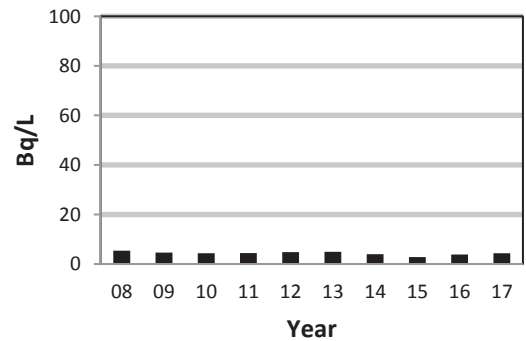


Figure 3-23: Toronto Harris WSP – Annual Average HTO in Water



## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>40 of 132</b>

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

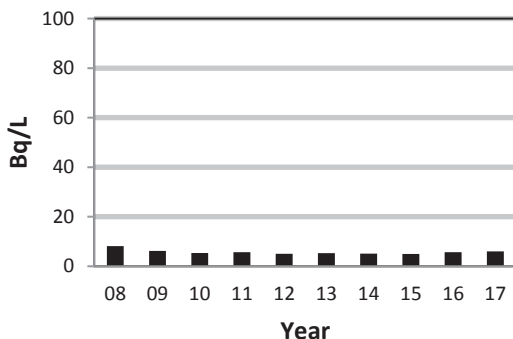


Figure 3-24: Whitby WSP – Annual Average HTO in Water

### Gross Beta

Annual average gross beta activity levels in samples from DN and PN area WSPs were 0.11 Bq/L. This is well below the gross beta activity screening level of 1 Bq/L, which is both an internal OPG level and a drinking water level recommended by Health Canada [R-16].

#### 3.3.4.2 Well Water

Monthly well water samples are collected from four wells around the DN area and two wells around the PN area. The wells sampled represent the potential critical groups for which the annual public dose is calculated under the EMP designs. Samples are analyzed monthly for HTO. Analytical results are provided in Appendix D, Table D8.

### Tritium Oxide

HTO concentrations in well water depend on the depth of the well and thus the amount of time it takes for precipitation to reach the aquifer from where the well draws its water. Radioactive decay of the tritium during its transit time to the aquifer determines the residual activity level in the well water. Deeper wells tend to have lower HTO concentrations. Well water HTO concentrations reflect the level of past atmospheric HTO releases because of the time it takes for precipitation to reach the well.

#### DN – Figure 3-25

The 2017 annual average HTO concentration observed in well water samples collected from the DN area was 10.6 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for DN HTO in well water.

#### PN – Figure 3-26

The 2017 annual average HTO concentration observed in well water samples collected from the PN area was 14.7 Bq/L. Based on the past 10 years of data, a

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>41 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend for PN HTO in well water.

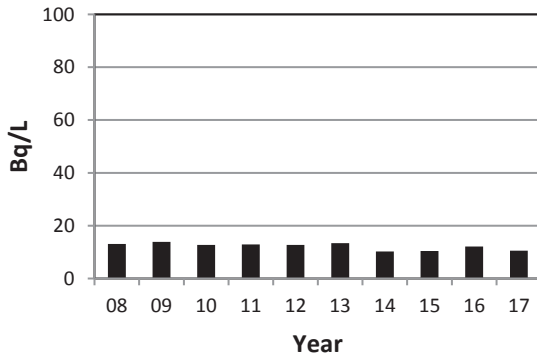


Figure 3-25: DN Annual Average HTO in Well Water

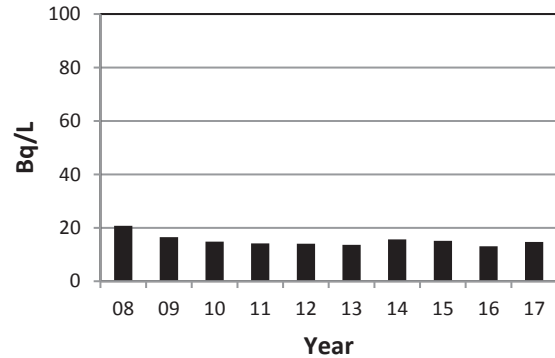


Figure 3-26: PN Annual Average HTO in Well Water

### 3.3.4.3 Lake Water

Lake water (non-drinking water) is sampled from two beaches in the vicinity of DN and three beaches in the vicinity of PN on a monthly basis and analysed for HTO. It is used to assess the water immersion dose exposure pathway from swimming in lake water. Sampling of lake water is not required during the winter months as it is not representative of public exposure. Analytical results are provided in Appendix D, Table D8.

#### DN – Figure 3-27

The 2017 annual average HTO concentration observed in lake water samples collected from two beaches in the DN area was 18.1 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend for DN HTO in lake water.

#### PN – Figure 3-28

The 2017 annual average HTO concentration observed in lake water samples collected from three beaches in the PN area was 21.2 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for PN HTO in lake water.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>42 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

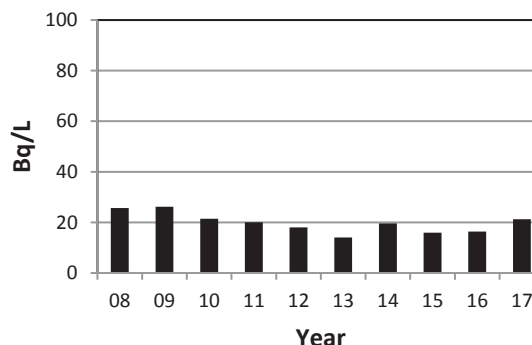
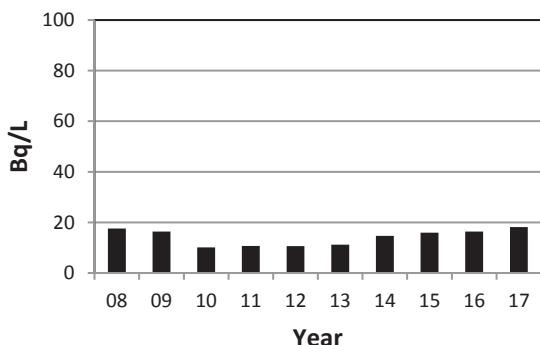


Figure 3-27: DN Annual Average HTO in Lake Water

Figure 3-28: PN Annual Average HTO in Lake Water

### 3.3.4.4 Fish

At the DN site, fish sampling takes place over the cooling water discharge diffuser. At the PN site, the sampling location is in the PN outfall. Background samples are taken from the Bay of Quinte area of Lake Ontario.

The target fish species to be collected at DN, PN, and at background locations is White Sucker, with Brown Bullhead as the backup species. Eight replicate fish samples are collected and analyzed at each location. A sample consists of the fish muscle tissue only, and excludes the head, skin, fins, and as many bones as possible. HTO, C-14, Co-60, Cs-134, Cs-137, and Potassium-40 (K-40) measurements are performed on each fish sample.

The results for fish are provided in Appendix D, Table D9.

#### Tritium Oxide

The HTO levels in fish change quickly in response to changes in water HTO levels from waterborne emissions. Thus, HTO concentrations measured in fish tissue reflect the HTO concentration in the water in the few hours before they were sampled. Long-term graphs of fish HTO levels for PN and DN are provided in Figures 3-29 and 3-30. In 2017, the HTO in Lake Ontario background fish samples averaged < 2.3 Bq/L.

#### DN – Figure 3-29

The HTO levels in the DN diffuser fish samples averaged 4.6 Bq/L. This value is slightly lower than levels observed in previous years. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for HTO in DN fish.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>43 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

PN – Figure 3-30

The HTO concentration in the PN outfall fish samples averaged 5.9 Bq/L. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level indicates a decreasing trend for HTO in PN fish.

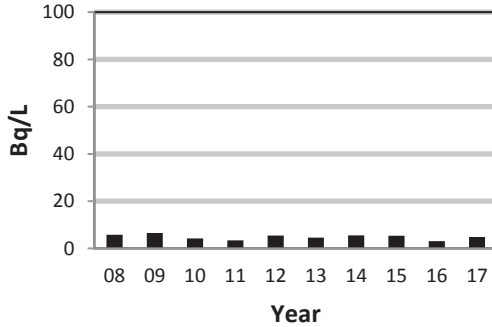


Figure 3-29: DN Annual Average HTO in Fish

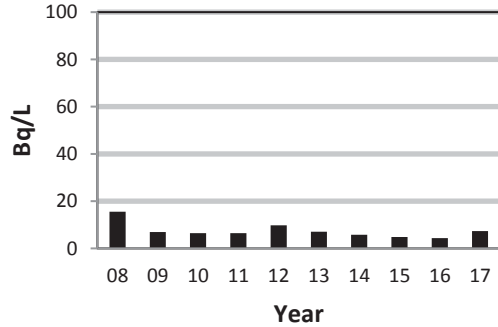


Figure 3-30: PN Annual Average HTO in Fish

**Carbon-14**

The average C-14 level in fish measured at a background Lake Ontario location was 230 Bq/kg-C in 2017.

The concentrations of C-14 in fish at both DN and PN are consistent with past years and comparable to background levels, as shown in Figures 3-31 and 3-32.

DN – Figure 3-31

The 2017 annual average C-14 level in DN fish was 240 Bq/kg-C. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for C-14 in DN fish.

PN – Figure 3-32

The 2017 annual average C-14 level in PN fish was 248 Bq/kg-C. Based on the past 10 years of data, a Mann-Kendall trend analysis at the 95% confidence level does not indicate any statistically significant trend for C-14 in PN fish.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>44 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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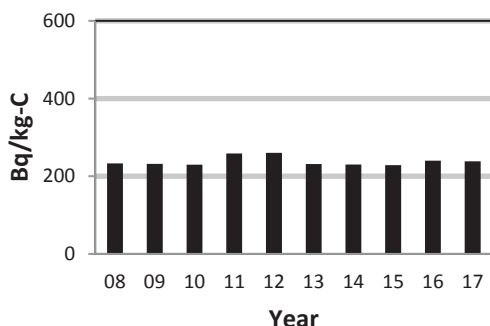


Figure 3-31: DN Annual Average C-14 in Fish

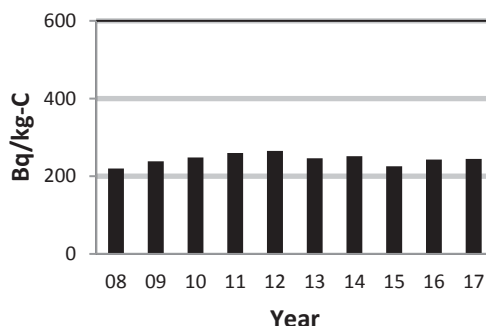


Figure 3-32: PN Annual Average C-14 in Fish

### Gamma Spectrometry

The majority of the gamma activity in fish is naturally occurring K-40. A small amount of Cs-137 is usually present which is primarily due to nuclear weapons testing and not reactor operation given that Cs-134 and Co-60, which are indicative of reactor operation, were not detected.

The average Cs-137 value for background Lake Ontario fish was 0.3 Bq/kg.

#### DN – Figure 3-33

The average Cs-137 value for DN fish was 0.1 Bq/kg. Given the level of uncertainty at such low concentrations, this is not distinguishable from background. Cs-134 and Co-60, which are indicative of reactor operation, were not detected in any fish samples at DN site in 2017.

#### PN – Figure 3-34

The average Cs-137 value for PN fish was 0.1 Bq/kg. Given the level of uncertainty at such low concentrations, this is not distinguishable from background. Cs-134 and Co-60, which are indicative of reactor operation, were not detected in any fish samples at PN site in 2017.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>45 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

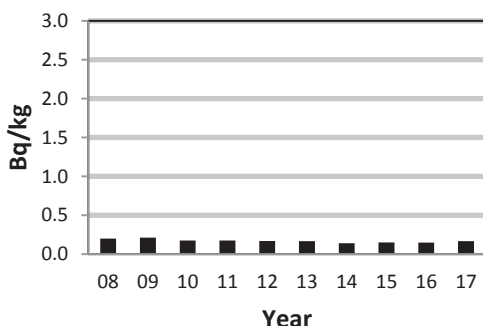


Figure 3-33: DN Annual Average Cs-137 in Fish

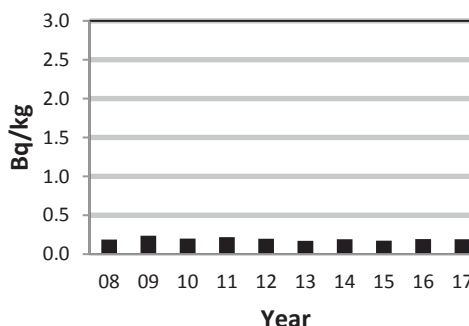


Figure 3-34: PN Annual Average Cs-137 in Fish

### 3.3.4.5 Beach Sand

Sand from three beaches around DN and three beaches around PN is collected annually to represent a potential pathway for external dose. Eight replicates are collected per sampling location. Gamma spectrometry is performed on these samples.

Beach sand samples were collected at Cobourg to determine the Cs-137 concentration in Lake Ontario background sand due to atmospheric weapons test fallout.

The results for beach sand are provided in Appendix D, Table D10.

#### Gamma Spectrometry

Background Cs-137 concentrations in beach sand samples measured at Cobourg averaged 0.4 Bq/kg in 2017. These values are consistent with values observed over the past five years.

#### DN

There was no Cs-137 detected in DN beach sand in 2017. Similar to previous years, there was no Co-60 or Cs-134 detected in any of the samples.

#### PN

The average concentration of Cs-137 measured at PN area beaches ranged from below detection (< 0.2) to 0.4 Bq/kg. Similar to previous years, there was no Co-60 or Cs-134 detected in the samples.

Wave action continuously moves the beach sand around, disturbing the original deposition patterns. This range of Cs-137 values is similar to the background values measured and, without the presence of other radionuclides such as Co-60 or Cs-134 that are more closely related to reactor operation, the Cs-137 measured along the shoreline cannot be confirmed to be the result of OPG operations.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>46 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

### 3.3.4.6 Sediment

Lake sediment is sampled every five years to identify possible radionuclide accumulation over time. The last sampling was conducted as part of a study commissioned by the CANDU Owners Group (COG) and took place in 2011 [R-17]. The 2011 results for sediment sampling are provided in the 2014 Results of Environmental Monitoring Programs report [R-18]. While EMP sediment sampling was scheduled to take place in 2016, it was deferred to 2019 as sampling was conducted only one year prior, in 2015, in support of the PN Safe Storage ERA [R-61]. The details and results of the 2015 sediment sampling is summarized in section 3.4.3 of this report. Station and background sediment sampling in support of the EMP will be conducted in 2019.

## 3.4 Supplementary Studies

CSA N288.4-10 specifies that supplementary studies can occasionally be conducted as part of the EMPs to achieve specific, well-defined objectives such as:

- providing the data required to reduce uncertainty and confounding factors in the risk assessment;
- increasing knowledge of the behaviour of contaminants and physical stressors in the environment (e.g., refining environmental transfer parameters);
- investigating specific EMP findings; and
- follow-up monitoring of mitigation activities implemented following an EA.

Supplementary studies are site-specific and as such may vary between nuclear facilities. These studies become part of the EMPs until the objective of the study has been achieved. At that time, the supplementary study is terminated.

In 2017, OPG conducted or finalized three supplementary studies in support of the PN EMP and three supplementary studies in support of the DN EMP.

- (a) Air Kerma Rate from the PWF at the PN Site
- (b) Contaminants of Potential Concern in Soil at PN
- (c) Sediment and Water Sampling of Non-radiological Contaminants at PN
- (d) Entrainment Study at DN
- (e) Conventional Effluent Characterization Study at DN
- (f) Benthic Study at DN

The following sections provide a description and the results of each study.

### 3.4.1 EMP Supplementary Study - Air Kerma Rate from the PWF at the PN Site

The PN EMP design document [R-59], identified a recommendation to measure direct gamma and skyshine dose from used fuel dry storage (UFDS) buildings to account for the changes in the amount of used fuel stored since the last gamma ray survey



## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>47 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

conducted in 2000 [R-51]. In 2017, a gamma ray survey was conducted on Lake Ontario to estimate the air kerma rate from the UFDS buildings.

### 3.4.1.1 Method

#### Sampling Plan

Direct measurements of gamma and skyshine dose are problematic because of the difficulties in detecting the used fuel radiation in the presence of the natural gamma background. Direct radiation and skyshine from used fuel storage buildings at PN was estimated by making a series of measurements at different distances, over the lake to screen out terrestrial gamma background (lake bottom terrestrial background radiation is shielded by the overlying water). Cosmic gamma background was measured by moving farther away from the PN site so background measurements could be subtracted, leaving only the site gamma dose contribution. Measurements of total gamma dose over the lake at specified distances from the source structures on site gives a dose rate versus distance relationships that can be applied to critical groups.

Health Canada conducted the gamma ray survey on June 7th and 8th, 2017 using a RS-250 spectrometer mounted on a boat. A series of measurements were taken at different distances from the shore along five transects. The location of two additional background transects were chosen sufficiently far from PN that the effect of radiation from the UFDS buildings could be neglected.

### 3.4.1.2 Results

The results demonstrate that the air kerma rate due to the waste storage facilities cannot be detected at distances greater than 400 m from the waste storage facilities. At a distance greater than 400 m from the storage facilities, the air kerma rate is less than the detection limit of 0.33 nGy/h. Data are summarized in Appendix H, Table H1.

When comparing the contributions from the storage facilities to the air kerma rate, they are slightly more elevated in 2017 than in 2000, due to increased storage of used fuel. This is approximately 0.06 % of the 30 nGy/y due to cosmic radiation and 30 nGy/y due to natural background radiation from the ground. [R-19]

### 3.4.1.3 Conclusions and Recommendations

The measurements around the waste storage facilities show that the highest air kerma rate contribution from the storage facilities at the closest data point to the UFDS buildings is 1.8 nGy/h. At 400 m distance, the air kerma rate contribution from the storage facility is approximately 0.1 nGy/h. [R-19]

The skyshine dose from this source is, therefore, not significant for potential critical groups outside the 1 km boundary, which are all the potential critical groups except the Fisher critical group, assumed to be located 500 m south of PN in Lake Ontario. [R-19]

**Report**

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>48 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**3.4.2 EMP Supplementary Study – Contaminants of Potential Concern in Soil at PN**

As recommended by the PN 2014 ERA, a supplementary monitoring study was conducted in 2015 to update soil data for the PN site. The goal of the soil monitoring program was to help reduce uncertainty regarding concentrations of contaminants of potential concern (COPCs) used in dose calculations for non-human biota. The results of this study are summarized as part of the most recent PN ERA [R-61], in 2017.

**3.4.2.1 Method**

**Sampling Plan**

Soil sampling occurred as part of the 2015 baseline environmental sampling program and focused on areas with historically elevated concentrations of tritium and metals. Soil was collected from areas with vegetation and organic soil cover that may represent potential habitat on site. Samples were collected in October 2015 from eight locations around the PN site.

**Screening for COPCs**

Maximum measured concentrations of parameters in soil were compared to the selected screening criteria to determine the list of COPCs to be assessed further. Radionuclides were carried forward for further assessment irrespective of screening criteria, as they were considered of public interest.

The following contaminants of potential concern were carried forward for further quantitative assessment in the Ecological Risk Assessment (EcoRA): Tritium, carbon-14, cesium-134, cesium-137, cobalt-60, cyanide, arsenic, copper, lead, zinc, and petroleum hydrocarbon F4.

**3.4.2.2 Results**

The EcoRA indicated that average concentrations in soil did not exceed acceptable risk levels for mammals or birds. Though the EcoRA indicated that localized effects to individual earthworms or plants may occur, the earthworm and terrestrial plant populations on site are not affected. Furthermore, species at risk are not impacted from site operations. Data are summarized in Appendix H, Table H2.

There were no exceedances of the radiation dose benchmark for terrestrial biota on the PN site.

**3.4.2.3 Conclusions and Recommendations**

In general, soils on site that exceed benchmark concentrations are localized. This suggests that the source of contaminants is from past industrial operations, and not deposition from atmospheric sources. Therefore, accumulation of COPCs in soil over time is not expected. [R-61]

### 3.4.3 EMP Supplementary Study – Sediment and Water Sampling of Non-radiological Contaminants at PN

As recommended by the PN 2014 ERA, a supplementary monitoring study was conducted in 2015 to collect sediment and water samples from the northern section of the Frenchman's Bay wetland. The goal of the study was to reduce uncertainty regarding the assessment of biota in the bay, since previously, biota were assessed at the mouth of the bay where sediment data were available, south of the wetland. The results of this study are summarized as part of the most recent PN ERA [R-61], in 2017.

#### 3.4.3.1 Method

##### Sampling Plan

Sediment and surface water sampling occurred as part of the 2015 baseline environmental sampling program. Ten sediment samples and three surface water samples were collected from the north and south ends of Frenchman's Bay in July 2015. As there was little variability in measurements, the data from both areas of the bay were pooled together.

##### Screening of COPCs

Samples were screened against relevant water and sediment quality guidelines to determine the list of COPCs to be assessed further. As expected, numerous COPCs exceeded screening levels, since Frenchman's Bay is impacted by urban runoff. Therefore, the proportion of overall risk to aquatic receptors attributed to PN was assessed.

The following COPCs in water were carried forward for further quantitative assessment in the EcoRA: total aluminum, copper, iron and sodium.

The following metals in sediment were carried forward for further quantitative assessment in the EcoRA: Aluminum, bismuth, boron, cadmium, calcium, chromium, copper, lead, manganese, nickel, phosphorous, thorium, tin, zinc.

#### 3.4.3.2 Results

The percent contributions from PN to water concentrations observed in Frenchman's Bay range from 0.3% to 22%. Overall, the contribution to the total concentration of metals at Frenchman's Bay from PN is low. Data are summarized in Appendix H, Tables H3 and H4.

#### 3.4.3.3 Conclusions and Recommendations

In general, the EcoRA showed that the exposure levels for non-radiological contaminants are below benchmark values. Where benchmark values were exceeded, the effects are highly localized and therefore the receptor populations are not expected

**Report**

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>50 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

to experience any adverse effects due to non-radiological releases from PN operations.

**3.4.4 EMP Supplementary Study – 2015/16 Entrainment Study for DN**

Following the completion of the DN refurbishment and continued operations EA and as committed in the DN Fisheries Act authorization, OPG completed a follow-up monitoring program to monitor entrainment at DN prior to the commencement of refurbishment in 2016 [R-62]. The goal of the program was to characterize the station’s entrainment of ichthyoplankton (i.e., fish eggs and larvae) and benthic invertebrates and to confirm no significant residual adverse effects to aquatic biota as a result of Condenser Cooling Water (CCW) operations. A final report was provided to the Department of Fisheries and Oceans Canada in December 2017 [R-63].

**3.4.4.1 Method**

**Sampling Plan**

Entrainment data was collected from December 2015 to November 2016. A total of 64 24-hour sampling events, spanning both daytime and nighttime periods (approximately 12 hours for each period), were completed. A total of 124 samples (daytime and nighttime) were collected. Water from the forebay was pumped through an ichthyoplankton net for sample collections. Flow rates were also measured to facilitate entrainment estimates. Densities of ichthyoplankton and benthic invertebrates were then extrapolated, based on station flows.

**3.4.4.2 Results**

A total of 2,613 fish eggs and 122 fish larvae were entrained over the 12 month sampling period. Alewife (*Alosa pseudoharengus*) comprised over 90% of fish eggs entrained, while Round Goby (*Neogobius melanostomus*) comprised 88.5% of total larvae entrained. It is estimated that a total of 94,482,521 eggs and 10,983,411 larvae were entrained over one year at DN. Data are summarized in Appendix H, Table H5.

Equivalent loss estimates of entrainment numbers were modeled and estimates were used to extrapolate the annual losses to equivalent numbers of Age 1 fish. The equivalent Age 1 biomass lost was estimated at 48 kg.

Approximately, 22,301 individual benthic invertebrates were entrained during the study. *Echinogammarus* and other amphipods (likely *Gammarus*) composed approximately 91% of the estimated benthic invertebrates entrained annually. It is estimated that a total of 1,548,288,043 benthic invertebrates were entrained over one year. Data are summarized in Appendix H, Table H6.

**3.4.4.3 Conclusions and Recommendations**

An estimated 589 kg of biomass was entrained during the study. This estimate is likely conservative as entrained eggs and larvae assume 100% mortality. The equivalent Age 1 biomass lost was estimated at 48 kg. Additionally, it is concluded that

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>51 of 132</b>

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

entrainment at DN does not negatively impacting local benthic invertebrate populations. [R-62]

### 3.4.5 EMP Supplementary Study – Effluent Characterization Study at DN

As identified in the DN refurbishment and continued operations EA, OPG completed an effluent characterization study at DN in 2016. The objective of the study was to confirm EA findings of non-significant impact from non-radiological effluent constituents on human and non-human biota [R-64]. The data from the study was used in the DN 2016 ERA [R-65].

#### 3.4.5.1 Method

##### Sampling Plan

Four effluent streams from facility operations to the CCW were sampled over six months for the purpose of this study. Sampled effluent streams included Active Liquid Waste, the Water Treatment Plant, Inactive Drainage, and Boiler Blowdown. As most streams are routinely sampled weekly, 24 samples from each effluent stream were collected. The sampling plan followed CSA N288.5 standard on effluent monitoring for Class 1 nuclear facilities and uranium mines and mills [R-66]

#### 3.4.5.2 Results

The chemical concentration data from the effluent characterization study was used in the most recent DN ERA to determine if any new COPCs warranted further assessment [R-65]. Data are summarized in Appendix H, Table H7 to H11.

#### 3.4.5.3 Conclusions and Recommendations

Analytical results were found to be acceptable and adequate for statistical characterization of effluent at DN. [R-64]

### 3.4.6 EMP Supplementary Study – 2016 Benthic Study for DN

Following the completion of the DN refurbishment and continued operations EA, OPG completed a follow-up monitoring program [R-67] to determine the baseline abundance and species diversity of benthic invertebrates in the area of the DN intake structure and to compare the results to nearshore studies completed in 2008 in the vicinity of the proposed New Nuclear at Darlington (NND) project infill area.

#### 3.4.6.1 Method

##### Sampling Plan

Benthic invertebrates on sediment (epifaunal) and in sediment (infaunal) were sampled in the vicinity of DN and at two reference locations; near Bond Head and Thickson Point. Epifauna samples were collected towing a paired benthic sled on three occasions (May, June and August) and infauna samples were collected using a diver

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>52 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

assisted airlift sampler in late August to early September. Twelve replicate samples were collected at each of the three locations, at 5m, 10m and 15m depths. Benthic samples were identified and counted and invertebrate abundance, diversity, community composition and ecological niche were calculated.

### 3.4.6.2 Results

#### Epifauna

On average, invertebrate abundance ranged from about 1,500 to 9,000 animals per benthic tow across the study area and taxa richness was in the range of 7 to 9. The range of Evenness and Simpson's Diversity scores were similar across the study area.

#### Infauna

At DN, mean invertebrate density ranged from 3,577 to 22,743 animals per m<sup>2</sup>, and taxa richness ranged from 11 to 23 taxa. Invertebrate density and richness at DN were within the ranges of what was measured at the reference locations at each depth. Evenness and Simpson's Diversity scores at DN were also within the range of those measured at the reference locations.

Data are summarized in Appendix H, Table H12.

### 3.4.6.3 Conclusions and Recommendations

Results of both the epifauna and infana sampling program demonstrate that the composition of the benthic invertebrate community at DN was within the range of variability observed in invertebrate community composition at the reference locations. Therefore, there is no effect on the benthic community related to CCW operations. Furthermore, the results of this study were similar to those of the nearshore studies completed in 2008 in support of the NND project EA. [R-67]

## 3.5 Other Studies

### 3.5.1 Potassium in Lake Water

Concentrations of potassium in lake water around PN and DN are monitored to support validation of the CSA N288.1-08 [R-20] default cesium bioaccumulation factor (Cs BAF) for fish of 3,500, which is used for the calculation of DRLs. The Cs-BAF value is based on an equation recommended by the International Atomic Energy Agency (IAEA) in the Technical Report Series (TRS)-472 report, which considers the relationship of the Cs BAF to lake water concentrations of potassium [R-21]. This study is conducted once every three years. The next potassium in lake water measurements will take place in 2019.

## 3.6 Areas of Regulatory Interest and Other Monitoring Programs

While the primary focus of this report is the results of 2017 monitoring conducted in support of the annual public dose calculation, the overall EMPs encompass several



## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>53 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

other OPG monitoring programs, which are described in Sections 3.6.1 to 3.6.3. Due to differences in reporting requirements and schedules, the information in the following sections is the most recent information available. Some 2017 information is based on preliminary data and/or reports as the finalized reports have not been issued at the time of this report's preparation.

### 3.6.1 Thermal Monitoring Program

The discharge of warm water through operation of the CCW system has the potential to impact the spawning success and larvae development of round whitefish. As a result of the CNSC's comments on a study completed in 2010 on the impact of PN thermal discharge on round whitefish spawning [R-22], a COG study on the effects of fixed and fluctuating temperatures on mortality and hatch of round whitefish and lake whitefish eggs was initiated and issued in 2014 [R-23]. This study prompted OPG to perform a re-assessment of the impacts of the thermal emissions from DN and PN on the survival of round whitefish eggs in Lake Ontario.

The COG study indicated that round whitefish are not as sensitive to thermal impact as previously suggested. Both station re-assessments concluded that the risk of thermal emissions on round whitefish is low and no further mitigation or offsetting is warranted. However, OPG made a commitment in the Darlington Refurbishment Environmental Assessment to monitor ambient substrate lake temperature during the winter months, and worked with Environment and Climate Change Canada and the CNSC to develop the monitoring program [R-24].

The program primarily uses the Darlington lake current monitor, with the Pickering lake current monitor as a backup. The objective is to examine the trend in winter water temperatures to inform an adaptive management program to protect round whitefish, should the potential effects of climate change cause significant increases in winter season lake bottom temperatures.

Whitefish spawn in late fall on coarse substrates (gravel or cobble) between the depths of 3 to 12 m. The embryos develop over the winter and hatch in spring. Suitable spawning habitat is present near the Darlington CCW discharge. Temperature impacts egg development in two ways: 1) increased temperature may lead to direct mortality of the eggs; and 2) increased temperature shortens the gestation period, leading to earlier hatch. The average winter temperature between December 1<sup>st</sup> and March 31<sup>st</sup> is compared to a threshold of 6.0° C, with the intention of implementing an adaptive management program if the 6.0° C threshold were to be exceeded [R-25].

The average lake temperature at the Darlington Lake Current Monitor between December 1<sup>st</sup> 2016 and March 31<sup>st</sup> 2017 was 3.0°C. Therefore, no additional actions are required. Long term trends are provided in Figure 3-35 below. There is no indication of a warming trend which would approach the threshold in the near term.



Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>54 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

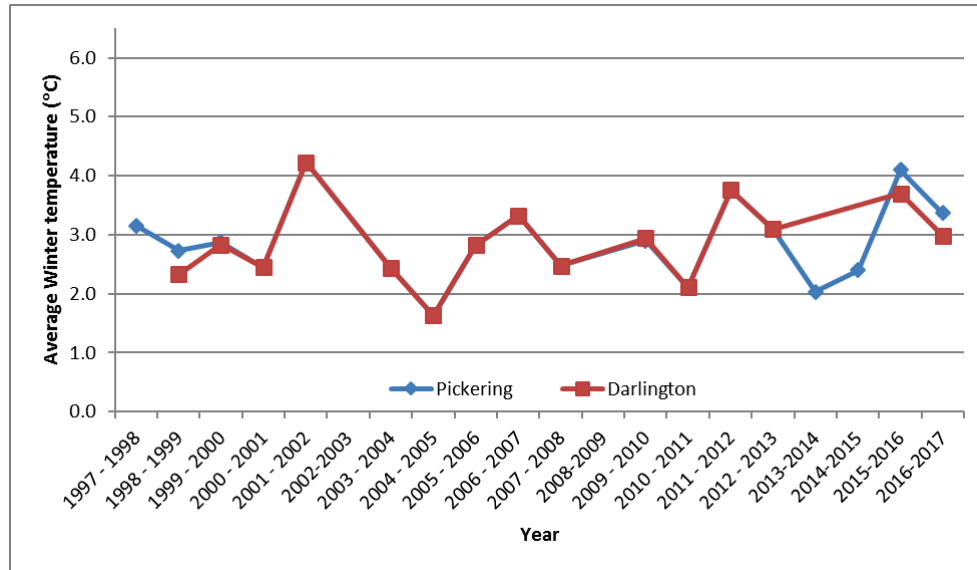


Figure 3-35: Long term trends in Lake Ontario winter temperatures (Dec 1st to March 31st)

Whitefish eggs may also be susceptible to temperature during the early phase of embryo development [R-26]. Environment and Climate Change Canada requested that OPG trend temperatures during the first month of egg development. Long term trends are provided in Figure 3-36 below. There is no indication of a warming trend approaching the threshold that would require adaptive management.

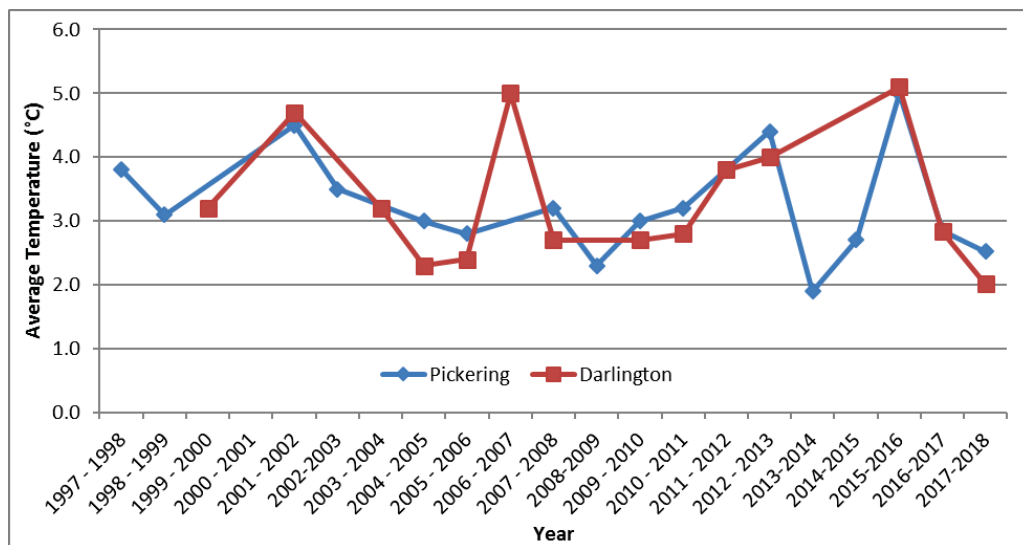


Figure 3-36: Long term trends in Lake Ontario winter temperatures (Dec 15th to January 15th)

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>55 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

### 3.6.2 Impingement and Entrainment Monitoring Program

Since 2009, OPG has seasonally installed a Fish Diversion System (FDS) at PN to reduce impingement of all fish species by 80%. Annual reporting of fish impingement is required by the CNSC to ensure ongoing compliance with reduction targets.

Results of the 2016 monitoring program are presented in the Pickering Nuclear 2016 Impingement Monitoring Report [R-27]. The biomass impinged in 2016 was estimated to be 1,035 kg, or 0.22 kg/million m<sup>3</sup> of station flow. This is the lowest annual biomass impingement rate since monitoring of the Fish Diversion System (FDS) began in 2010.

The preliminary estimate of biomass impinged in 2017 is 1,217 kg. In addition to this amount, an impingement event occurred on November 16-18, 2017 during which an estimated 24,000 kg of Alewife were impinged. OPG is currently investigating this event, and the result of the investigation will be discussed further in the Pickering Nuclear 2017 Impingement Monitoring Report.

In addition, with respect to the wetland improvement project initiated in 2014, OPG retained the Toronto and Region Conservation Authority (TRCA) to upgrade a portion of the Duffins Creek wetland in order to offset residual impingement losses. The project design phase was initiated in 2015 and will include the construction of a levee, fish gate and water control structure at Simcoe Point Wetland. In 2017, per Section 35 of the Federal Fisheries Act, OPG obtained a Fisheries Act Authorization for the Duffins Creek wetland project and the residual impingement from the operation of Pickering Nuclear.

### 3.6.3 Groundwater Monitoring Program

In 2017, PN and DN completed annual groundwater monitoring programs to evaluate groundwater quality and flow across the sites and to detect any emergent issues.

The groundwater monitoring programs occur from January 1 to December 31 of each year with 213 groundwater monitoring wells sampled in 2017 for tritium, the key contaminant of concern. Annual water level measurement events were also conducted for each site. Within certain areas, samples were also analyzed for select hazardous substances, such as petroleum hydrocarbons (PHCs) and volatile organic compounds (VOCs).

In general, tritium trends over time show levels for the most part that have remained nearly steady or have decreased, indicating stable or improved environmental performance. There are isolated cases where tritium concentrations have shown increases. Expected increases occur when tritium is migrating as a plume. Where unexpected tritium concentrations are identified, investigations are completed to determine the root cause and to implement corrective measures. Ongoing results confirm that tritium in groundwater is mainly localized within the station protected area and the site perimeter tritium concentrations remain low.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>56 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

#### 4.0 ASSESSMENT OF RADIOLOGICAL DOSE TO THE PUBLIC

This section contains an assessment of doses to the public resulting from the operation of OPG’s PN and DN sites. The effective dose limit for members of the public as set out in the Radiation Protection Regulations [R-28] is 1,000 µSv/year. The environmental samples collected and analysed through the PN and DN EMPs are used to produce realistic estimates of radiation doses to the public resulting from the operation of PN and DN sites, and to demonstrate that these doses remain below the regulatory limit.

The doses are heavily based on environmental concentrations of radionuclides measured at the potential critical group locations and surrounding environment. For the radionuclides and pathways where environmental measurements are not available, dose is modeled from emissions.

The dose calculation follows the method described in OPG’s Methodology for Data Analysis and Public Dose Determination for the Environmental Monitoring Program [R-29]. Assumptions, model parameters, and mean intake rates are used in accordance with CSA N288.1-08 [R-20]. Annual average meteorological data are used along with local intake fractions and representative locations for potential critical groups identified in the site-specific survey reviews [R-30] [R-31]. Appendix F provides details on how the data are used.

Figure 4-1 represents the model of exposure pathways to human receptors used for public dose calculation.

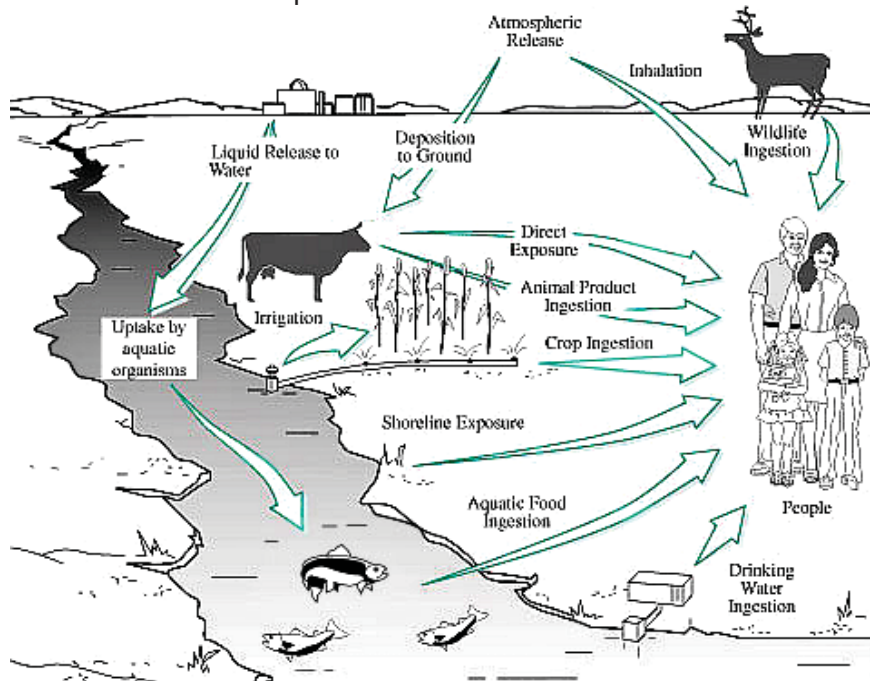


Figure 4-1: Model of Exposure Pathways from Site Emissions

Source: Based on United States Department of Energy/Hanford Site

Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>57 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****4.1 Modelling****4.1.1 Integrated Model for Probabilistic Assessment of Contaminant Transport (IMPACT)**

The IMPACT version 5.4.0 software was used to calculate doses to the potential critical groups using 2016 environmental monitoring data. Where measured environmental data is not available, IMPACT calculates the doses from emissions. IMPACT 5.4.0 is consistent with the method of dose calculation described in the CSA N288.1-08 standard [R-20].

IMPACT 5.5.1, an updated version of IMPACT 5.4.0, has been released and is consistent with the revised CSA N288.1-14 standard [R-56]. IMPACT 5.5.1 has been used to update the DN and PN DRLs. Following implementation of these updated DRLs, IMPACT 5.5.1 will be used for public dose calculations.

**4.1.2 Calculated Atmospheric Dispersion Factors**

Atmospheric dispersion factors (Ka) provide a measure of the dilution of station radiological stack emissions to the atmosphere. Ka values are used to estimate radionuclide concentrations in air at the boundary monitor locations when local measured values are not available. Details of how and when the Ka values are used are provided in Appendix F, Dose Calculation Procedure and Concentrations.

Factors influencing atmospheric dispersion at a specific location include wind speed and direction, as well as the level of turbulence in the atmosphere.

Ka values are calculated from the measured HTO in air concentrations and HTO emissions using the relationship:

$$K_a = C/Q \text{ (s/m}^3\text{)}$$

Where C is the annual average HTO in air concentration (Bq/m<sup>3</sup>) above background measured outside the site boundary, and Q is the average annual HTO release rate (Bq/s) measured by stack monitors at the point of release. The release rate is determined by dividing the total annual emission of HTO as given in Table 2-1 by  $3.16 \times 10^7$  seconds per year.

Ka values have been calculated using HTO in air concentrations from the active samplers at the boundary locations. These values are listed in Tables 4-1 and 4-2 for DN and PN, respectively.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>58 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table 4-1: Darlington Nuclear Annual Boundary Dispersion Factors – 2017**

INDICATOR SITES	Measured Average	
	Airborne Tritium Concentration (Bq/m <sup>3</sup> )	Measured Ka (s/m <sup>3</sup> )
D1 – Southeast Fence	1.27	1.6E-07
D2 – East Fence	1.37	1.8E-07
D5 – Knight Road	0.40	4.8E-08
D9- Courtice WPCP	0.53	6.54E-08
D10 – Holt Road	0.21	2.31E-08
<b>Average</b>		<b>9.5E-08</b>

NOTE: The measured annual HTO to air emission is used together with the measured levels of HTO in the environment to calculate Ka.

**Table 4-2: Pickering Nuclear Annual Boundary Dispersion Factors – 2017**

INDICATOR SITES	Measured Average	
	Airborne Tritium Concentration (Bq/m <sup>3</sup> )	Measured Ka (s/m <sup>3</sup> )
P2 – Montgomery Park Rd.	12.64	5.8E-07
P3 – Sandy Beach Rd.	2.58	1.2E-07
P4 – Liverpool Rd.	1.17	5.2E-08
P6 – East Boundary	6.83	3.1E-07
P10 – Central Maintenance –East	9.10	4.1E-07
P11 – Alex Robertson Park	2.65	1.2E-07
<b>Average</b>		<b>2.6E-07</b>

NOTE: The measured annual HTO to air emission is used together with the measured levels of HTO in the environment to calculate Ka.

#### 4.1.3 Meteorological Data

Wind speed, direction and frequency are measured continuously at meteorological towers at each nuclear site. The average annual wind frequencies at a 10 m height in 2017 for the DN and PN sites are presented in Table 4-3 for 16 wind sectors.

The meteorological data are used in the IMPACT program to model radionuclide concentrations at the potential critical group locations where measured data is not available (such as pathways for I(mfp), Co-60, Cs-137+ and HT). In 2017, the landward wind sector which the wind predominantly blew towards was NW for DN and NNE for PN. Table 4-3 indicates the wind frequencies blowing from each direction.

## Public Information

Document Number:

N-REP-03443-10017

Usage Classification:

Information

Sheet Number:

N/A

Revision Number:

R000

Page:

59 of 132

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

Table 4-3: Darlington and Pickering Nuclear – 2017 Annual Average Wind Frequency by Direction (at 10 m height)

Direction Wind Blowing From	Darlington Nuclear Wind Frequency (%)	Pickering Nuclear Wind Frequency (%)
N	10.27	9.65
NNE	7.06	6.22
NE	3.59	3.82
ENE	2.96	5.48
E	5.90	7.78
ESE	8.02	5.01
SE	<b>8.41</b>	3.38
SSE	3.11	1.49
S	2.38	2.42
SSW	2.23	<b>10.32</b>
SW	1.46	7.48
WSW	4.65	6.46
W	10.66	6.58
WNW	11.42	7.43
NW	8.58	8.80
NNW	9.29	7.68
Total	100.00	100.00

Note: Shaded fields indicate landward wind sectors.  
 Bolded values indicate landward wind sectors with the highest wind frequency.

## 4.2 Critical Group Dose

The calculation of public dose in this report is intended to be realistic, using the potential critical group lifestyles and attributes collected in the DN and PN site-specific surveys [R-30] [R-31] [R-32] [R-33]. The site specific surveys identify the potential critical groups for DN and PN as discussed in Appendix E. Every five years the site specific surveys and pathway analyses are reviewed to ensure the public dose accurately represents the public living near the nuclear generating stations.

Current EMP designs are based on the 2006 site specific survey information. Site specific surveys were updated in 2012 and pathway analyses were updated in 2014, however these did not identify any significant changes with the potential to substantially alter the predictions of the ERAs or the implementation of the EMPs. Therefore, in accordance with CSA N288.4-10 Clause 5.3, no immediate action or change is required to the EMP designs. Recommendations from these studies will be incorporated during the next EMP design revisions.

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>60 of 132</b>

Title:

### **2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

In public dose assessments, “critical groups” are used to estimate the mean realistic impacts of emissions on the most affected individuals. An individual with the average characteristics of the group is known as the “Representative Person” as described in CSA N288.1-08 [R-20]. Dose estimates are calculated for a number of potential critical groups for each site, and for three age classes within each potential critical group; adult, child, and infant. The group and age class with the highest dose is reported as the site public dose for the given year.

Doses are reported for each of the top three potential critical groups at DN and PN, i.e. the three potential critical groups for each site which yield the highest dose estimates based on the pathway analyses. For DN these are the Dairy Farm, the Farm, and the Rural Resident. For PN these are the Industrial/Commercial Worker, the Urban Resident, and the occupants of a Correctional Institute. Additionally, the annual public dose is also calculated for the PN Dairy Farm potential critical group as this group is exposed to the most media types and pathways. Including the Dairy Farm assures that any future changes in emissions, environmental transfer factors, exposure factors, and dosimetry, and changes in the distribution of radionuclides released will be captured. The EMP sampling plan is designed to monitor for these potential critical groups.

For groups that occupy a relatively small geographic area, radionuclide measurements taken at that location are used in the potential critical group calculations. For groups such as the Farm, Dairy Farm or Urban Resident that are spread over much wider geographic areas, air concentrations are determined for a single conservative representative location, and group average values are used for terrestrial samples and water sources.

A small fraction of the adult residents living near DN or PN also work within 5 km of the stations, thereby receiving a different dose while at work and at home. Similarly, a small fraction of the Industrial/Commercial potential critical group workers live near DN or PN station and continue to receive a dose while at home. As a result, the dose estimates for these potential critical groups have been adjusted to account for this portion of the population.

The following sections provide the basis for the dose calculation, results, and interpretation of the public dose for DN and PN. Details on the calculations, how the radionuclide concentrations are determined, background subtractions, and whether data is measured or modeled are provided in Appendix F. Tables of doses calculated for all the potential critical groups are provided in Appendix G, Tables of Public Doses by Radionuclide, Pathway and Age Group for Darlington Nuclear and Pickering Nuclear Potential Critical Groups.

#### **4.2.1 Exposure Pathways**

The dose calculations include all pathways of radionuclide uptake or external exposure by humans, as illustrated previously in Figure 4-1. The dose contribution from each pathway was estimated with IMPACT 5.4.0 either using direct measurements in the environment or by modelling from emissions.



## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>61 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

### 4.2.2 Age Classes

In accordance with CSA N-288.1-08 [R-20], three age classes are used for estimating annual dose to the representative person. The three age classes are 0-5 years (infant), 6-15 years (child), and 16-70 years (adult). The dose estimates to these three age groups are sufficient to characterize doses to the public. For practical implementation in dose calculations, the dose coefficients and characteristics for a one-year old infant, a 10-year old child, and an adult are used to represent the three age classes [R-34].

### 4.2.3 Basis of Dose Calculation

- For each potential critical group, the annual average concentration of each environmental medium sampled from that group is used for the dose calculation with the background subtracted.
- OBT doses from terrestrial animals, plants, and fish are modeled from measured HTO concentrations in terrestrial media and fish.
- Doses from HTO, noble gases, and C-14 in air (where samplers are not at potential critical group locations) are estimated based on measurements at the fence line boundary and applying a calculated air dispersion ratio for the potential critical group location.
- Doses from the remaining radionuclide pathways for I(mfp), Co-60, and HT, are modeled from emissions applying the Ka dispersion factor as well as the calculated air dispersion ratio for the potential critical group location (see Appendix F)

### 4.2.4 Uncertainty in Dose Calculation

As described previously, the public dose estimates use a combination of measured and modeled environmental concentrations of radionuclides. A study was completed through COG to quantify the uncertainties associated with public dose estimation. This study concluded that dose estimates which start with concentration measurements in environmental media for the important exposure pathways, such as OPG's EMP dose estimates, tend to have uncertainties in the order of  $\pm 30\%$  [R-35].

## 4.3 Darlington Nuclear Public Dose

### 4.3.1 Darlington Nuclear Potential Critical Groups

The three potential critical groups at DN for which doses are calculated in this report are shown in Figure C1, Appendix C and are described in Appendix E, Potential Critical Group Descriptions. The potential critical groups and their representative locations are primarily based on the DN site-specific survey review [R-30] and modified, if required, if significant changes occur ahead of the next site-specific survey review.

## Public Information

Document Number:

N-REP-03443-10017

Usage Classification:

Information

Sheet Number:

N/A

Revision Number:

R000

Page:

62 of 132

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

## 4.3.2 Dose Calculation Results

For 2017, the limiting critical group at DN was the Dairy Farm infant, with a dose of 0.7  $\mu\text{Sv}/\text{year}$ , as indicated in Table 4-4.

The Dairy Farm critical group represents dairy farms located within approximately 10 km of the DN site. The closest dairy farm is in the North wind sector about 3 km from the site. Members of this group obtain their water supply from wells and use it for drinking, bathing, irrigation, and livestock watering. They also obtain a large fraction of their annual fruit, vegetable and animal product consumption, including fresh cow's milk, from local sources. Members also consume some locally caught fish and are externally exposed to beach sand at local beaches. The results of the 2017 DN public dose calculation are presented in Table 4-4.

Table 4-4: 2017 Annual Darlington Nuclear Critical Group Doses

Potential Critical Group	Dose per Age Class (microsieverts)		
	Adult	Child (10-year old)	Infant (One-year old)
Dairy Farm Residents	0.4	0.5	<b>0.7</b>
Farm Residents	0.6	0.5	0.4
Rural Residents	0.3	0.2	0.2

Table 4-5 illustrates the dose contribution from each radionuclide for the Dairy Farm infant and percent contribution to the total dose. C-14, HTO, and noble gases contribute over 96% of the total dose.

Table 4-5: 2017 Darlington Nuclear Public Dose

Radionuclide	Dose ( $\mu\text{Sv}/\text{a}$ )	% Dose Contribution
C-14	5.7E-01	75.9%
Co-60	7.7E-04	0.1%
Cs-137+	0.0E+00	0.0%
HT	9.2E-07	0.0%
HTO	8.3E-02	11.1%
Noble Gases	7.2E-02	9.7%
OBT	9.9E-03	1.3%
I (mfp)	1.4E-02	1.9%
Total	7.5E-01	100%

NOTE: "+" indicates that contributions from progeny are included.

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>63 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

This distribution of dose by radionuclides reflects the characteristics of the Dairy Farm group. C-14 dose is mostly from ingestion of terrestrial plants and animal products. The Dairy Farm one-year old infant consumes animal products that are almost entirely from local sources, including milk from local cows, as well as a portion of its fruits and vegetables. Dose from HTO is attributed to air inhalation and ingestion of local terrestrial plants and animal products. The public dose trend for DN is presented on a logarithmic scale in Figure 4-2. The DN dose remains essentially unchanged over the last ten years and is below 1% of the legal limit.

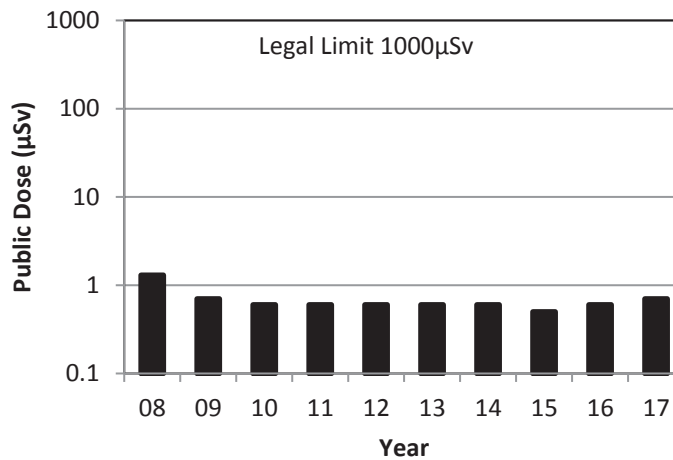


Figure 4-2: Darlington Nuclear Annual Public Dose Trend

**4.3.3 Discussion of Results**

The 2017 DN site public dose of 0.7 µSv, as represented by the Dairy Farm infant, is about 0.1% of the 1,000 µSv/year legal limit for a member of the public. The DN dose for 2017 is essentially unchanged from the 2016 site dose of 0.6 µSv for the Dairy Farm infant. The critical group has remained unchanged.

The DN dose for 2017 is less than 0.1% of the estimated average background dose around DN, from naturally occurring and anthropogenic (man-made) radiation, of about 1,400 µSv/year (excluding medical doses, refer to Section 4.5). Figure 4-3 is a graphical representation of critical group dose compared to background radiation around DN. As an additional source of comparison, Table 4-8 provides examples of typical doses from exposure to natural and anthropogenic sources.

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>64 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

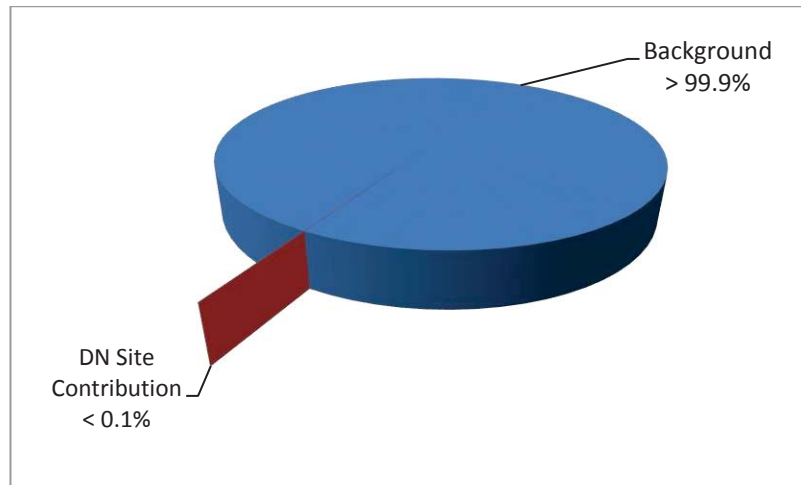


Figure 4-3: Comparison of Darlington Nuclear Public Dose to Background Dose

#### 4.4 Pickering Nuclear Public Dose

##### 4.4.1 Pickering Nuclear Potential Critical Groups

The four potential critical groups at PN for which doses are calculated in this report are shown in Figure C2, Appendix C and are described in Appendix E. The potential critical groups and their representative locations are primarily based on the site-specific survey review conducted in 2005 [R-31] and modified, if required, if significant changes occur ahead of the next site-specific review cycle.

##### 4.4.2 Dose Calculation Results

For 2017, the limiting critical group at PN was the Urban Resident adult, with a dose of 1.8  $\mu$ Sv/year, as indicated in Table 4-6.

The Urban Resident critical group consists of Pickering and Ajax residents in the areas surrounding the PN site. Members of this group drink mostly water from Ajax WSP and also consume a diet comprised in part of locally grown produce and some locally caught fish. Members of this group are also externally exposed to beach sand at local beaches.

A fraction of adult residents within the Urban Resident critical group also work within 5 km of PN station and receive some dose from the station while at work. The average dose for the Urban Resident Adult has been adjusted to account for these residents.

The results of the 2017 PN public dose calculation are presented in Table 4-6.

## Public Information

Document Number:

N-REP-03443-10017

Usage Classification:

Information

Sheet Number:

N/A

Revision Number:

R000

Page:

65 of 132

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

Table 4-6: 2017 Annual Pickering Nuclear Critical Group Doses

Potential Critical Group	Dose per Age Class (microsieverts)		
	Adult	Child (10-year old)	Infant (One-year old)
Dairy Farm Residents	0.6	0.6	0.8
Urban Residents	1.8	1.7	1.7
C2 Correctional Institution	1.2	1.3	
Industrial Workers	1.5		

Table 4-7 illustrates the dose from each radionuclide and percent contribution to the total dose. HTO and noble gases contribute over 95% of the total dose.

Table 4-7: 2017 Pickering Nuclear Public Dose

Radionuclide	Dose ( $\mu\text{Sv/a}$ )	% Dose Contribution
C-14	1.1E-02	0.6%
Co-60	4.1E-02	2.3%
Cs-137+	3.1E-02	1.7%
HTO	6.1E-01	33.8%
Noble Gases	1.1E+00	61.6%
OBT	2.2E-03	0.1%
I (mfp)	3.2E-05	0.0%
Total	1.8E+00	100%

NOTE: "+" indicates that contributions from progeny are included.

This distribution of dose by radionuclides reflects the characteristics of the Urban Resident group since their exposure is mainly from inhalation of HTO and external exposure to noble gases. The public dose trend for PN is presented on a logarithmic scale in Figure 4-4. The PN dose remains below 1% of the legal limit.

The reduction in dose from 2008 to 2009 is primarily attributed to changes in methodology and transfer parameters specified by CSA N288.1-08 [R-36].

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>66 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

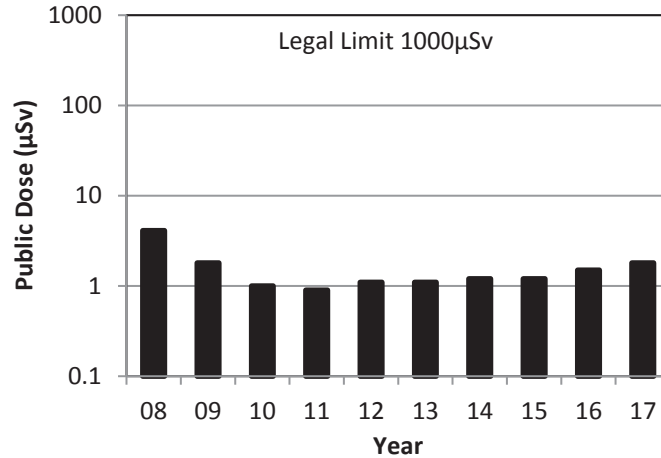


Figure 4-4: Pickering Nuclear Annual Public Dose Trend

4.4.3 Discussion of Results

The 2017 PN site public dose of 1.8 µSv, as represented by the Urban Resident adult, is 0.2% of the 1,000 µSv/year legal limit for a member of the public. The PN dose for 2017 is slightly higher than the 2016 site dose, as consistent with emissions trends. The critical group has remained unchanged.

The PN dose for 2017 was equivalent to 0.1% of the estimated background dose around PN of 1,400 µSv/year, from naturally occurring and anthropogenic (man-made) radiation (excluding medical doses, refer to Section 4.5). Figure 4-5 is a graphical representation of critical group dose compared to background radiation around PN. As an additional source of comparison, Table 4-8 provides examples of typical doses from exposure to natural and anthropogenic sources.

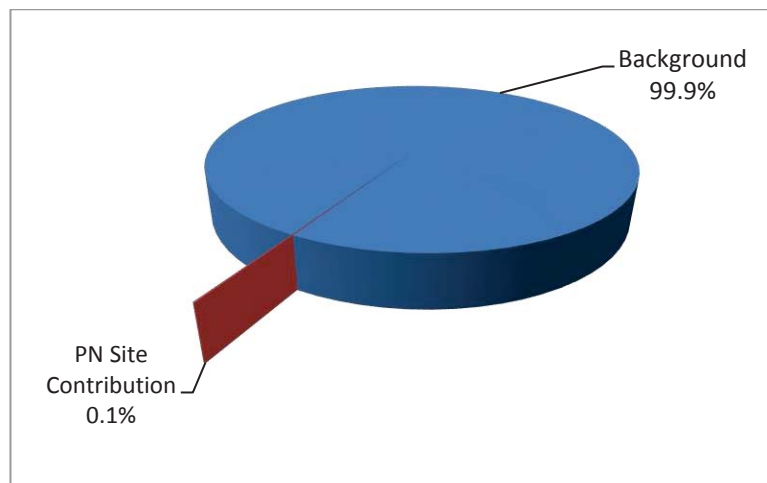


Figure 4-5: Comparison of Pickering Nuclear Public Dose to Background Dose

## Report

Public Information		
Document Number:	<b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>
Sheet Number:	<b>N/A</b>	Revision Number: <b>R000</b>
		Page: <b>67 of 132</b>

Title:

## 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

### 4.5 Natural and Anthropogenic Data

Table 4-8 provides some typical doses received by members of the public from exposure to natural and anthropogenic sources.

**Table 4-8: Typical Doses from Exposure to Natural and Anthropogenic Sources**

Source of Exposure	Effective Dose ( $\mu\text{Sv}$ )
Annual External Exposure during Precipitation Events (Gamma Radiation from Naturally Occurring Radon Gas Decay Products) [R-37]	4
Chest X-Ray (single film) [R-38]	10
Airplane Travel (two hour flight) [R-39]	12

Information on Canadian public doses from naturally occurring sources, including data from ground gamma surveys in four major Canadian cities, was provided in 2002 [R-40] [R-41]. Results are summarized in Table 4-9, where it can be seen that most of the variation is due to the inhalation dose from Radon-222 (Rn-222).

**Table 4-9: Naturally Occurring Annual Public Effective Doses**

Radiation Source	Worldwide Average ( $\mu\text{Sv}$ )	Canada ( $\mu\text{Sv}$ )	Toronto ( $\mu\text{Sv}$ )	Montreal ( $\mu\text{Sv}$ )	Winnipeg ( $\mu\text{Sv}$ )	Pickering Nuclear Site ( $\mu\text{Sv}$ )	Darlington Nuclear Site ( $\mu\text{Sv}$ )
Cosmic	380	318	313	313	315	313	313
Internal	306	306	306	306	306	306	306
Inhalation <sup>(a)</sup>	1,256	926	757	667	3,225	565	565
External	480	219	178	278	176	154	154
<b>Total<sup>(b)</sup></b>	<b>2,400</b>	<b>1,800</b>	<b>1,600</b>	<b>1,600</b>	<b>4,000</b>	<b>1,300</b>	<b>1,300</b>

(a) Mostly from Rn-222.

(b) Total doses have been rounded to two significant figures to reflect the inherent uncertainty. Some components are based on direct measurements and others are estimated from related measurements.

In addition to naturally occurring radiation, the public also receives about 70  $\mu\text{Sv}/\text{year}$  effective dose from anthropogenic sources such as nuclear weapon test fallout, and exposures from technological processes and consumer products and services, excluding medical sources. Thus, the total background dose around PN and DN from naturally occurring and anthropogenic sources is 1,400  $\mu\text{Sv}/\text{year}$ . Furthermore, the average Canadian dose from medical sources averages 1,100  $\mu\text{Sv}/\text{year}$  per person. The legal limit of 1,000  $\mu\text{Sv}$  per year from licensed industrial practices is over and above the dose the public already receives from the natural environment and from medical procedures [R-42].

### 5.0 QUALITY ASSURANCE AND PERFORMANCE

The Quality Assurance (QA) program for the EMPs encompasses all activities from sample collection, laboratory analysis, laboratory quality control and external laboratory comparison, to program audits, self-assessments, and dose verifications. The objectives include ensuring that EMP samples are representative and their

Public Information		
Document Number: <b>N-REP-03443-10017</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>68 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

analytical results are accurate such that best estimates of radiation doses to the public can be provided, as well as complying with procedures and program quality requirements. This section provides an overview of quality assurance activities that are critical to ensuring the quality of the EMP data and processes.

**5.1 Laboratory Quality Assurance and Quality Control**

The OPG Health Physics Laboratory (HPL) is accredited for radioanalysis of drinking water and soil by the Canadian Association for Laboratory Accreditation (CALA). The accreditation is based on demonstrated compliance with ISO 17025, General Requirements for the Competence of Testing and Calibration Laboratories. HPL is also licensed for radioanalysis of drinking water by the Province of Ontario’s Ministry of Environment and Climate Change. HPL performs laboratory activities in accordance with the OPG Dosimetry and Radiological Environmental Measurement Services Quality Assurance Manual [R-43].

**5.1.1 Laboratory Quality Control**

Quality control (QC) samples are used to estimate the precision and accuracy of analytical results and to examine any sources of error introduced by laboratory practices which require corrective actions. Two types of QC samples are used to accompany the analyses of the environmental samples collected for the EMPs:

- (a) Process control samples are ‘dead water’ (radiation-free water/blank) samples that go through the same handling process as the real samples.
- (b) QC standards are samples with predetermined values (usually traceable standards) that go through the same handling process as the real samples. The analysis of the environmental sample is considered valid when the results of the accompanying QC samples are within the expected set limits, depending on the analysis type.

For 2017, the results for the QC samples were all within the required range. These results provide confidence in the quality of data for the program and the consistency of laboratory measurements.

**5.1.2 Laboratory Performance Testing**

The main purpose of the laboratory performance testing programs is to provide assurance to OPG Nuclear and the CNSC of the laboratory’s analytical proficiency (i.e., the accuracy of the measurements). The testing programs provide a quality check to laboratory operations including equipment calibration, analytical procedures, data review and internal QC. These testing programs are a crucial part of the laboratory QA program to demonstrate that the laboratory is performing within the acceptable limits as measured against external unbiased standards.

For 2017, OPG Nuclear participated in a laboratory performance testing program that included the measurement of tritium in water, gross beta in water, and gamma emitters in water, soil and milk.



## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>69 of 132</b>

Title:

### 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

QA test samples were supplied on a quarterly basis by Eckert and Ziegler Analytics [R-44]. Results of analyses were reported back to Eckert and Ziegler Analytics who then provide performance reports for each of the analytical types. The performance test limits were as follows:

$$-25\% \leq \text{Relative Difference} \leq +50\%$$

$$\text{Relative Precision} \leq 40\%$$

These test limits are adapted from the *in vitro* accuracy specifications of the CNSC's Regulatory Standard S-106 Revision 1, Technical and Quality Assurance Requirements for Dosimetry Services [R-45].

All QA performance test results in 2017 met the specified limits. The maximum and minimum Relative Difference and Relative Precision are summarized for each sample type and presented in Table 5-1.

**Table 5-1: Summary of Analytics Performance Test Results – 2017**

Sample Types	Relative Difference (%)		Relative Precision (%)	
	High	Low	High	Low
Tritium in Water	1	-2	2	0
Gross Beta in Water	5	-4	9	7
Gamma in Water	35	0	12	2
Gamma in Soil	14	-7	7	1
Gamma in Milk	22	-15	5	1

## 5.2 Equipment Calibrations/Maintenance

Equipment calibrations and maintenance are conducted in accordance with the Environmental Monitoring Program Equipment Maintenance Manual [R-46].

In addition, annual sensitivity checks are performed on the noble gas detectors to quantify the sensitivity of the sodium iodide crystal in each detector. The 2017 results indicate that detectors are functioning at acceptable levels of sensitivity [R-47].

## 5.3 Program Quality Assurance

### 5.3.1 Audits

An independent audit, also referred to as a performance based assessment, of the EMPs is conducted once every five years in accordance with CSA N288.4-10 [R-2]. The last audit of the EMPs was performed in 2014 by OPG's Nuclear Oversight department.

The OPG HPL also has a commitment to perform a minimum of one independent audit each year of the quality system used for dosimetry and environmental measurement

## Report

Public Information		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>70 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

services. These may not always be related to the EMPs. In 2017, an HPL QA audit was conducted record handling in the Environment duty area. There were no significant adverse findings or conditions arising from this self-assessment that affected the quality of results and measurements in the dosimetry and environmental laboratory. Minor recommendations have been assigned and will be tracked to completion [R-44].

### 5.3.2 Self-Assessments

In 2017, two self-assessments were performed on different elements of the EMPs.

#### (a) Field Verification of both Monthly and Quarterly Sample Collection

The focus of this self-assessment was to observe OPG HPL staff collect monthly and quarterly sample collection for EMP in accordance with approved program procedures. The field verification was conducted via direct observation of sample collection in the field, and confirmed that HPL's procedural use and adherence to collection procedures is satisfactory. This self-assessment is documented in the OPG Self Assessment Database under NO17-001843-SA.

#### (b) Annual Performance Assessment

Self-assessment LEC17-001852-SA was completed for the EMP Annual Performance Assessment. The assessment confirmed that all EMP design objectives were met. Required equipment repairs and maintenance to EMP stations have been carried out. Revisions required for various EMP procedures and documents were completed.

### 5.4 Third-Party Verification of Annual EMP Report

An independent third-party verification of the annual dose calculations and this report was carried out by EcoMetrix Incorporated. Verification was done on the methodology used, assumptions made, input parameter values and data used. This involved checking the dose calculations, IMPACT scenarios, and performing independent replicate IMPACT model runs and hand calculations to validate the results obtained by OPG. Any necessary changes identified by the third-party verification have been addressed and incorporated in this report.

### 5.5 Program Performance

#### 5.5.1 Sample Unavailability

A total of 979 laboratory analyses were performed for the 2017 dose calculation. The analyses covered HTO, C-14, and gamma scan. The PN site accounted for 35% of these sample analyses, while the DN and provincial-background programs accounted for 50% and 15% respectively. Table 5-2 shows the sample types, number of locations, number of samples used for the dose calculation, and the unavailability of each sample type.

**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-10017</b>		Usage Classification: <b>Information</b>
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>71 of 132</b>

<b>Title:</b> <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
---

The unavailability indicator tracks the performance of sample collection and analysis for the EMPs. The sampling portion of the EMPs is designed to collect representative field samples from selected pathways near each nuclear site and from background locations, in order to meet the program objectives as defined in Section 1.1. The sample unavailability percentage is determined by dividing the number of missed or invalid sample analyses by the number of planned sample analyses for each EMP site.

An important objective of the EMP is to estimate the doses to the public based on environmental data measured in the public domain. In accordance with the EMP governing document [R-48], the requirement to meet unavailability limits is specific to the analysis of samples used in the dose calculation. These limits are applied to the PN, DN and provincial-background EMPs separately.

The unavailability limits for samples used in the dose calculation are provided in Table 5-2 and range from 10 to 25%. The unavailability limits were derived based on the relative contributions to total dose, therefore higher dose contributors have a lower unavailability limit. The overall unavailability for PN, DN and provincial-background EMPs was 6%, 5% and 5%, respectively.

The unavailability limit of 20% for PN fruits was exceeded in 2017. Available samples adequately represented the Dairy Farm and Farm critical groups. Urban Resident critical group did not have a representative sample of fruit, as over 75% of the planned samples were unavailable. For dose calculation purposes, PN dose from fruit was conservatively modeled in IMPACT in 2017 for the Urban Resident critical group.

The unavailability limit of 20% for DN vegetables was exceeded in 2017 due to several locations not growing vegetable gardens in 2017, or vegetables samples not being available due to poor growing season. In 2017, available samples were not adequate to represent critical groups. Therefore, for purpose of the dose calculation, DN dose from vegetables was conservatively modeled in IMPACT in 2017.

Though not exceeded, the unavailability limit of 20% for provincial background fruits and vegetables was reached in 2017. This was due to the fact that fruit and vegetable samples were not available at the Barrie location in 2017. Eighty percent of provincial fruit and vegetables were available and considered representative of background locations. Therefore, these samples were used in the dose calculation.

Report

Public Information			
Document Number: <b>N-REP-03443-1001</b>		Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>		Page: <b>72 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table 5-2: Unavailability of EMP Sample Data Used for Dose Calculation Purposes**

Sample Types	Collection Frequency	Pickering Nuclear				Darlington Nuclear				Provincial Background				Unavailability Limit <sup>(d)</sup>
		Locations	Planned Analyses	Actual Analyses	Unavailability	Locations	Planned Analyses	Actual Analyses	Unavailability	Locations	Planned Analyses	Actual Analyses	Unavailability	
<b>Tritium</b>														
Tritium in Air (Molecular Sieve)	Monthly/Quarterly	6	72	72	0%	6	72	70	3%	1	12	12	0%	10%
Water Supply Plants	Weekly Composite	1	48	48	0%	2	96	96	0%					15%
Residential Wells	Monthly	2	24	24	0%	4	48	47	2%					15%
Milk	Monthly	2	24	24	0%	3	36	36	0%					25%
Milk	Quarterly									1	12	12	0%	25%
Lake Water	Monthly <sup>(a)</sup>	3	24	24	0%	2	16	16	0%					25%
Fruits	Annual	5	15	7	53%	5	15	15	0%	5	10	8	20%	20%
Vegetables	Annual	5	15	15	0%	7	21	10	52%	5	10	8	20%	20%
Animal Feed	Annual	1	8	8	0%	4	16	16	0%	1	8	8	0%	25%
Poultry	Annual					1	8	8	0%	1	8	8	0%	25%
Eggs	Quarterly					1	12	12	0%	1	12	12	0%	25%
Fish	Annual	1	8	8	0%	1	8	8	0%					25%
<b>Carbon-14</b>														
Carbon-14 in Air	Quarterly	4	16	16	0%	4	16	16	0%	1	4	4	0%	25%
Milk	Monthly	2	24	24	0%	3	36	36	0%					10%
Milk	Quarterly									1	12	12	0%	25%
Fruits	Annual	5	15	7	53%	5	15	15	0%	5	10	8	20%	20%
Vegetables	Annual	5	15	15	0%	7	21	10	52%	5	10	8	20%	20%
Animal Feed	Annual	1	8	8	0%	4	16	16	0%	1	8	8	0%	25%
Poultry	Annual					1	8	8	0%	1	8	8	0%	25%
Eggs	Quarterly					1	12	12	0%	1	12	12	0%	25%
Fish	Annual	1	8	8	0%	1	8	8	0%	1	8	8	0%	25%
<b>Noble Gases</b>														
External Gamma (Noble Gas Monitors) <sup>(b)</sup>	Continuous	6	NA	NA	1%	5	NA	NA	2%					25%
<b>Gamma</b>														
Fish	Annual	1	8	8	0%	1	8	8	0%	1	8	8	0%	25%
Beach Sand	Annual	3	24	24	0%	3	24	24	0%	1	8	8	0%	25%
<b>Overall dose sample Unavailability <sup>(c)</sup></b>			356	340	6%		512	487	5%		160	152	5%	

Notes: NA = Not Applicable.

(a) For safety considerations, samples are not required during the winter months (Dec. - Mar.).

(b) Noble gas detector unavailability is based on an average of actual run time of all monitors for PN and DN.

(c) Unavailability defined as an average of the percent unavailability of all sample types.

(d) Unavailability limit for all Provincial samples types is 25%.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>73 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****5.6 Annual Assessment of the EMPs**

The annual assessment of OPG's 2017 EMPs is summarized as follows:

- Overall, the EMPs met their objectives as defined in Section 1.1.
- A total of 979 environmental data analyses were completed for samples collected around DN and PN sites and at various Ontario background locations in support of the radiological dose calculations. The overall unavailabilities were 6%, 5%, and 5% for the PN, DN, and provincial-background EMPs, respectively.
- Two self assessments were completed this year for the EMPs. No significant findings were identified.
- An independent third-party verification of the annual dose calculations and this report was carried out by EcoMetrix Incorporated.

**5.6.1 Summary of Darlington Results**

- Site emissions remained at very small fractions of their respective DRLs.
- Boundary noble gas detector dose rates remained below detection limits.
- As indicated in Figure 2-1, the HT emissions from DN have remained at low levels. In 2017, the HT emissions were  $1.4 \times 10^{14}$  becquerels (Bq). As of 2017, these emissions include HT emissions from the powerhouse. However, the increase in elemental tritium emissions observed in 2017 is primarily attributed to a valve that was inadvertently opened and vented to the TRF stack on May 8, 2017. Corrective actions were immediately taken and a procedure update was subsequently initiated to rectify the deficiencies.
- A follow-up monitoring program to monitor entrainment at DN was completed in 2016. A conservative estimate of 589 kg of biomass was entrained during the study. Additionally, the study concluded that DN does not negatively impact local benthic invertebrate populations.
- In 2016, OPG completed an effluent characterization study at DN to confirm EA findings of non-significant impact from non-radiological effluent constituents on human and non-human biota. The results adequately characterized effluent at DN and were used in the 2016 DN ERA.
- Annual average tritium concentrations in drinking water from the nearby water supply plants were well below OPG's commitment of 100 Bq/L. The annual average HTO activity in well water was 10.6 Bq/L.
- Soil samples had Cs-137 at levels within the range of values seen at the background locations and had no detectable Co-60 or Cs-134. There is no indication of buildup of activity in soil.

## Report

Public Information		
Document Number:	Usage Classification:	
<b>N-REP-03443-1001</b>	<b>Information</b>	
Sheet Number:	Revision Number:	Page:
<b>N/A</b>	<b>R000</b>	<b>74 of 132</b>

Title:

### **2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- Concentrations of HTO and C-14 in air, vegetation, milk, and fish and Cs-137 in fish were in line with levels seen over the last ten years. Eggs and poultry sampling resulted in concentrations for HTO and C-14 that were similar to those in 2016.
- The 2017 public dose for the DN site was 0.7  $\mu$ Sv and was represented by the infant of the Dairy Farm critical group. The 2017 site public dose remains a small fraction of both the annual legal dose limit and the annual natural background radiation in the area.

#### **5.6.2 Summary of Pickering Results**

- Site emissions remained at a very small fraction of their respective DRLs.
- The PN waterborne HTO emissions remain stable. The PN tritium to water emission in 2017 was  $3.8 \times 10^{14}$  Bq. The slight increase observed in 2017 is attributed to a leak in the Unit 5 moderator pit. Tritiated water from the moderator room was processed and discharged through the active liquid waste system. Sealing and repair work to the moderator pit was completed in April 2017.
- A supplementary study was conducted which confirmed that the air kerma rate due to the waste storage facilities cannot be detected at distances greater than 400 m from the facilities and that the skyshine dose from this source is, therefore, not significant for potential critical groups outside the 1 km boundary.
- A supplementary study on soil was conducted to help reduce uncertainty regarding concentration of COPCs used in dose calculations for non-human biota at PN. The study concluded that accumulation of COPCs in soil over time is not expected.
- A supplementary study was conducted in 2015 to collect sediment and water samples from the northern section of Frenchman's Bay wetland. The study aimed to reduce uncertainty regarding the assessment of biota in the bay and concluded that receptor populations are not expected to experience any adverse effects due to non-radiological releases from PN operations.
- Annual average tritium concentrations in drinking water from the nearby water supply plants were below OPG's commitment of 100 Bq/L. The annual average HTO activity in well water was 14.7 Bq/L.
- Soil samples had Cs-137 at levels within the range of values seen at the background locations and had no detectable Co-60 or Cs-134. There is no indication of buildup of activity in soil.
- Concentrations of HTO and C-14 in air, vegetation, milk, and fish, and Cs-137 in fish were in line with levels seen over the last ten years.

## Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>75 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

- The 2017 public dose for the PN site was 1.8 µSv and was represented by the adult of the Urban Resident group. The 2017 site public dose remains a small fraction of both the annual legal dose limit and the annual natural background radiation in the area.

### 6.0 OUTLOOK FOR 2018

The most recent program design reviews on the PN and DN EMPs were issued in 2015. The design reviews incorporated the most recent ERA results, updated pathway analyses, and the results of the latest site specific surveys. However these reviews did not identify any significant changes with the potential to substantially alter the predictions of the ERAs or the implementation of the EMPs. Therefore, in accordance with N288.4-10 Clause 5.3, no immediate action or change is required to the EMP designs. Recommendations from these studies will be incorporated into the EMPs following implementation of the revised DRLs and incorporation of CSA N288.1-14 into the public dose calculations. CSA N288.1-14 compliant PN and DN DRLs were calculated in 2016 and are undergoing review.

An updated DN ERA was completed in 2016 and an updated PN ERA was completed in 2017 in support of PN's licence application. Changes to the EMP as a result of the latest ERA will be identified and captured in the next EMP design review, which will be undertaken in 2018.



## Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>76 of 132</b>

Title:

### 2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS

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**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>77 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

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Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>78 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

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Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>79 of 132</b>

Title:

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<b>Public Information</b>		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>80 of 132</b>

Title:

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Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>81 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

### Appendix A: Radiological Units and Conversions

#### Absorbed Dose

1 gray (Gy) = 1 joule/kg

#### Effective Dose

1 sievert (Sv) = 100 rem  
1 millisievert (mSv) = 100 millirem (mrem)  
1 microsievert ( $\mu$ Sv) = 0.1 millirem (mrem)

#### Quantity of Radionuclide

1 becquerel (Bq) = 1 disintegration per second  
1 curie (Ci) =  $3.7 \times 10^{10}$  Bq  
1 mCi/(km<sup>2</sup>·month) = 37 Bq/(m<sup>2</sup>·month)

**Report**

Public Information		
Document Number:	Usage Classification:	
<b>N-REP-03443-1001</b>	<b>Information</b>	
Sheet Number:	Revision Number:	Page:
<b>N/A</b>	<b>R000</b>	<b>82 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS****Appendix B: Glossary of Acronyms and Symbols****Radionuclides and Units of Measure**

<b>Ar-41</b>	Argon-41
<b>C-14</b>	Carbon-14
<b>CO<sub>2</sub></b>	Carbon Dioxide
<b>Co-60</b>	Cobalt-60
<b>Cs-134</b>	Cesium-134
<b>Cs-137</b>	Cesium-137
<b>Cs-137+</b>	Cesium-137 including progeny
<b>HT</b>	Elemental Tritium
<b>HTO</b>	Tritium Oxide
<b>I(mfp)</b>	Mixed Fission Products Radioiodines
<b>I-131</b>	Iodine-131
<b>Ir-192</b>	Iridium-192
<b>K-40</b>	Potassium-40
<b>Rn-222</b>	Radon-222
<b>Xe-133</b>	Xenon-133
<b>Xe-135</b>	Xenon-135
<b>µGy</b>	microgray
<b>µSv</b>	microsievert
<b>Bq</b>	becquerel
<b>Bq/kg-C</b>	becquerels per kilogram carbon
<b>Ci</b>	Curie
<b>Gy</b>	Gray
<b>kg</b>	kilogram
<b>L</b>	Litre
<b>mGy</b>	milligray
<b>mSv</b>	millisievert
<b>nGy</b>	nanogray
<b>Sv</b>	Sievert

**Acronyms and Abbreviations**

<b>ACU</b>	Air Conditioner Unit
<b>BAF</b>	Bioaccumulation Factor
<b>CALA</b>	Canadian Association for Laboratory Accreditation
<b>CANDU</b>	Canada Deuterium Uranium
<b>CCW</b>	Condenser Cooling Water
<b>CNSC</b>	Canadian Nuclear Safety Commission
<b>COG</b>	CANDU Owners Group
<b>CSA</b>	Canadian Standards Association
<b>DN</b>	Darlington Nuclear
<b>DRL</b>	Derived Release Limit
<b>DWMF</b>	Darlington Waste Management Facility
<b>E</b>	East wind sector
<b>EA</b>	Environmental Assessment
<b>EcoRA</b>	Ecological Risk Assessment
<b>EMP</b>	Environmental Monitoring Program

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>83 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

<b>ENE</b>	East North East wind sector
<b>EPA</b>	Environmental Protection Agency
<b>ERA</b>	Environmental Risk Assessment
<b>ESE</b>	East South East wind sector
<b>FDS</b>	Fish Diversion System
<b>FPS</b>	Fixed Point Surveillance
<b>HC</b>	Health Canada
<b>HPL</b>	Health Physics Laboratory
<b>IAEA</b>	International Atomic Energy Agency
<b>ICRP</b>	International Commission on Radiological Protection
<b>IMPACT</b>	Integrated Model for Probabilistic Assessment of Contaminant Transport
<b>ISO</b>	International Organization for Standardization
<b>Ka</b>	Atmospheric Dispersion Factor ( $s/m^3$ )
<b>Kerma</b>	Kinetic Energy Released in Matter
<b>Lc</b>	Critical Level ( $\approx 0.5Ld$ )
<b>Ld</b>	Limit of Detection
<b>MOECC</b>	Ministry of Environment and Climate Change
<b>MOEE</b>	Ministry of Environment and Energy
<b>MOU</b>	Memorandum of Understanding
<b>MW</b>	Megawatts
<b>N</b>	North wind sector
<b>NaI</b>	Sodium Iodide
<b>NE</b>	North East wind sector
<b>NND</b>	New Nuclear at Darlington
<b>NNE</b>	North North East wind sector
<b>NNW</b>	North North West wind sector
<b>NW</b>	North West wind sector
<b>OBT</b>	Organically Bound Tritium
<b>ODS</b>	Ozone Depleting Substances
<b>OPG</b>	Ontario Power Generation
<b>PHC</b>	Petroleum Hydrocarbon
<b>PN</b>	Pickering Nuclear
<b>PWMF</b>	Pickering Waste Management Facility
<b>PWQO</b>	Provincial Water Quality Objective
<b>QA</b>	Quality Assurance
<b>QC</b>	Quality Control
<b>QOR</b>	Quarterly Operations Report
<b>REMP</b>	Radiological Environmental Monitoring Program
<b>S</b>	South wind sector
<b>SE</b>	South East wind sector
<b>SOR</b>	Statement of Requirements
<b>SSE</b>	South South East wind sector
<b>SSW</b>	South South West wind sector
<b>SW</b>	South West wind sector
<b>TOC</b>	Total Organic Carbon
<b>TRC</b>	Total Residual Chlorine
<b>TRCA</b>	Toronto and Region Conservation Authority
<b>TRF</b>	Tritium Removal Facility



**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>84 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- TRS**            Technical Report Series
- TRV**            Toxicity Reference Value
- TWh**            Terawatt Hour
- UFDS**           Used Fuel Dry Storage
- VOC**            Volatile Organic Compounds
- VBO**            Vacuum Building Outage
- W**                West wind sector
- WNW**           West North West wind sector
- WPCP**          Water Pollution Control Plant
- WSP**            Water Supply Plant

**Report**

<b>Public Information</b>		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>85 of 132</b>

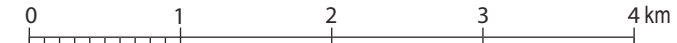
Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Appendix C: Maps of Environmental Monitoring and Critical Group Locations**

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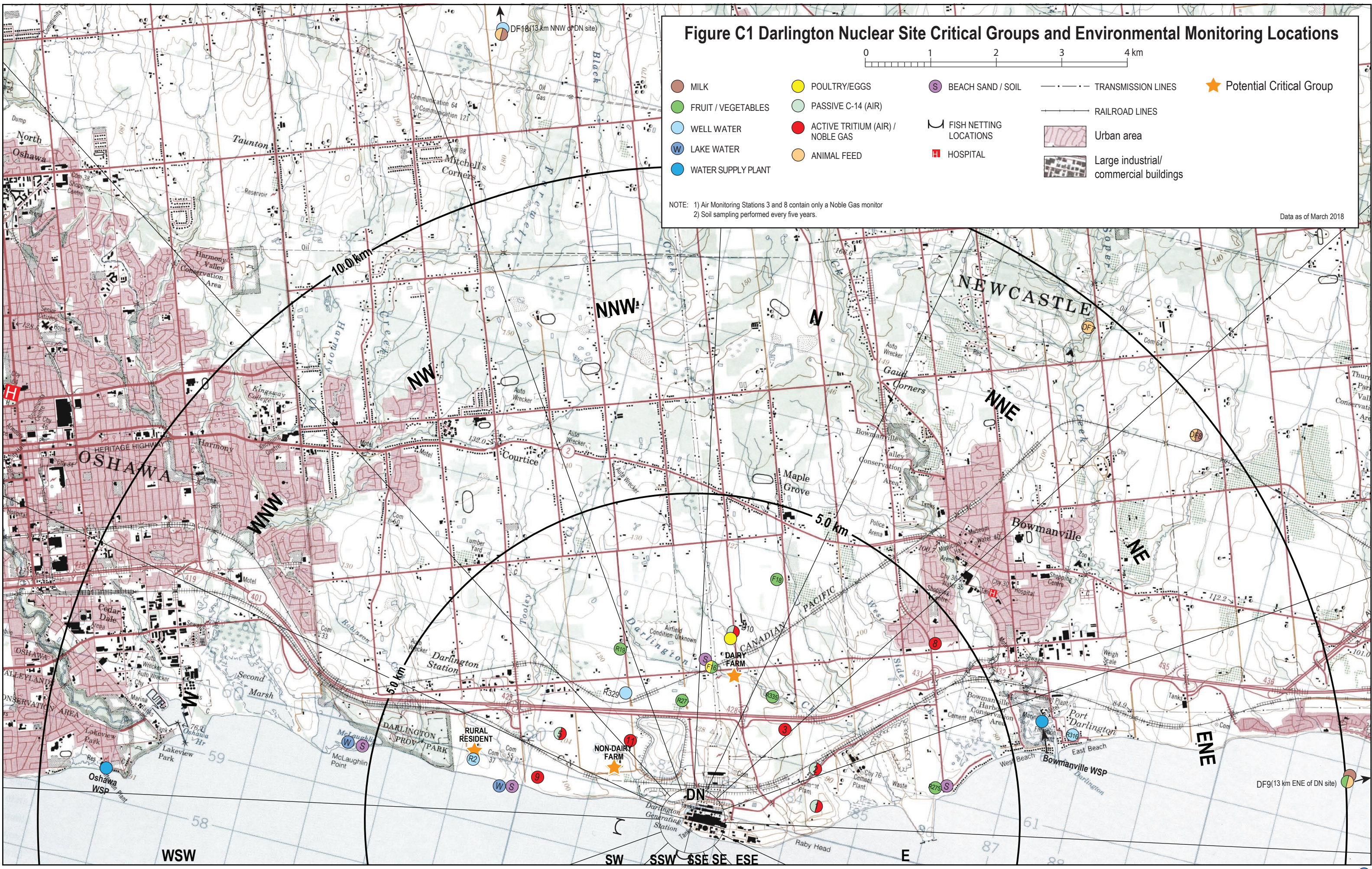
**Figure C1 Darlington Nuclear Site Critical Groups and Environmental Monitoring Locations**



- MILK
- POULTRY/EGGS
- BEACH SAND / SOIL
- TRANSMISSION LINES
- Potential Critical Group
- FRUIT / VEGETABLES
- PASSIVE C-14 (AIR)
- ACTIVE TRITIUM (AIR) / NOBLE GAS
- RAILROAD LINES
- WELL WATER
- ANIMAL FEED
- LAKE WATER
- HOSPITAL
- WATER SUPPLY PLANT
- FISH NETTING LOCATIONS
- Urban area
- Large industrial/commercial buildings

NOTE: 1) Air Monitoring Stations 3 and 8 contain only a Noble Gas monitor  
 2) Soil sampling performed every five years.

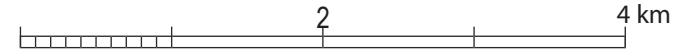
Data as of March 2018



Newcastle WSP (13 km ENE of DN site)



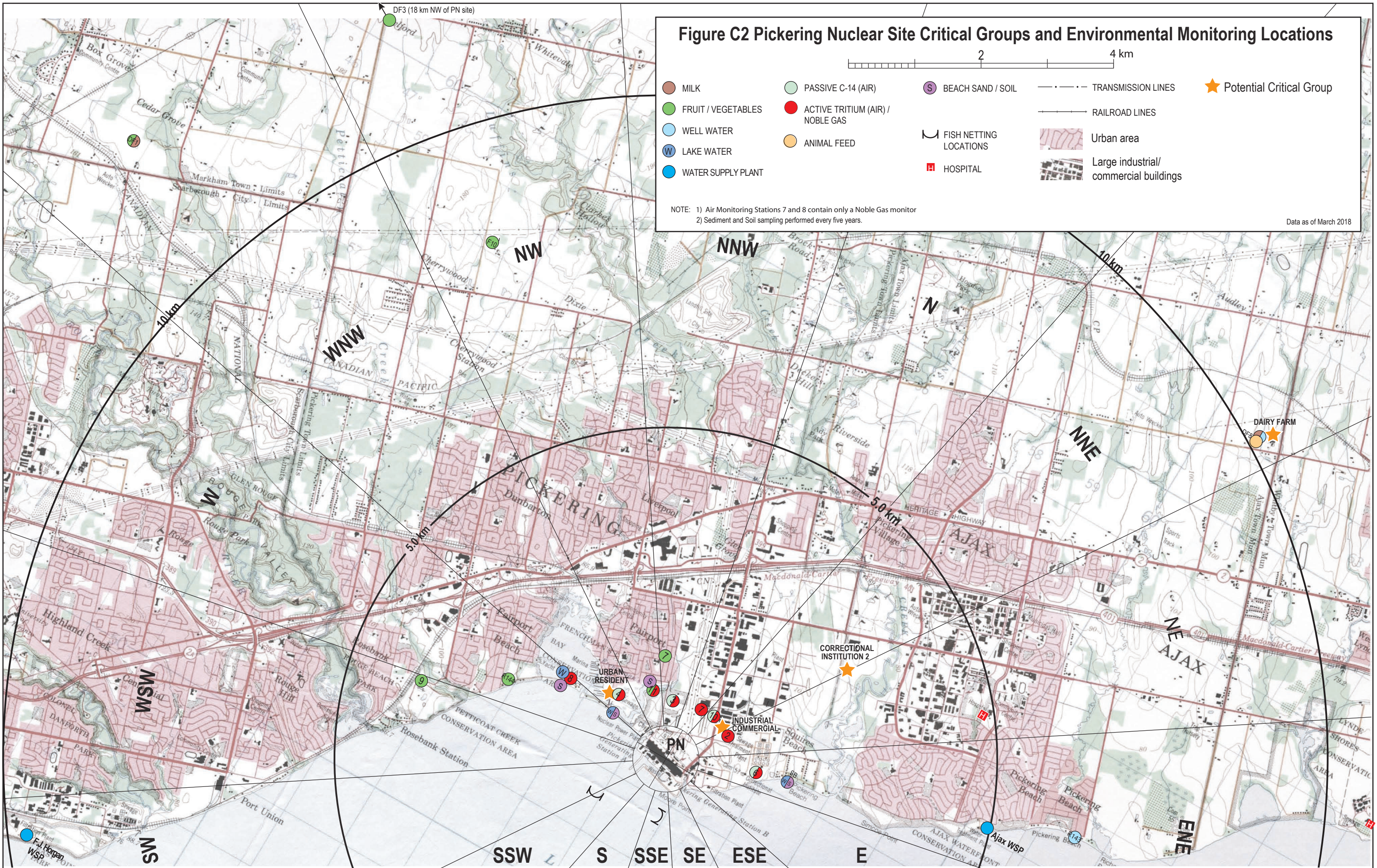
**Figure C2 Pickering Nuclear Site Critical Groups and Environmental Monitoring Locations**



- |                    |                                  |                    |                        |                                       |
|--------------------|----------------------------------|--------------------|------------------------|---------------------------------------|
| MILK               | PASSIVE C-14 (AIR)               | BEACH SAND / SOIL  | TRANSMISSION LINES     | Potential Critical Group              |
| FRUIT / VEGETABLES | ACTIVE TRITIUM (AIR) / NOBLE GAS | ANIMAL FEED        | RAILROAD LINES         | Urban area                            |
| WELL WATER         | LAKE WATER                       | WATER SUPPLY PLANT | FISH NETTING LOCATIONS | Large industrial/commercial buildings |
| LAKE WATER         |                                  | HOSPITAL           |                        |                                       |

NOTE: 1) Air Monitoring Stations 7 and 8 contain only a Noble Gas monitor  
 2) Sediment and Soil sampling performed every five years.

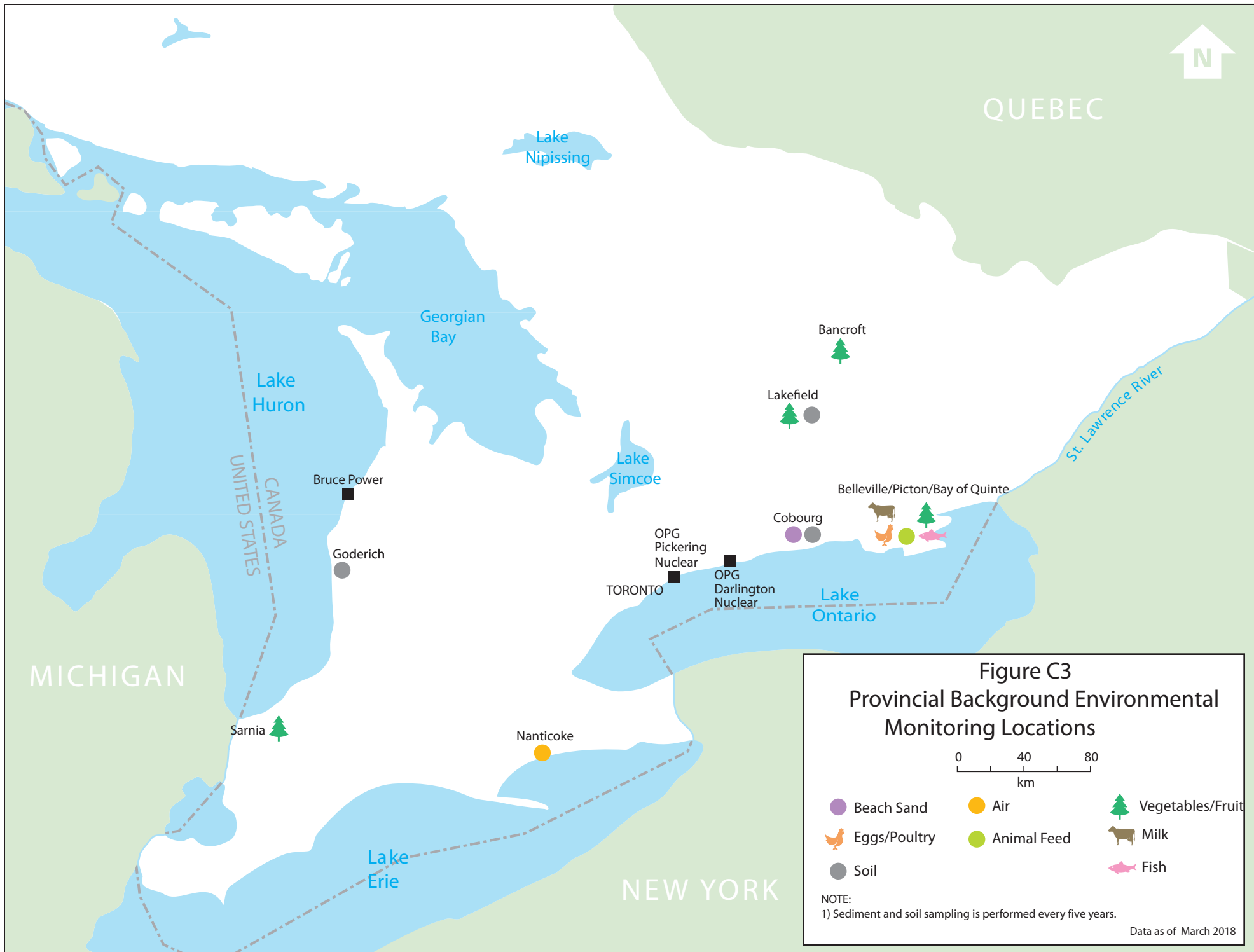
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R.C. Harris WSP (22 km WSW of PN site)

Whitby WSP (12 km ENE of PN site)





**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>89 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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**Appendix D: Environmental Monitoring Data**

**Table D-1: Annual Average Concentrations of Tritium-in-Air – 2017**

Molecular Sieve Tritium-in-Air											
DN EMP Locations	N	Location Average (Bq/m <sup>3</sup> ) <sup>(a)</sup>	Uncertainty (±2σ) <sup>(b)</sup>	PN EMP Locations	N	Location Average (Bq/m <sup>3</sup> ) <sup>(a)</sup>	Uncertainty (±2σ) <sup>(b)</sup>	Background Locations	N	Location Average (Bq/m <sup>3</sup> ) <sup>(a)</sup>	Uncertainty (±2σ) <sup>(b)</sup>
D1	12	1.3	1.6	P10	12	9.1	11.0	Nanticoke	12	<0.1	
D2	12	1.4	2.5	P11	12	2.6	2.6				
D5	12	0.4	0.4	P2	12	12.6	15.4				
D9	12	0.5	0.5	P3	12	2.6	2.3				
D10	11	0.2	0.3	P4	12	1.2	1.3				
D11	11	0.5	0.8	P6	12	6.8	6.2				
Annual Average <sup>(c)</sup>	70	0.7	1.6	Annual Average <sup>(c)</sup>	72	5.8	11.5				

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Molecular Sieve Tritium Ld = 0.2 Bq/m<sup>3</sup> and Lc = 0.1 Bq/m<sup>3</sup>.

(b) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(c) Annual averages are calculated using the entire dataset.

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>90 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table D-2: Annual Average Concentrations of Carbon-14 in Air – 2017**

Passive Sampler C-14 in Air											
DN EMP Locations	N	Location Average (Bq/kg-C) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(b)</sup>	PN EMP Locations	N	Location Average (Bq/kg-C) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(b)</sup>	Background Locations	N	Location Average (Bq/kg-C) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(b)</sup>
D1	4	274	82	P10	4	425	211	Nanticoke	4	238	10
D2	4	271	55	P3	4	292	59				
D5	4	256	30	P4	4	289	50				
D10	4	234	29	P6	4	351	54				
Annual Average <sup>(c)</sup>	16	259	58	Annual Average <sup>(c)</sup>	16	339	154				

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples.

(a) Bq/kg-C (Bq per kg of carbon). Ld for C-14 = 40 Bq/kg-C.

(b) Averages of datasets are reported.  $2\sigma$  denotes two times the standard deviation of the dataset.

(c) Annual averages are calculated using the entire dataset.



Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>91 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-3: Annual Average Dose Rates of Noble Gases and Ir-192 Skyshine in Air – 2017

DN EMP	N	Air Kerma Rates							
		Ar-41 <sup>(c)</sup>		Ir-192		Xe-133		Xe-135	
		Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ )	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ )	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ )	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ )
D1	12	6*	< 1	ND	NA	< 3	NA	< 3	NA
D2	12	6*	< 1	ND	NA	< 3	NA	< 3	NA
D3	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
D5	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
D8	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
D9	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
D10	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
D11	12	< 6	NA	ND	NA	< 3	NA	< 3	NA
Annual Average <sup>(b)</sup>	96	6	0	ND	NA	< 3	NA	< 3	NA
PN EMP	N	Ar-41		Ir-192		Xe-133 <sup>(c)</sup>		Xe-135	
		Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ ) <sup>(a)</sup>	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ )	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ ) <sup>(a)</sup>	Location Average (nGy/month)	Uncertainty ( $\pm 2\sigma$ ) <sup>(a)</sup>
		P2	12	420	630	ND	NA	12	20
P3	12	182	259	ND	NA	5*	4	< 3	NA
P4	12	118	156	ND	NA	4*	2	< 3	NA
P6	12	254	250	ND	NA	8*	6	< 3	NA
P7	12	394	700	ND	NA	11*	23	< 3	NA
P8	12	109	156	ND	NA	4*	3	< 3	NA
P10	12	651	1160	ND	NA	21	38	< 3	NA
P11	12	149	212	ND	NA	4*	4	< 3	NA
Annual Average <sup>(b)</sup>	96	285	642	ND	NA	9	20	< 3	NA

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples.

"<" indicates less than Ld. NA= Not Applicable. ND = Not Detected.

\* indicates that dataset contains both detected and censored non-detected values

(a) Averages of datasets are reported.  $2\sigma$  denotes two times the standard deviation of the dataset.

(b) Annual averages are calculated using the entire dataset.

(c) For datasets partially composed of values censored at the Ld, the Kaplan-Meier methodology is used to determine the mean and standard deviation of the dataset.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>92 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-4: Fruits and Vegetables – 2017

Darlington EMP						
Location	Sample Type	N	HTO (Bq/L) <sup>(a)</sup>		C-14 (Bq/kg-C) <sup>(a)</sup>	
			Location Average	Uncertainty (±2σ) <sup>(c)(d)</sup>	Location Average	Uncertainty (±2σ) <sup>(c)(d)</sup>
DF9	Fruit	3	6.9	1.5	239	23
F18	Fruit	3	12.4	2.2	245	9
R19	Fruit	3	23.3	12.1	242	11
R27	Fruit	3	34.8	21.8	267	31
R335	Fruit	3	18.5	4.3	244	32
<b>Annual Average<sup>(b)</sup></b>	Fruit	15	19.2	22.0	247	28
DF7	Vegetables	0	NA	NA	NA	NA
F16	Vegetables	3	18.1	5.2	243	22
R19	Vegetables	3	22.4	6.6	256	16
R275	Vegetables	1	21.9	3.3	249	22
R335	Vegetables	3	16.3	3.9	237	10
<b>Annual Average<sup>(b)</sup></b>	Vegetables	10	19.2	7.0	246	21

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NA= not applicable.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Annual averages are calculated using the entire dataset.

(c) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(d) For datasets of a single measured value, associated uncertainty is the laboratory analytical uncertainty for that sample

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>93 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-4: Fruits and Vegetables – 2017 (Continued)

Pickering EMP								
Location	Sample Type	N	HTO (Bq/L) <sup>(a)</sup>		C-14 (Bq/kg-C) <sup>(a)</sup>		OBT (Bq/L (w.e.)) <sup>(d)</sup>	
			Location Average	Uncertainty (±2σ) <sup>(c)</sup>	Result	Uncertainty (±2σ) <sup>(c)</sup>	Result	Uncertainty (±2σ) <sup>(b)</sup>
DF3	Fruit	2	10.3	10.0	235	38		
F10	Fruit	3	22.1	23.9	248	32	41.6	3.4
LOC10	Fruit	0	NA	NA	NA	NA		
LOC35	Fruit	0	NA	NA	NA	NA		
LOC7	Fruit	2	117.0	19.8	301	41		
Annual Average <sup>(b)</sup>	Fruit	7	45.9	99.0	257	70	41.6	3.4
DF1	Vegetables	3	15.1	8.5	229	8		
DF3	Vegetables	2	10.5	2.3	226	33		
P11	Vegetables	3	168.5	19.0	327	42		
P9	Vegetables	3	88.7	6.5	247	6		
R144	Vegetables	3	110.2	5.9	272	69		
Annual Average <sup>(b)</sup>	Vegetables	15	78.6	124.1	262	82		

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NA = not applicable.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Annual averages are calculated using the entire dataset.

(c) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(d) w.e. = water equivalent.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>94 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-4: Fruits and Vegetables – 2017 (Continued)

Background Locations								
Location	Sample Type	N	HTO (Bq/L)(a)		C-14 (Bq/kg-C) <sup>(a)</sup>		OBT (Bq/L (w.e.)) <sup>(d)</sup>	
			Result	Uncertainty (±2σ) <sup>(b)</sup>	Result	Uncertainty (±2σ) <sup>(b)</sup>	Result	Uncertainty (±2σ) <sup>(b)</sup>
F1   Bancroft- Sample A	Fruit	1	<1.6	2.4	244	22	NR	NR
F1   Bancroft- Sample B	Fruit	1	<1.7	2.4	224	20		
F2   Lakefield- Sample A	Fruit	1	4.7	2.5	232	21		
F2   Lakefield- Sample B	Fruit	1	<b>2.8</b>	2.4	268	23		
F3   Picton- Sample A	Fruit	1	<0.5	2.3	228	20		
F3   Picton- Sample B	Fruit	1	<1.5	2.4	220	20		
F4   Sarnia- Sample A	Fruit	1	<1.8	2.4	231	21		
F4   Sarnia- Sample B	Fruit	1	<1.0	2.3	231	20		
Annual Average <sup>(c)</sup>		8	<1.9	2.6	235	30		
F1   Bancroft- Sample A	Vegetables	1	<2.2	2.3	224	21	20.0	2.9
F1   Bancroft- Sample B	Vegetables	1	<1.4	2.3	208	20	NR	NR
F2   Lakefield- Sample A	Vegetables	1	<b>3.9</b>	2.4	236	21	19.5	2.8
F2   Lakefield- Sample B	Vegetables	1	<b>3.2</b>	2.4	221	21	NR	NR
F3   Picton- Sample A	Vegetables	1	<0.7	2.2	227	21	16.4	2.8
F3   Picton- Sample B	Vegetables	1	<1.8	2.3	213	21	NR	NR
F4   Sarnia- Sample A	Vegetables	1	<b>3.5</b>	2.4	215	21	18.9	2.8
F4   Sarnia- Sample B	Vegetables	1	<0.9	2.2	216	21	NR	NR
Annual Average <sup>(c)</sup>		8	<2.2	2.4	220	18	18.7	3.2

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NA = not applicable. NR = not required by program.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Individual analytical results are reported. 2σ denotes the laboratory uncertainty of the individual sample.

(c) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(d) w.e. = water equivalent.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>95 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-5: Animal Feed – 2017

Animal Feed <sup>(b)</sup>							
Location	Sample Type	N	HTO (Bq/L) <sup>(a)</sup>		N	C-14 (Bq/kg-C) <sup>(a)</sup>	
			Location Average	Uncertainty ( $\pm 2\sigma$ ) <sup>(d)</sup>		Location Average	Uncertainty ( $\pm 2\sigma$ ) <sup>(d)</sup>
<b>Darlington EMP</b>							
DF18	Generic Feed	1	5.9	2.7	1	241	21
DF18	Generic Feed	1	6.9	2.7	1	247	21
DF7	Generic Feed	1	9.5	2.9	1	252	21
DF7	Generic Feed	1	10.6	2.9	1	242	21
DF8	Generic Feed	1	20.1	3.4	1	248	21
DF8	Generic Feed	1	18.0	3.3	1	221	20
DF9	Generic Feed	1	28.2	3.7	1	224	20
DF9	Generic Feed	1	31.8	3.9	1	242	21
Annual Average <sup>(c)</sup>	Generic Feed	8	16.4	19.6	8	239	22
DF18	Forage	1	8.2	2.8	1	232	21
DF18	Forage	1	6.7	2.8	1	227	21
DF7	Forage	1	11.0	3.0	1	255	22
DF7	Forage	1	14.8	3.2	1	255	23
DF8	Forage	1	7.0	2.8	1	258	23
DF8	Forage	1	6.4	2.8	1	263	22
DF9	Forage	1	11.5	3.0	1	250	22
DF9	Forage	1	12.7	3.1	1	238	21
Annual Average <sup>(c)</sup>	Forage	8	9.8	6.3	8	247	27
<b>Pickering EMP</b>							
DF8	Generic Feed	4	20.7	2.2	4	252	28
DF8	Forage	4	33.7	1.7	4	256	14
<b>Background Locations</b>							
Belleville	Generic Feed	4	5.7	1.6	4	236	17
Belleville	Forage	4	6.6	7.5	4	233	7

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NA= not applicable.

Generic Feed = dry feed, Forage = wet feed

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Animal feed is collected semi-annually. This table depicts the average of the results for each sampling location.

(c) Annual averages are calculated using the entire dataset.

(d) Averages of datasets are reported.  $2\sigma$  denotes two times the standard deviation of the dataset. However, where N < 3, individual sample results are reported and  $2\sigma$  denotes the laboratory uncertainty of the individual sample.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>96 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-6: Annual Average Concentrations in Milk – 2017

Location	N	HTO (Bq/L) <sup>(a)</sup>		C-14 (Bq/kg-C) <sup>(a)</sup>		OBT (Bq/L w.e.)	
		Location Average	Uncertainty (±2σ) <sup>(b)</sup>	Location Average	Uncertainty (±2σ) <sup>(b)</sup>	Location Average	Uncertainty (±2σ) <sup>(b)</sup>
<b>DN EMP</b>							
DF18	12	5.8	2.9	229	18		
DF9	12	<b>4.4</b>	5.3	234	20		
DF8	12	6.5	4.6	237	22		
Annual Average <sup>(c)</sup>	36	5.6	4.6	233	20		
<b>PN EMP</b>							
DF1	12	13.0	3.3	231	16	NR	NR
DF8	12	16.2	5.0	236	19	23.9	7.8
Annual Average <sup>(c)</sup>	24	14.6	5.3	233	18	23.9	7.8
<b>Background Locations</b>							
Belleville	12	<2.1	3.8	243	25	NR	NR

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NA = not applicable. NR = not required by program.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(c) Annual averages are calculated using the entire dataset.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>97 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-7: Annual Average Concentrations in Eggs and Poultry – 2017

Location	Sample Type	N	HTO (Bq/L) <sup>(a)</sup>		C-14 (Bq/kg-C) <sup>(a)</sup>	
			Location Average	Uncertainty (±2σ) <sup>(b)</sup>	Location Average	Uncertainty (±2σ) <sup>(b)</sup>
<b>Darlington EMP</b>						
F16	Poultry	8	8.9	2.3	235	11
D10	Eggs	12	4.0	5.9	233	16
<b>Background</b>						
Picton	Poultry	8	<b>2.9</b>	1.8	236	20
Picton	Eggs	12	<b>2.6</b>	4.2	233	19

NOTES:

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

Egg and poultry sampling not required for PN EMP.

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C.

(b) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

# Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>98 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table D-8: Annual Average Drinking Water and Lake Water Concentrations – 2017**

DN EMP							PN EMP						
Location	Tritium Concentration			Gross Beta Activity Concentration			Location	Tritium Concentration			Gross Beta Activity Concentration		
	N	Location Average (Bq/L) <sup>(b)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	N	Location Average (Bq/L) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>		N	Location Average (Bq/L) <sup>(b)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	N	Location Average (Bq/L) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>
<b>WSP</b>							<b>WSP</b>						
Bowmanville WSP	48	6.8	6.2	12	0.10	0.03	Ajax WSP	48	7.0	8.5	12	0.10	0.05
Newcastle WSP	48	6.4	7.5	12	0.11	0.03	F. J. Horgan WSP	48	5.3	3.8	12	0.10	0.03
Oshawa WSP	48	10.8	13.9	12	0.11	0.03	R.C. Harris WSP	48	4.6	3.2	12	0.11	0.05
							Whitby WSP	48	8.3	10.7	12	0.11	0.03
Annual Average <sup>(d)</sup>	144	8.0	10.5	36	0.11	0.03	Annual Average <sup>(d)</sup>	192	6.3	7.8	48	0.11	0.04
<b>Well Water</b>							<b>Well Water</b>						
DF18	12	<b>3.6</b>	2.7				DF8	12	11.3	3.2			
R2	12	18.6	2.8				R143	12	18.1	4.4			
R316	11	7.6	2.7										
R329	12	12.2	7.5										
Annual Average <sup>(d)</sup>	47	10.6	12.2				Annual Average <sup>(d)</sup>	24	14.7	8.0			
<b>Lake Water</b>							<b>Lake Water</b>						
Courtice Road Beach	8	10.1	8.3				Beachfront Park	8	25.7	26.8			
McLaughlin Bay	8	26.2	5.2				Frenchman's Bay	8	25.9	26.4			
							Squires Beach	8	12.1	15.6			
Annual Average <sup>(d)</sup>	16	18.1	18.0				Annual Average <sup>(d)</sup>	24	21.2	26.1			

**NOTES:**

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NR = not required by program.

Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

- (a) Ld for gross beta = 0.03 Bq/L and Lc = 0.02 Bq/L.
- (b) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L.
- (c) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.
- (d) Annual averages are calculated using the entire dataset.
- (e) Samples are not required during the winter months.



# Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>99 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table D-9: Lake Fish – 2017**

	Sample Type	N	HTO		C-14		Co-60	Cs-134	Cs-137		K-40		OBT composite <sup>(e)</sup>	
			Result (Bq/L) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	Result (Bq/kg-C) <sup>(a)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	Result (Bq/kg fw) <sup>(b)</sup>	Result (Bq/kg fw) <sup>(b)</sup>	Result (Bq/kg fw) <sup>(b)(d)</sup>	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	Result (Bq/kg fw)	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>	Result (Bq/L) w.e.	Uncertainty ( $\pm 2\sigma$ ) <sup>(c)</sup>
<b>DN EMP - Locations</b>														
Darlington Diffuser	White sucker	8	4.6	1.3	240	15	<0.1	<0.1	0.1*	<0.1	130	12	19.2	2.8
<b>PN EMP - Locations</b>														
Pickering 5-8 Outfall	White sucker	8	5.9	2.6	248	22	<0.1	<0.1	0.1*	0.1	123	17	21.8	2.9
<b>Background Locations</b>														
Lake Ontario (USA) Far Field	White sucker	8	<2.3	1.9	230	11	<0.1	<0.1	0.3	0.2	123	8	14.3	2.7

**NOTES:**

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples

fw = fresh weight

w.e. = water equivalent

(a) Ld for tritium = 4.5 Bq/L and Lc = 2.3 Bq/L. Ld for C-14 = 40 Bq/kg-C. Bolded values are greater than Lc but less than Ld. "<" indicates less than Lc.

(b) For gamma analysis (Co-60, Cs-134, Cs-137, K-40), "<" indicates less than Ld.

(c) Averages of datasets are reported.  $2\sigma$  denotes two times the standard deviation of the dataset.

(d) For datasets partially composed of values censored at the Ld, the Kaplan-Meier methodology is used to determine the mean and standard deviation of the dataset. This is indicated by "".

(e) Where individual analytical results are reported,  $2\sigma$  denotes the laboratory uncertainty of the individual sample.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>100 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table D-10: Beach Sand – 2017

Beach Sand	N	Gamma Analysis (Bq/kg dw) <sup>(a)</sup>					
		Co-60 Result	Cs-134 Result	Cs-137 <sup>(c)</sup>		K-40	
				Result	Uncertainty (±2σ) <sup>(b)</sup>	Result	Uncertainty (±2σ) <sup>(b)</sup>
<b>DN EMP - Locations</b>							
Courtice Road Beach	8	< 0.1	< 0.1	< 0.1	NR	268	13
McLaughlin Bay	8	< 0.1	< 0.2	< 0.1	NR	293	27
West/East Beach	8	< 0.1	< 0.2	< 0.1	NR	333	54
<b>PN EMP - Locations</b>							
Beachfront Park	8	< 0.1	< 0.1	0.4	0.1	297	18
Beachpoint Promenade	8	< 0.1	< 0.1	0.4	0.1	372	25
Squire Beach	8	< 0.1	< 0.2	< 0.2	NR	228	71
<b>Background Locations</b>							
Cobourg	8	< 0.1	< 0.1	0.4	0.1	262	18

Refer to Section 3.3.1 for complete list of reporting conventions.

N = number of samples. NR = not required by program.

\* indicates that dataset contains both detected and censored non-detected values

(a) For gamma analysis "<" indicates less than Ld.

(b) Averages of datasets are reported. 2σ denotes two times the standard deviation of the dataset.

(c) For datasets partially composed of values censored at the Ld, the Kaplan-Meier methodology is used to determine the mean and standard deviation of the dataset.

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>101 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table D-11: Soil – 2017**

Location	Sample Type	Gamma Analysis (Bq/kg dw) <sup>(a)</sup>					
		Co-60 Result	Cs-134 Result	Cs-137 Result	Cs-137 Uncertainty (±2σ)	K-40 Result	K-40 Uncertainty (±2σ)
<b>DN EMP</b>							
F16	Soil - Undisturbed	<0.3	<0.4	7.2	0.3	691.0	9.3
F16	Soil - Irrigated	<0.2	<0.2	5.8	0.2	647.1	7.6
R316	Soil - Irrigated	<0.2	<0.3	5.1	0.2	540.1	7.1
<b>PN EMP</b>							
P11	Soil - Undisturbed	<0.2	<0.3	4.5	0.3	724.2	7.8
P11	Soil - Irrigated	<0.2	<0.3	3.0	0.3	499.7	8.2
<b>Background Locations</b>							
Lakefield (A)	Soil - Undisturbed	<0.2	<0.3	5.5	0.2	761.1	8.3
Lakefield (B)	Soil - Undisturbed	<0.2	<0.2	5.6	0.3	745.6	8.2
Cobourg (A)	Soil - Undisturbed	<0.2	<0.3	8.7	0.3	590.2	8.4
Cobourg (B)	Soil - Undisturbed	<0.2	<0.3	9.0	0.3	585.4	8.6
Goderich (A)	Soil - Undisturbed	<0.2	<0.3	1.8	0.2	393.5	7.0
Goderich (B)	Soil - Undisturbed	<0.1	<0.2	1.7	0.1	394.4	5.9

NOTES:

(a) For gamma analysis "<" indicates less than Ld.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>102 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

## Appendix E: Potential Critical Group Descriptions

### E.1.0 DARLINGTON NUCLEAR POTENTIAL CRITICAL GROUPS

Nine potential critical groups are identified for Darlington Nuclear. The annual public dose is calculated for the top three DN potential critical groups only, which have yielded the highest dose estimates in recent years. These are the Dairy Farm, the Farm, and the Rural Resident, as shown in Figure C1 (see Appendix C, Maps of Environmental Monitoring and Critical Group Locations). The EMP sampling plan is structured around monitoring for these three potential critical groups. These groups can change based on the updated pathway analysis results. For informational purposes, descriptions for all nine potential critical groups considered are provided below.

All of the potential critical groups, with the exception of the Industrial/Commercial group, consume some locally caught fish near the DN diffuser. All potential critical groups with the exception of the Sport Fisher and Industrial/Commercial groups are assumed to be exposed to local beach sand. The one-year old infant is assumed to drink cow's milk and water (not infant formula). For all potential critical groups except the dairy farm infant, who drinks fresh local cow's milk, the assumption is made that the milk consumed is a composite from dairy farms all over Ontario which are not affected by station operations.

Based on the site-specific survey review [R-30], a small fraction of residents from the Oshawa/Courtice, Bowmanville, West/East Beach, and Rural Resident potential critical groups work within 5 km of DN. In addition, a small fraction of the Industrial/Commercial potential critical group resides close to DN. Therefore, the average Adult dose for the Rural Resident potential critical group has been adjusted to account for the exposure this portion of the population receives while at work and at home.

The DN potential critical groups are described as follows:

- (c) The **Oshawa/Courtice** potential critical group consists of urban residents in Oshawa and in the community of Courtice within the Municipality of Clarington located to the W and WNW of the site starting at about 6 km from the site. These residents obtain drinking water from the Oshawa WSP, and grow a small percentage of their annual fruit and vegetable consumption in gardens.
- (d) The **Bowmanville** potential critical group consists of urban residents located to the NE and NNE of the site at distances from 4 to 7 km from DN. These residents obtain drinking water from the Bowmanville WSP, and grow a small percentage of their annual fruit and vegetable consumption in gardens. They also purchase a small percentage of their annual meat, poultry and egg consumption from local farms.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>103 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- (e) The **West/East Beach** potential critical group consists of urban residents located to the ENE of the site at distances from 3.5 km to 7 km. These residents obtain their drinking water from both wells and the Bowmanville WSP, and grow a small percentage of their annual fruit and vegetable consumption in gardens. They also purchase a small percentage of their annual poultry and egg consumption from local farms.
- (f) The **Farm** potential critical group consists of agricultural farms (but not dairy farms) located in all landward wind sectors around the DN site at distances from 1.5 km to 10 km. The closest is in the WNW wind sector. Members of this group obtain their water supply mostly from wells and use it for drinking, bathing, irrigation and watering livestock. They also obtain a large fraction of their annual fruit, vegetable and animal product consumption from locally grown products.
- (g) The **Dairy Farm** potential critical group consists of dairy farms located in all landward wind sectors around the DN site at distances from 3 km to over 10 km. The closest is in the N wind sector. Members of this group obtain their water supply from wells and use it for drinking, bathing, irrigation, and livestock watering. They also obtain a large fraction of their annual fruit, vegetable and animal product consumption, including fresh cow's milk, from locally grown products.
- (h) The **Rural Residents** potential critical group consists of residents in rural areas in all landward wind sectors around the site at distances of about 2 km to 5 km. Members of this group obtain about half of their water supply from wells and half from the Bowmanville WSP, and use it for drinking, bathing, and irrigation. They obtain a moderate fraction of their annual fruits, vegetables, poultry and eggs from locally grown products.
- (i) The **Industrial/Commercial** potential critical group consists of adult workers whose work location is close to the nuclear site. The closest location for this group is the St. Mary's cement plant about 1.8 km NE of the site, however, the most affected location due to updated meteorological data is the Courtice Water Pollution Control Plant about 2 km W of DN. Members of this group are typically at this location about 23% of the time. They consume water from the Bowmanville WSP.
- (j) The **Sport Fisher** potential critical group is comprised of non-commercial individuals fishing near the DN site discharge, about 0.5 km S of the DN site. Members of this group were conservatively assumed to obtain their entire amount of fish for consumption from the vicinity of the DN site and spend 1% of their time at the discharge location where atmospheric exposure occurs.
- (k) The **Camper** potential critical group consists of campers at the Darlington Provincial Park, located from 4 to 6 km W of the site at the lakeshore, and includes McLaughlin Bay, a shallow water body where some fishing takes place. The campers are assumed to be in the park no more than six months of the year. They consume drinking water from the Oshawa WSP, and purchase a small fraction of their annual fruits, vegetables, meat, poultry, and eggs from locally grown sources.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>104 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

## E.2.0 PICKERING NUCLEAR POTENTIAL CRITICAL GROUPS

Six potential critical groups are identified for Pickering Nuclear. Note that the annual public dose is calculated for the top three PN potential critical groups, which have yielded the highest dose estimates in recent years. These are the Industrial Worker, the Urban Resident, and the inhabitants of a Correctional Institution. In addition, PN dose is calculated for the Dairy Farm potential critical group since it is exposed to the most media/pathways. Including the Dairy Farm group assures that any future changes in emissions, environmental transfer factors, exposure factors, and dosimetry, and changes in the distribution of radionuclides released will be captured. Refer to Figure C2 in Appendix C, Maps of Environmental Monitoring and Critical Group Locations.

The annual sampling plan is structured around monitoring for these four potential critical groups. These groups can change based on the updated pathway analysis results. For informational purposes, descriptions for all six potential critical groups considered are provided below.

The one-year old infant is assumed to drink cow's milk and water (not infant formula). For all potential critical groups except the dairy farm infant, who drinks fresh local cow's milk, the assumption is made that the milk is a composite from dairy farms all over Ontario which are not affected by station operations.

Based on the site-specific survey [R-31], a small fraction of Industrial/Commercial workers reside close to PN. Similarly, a fraction of residents from the Urban Resident potential critical group work within 5 km of PN. Therefore, the average Adult doses for these groups have been adjusted to account for the exposure this portion of the population receives while at work and at home.

The PN potential critical groups are described as follows:

- (a) The **C2** potential critical group consists of inhabitants at a correctional institute, located approximately 3 km NNE of the PN Site. The C2 group obtains drinking water from the Ajax WSP and does not consume locally grown fruits or vegetables. The C2 resident is conservatively assumed to be at this location 100 percent of the time over the full year.
- (b) The **Industrial/Commercial** potential critical group consists of adult workers whose work location is close to the nuclear site. Members of this group are typically at this location about 23% of the time. They consume water from the Ajax WSP. The closest location for this group is about 1 km NNE of the site.
- (c) The **Urban Residents** potential critical group consists of Pickering and Ajax area residents which surround the PN Site (e.g., Fairport, Fairport Beach, Rosebank, Liverpool, Pickering Village, etc.). The members of this group mostly consume water from the Ajax WSP and also consume a diet composed in part of locally grown produce and some locally caught fish. Members of this potential critical group are also externally exposed to beach sand at local beaches (Beachpoint Promenade, Beachfront Park, or Squires Beach).

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>105 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- (d) The **Farm** potential critical group consists of residents of agricultural farms (but not dairy farms) within a 10 km radius of the PN Site. Members of this group obtain most of their water supply from wells but also a portion from the Ajax WSP. Members of this potential critical group consume locally grown produce and animal products, as well as locally caught fish. They are also externally exposed to beach sand at local beaches (Beachpoint Promenade, Beachfront Park, or Squires Beach).
- (e) The **Dairy Farm** potential critical group consists of residents of dairy farms within a 20 km radius of the PN Site. This group obtains most of their water supply from local wells. They also consume locally grown fruit and vegetables and locally produced animal products, including fresh cow's milk. Members of this potential critical group are also externally exposed to beach sand at local beaches (Beachpoint Promenade, Beachfront Park, or Squires Beach).
- (f) The **Sport Fisher** potential critical group is comprised of non-commercial individuals fishing near the PN site outfalls, 0.5 km S of the PN site. Members of this group were conservatively assumed to obtain their entire amount of fish for consumption from the vicinity of the PN site and spend 1% of their time at the outfall location where atmospheric exposure occurs.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>106 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

## Appendix F: Dose Calculation Procedure and Concentrations

### F.1.0 CRITICAL GROUP DOSE CALCULATION PROCEDURE

The dose calculations were performed according to N-INS-03443-00001, Methodology for Data Analysis and Public Dose Determination for the Environmental Monitoring Program [R-29]. Deviations from this methodology are listed below. The methodology used and software used for dose calculation, IMPACT 5.4.0, are consistent and compliant with CSA N288.1-08 [R-20]. As mentioned in Section 4.1.1, the recently released IMPACT 5.5.1, which is compliant with CSA N288.1-14 [R-56], will be used first to update DN and PN DRLs before it is applied to public dose calculations.

- An update to N288.1-08 was issued in 2011. Only one change in this update affects the dose calculation and it is related to the parameters used for beef cattle consuming dry feed. Given that the use of the existing parameters produces a conservative dose estimate, this change will not be applied at this time.
- OBT doses from terrestrial animals and terrestrial plants were modeled using HTO concentrations measured in terrestrial samples at the potential critical groups. OBT doses from fish were modeled from HTO concentrations in fish.
- HTO and C-14 concentrations in terrestrial animal products other than milk, eggs, and poultry are modeled from measured concentrations of HTO and C-14 in animal feed, forage, air and water. The concentrations are used to calculate the dose from ingestion of animal products. The dose resulting from I(mfp) and particulate is modeled from emissions and empirical Ka values and the ratio of modeled Ka values for the boundary monitor location and the potential critical group location.
- Location specific measures of each radionuclide were used in the potential critical group calculations where the group occupied a relatively small geographic location. Some groups such as the Farm, Dairy Farm or Urban Resident are spread over much wider geographic areas, and for these groups air concentrations were determined for a single conservative representative location, and group average values were used for terrestrial samples and water sources.
- Only dairy farm residents ingest local cow's milk.
- People are generally assumed to be at the potential critical group location 100% of the time, with the exception of the Industrial/Commercial group. Details are provided in Appendix E. Based on the site specific surveys, a small fraction of residential potential critical group members at both PN and DN work within 5 km of the station. In addition, a small fraction of Industrial/Commercial workers reside close to the station at both PN and DN. Therefore, the average Adult doses for these groups have been adjusted at both PN and DN to account for the exposure this portion of the population receives while at work and at home.
- No local grain products are consumed by humans.



## Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>107 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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### F.2.0 PROVINCIAL-BACKGROUND DATA

Treatment of provincial-background data for public dose calculation purposes is as follows:

- If the mean (arithmetic or Kaplan-Meier) is below the Lc, a concentration of 0 (zero) is used for the dose calculation in order to be conservative, i.e. no background concentration is subtracted from the concentration measured around PN or DN.
- If all values in a dataset are below the Ld, a concentration of 0 (zero) is used for the dose calculation in order to be conservative.
- If there are not enough samples collected in a given year to accurately reflect the background dose in a particular sample media, 0 (zero) is used for HTO and gamma in order to be conservative. Previous sampling years may be consulted to arrive at an estimated C-14 concentration in the affected media as background values are not expected to vary significantly from year to year.

### F.3.0 POTENTIAL CRITICAL GROUP RADIONUCLIDE CONCENTRATIONS AND BACKGROUND SUBTRACTIONS

The following section details how the radionuclide concentrations are determined, whether they are measured or modeled, and any calculations made to obtain results.

A summary on the radionuclides and pathways measured and modeled in the dose calculation is presented in Table F-1. DRL Guidance document [R-49] provides a description of each pathway.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>108 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table F-1: Radionuclides and Pathways Measured and Modeled in the Dose Calculation

Pathway	Radionuclide	Modeled <sup>(a)</sup>	Measured
Air Inhalation	HTO	√(Fisher)	√ <sup>(c)</sup>
	HT	√ <sup>(b)</sup>	
	C-14	√ <sup>(b)</sup>	√
	I(mfp)	√ <sup>(b)</sup>	
	Co-60	√ <sup>(b)</sup>	
Air External Exposure	Noble Gas		√ <sup>(c)</sup>
	C-14	√ <sup>(b)</sup>	√
	I(mfp)	√ <sup>(b)</sup>	
	Co-60	√ <sup>(b)</sup>	
Soil External Exposure	C-14	√	
	I(mfp)	√	
	Cs-137+, Co-60	√	
Sand External Exposure	C-14	√	
	Cs-137+		√
Water External Exposure (Lakes, WSPs, Wells)	HTO	√ (wells)	√
	C-14	√	
	I(mfp)	√	
	Cs-137+	√	
Terrestrial Animals Ingestion	HTO	√	√ (milk, eggs, poultry)
	C-14	√	√ (milk, eggs, poultry)
	I(mfp)	√	
	Cs-137+, Co-60	√	
	OBT	√ <sup>(d)</sup>	
Terrestrial Plants Ingestion	HTO		√
	C-14		√
	I(mfp)	√	
	Cs-137+, Co-60	√	
	OBT	√ <sup>(d)</sup>	
Aquatic Animals Ingestion	HTO		√
	C-14		√
	I(mfp)	√	
	Cs-137+		√
	OBT	√ <sup>(d)</sup>	
Sand and Soil Incidental Ingestion	HTO	√	
	C-14	√	
	I(mfp)	√ (soil)	
	Cs-137+, Co-60	√	√ (sand)
Water Ingestion (WSPs, Wells)	HTO		√
	C-14	√	
	I(mfp)	√	
	Cs-137+	√	

"+" indicates that contributions from progeny are included.

- (a) Modeling is based on emissions or from local air measurements where they are available.
- (b) Concentrations are modeled from emissions and adjusted using empirical Ka determined for each potential critical group location.
- (c) Doses are measured directly at the site boundary and adjusted to potential critical group locations using the ratio of modeled air dispersion factors for the boundary monitor and potential critical group.
- (d) OBT dose is modeled from HTO concentration in terrestrial plants, terrestrial animals, or fish respectively.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>109 of 132</b>
Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>		

### F.3.1 Tritium

For the purpose of estimating the critical group dose, the concentrations used in the corresponding pathways were determined as follows:

**Air** – Tritium-in-air is measured at boundary locations with measured background tritium-in-air subtracted, and these values are used to estimate concentrations at each potential critical group location using the ratio of modeled atmospheric dispersion factors for the boundary monitor location and the potential critical group location (except for the Fisher potential critical group where it is modeled from emissions).

Concentrations of radionuclides in air that are not monitored at boundary sites or potential critical groups are obtained for the potential critical group location as follows:

The concentrations at the boundary monitor sites are estimated using their emissions data and empirical Ka values obtained from HTO emissions and HTO boundary monitor measurements. The concentrations at potential critical group locations are modeled from the empirically estimated boundary location concentration by using the ratio of modeled air dispersion factors for the boundary monitor location and the potential critical group location.

- **Water** – Drinking water is sampled and measured at the local WSPs and also at wells where local residents obtain their water. For the WSPs, the annual average concentration is used with background tritium concentration subtracted. The background tritium concentration is calculated for natural and weapons fallout contributions using the Great Lakes Time-Concentration Tritium Model [R-15]. For wells, the average concentration found at each potential critical group is used and background is assumed to be zero. Tritium concentration in wells used for purposes other than drinking water is modeled. Lake water HTO concentrations are measured monthly and used to calculate the dose from water immersion. Background HTO concentrations from the Great Lakes Time-Concentration Tritium model [R-15], are subtracted.
- **Milk** – Milk from local dairy farms is sampled on a monthly basis. The annual average of all the dairy farms is used for the dose calculation, with background tritium in milk concentration subtracted. Only dairy farm residents drink local milk since it is illegal to sell unprocessed milk.
- **Poultry** – Poultry from a local farm is sampled on an annual basis. The annual average is used for the dose calculation, with background values subtracted. Since the farm where poultry is sampled is located in close proximity to the dairy farm, it is assumed that there is not a large difference in radionuclide concentrations in poultry obtained from the local farm vs. the local dairy farm. Therefore, the poultry samples taken are applied to both the Farm and Dairy Farm potential critical groups.

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>110 of 132</b>

Title:

**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- **Eggs** – Eggs from a local farm are sampled on a quarterly basis. The annual average is used for the dose calculation, with background values subtracted. Since the farm where eggs are sampled is located in close proximity to the dairy farm, it is assumed that there is not a large difference in radionuclide concentrations in eggs obtained from the local farm vs. the local dairy farm. Therefore, the egg samples taken are applied to both the Farm and Dairy Farm potential critical groups.
- **Fruits and Vegetables** – Fruit and vegetable tritium concentrations are measured at each potential critical group location and the background tritium concentration is subtracted. The average concentration from all samples measured for each potential critical group is used in the dose calculation.
- **Animal Feed** – The animal feed (wet and dry) is collected from dairy farms bi-annually and is usually from the previous year's harvest. The annual averages of wet and dry feed are used for the dose calculation with background values subtracted.
- **Fish** – The radionuclide concentrations used for locally caught fish are the average measured values in the fish samples, minus background tritium in water. The background tritium in water concentration is for natural and weapons fallout contributions only, as calculated using the Great Lakes Time-Concentration Tritium Model [R-15].

**F.3.2 Carbon-14**

For the purpose of estimating the critical group dose, the concentrations used in the corresponding pathways were determined as follows:

- Air** – C-14 via air inhalation is monitored at boundary locations for about half the landward wind sectors. Where C-14 in air measurements are available, the concentration of C-14 in air is based on the annual average of measurements for each potential critical group location. If more than one sample location is used to represent one potential critical group, then the maximum of the annual averages is taken. Where C-14 in air measurements are not available C-14 in air is modeled from emissions and adjusted using the empirical Ka as described in Section 4.1.2. For all measurements, the average background C-14 concentration in air is subtracted.
- Water** – Concentrations of C-14 in well water are modeled from measured local air concentrations at each potential critical group location, and concentrations in the WSPs and lake water are modeled from site waterborne emissions.
- Terrestrial media** – The concentrations of C-14 in terrestrial media (plants, milk, animal feed, eggs, and poultry) are based on the average of the measurements for each sample type for each potential critical group, minus the average C-14 concentration measured in background media. Where average measurements for a sample type are less than average concentrations in background media, C-14 is conservatively modeled.

## Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>111 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

- (d) **Fish** – For fish, the average C-14 concentration of all samples per site is used, minus the average concentration of C-14 in Lake Ontario fish measured in background locations.

### F.3.3 Noble Gases and Skyshine

The noble gas detectors measure the air kerma rate, which is converted to effective dose using appropriate age-specific conversion factors (effective dose/air kerma rate) [R-50] and standard occupancy and shielding factors for air immersion dose as described in CSA N288.1-08 [R-20].

Noble gas dose is measured directly in most landward wind sectors around the DN and PN site boundaries, and adjusted to the potential critical group location using calculated air dispersion ratios.

The air kerma rate from the PWMF at the PN site was measured in June 2017 over water on Lake Ontario [R-19]. The results showed a rapid drop in the measured air kerma rate with distance, such that it is below the detection limit (0.33 nGy/h) at a distances greater than 400 m from the storage areas. At 1 km distance, the air kerma rate is estimated to be negligible. The skyshine dose from this source is, therefore, not significant for potential critical groups outside the 1 km boundary, which are all the potential critical groups except the Fisher which is assumed to be located 500 m south of PN in Lake Ontario. Skyshine doses from the PWMF are estimated and included in the total noble gas dose for all potential critical groups. Skyshine doses from the DWMF are negligible as all potential critical groups are located beyond 1 km from the DWMF.

Ir-192 skyshine doses from radiography conducted at DN and PN stations are estimated and included in the potential critical group noble gas doses. Skyshine doses are found to be negligible for all potential critical groups.

### F.3.4 Radioiodines

Radioiodine emissions are assumed to have an equilibrium mixture of radioiodines based on I(mfp). This is to account for short-lived radioiodines which may be emitted along with I-131. Emissions for each short-lived radioiodine are incorporated into the dose model based on its equilibrium ratio to the measured I-131 emission. Doses are modeled for the individual radioiodines and summed for the total I(mfp) dose. Due to the very short half-lives of some of these radioiodines, this calculation may overestimate the doses.

Radioiodines are an airborne emission and concentrations at potential critical group locations are modeled using emissions, the empirical Ka at each potential critical group location and modeled atmospheric dispersion factors.

### F.3.5 Particulates and Gross-Beta Gamma

Both airborne particulates and waterborne gross-beta emissions represent a mixture of beta and gamma emitting radionuclides. In order to obtain conservative doses for

## Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>112 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

these mixtures, they are represented by the most limiting radionuclides typically found in the mixtures. According to the pathway analyses [R-53][R-54], the most limiting radionuclide for atmospheric particulate emissions is Co-60 and for liquid effluent beta-gamma emissions it is Cs-137. There was no analysis for alpha radioactivity because alpha radionuclide emissions from the stations are extremely low [R-55].

For airborne particulates, concentrations in air are modeled using emissions, the empirical Ka at each potential critical group location and modeled atmospheric dispersion factors. Concentrations in terrestrial media are subsequently modeled from the airborne concentrations. These concentrations are used to calculate doses to potential critical groups.

For waterborne gross-beta gamma, potential critical group doses are directly modeled from emissions in aquatic media where no local measurements are available. The only pathways used for dose calculation in which gross beta-gamma activity is measured in environmental samples are fish and beach sand. Background values of activity in Lake Ontario fish and beach sand are subtracted from these measurements.

### F.3.6 Elemental Tritium

For HT, the inhalation pathway is the only direct pathway to humans resulting in dose. Concentrations in air are modeled using emissions, the empirical Ka at each potential critical group location and modeled atmospheric dispersion factors. HT converts into HTO through interaction with microbes in the soil. The resultant HTO is routinely measured in air and local biota around nuclear sites.

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>113 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Appendix G: Tables of Public Doses by Radionuclide, Pathway and Age Group for Darlington Nuclear and Pickering Nuclear Potential Critical Groups**

**Table G-1: Darlington Nuclear – Farm Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total
Adult	C-14	uSv/a	2.60E-04	2.99E-07	4.07E-06	5.76E-11	0.00E+00	0.00E+00	1.42E-10	9.10E-12	0.00E+00	1.63E-04	6.59E-02	6.65E-02	1.33E-01
	Co-60	uSv/a	4.63E-06	1.76E-07	1.01E-07	1.36E-08	4.49E-09	4.65E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.19E-05	2.46E-06	4.71E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	2.79E-05	4.09E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.20E-05
	HT	uSv/a	4.99E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.99E-06
	HTO	uSv/a	1.28E-01	0.00E+00	1.02E-01	2.79E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.15E-05	1.02E-01	7.13E-03
	NobleGases	uSv/a	0.00E+00	1.24E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.24E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.93E-06	1.59E-02	4.74E-03
	I (mfp)	uSv/a	8.21E-05	7.10E-06	6.79E-07	3.55E-09	4.23E-10	1.86E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.12E-03	3.45E-03
	Total	uSv/a	1.29E-01	1.24E-01	1.02E-01	2.80E-03	4.91E-09	4.67E-03	1.42E-10	9.10E-12	0.00E+00	1.79E-04	1.86E-01	7.96E-02	6.28E-01
	Child-10y	C-14	uSv/a	3.71E-04	2.99E-07	2.89E-06	5.76E-11	0.00E+00	0.00E+00	7.85E-10	9.10E-12	0.00E+00	9.63E-05	5.04E-02	4.38E-02
Co-60		uSv/a	6.61E-06	1.76E-07	1.67E-07	1.36E-08	5.80E-08	4.65E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	8.43E-05	4.83E-06	4.74E-03
Cs-137+		uSv/a	0.00E+00	0.00E+00	1.10E-05	4.09E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.51E-05
HT		uSv/a	5.93E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.93E-06
HTO		uSv/a	1.52E-01	0.00E+00	6.56E-02	2.33E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.15E-06	6.77E-02	4.29E-03
NobleGases		uSv/a	0.00E+00	1.24E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.24E-01
OBT		uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.89E-06	1.21E-02	2.92E-03
I (mfp)		uSv/a	1.86E-04	7.10E-06	8.24E-07	3.55E-09	3.99E-09	1.86E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.50E-03	4.79E-03
Total		uSv/a	1.53E-01	1.24E-01	6.56E-02	2.33E-03	6.20E-08	4.67E-03	7.85E-10	9.10E-12	0.00E+00	1.05E-04	1.33E-01	5.31E-02	5.36E-01
Infant_1y		C-14	uSv/a	2.53E-04	2.99E-07	0.00E+00	1.82E-11	0.00E+00	0.00E+00	1.57E-09	9.10E-12	0.00E+00	5.67E-05	4.09E-02	3.94E-02
	Co-60	uSv/a	4.84E-06	2.28E-07	0.00E+00	1.77E-08	1.42E-07	6.04E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	8.19E-05	8.20E-06	6.14E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00
	HT	uSv/a	4.06E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.06E-06
	HTO	uSv/a	1.04E-01	0.00E+00	0.00E+00	8.76E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.84E-06	5.74E-02	4.27E-03
	NobleGases	uSv/a	0.00E+00	1.53E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.53E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.76E-06	9.72E-03	2.39E-03
	I (mfp)	uSv/a	2.17E-04	9.23E-06	0.00E+00	4.62E-09	1.40E-08	2.42E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.47E-03	5.99E-03
	Total	uSv/a	1.05E-01	1.53E-01	0.00E+00	8.76E-04	1.56E-07	6.07E-03	1.57E-09	9.10E-12	0.00E+00	6.23E-05	1.12E-01	5.21E-02	4.29E-01

Report

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Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>114 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table G-2: Darlington Nuclear – Dairy Farm Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total
Adult	C-14	uSv/a	0.00E+00	0.00E+00	0.00E+00	1.39E-11	0.00E+00	0.00E+00	1.42E-10	9.10E-12	0.00E+00	3.24E-05	2.15E-02	2.07E-01	2.28E-01
	Co-60	uSv/a	1.05E-06	3.99E-08	0.00E+00	0.00E+00	5.35E-10	5.54E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.92E-05	3.12E-06	5.77E-04
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	4.09E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.09E-06
	HT	uSv/a	1.13E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.13E-06
	HTO	uSv/a	2.91E-02	0.00E+00	3.66E-02	1.54E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.28E-06	3.84E-02	1.29E-02	1.19E-01
	NobleGases	uSv/a	0.00E+00	5.88E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.88E-02
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	9.80E-07	5.99E-03	3.19E-03
	I (mfp)	uSv/a	1.87E-05	1.58E-06	0.00E+00	0.00E+00	9.36E-11	4.14E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	8.21E-04	1.73E-03
	Total	uSv/a	2.91E-02	5.88E-02	3.66E-02	1.54E-03	6.28E-10	5.58E-04	1.42E-10	9.10E-12	0.00E+00	3.57E-05	6.67E-02	2.24E-01	4.18E-01
	Child-10y	C-14	uSv/a	0.00E+00	0.00E+00	0.00E+00	1.39E-11	0.00E+00	0.00E+00	7.85E-10	9.10E-12	0.00E+00	1.92E-05	1.64E-02	2.64E-01
Co-60		uSv/a	1.50E-06	3.99E-08	0.00E+00	0.00E+00	6.92E-09	5.54E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.11E-05	9.53E-06	5.96E-04
Cs-137+		uSv/a	0.00E+00	0.00E+00	0.00E+00	4.09E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.09E-06
HT		uSv/a	1.35E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.35E-06
HTO		uSv/a	3.46E-02	0.00E+00	2.35E-02	1.28E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.22E-06	2.54E-02	1.62E-02	1.01E-01
NobleGases		uSv/a	0.00E+00	5.88E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.88E-02
OBT		uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.75E-07	4.54E-03	3.34E-03	7.88E-03
I (mfp)		uSv/a	4.24E-05	1.58E-06	0.00E+00	0.00E+00	8.84E-10	4.14E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	9.66E-04	3.95E-03	4.96E-03
Total		uSv/a	3.46E-02	5.88E-02	2.35E-02	1.29E-03	7.80E-09	5.58E-04	7.85E-10	9.10E-12	0.00E+00	2.10E-05	4.73E-02	2.87E-01	4.53E-01
Infant_1y		C-14	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.57E-09	9.10E-12	0.00E+00	1.13E-05	1.34E-02	5.54E-01
	Co-60	uSv/a	1.10E-06	5.19E-08	0.00E+00	0.00E+00	1.70E-08	7.20E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.02E-05	2.26E-05	7.74E-04
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00
	HT	uSv/a	9.23E-07	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	9.23E-07
	HTO	uSv/a	2.37E-02	0.00E+00	0.00E+00	2.01E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.63E-07	2.15E-02	3.76E-02	8.30E-02
	NobleGases	uSv/a	0.00E+00	7.23E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.23E-02
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.49E-07	3.65E-03	6.30E-03	9.95E-03
	I (mfp)	uSv/a	4.96E-05	2.05E-06	0.00E+00	0.00E+00	3.10E-09	5.38E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.34E-03	1.27E-02	1.40E-02
	Total	uSv/a	2.38E-02	7.23E-02	0.00E+00	2.01E-04	2.01E-08	7.26E-04	1.57E-09	9.10E-12	0.00E+00	1.24E-05	4.00E-02	6.11E-01	7.47E-01



Report

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Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>115 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table G-3: Darlington Nuclear – Rural Resident Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total
Adult	C-14	uSv/a	1.48E-04	1.70E-07	1.75E-06	3.62E-11	4.20E-14	7.84E-13	1.40E-10	8.93E-12	0.00E+00	1.09E-04	2.15E-02	6.74E-03	2.85E-02
	Co-60	uSv/a	2.25E-06	8.54E-08	5.71E-08	4.37E-09	2.03E-09	2.10E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.50E-05	5.48E-07	2.12E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	1.05E-04	4.25E-06	5.53E-09	3.61E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.22E-06	1.66E-07	4.74E-04
	HT	uSv/a	2.42E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.42E-06
	HTO	uSv/a	6.23E-02	0.00E+00	8.82E-02	1.81E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.70E-06	3.00E-02	8.00E-04	1.83E-01
	NobleGases	uSv/a	0.00E+00	4.44E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.44E-02
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.31E-06	4.63E-03	4.61E-04
	I (mfp)	uSv/a	3.99E-05	3.10E-06	4.11E-07	1.22E-09	2.04E-10	9.02E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.20E-04	1.78E-04
	Total	uSv/a	6.25E-02	4.44E-02	8.83E-02	1.81E-03	7.77E-09	2.47E-03	1.40E-10	8.93E-12	0.00E+00	1.20E-04	5.68E-02	8.18E-03	2.65E-01
	Child-10y	C-14	uSv/a	2.02E-04	1.63E-07	1.24E-06	3.69E-11	2.36E-13	7.99E-13	7.85E-10	9.10E-12	0.00E+00	6.58E-05	1.68E-02	5.08E-03
Co-60		uSv/a	3.13E-06	8.32E-08	9.67E-08	4.45E-09	2.56E-08	2.05E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.49E-05	1.09E-06	2.08E-03
Cs-137+		uSv/a	0.00E+00	0.00E+00	3.79E-05	4.32E-06	1.73E-08	3.68E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.33E-06	6.06E-08	4.12E-04
HT		uSv/a	2.80E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.80E-06
HTO		uSv/a	7.22E-02	0.00E+00	5.71E-02	1.53E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.20E-06	2.03E-02	5.24E-04	1.52E-01
NobleGases		uSv/a	0.00E+00	4.32E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.32E-02
OBT		uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.98E-06	3.58E-03	3.01E-04
I (mfp)		uSv/a	8.82E-05	3.02E-06	5.09E-07	1.25E-09	1.88E-09	8.79E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.45E-04	3.58E-04	1.20E-03
Total		uSv/a	7.25E-02	4.32E-02	5.71E-02	1.54E-03	4.48E-08	2.43E-03	7.85E-10	9.10E-12	0.00E+00	7.20E-05	4.15E-02	6.26E-03	2.25E-01
Infant_1y		C-14	uSv/a	1.38E-04	1.63E-07	0.00E+00	9.17E-12	4.73E-13	7.99E-13	1.57E-09	9.10E-12	0.00E+00	3.88E-05	1.42E-02	6.15E-03
	Co-60	uSv/a	2.30E-06	1.08E-07	0.00E+00	5.78E-09	6.28E-08	2.66E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.43E-05	2.05E-06	2.69E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	3.12E-07	2.08E-08	4.80E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.98E-07	3.62E-08	4.81E-04
	HT	uSv/a	1.92E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.92E-06
	HTO	uSv/a	4.95E-02	0.00E+00	0.00E+00	3.79E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.62E-06	1.78E-02	7.35E-04	6.84E-02
	NobleGases	uSv/a	0.00E+00	5.31E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.31E-02
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.20E-06	2.95E-03	3.59E-04
	I (mfp)	uSv/a	1.03E-04	3.92E-06	0.00E+00	1.62E-09	6.58E-09	1.14E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.04E-03	1.26E-03	2.42E-03
	Total	uSv/a	4.97E-02	5.31E-02	0.00E+00	3.79E-04	9.02E-08	3.16E-03	1.57E-09	9.10E-12	0.00E+00	4.26E-05	3.60E-02	8.51E-03	1.51E-01

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Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>116 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table G-4: Pickering Nuclear – Dairy Farm Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total
Adult	C-14	uSv/a	1.13E-04	1.30E-07	1.95E-06	3.04E-10	0.00E+00	0.00E+00	2.93E-09	1.87E-10	0.00E+00	0.00E+00	2.09E-02	3.26E-01	3.47E-01
	Co-60	uSv/a	8.40E-06	3.18E-07	0.00E+00	3.00E-09	4.59E-09	4.76E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.72E-05	9.98E-06	4.83E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	1.04E-04	7.67E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.87E-07	1.81E-04
	HTO	uSv/a	7.58E-02	0.00E+00	7.63E-02	1.95E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.14E-02	1.78E-02	1.83E-01
	NobleGases	uSv/a	0.00E+00	1.02E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.02E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.85E-03	6.42E-03
	I (mfp)	uSv/a	1.47E-06	7.77E-08	0.00E+00	1.33E-11	7.40E-12	3.22E-07	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.50E-05	3.43E-05	6.12E-05
Total	uSv/a	7.59E-02	1.02E-01	7.64E-02	2.03E-03	4.60E-09	4.76E-03	2.93E-09	1.87E-10	0.00E+00	0.00E+00	3.42E-02	3.50E-01	6.46E-01	
Child-10y	C-14	uSv/a	1.61E-04	1.30E-07	1.38E-06	3.04E-10	0.00E+00	0.00E+00	1.62E-08	1.87E-10	0.00E+00	0.00E+00	1.54E-02	3.23E-01	3.38E-01
	Co-60	uSv/a	1.20E-05	3.18E-07	0.00E+00	3.00E-09	5.94E-08	4.76E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.61E-05	2.48E-05	4.87E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	4.09E-05	7.67E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.44E-07	1.18E-04
	HTO	uSv/a	9.01E-02	0.00E+00	4.90E-02	1.63E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.56E-03	1.87E-02	1.67E-01
	NobleGases	uSv/a	0.00E+00	1.02E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.02E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.42E-03	5.16E-03
	I (mfp)	uSv/a	3.33E-06	7.77E-08	0.00E+00	1.33E-11	7.00E-11	3.22E-07	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.94E-05	6.89E-05	1.02E-04
Total	uSv/a	9.03E-02	1.02E-01	4.91E-02	1.71E-03	5.95E-08	4.76E-03	1.62E-08	1.87E-10	0.00E+00	0.00E+00	2.44E-02	3.47E-01	6.19E-01	
Infant_1y	C-14	uSv/a	1.10E-04	1.30E-07	0.00E+00	7.62E-12	0.00E+00	0.00E+00	3.23E-08	1.87E-10	0.00E+00	0.00E+00	1.15E-02	5.68E-01	5.80E-01
	Co-60	uSv/a	8.78E-06	4.14E-07	0.00E+00	3.90E-09	1.46E-07	6.19E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.09E-05	5.67E-05	6.33E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.29E-07	1.29E-07
	HTO	uSv/a	6.18E-02	0.00E+00	0.00E+00	3.13E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.06E-03	4.03E-02	1.08E-01
	NobleGases	uSv/a	0.00E+00	1.24E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.24E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.09E-03	7.96E-03
	I (mfp)	uSv/a	3.90E-06	1.01E-07	0.00E+00	1.72E-11	2.45E-10	4.19E-07	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.92E-05	2.42E-04	2.85E-04
Total	uSv/a	6.19E-02	1.24E-01	0.00E+00	3.13E-04	1.46E-07	6.19E-03	3.23E-08	1.87E-10	0.00E+00	0.00E+00	1.88E-02	6.17E-01	8.28E-01	

**Table G-5: Pickering Nuclear – Industrial/Commercial Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total
Adult	C-14	uSv/a	6.58E-04	7.57E-07	3.89E-06	2.75E-11	2.20E-13	4.10E-12	1.80E-10	1.15E-11	0.00E+00	2.43E-07	6.72E-04	6.17E-07	1.34E-03
	Co-60	uSv/a	5.43E-05	2.06E-06	4.81E-295	1.68E-10	2.53E-09	2.62E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.76E-06	1.69E-10	2.68E-03
	Cs-137+	uSv/a	0.00E+00	0.00E+00	1.21E-03	7.36E-06	2.64E-08	1.72E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.95E-06	8.98E-11	2.95E-03
	HTO	uSv/a	5.47E-01	0.00E+00	1.63E-02	1.52E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.35E-08	8.96E-04	2.10E-07	5.64E-01
	NobleGases	uSv/a	0.00E+00	8.94E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	8.94E-01
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.79E-09	1.41E-04	1.41E-04
	I (mfp)	uSv/a	1.14E-05	8.06E-07	0.00E+00	6.37E-13	3.42E-12	1.52E-07	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.08E-06	2.43E-09	1.34E-05
Total	uSv/a	5.48E-01	8.94E-01	1.75E-02	1.60E-04	2.89E-08	4.34E-03	1.80E-10	1.15E-11	0.00E+00	2.62E-07	1.71E-03	9.45E-07	1.47E+00	

Report

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Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>117 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table G-6: Pickering Nuclear – Correctional Institute (C2) Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total	
Adult	C-14	uSv/a	7.32E-04	8.42E-07	1.38E-05	4.14E-11	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.47E-04	
	Co-60	uSv/a	5.53E-05	2.10E-06	0.00E+00	0.00E+00	2.92E-08	3.02E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.03E-02	
	Cs-137+	uSv/a	0.00E+00	0.00E+00	4.31E-03	1.11E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.32E-03	
	HTO	uSv/a	4.93E-01	0.00E+00	5.75E-02	3.62E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.51E-01	
	NobleGases	uSv/a	0.00E+00	6.51E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.51E-01	
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	
	I (mfp)	uSv/a	9.59E-06	6.46E-07	0.00E+00	0.00E+00	4.78E-11	2.15E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.24E-05	
	Total	uSv/a	4.93E-01	6.51E-01	6.19E-02	3.74E-04	2.92E-08	3.02E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.24E+00	
	Child-10y	C-14	uSv/a	1.05E-03	8.42E-07	9.78E-06	4.14E-11	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.06E-03
		Co-60	uSv/a	7.89E-05	2.10E-06	0.00E+00	0.00E+00	3.77E-07	3.02E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.03E-02
Cs-137+		uSv/a	0.00E+00	0.00E+00	1.71E-03	1.11E-05	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.72E-03	
HTO		uSv/a	5.86E-01	0.00E+00	3.70E-02	3.02E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.23E-01	
NobleGases		uSv/a	0.00E+00	6.51E-01	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.51E-01	
OBT		uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	
I (mfp)		uSv/a	2.18E-05	6.46E-07	0.00E+00	0.00E+00	4.52E-10	2.15E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.46E-05	
Total		uSv/a	5.87E-01	6.51E-01	3.87E-02	3.13E-04	3.78E-07	3.02E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.31E+00	

**Table G-7: Pickering Nuclear – Urban Resident Doses – 2017**

HumanType	Radionuclide	Unit	Air (inhalation)	Air (external)	Water (ingestion)	Water (external)	Soil (ingestion)	Soil (external)	Sediment (ingestion)	Sediment (external)	Aquatic plants	Aquatic animals	Terrestrial plants	Terrestrial animals	Total	
Adult	C-14	uSv/a	8.41E-04	9.68E-07	1.17E-05	4.29E-10	3.44E-12	6.41E-11	2.82E-09	1.80E-10	0.00E+00	3.80E-06	1.05E-02	9.65E-06	1.14E-02	
	Co-60	uSv/a	4.86E-05	1.84E-06	7.52E-294	2.63E-09	3.95E-08	4.09E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.76E-05	2.64E-09	4.10E-02	
	Cs-137+	uSv/a	0.00E+00	0.00E+00	3.61E-03	1.15E-04	4.13E-07	2.70E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	6.18E-05	1.40E-09	3.08E-02	
	HTO	uSv/a	5.48E-01	0.00E+00	4.95E-02	2.38E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.11E-07	1.40E-02	3.28E-06	6.14E-01	
	NobleGases	uSv/a	0.00E+00	1.12E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.12E+00	
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	9.05E-08	2.20E-03	2.20E-03	
	I (mfp)	uSv/a	1.21E-05	9.20E-07	0.00E+00	9.96E-12	5.35E-11	2.38E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.68E-05	3.81E-08	3.23E-05	
	Total	uSv/a	5.49E-01	1.12E+00	5.32E-02	2.50E-03	4.52E-07	6.79E-02	2.82E-09	1.80E-10	0.00E+00	4.10E-06	2.68E-02	1.48E-05	1.82E+00	
	Child-10y	C-14	uSv/a	1.10E-03	8.88E-07	8.22E-06	4.46E-10	1.97E-11	6.65E-11	1.62E-08	1.87E-10	0.00E+00	2.33E-06	8.04E-03	9.84E-06	9.17E-03
		Co-60	uSv/a	5.98E-05	1.59E-06	1.30E-293	2.73E-09	5.31E-07	4.25E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.62E-05	7.96E-09	4.26E-02
Cs-137+		uSv/a	0.00E+00	0.00E+00	1.42E-03	1.19E-04	1.32E-06	2.80E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.59E-05	5.42E-10	2.96E-02	
HTO		uSv/a	5.74E-01	0.00E+00	3.17E-02	2.06E-03	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.17E-07	9.67E-03	2.47E-06	6.17E-01	
NobleGases		uSv/a	0.00E+00	1.02E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.02E+00	
OBT		uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	5.51E-08	1.74E-03	1.74E-03	
I (mfp)		uSv/a	2.45E-05	8.30E-07	0.00E+00	1.03E-11	5.25E-10	2.47E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.05E-05	8.50E-08	4.85E-05	
Total		uSv/a	5.75E-01	1.02E+00	3.31E-02	2.18E-03	1.85E-06	7.05E-02	1.62E-08	1.87E-10	0.00E+00	2.50E-06	1.96E-02	1.37E-05	1.72E+00	
Infant_1y		C-14	uSv/a	7.52E-04	8.88E-07	0.00E+00	4.18E-11	3.94E-11	6.65E-11	3.23E-08	1.87E-10	0.00E+00	1.37E-06	6.51E-03	1.80E-05	7.29E-03
		Co-60	uSv/a	4.38E-05	2.06E-06	0.00E+00	3.55E-09	1.30E-06	5.52E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	4.38E-05	2.06E-08	5.53E-02
	Cs-137+	uSv/a	0.00E+00	0.00E+00	0.00E+00	1.42E-05	1.58E-06	3.65E-02	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.27E-05	3.84E-10	3.65E-02	
	HTO	uSv/a	3.93E-01	0.00E+00	0.00E+00	4.07E-04	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	7.32E-08	8.40E-03	3.98E-06	4.02E-01	
	NobleGases	uSv/a	0.00E+00	1.25E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	1.25E+00	
	OBT	uSv/a	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	0.00E+00	3.35E-08	1.42E-03	1.42E-03	
	I (mfp)	uSv/a	2.87E-05	1.08E-06	0.00E+00	1.34E-11	1.84E-09	3.21E-06	0.00E+00	0.00E+00	0.00E+00	0.00E+00	2.79E-05	3.15E-07	6.12E-05	
	Total	uSv/a	3.94E-01	1.25E+00	0.00E+00	4.21E-04	2.89E-06	9.17E-02	3.23E-08	1.87E-10	0.00E+00	1.48E-06	1.64E-02	2.41E-05	1.75E+00	

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Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>118 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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**Appendix H: Supplementary Study Data**

**Table H-1: Air Kerma Rates from the PWF at the PN Site**

PWF Transects				Background Transects			
ID	Northing (UTM 17T)	Easting (UTM 17T)	Air Kerma Rate [nGy.h <sup>-1</sup> ]	ID	Northing (UTM 17T)	Easting (UTM 17T)	Air Kerma Rate [nGy.h <sup>-1</sup> ]
A1	4852151.9	655754.5	1.83	BG-A1	4852067.3	652818.4	-0.74
A2	4852081.5	655706.3	0.43	BG-A2	4851982.6	652900.4	-0.86
A3	4852008.3	655649.2	-0.03	BG-A3	4851909.3	652972.1	-0.85
A4	4851919.6	655587.1	-0.18	BG-A4	4851866.6	653037.2	-0.66
A5	4851829.3	655527.6	-0.13	BG-A5	4851785.3	653118.6	-0.17
A6	4851756.8	655467	-0.1	BG-A6	4851739	653185.2	-0.24
A8	4851607.1	655349.5	-0.04	BG-A8	4851597.3	653335.5	0.16
A10	4851434.5	655228.9	0.03	BG-A10	4851449.7	653493.1	0.23
B1	4852104.8	655908.5	1.53	BG-A15	4851065.6	653826.5	-0.08
B2	4852003.2	655897.8	0.14	BG-A20	4850711.4	654151.8	-0.45
B3	4851905.7	655896.5	-0.19	BG-B1	4853194.7	658883.7	0.08
B4	4851810.4	655890.1	-0.19	BG-B2	4853104.6	658888.1	-0.06
B5	4851699.5	655895.6	-0.11	BG-B3	4853007.5	658890.3	-0.14
B6	4851605.5	655903.4	-0.08	BG-B4	4852909.3	658884.9	-0.1
B8	4851404.6	655907.7	0.02	BG-B5	4852804.4	658885	0.01
B10	4851209.1	655908.7	-0.07	BG-B6	4852705.9	658885.2	-0.04
C1	4852377.6	656295.9	1.17	BG-B8	4852499.3	658886.9	0.03
C2	4852282.7	656308	0.12	BG-B10	4852303.8	658885.5	-0.07
C3	4852181.7	656312.8	-0.05	BG-B15	4851806.5	658875.4	-0.08
C4	4852068	656312.6	-0.05	BG-B20	4851304.4	658875.8	-0.02
C5	4851965.5	656307.4	-0.18				
C6	4851866.7	656299.2	-0.08				
C8	4851655.8	656302.1	-0.03				
C10	4851474.1	656300.5	-0.11				
D1	4852432	656406.8	1.2				
D2	4852339	656463.6	-0.08				
D3	4852251.2	656528.9	-0.2				
D4	4852164.7	656576	-0.18				
D5	4852094.1	656632.2	-0.13				
D6	4851999.9	656678.9	-0.17				
D8	4851831.7	656797.5	0.04				
D10	4851678.2	656907.2	-0.05				
E1	4852473	656501.9	0.76				
E2	4852443.4	656593.2	-0.18				
E3	4852421.8	656689.4	-0.22				
E4	4852387.1	656793.6	-0.23				
E5	4852368.3	656894.6	-0.27				
E6	4852338	656991	-0.31				
E8	4852275.9	657177.3	-0.01				
E10	4852233.3	657383	-0.04				

Note: Values in red are notable values above detection.  
 Sample points increase in distance from shoreline along each transect.  
 Source: Table 5 of N-REP-03443-10018 [R-19]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>119 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table H-2: Screening of Soil COPCs for Ecological Risk Assessment**

Parameter	Unit	Detection Limit	MOE2011 (Table 3 - Industrial)	CCMESQG (Industrial)	US EPA Region 5 RCRA	Soil Screening Level	Ref	Max Soil Conc.	Carried Forward as COPC?	Notes
<b>Inorganics</b>										
Conductivity	ms/cm	0.002	1.4			1.4	(1)	1.2	No	
Cyanide (free)	ug/g	0.01	0.051	8	1.3	0.051	(1)	0.33	Yes	
Moisture, Percent	%	1						20	No	
pH	pH units							8.07	No	
Sodium Adsorption Ratio	-		12	12		12	(1)	11	No	
<b>Metals</b>										
Antimony	ug/g	0.2	40	40		40	(1)	9.7	No	
Arsenic	ug/g	1	18	12		12	(2)	58	Yes	
Barium	ug/g	0.5	670	2000		670	(1)	140	No	
Beryllium	ug/g	0.2	8	8		8	(1)	0.79	No	
Boron	ug/g	5				0	(1)	12	No	
Boron, Hot Water Soluble	ug/g	0.1	2			2	(1)	0.42	No	
Cadmium	ug/g	1	1.9	22		1.9	(1)	1.7	No	
Chromium	ug/g	0.1	160	87		87	(2)	44	No	
Cobalt	ug/g	1	80	300		80	(1)	67	No	
Copper	ug/g	0.5	230	91		91	(2)	830	Yes	
Hexavalent Chromium	ug/g	0.2	8	1.4		1.4	(2)	<0.2	No	
Lead	ug/g	1	120	600		120	(1)	230	Yes	
Mercury	ug/g	0.05	3.9	50			(1)	<0.05	No	
Molybdenum	ug/g	0.5	40	40		40	(1)	16	No	
Nickel	ug/g	0.5	270	89		89	(2)	23	No	
Selenium	ug/g	0.5	5.5	2.9		2.9	(2)	2	No	
Silver	ug/g	0.2	40	40		40	(1)	0.53	No	
Thallium	ug/g	0.05	3.3	1		1	(2)	0.22	No	
Uranium	ug/g	0.05	33	300		33	(1)	0.89	No	
Vanadium	ug/g	5	86	130		86	(1)	41	No	
Zinc	ug/g	5	340	360		340	(1)	3200	Yes	
<b>Petroleum Hydrocarbons</b>										
0										
Benzene	ug/g	0.02	0.32			0.32	(1)	< 0.020	No	
Ethylbenzene	ug/g	0.02	9.5			9.5	(1)	< 0.020	No	
Toluene	ug/g	0.05	68			68	(1)	< 0.020	No	
Xylenes, Total	ug/g	0.02	26			26	(1)	< 0.020	No	
Petroleum Hydrocarbons - F1 (C6-C10)	ug/g	10	55			55	(1)	< 10	No	
Petroleum Hydrocarbons - F1 (C6-C10)-BTEX	ug/g	10	55			55	(1)	< 10	No	
Petroleum Hydrocarbons - F2 (C10-C16)	ug/g	10	230			230	(1)	< 10	No	
Petroleum Hydrocarbons - F3 (C16-C34)	ug/g	1000	1700			1700	(1)	1100	No	
Petroleum Hydrocarbons - F4 (C34-C50)	ug/g	1000	3300			3300	(1)	1500	No	
Petroleum Hydrocarbons - F4 Gravimetric	ug/g	1000	3300			3300	(1)	5700	Yes	
Reached Baseline at C50	ug/g					0		-	-	

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>120 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table H-2: Screening of Soil COPCs for Ecological Risk Assessment (Continued)

Parameter	Unit	Detection Limit	MOE2011 (Table 3 - Industrial)	CCMESQG (Industrial)	US EPA Region 5 RCRA	Soil Screening Level	Ref	Max Soil Conc.	Carried Forward as COPC?	Notes
VOCs						0				
1,1,1,2-Tetrachloroethane	ug/g	0.05	0.087			0.087	(1)	< 0.050	No	
1,1,1-Trichloroethane	ug/g	0.05	6.1	50		6.1	(1)	< 0.050	No	
1,1,2,2-Tetrachloroethane	ug/g	0.05	0.05	0.6		0.05	(1)	< 0.050	No	
1,1,2-Trichloroethane	ug/g	0.05	0.05	50		0.05	(1)	< 0.050	No	
1,1-Dichloroethane	ug/g	0.05	17	50		17	(1)	< 0.050	No	
1,1-Dichloroethene	ug/g	0.05	0.064	50		0.064	(1)	< 0.050	No	
1,2-Dibromoethane	ug/g	0.05	0.05			0.05	(1)	< 0.050	No	
1,2-Dichlorobenzene	ug/g	0.05	6.8			6.8	(1)	< 0.050	No	
1,2-Dichloroethane	ug/g	0.05	0.05			0.05	(1)	< 0.050	No	
1,2-Dichloropropane	ug/g	0.05	0.16			0.16	(1)	< 0.050	No	
1,3-Dichlorobenzene	ug/g	0.05	9.6			9.6	(1)	< 0.050	No	
1,3-Dichloropropene, Total	ug/g	0.05	0.18			0.18	(1)	< 0.050	No	
1,4-Dichlorobenzene	ug/g	0.05	0.2	10		0.2	(1)	< 0.050	No	
2-Butanone	ug/g	0.5	70			70	(1)	< 0.50	No	
4-Methyl-2-pentanone	ug/g	0.5	31			31	(1)	< 0.50	No	
Acetone	ug/g	0.5	16			16	(1)	< 0.50	No	
Benzene	ug/g	0.5	0.32			0.32	(1)	< 0.020	No	
Bromodichloromethane	ug/g	0.02	18			18	(1)	< 0.050	No	
Bromoform	ug/g	0.05	0.61			0.61	(1)	< 0.050	No	
Bromomethane	ug/g	0.05	0.05			0.05	(1)	< 0.050	No	
Carbon Tetrachloride	ug/g	0.05	0.21	50		0.21	(1)	< 0.050	No	
Chlorobenzene	ug/g	0.05	2.4			2.4	(1)	< 0.050	No	
Chloroform	ug/g	0.05	0.47	50		0.47	(1)	< 0.050	No	
cis-1,2-Dichloroethene	ug/g	0.05	55			55	(1)	< 0.050	No	
cis-1,3-Dichloropropene	ug/g	0.03		50	0.398	0.398	(3)	< 0.030	No	
Dibromochloromethane	ug/g	0.05	13			13	(1)	< 0.050	No	
Dichlorodifluoromethane	ug/g	0.05	16			16	(1)	< 0.050	No	
Ethylbenzene	ug/g	0.02	9.5			9.5	(1)	< 0.020	No	
Methyl tert-Butyl Ether	ug/g	0.05	11			11	(1)	< 0.050	No	
Methylene Chloride	ug/g	0.05	1.6	50		1.6	(1)	< 0.050	No	
n-Hexane	ug/g	0.05	46			46	(1)	< 0.050	No	
Styrene	ug/g	0.05	34	50		34	(1)	< 0.050	No	
Tetrachloroethene	ug/g	0.05	4.5	0.6		0.6	(2)	< 0.050	No	
Toluene	ug/g	0.02	68			68	(1)	< 0.020	No	
trans-1,2-Dichloroethene	ug/g	0.05	1.3			1.3	(1)	< 0.050	No	
trans-1,3-Dichloropropene	ug/g	0.04			0.398	0.398	(3)	< 0.040	No	
Trichloroethene	ug/g	0.05	0.91			0.91	(1)	< 0.050	No	
Trichlorofluoromethane	ug/g	0.05	4			4	(1)	< 0.050	No	
Vinyl Chloride	ug/g	0.02	0.032			0.032	(1)	< 0.020	No	
Xylenes, Total	ug/g	0.02	26			26	(1)	< 0.020	No	



**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>121 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table H-2: Screening of Soil COPCs for Ecological Risk Assessment (Continued)**

Parameter	Unit	Detection Limit	MOE 2011 (Table 3 - Industrial)	CCME SQG (Industrial)	US EPA Region 5 RCRA	Soil Screening Level	Ref	Max Soil Conc.	Carried Forward as COPC?	Notes
<b>Semi-VOCs</b>										
1- & 2-Methylnaphthalene	ug/g	0.0071	76			76	(1)	0.018	No	
1-Methylnaphthalene	ug/g	0.005	76			76	(1)	0.0094	No	
2-Methylnaphthalene	ug/g	0.005	76			76	(1)	0.0083	No	
4-Methyl-2-pentanone	ug/g	0.005	31			31	(1)	< 0.50	No	
Acenaphthene	ug/g	0.005	96			96	(1)	< 0.0050	No	
Acenaphthylene	ug/g	0.005	0.15			0.15	(1)	< 0.0050	No	
Anthracene	ug/g	0.005	0.67			0.67	(1)	0.012	No	
Benzo [b,j] fluoranthene	ug/g	0.005	0.96			0.96	(1)	0.11	No	
Benzo[a]anthracene	ug/g	0.005	0.96			0.96	(1)	0.065	No	
Benzo[a]pyrene	ug/g	0.005	0.3			0.3	(1)	0.052	No	
Benzo[g,h,i]perylene	ug/g	0.005	9.6			9.6	(1)	0.031	No	
Benzo[k]fluoranthene	ug/g	0.005	0.96			0.96	(1)	0.022	No	
Chrysene	ug/g	0.005	9.6			9.6	(1)	0.064	No	
Dibenzo[a,h]anthracene	ug/g	0.005	0.1			0.1	(1)	0.0066	No	
Fluoranthene	ug/g	0.005	9.6			9.6	(1)	0.17	No	
Fluorene	ug/g	0.005	62			62	(1)	< 0.0050	No	
Indeno[1,2,3-cd]pyrene	ug/g	0.005	0.76			0.76	(1)	0.036	No	
Naphthalene	ug/g	0.005	9.6			9.6	(1)	< 0.0050	No	
Phenanthrene	ug/g	0.005	12			12	(1)	0.068	No	
Pyrene	ug/g	0.005	96			96	(1)	0.12	No	
Styrene	ug/g	0.05	34	50		34	(1)	< 0.050	No	
Diethylene Glycol	mg/kg	10	-		6200	6200	(4)	< 10	No	
Ethylene Glycol	mg/kg	10	-	960		960	(2)	< 10	No	
Propylene Glycol	mg/kg	10	-	insufficient info	100000	100000	(4)	< 10	No	
Total Glycols	mg/kg	10	-					< 10	No	No toxicity data, non-detect
<b>Radionuclides</b>										
Carbon-14	Bq/kg							-	No	Assessed quantitatively for public interest purposes (H-3, C-14, Cs-134, Cs-137, Co-60)
Carbon-14	Bq/kg-C							557	No	
Cobalt-60	Bq/kg							<1.00	No	
Cesium-134	Bq/kg							<1.00	No	
Cesium-137	Bq/kg							<1.00	No	
Potassium-40	Bq/kg							663	No	
Tritium	Bq/kg							92.4	No	

Notes:

- (1) MOE (2011), Table 3 Standards, Industrial, Coarse soil
- (2) CCME Soil Quality Guidelines, Industrial
- (3) US EPA Region 5 RCRA Ecological Screening Levels, <http://www.epa.gov/Region5/waste/cars/pdfs/ecological-screening-levels-200308.pdf>
- (4) B.C. Reg. 375/96 Contaminated Sites Regulation

Source: Table A13 from P-REP-07701-00001 [R-61]

Report

Public Information		
Document Number:	Usage Classification:	
<b>N-REP-03443-1001</b>	<b>Information</b>	
Sheet Number:	Revision Number:	Page:
<b>N/A</b>	<b>R000</b>	<b>122 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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**Table H-3: Exposure Concentrations for Frenchman’s Bay Exposure Assessment**

Media	VEC Category	COPC	Units	Maximum Concentration	Mean Concentration
Water	Aquatic invertebrate	Aluminum	mg/L	2.70E-01	1.40E-01
	Riparian birds	Bismuth		<1.00E-03	<1.00E-03
	Amphibians	Boron		4.20E-02	3.48E-02
	Riparian mammals	Cadmium		1.00E-05	<1.00E-05
	Aquatic Plants	Calcium		6.40E+01	5.00E+01
		Chromium		<5.00E-03	<5.00E-03
		Copper		2.10E-03	1.69E-03
		Iron		5.60E-01	2.84E-01
		Lead		9.20E-04	<6.09E-04
		Manganese		8.00E-02	4.55E-02
		Nickel		1.30E-03	<1.04E-03
		Sodium		9.10E+01	5.70E+01
		Thallium		<5.00E-05	<5.00E-05
		Tin		<1.00E-03	<1.00E-03
Zinc	7.40E-03	<5.31E-03			
Sediment	Aquatic invertebrate	Aluminum	mg/kg dw	1.30E+04	8.86E+03
	Riparian birds	Bismuth		<1.00E+00	<1.00E+00
	Amphibians	Boron		2.50E+01	7.78E+00
	Riparian mammals	Cadmium		7.50E-01	4.82E-01
	Aquatic Plants	Calcium		1.30E+05	9.07E+04
		Chromium		3.10E+01	2.23E+01
		Copper		7.40E+01	4.53E+01
		Iron		2.10E+04	1.61E+04
		Lead		4.30E+01	2.98E+01
		Manganese		6.60E+02	4.39E+02
		Nickel		2.30E+01	1.64E+01
		Phosphorus		1.50E+03	9.84E+02
		Sodium		5.90E+02	3.80E+02
		Thallium		2.60E-01	1.92E-01
Tin	<5.00E+00	<5.00E+00			
Zinc	2.30E+02	1.65E+02			

Note: Exposure point concentrations are based on measured data from July 2015.  
 VEC = Valued Ecosystem Component  
 COPC = Contaminant of Potential Concern  
 Source: Table E2 from P-REP-07701-00001 [R-61]

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>123 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

**Table H-4: Measured Water Concentrations at Frenchman’s Bay Compared to PN Contribution**

COPC	Frenchman’s Bay Measured Concentration (mg/L)		Pickering Measured Concentration (mg/L)		Estimated Pickering Contribution at Frenchman’s Bay (mg/L)		% Contribution from PN	
	Max	Mean	Max	Mean	Max	Mean	Max %	Mean %
Aluminum	2.70E-01	1.40E-01	9.60E-03	7.27E-03	1.12E-03	8.46E-04	0.41%	0.60%
Bismuth	1.00E-03	1.00E-03	<0.001	<0.001	1.16E-04	1.16E-04	11.64%	11.64%
Boron	4.20E-02	3.48E-02	2.7E-02	2.49E-02	3.14E-03	2.89E-03	7.48%	8.31%
Cadmium	1.00E-05	1.00E-05	1.90E-05	<0.000010	2.21E-06	1.15E-06	22.12%	11.47%
Calcium	6.40E+01	5.00E+01	3.50E+01	3.39E+01	4.07E+00	3.94E+00	6.37%	7.88%
Chromium	5.00E-03	5.00E-03	<0.0050	<0.0050	5.82E-04	5.82E-04	11.64%	11.64%
Copper	2.10E-03	1.69E-03	2.00E-03	1.31E-03	2.33E-04	1.53E-04	11.09%	9.05%
Iron	5.60E-01	2.84E-01	1.00E-01	1.00E-01	1.16E-02	1.16E-02	2.08%	4.10%
Lead	9.20E-04	6.09E-04	<0.0005	<0.0005	5.82E-05	5.82E-05	6.33%	9.56%
Manganese	8.00E-02	4.55E-02	<0.0020	<0.0020	2.33E-04	2.33E-04	0.29%	0.51%
Nickel	1.30E-03	1.04E-03	1.20E-03	1.06E-03	1.40E-04	1.23E-04	10.74%	11.83%
Sodium	9.10E+01	5.70E+01	1.40E+01	1.40E+01	1.63E+00	1.63E+00	1.79%	2.86%
Thallium	5.00E-05	5.00E-05	<0.000050	<0.000050	5.82E-06	5.82E-06	11.64%	11.64%
Tin	1.00E-03	1.00E-03	<0.0010	<0.0010	1.16E-04	1.16E-04	11.64%	11.64%
Zinc	7.40E-03	5.31E-03	5.50E-03	2.07E-03	6.40E-04	5.90E-04	8.65%	11.12%

Source: Table E9 from P-REP-07701-00001 [R-61]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>124 of 132</b>

Title: **2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table H-5: Ichthyoplankton Taxa Collected by Month at DN Based on Counts per Sample**

Scientific Name	Common Name	Dec-15	Jan-16	Feb-16	Mar-16	Apr-16	May-16	Jun-16	Jul-16	Aug-16	Sep-16	Oct-16	Nov-16	Total	Percent of Total Number Entrained
<b>Sample Volume (m<sup>3</sup>)</b>		5002	2075	3894	6128	8960	10875	9225	8604	4182	4435	2279	2392	67497	-
<b>Eggs</b>															
<i>Cottus cognatus/Cottus bairdii</i>	Slimy/Mottled Sculpin	0	0	0	0	2	7	0	0	0	0	0	0	9	0.3%
<i>Neogobius melanostomus</i>	Round Goby	0	0	0	0	3	7	0	0	0	0	0	0	10	0.4%
<i>Sander vitreus</i>	Walleye	0	0	0	0	69	172	0	0	0	0	0	0	241	9.2%
<i>Alosa pseudoharengus</i>	Alew ife	0	0	0	0	0	0	970	1322	59	0	0	0	2351	90.0%
Unidentified	Unidentified	0	0	0	0	0	2	0	0	0	0	0	0	2	0.1%
<b>Total Fish Eggs</b>		0	0	0	0	74	188	970	1322	59	0	0	0	2613	<b>100%</b>
% Entrainment per Month		0.0%	0.0%	0.0%	0.0%	2.8%	7.2%	37.1%	50.6%	2.3%	0.0%	0.0%	0.0%	<b>100%</b>	
Total Fish Eggs Excluding Round Goby		0	0	0	0	71	181	970	1322	59	0	0	0	2603	
% Entrainment per Month Excluding Round Goby		0.0%	0.0%	0.0%	0.0%	2.7%	7.0%	37.3%	50.8%	2.3%	0.0%	0.0%	0.0%	<b>100%</b>	
<b>Larvae</b>															
<i>Neogobius melanostomus</i>	Round Goby	1	0	0	0	10	19	4	1	46	24	2	1	108	88.5%
<i>Myoxocephalus thompsonii</i>	Deepwater Sculpin	0	2	3	3	1	0	0	0	0	0	0	0	9	7.4%
<i>Lota lota</i>	Burbot	0	0	0	0	0	1	0	0	0	0	0	0	1	0.8%
<i>Alosa pseudoharengus</i>	Alew ife	0	0	0	0	0	0	0	0	0	1	0	0	1	0.8%
<i>Osmerus mordax / Alosa pseudoharengus</i>	Rainbow Smelt / Alew ife	0	0	0	0	0	0	2	0	0	0	0	0	2	1.6%
Unidentified	Unidentified	0	0	1	0	0	0	0	0	0	0	0	0	1	0.8%
<b>Total Fish Larvae</b>		1	2	4	3	11	20	6	1	46	25	2	1	122	<b>100%</b>
% Entrainment per Month		0.8%	1.6%	3.3%	2.5%	9.0%	16.4%	4.9%	0.8%	37.7%	20.5%	1.6%	0.8%	<b>100%</b>	
Total Fish Larvae Excluding Round Goby		0	2	4	3	1	1	2	0	0	1	0	0	14	
% Entrainment per Month Excluding Round Goby		0.0%	14.3%	28.6%	21.4%	7.1%	7.1%	14.3%	0.0%	0.0%	7.1%	0.0%	0.0%	<b>100%</b>	
<b>Eggs &amp; Larvae</b>															
<b>Total Ichthyoplankton</b>		1	2	4	3	85	208	976	1323	105	25	2	1	2735	

Source: Table 3.1 from NK38-REP-07260-00005 [R-62]

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>125 of 132</b>

Title:  
**2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS**

**Table H-6: Benthic Invertebrate Taxa Collected by Month at DN (December 7<sup>th</sup>, 2015 to November 22<sup>nd</sup>, 2016)**

Scientific Name	Feb-16	Mar-16	Apr-16	May-16	Jun-16	Jul-16	Aug-16	Sep-16	Oct-16	Nov-16	Total	%of Total Entrained
<b>Sample Volume (m³)</b>	3894	6128	8960	10875	9225	8604	4182	4435	2279	2392	67497	-
<i>Echinogammarus</i>	1025	1905	3674	2414	0	0	0	0	770	2291	12079	54.2%
Other Amphipods (likely <i>Gammarus</i> )	0	0	0	810	659	782	614	2836	1273	0	6974	31.3%
Chironomids (all life stages)	47	916	1159	684	61	19	57	22	28	3	2996	13.4%
<i>Hemimysis</i>	27	15	58	57	7	0	0	14	20	15	213	1.0%
<i>Asellus</i>	0	0	5	0	0	0	0	26	0	0	31	0.14%
Other Isopods	0	1	3	0	0	0	1	2	0	0	7	0.03%
<i>Chaoborus</i>	0	0	1	0	0	0	0	0	0	0	1	0.004%
<b>Total Benthic Invertebrates</b>	1099	2837	4900	3965	727	801	672	2900	2091	2309	22301	100.0%
<b>% Entrainment per Month</b>	4.9%	12.7%	22.0%	17.8%	3.3%	3.6%	3.0%	13.0%	9.4%	10.4%	100.0%	

Source: Table 3.19 from NK38-REP-07260-00005 [R-62]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>126 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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Table H-7: Active Liquid Waste COPC Summary Statistics with Outliers Removed

Active Liquid Waste (ALW)	Units	No. of Outliers	N	NDL	Min	Percentile			Max	Mean	%RSD	
						25th	75th	95th				
Organics	Ethylene Glycol	mg/L	0	24	24	5	5	5	5	20	5.6	54
	Propylene Glycol	mg/L	0	24	24	5	5	5	5	20	5.63	54
	F1 (C6-C10)	µg/L	0	24	24	20	25	25	25	25	25	12
	F2 (C10-C16)	µg/L	0	24	24	100	100	100	100	100	100	3
	Hydrocarbons)											
	F3 (C16-C34)	µg/L	0	24	24	100	200	200	200	200	192	2
	Hydrocarbons)											
	F4 (C34-C50)	µg/L	0	24	24	100	200	200	200	200	192	2
	Hydrocarbons)											
	AEO C8-C9	µg/L	0	23	23	0.03	0.03	0.03	0.03	0.03	0.03	0
	AEO C10-C11	µg/L	0	23	23	0.03	0.03	0.03	0.03	0.03	0.03	0
	AEO C12-C13	µg/L	0	23	2	0.03	1.14	5.73	13	19	4.29	112
	AEO C14-C15	µg/L	0	23	4	0.03	0.1	1.92	2.86	2.93	1.14	7
	AEO C16-C18	µg/L	0	23	23	0.03	0.03	0.03	0.03	0.03	0.03	0
	TOTAL AEOs	µg/L	0	23	2	0.03	1.17	7.91	16	21	5.41	102
	LAS C10	µg/L	0	23	10	0.06	0.06	0.2	0.2	0.2	0.14	51
LAS C12	µg/L	0	23	7	0.06	0.06	4.7	7.81	16	3.18	118	
TOTAL LAS	µg/L	0	23	7	0.06	0.06	4.73	7.88	16	3.21	118	
NP1EC	µg/L	0	23	20	0.01	0.01	0.01	0.01	0.01	0.01	0	
Metals	Mercury (Hg)	µg/L	0	24	22	0.01	0.01	0.01	0.1	0.1	0.02	136
	Total Aluminum (Al)	µg/L	1	23	0	9.72	14.5	22	26	34	18.3357	33
	Total Cadmium (Cd)	µg/L	2	22	2	0.06	0.07	0.10	0.16	0.20	0.09	39
	Total Chromium (Cr)	µg/L	2	22	2	0.1	0.1625	0.33	0.4095	0.42	0.25	42
	Total Copper (Cu)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Extractable Gadolinium (Gd)	µg/L	1	23	9	0.1	2	4.15	9.66	12	3.46	81
	Total Iron (Fe)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Lead (Pb)	µg/L	1	23	0	0.91	1.615	3.25	6.98	7.85	2.76	71
	Total Lithium (Li)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Molybdenum (Mo)	µg/L	1	23	0	0.44	0.85	1.54	2.17	3.52	1.24	55
	Total Nickel (Ni)	µg/L	3	21	0	0.72	1.30	4.65	9.10	13.00	3.59	88
	Total Selenium (Se)	µg/L	2	22	4	0.04	0.05	0.08	0.09	0.09	0.07	25
Total Zinc (Zn)	µg/L	-	-	-	-	-	-	-	-	-	-	
Nutrients	Nitrate	mg/L	-	-	-	-	-	-	-	-	-	-
	Total Phosphorus	mg/L	-	-	-	-	-	-	-	-	-	-
Other	Morpholine	µg/L	0	24	24	1.0	4.0	4.0	4.0	4.0	3.75	82

Note:

AEO= alcohol ethoxylates

LAS= linear alkylbenzene sulphonates

NP1EC= nonylphenol ethoxycarboxylate

NDL= number of values below detection limit (included in statistics at value of detection limit).

%RSD= percent relative standard deviation

Source: Table B2 from NK38-REP-03480-10033 [R-64]



Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>127 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table H-8: Water Treatment Plant COPC Summary Statistics with Outliers Removed

Water Treatment Plant (WTP)	Units	No. of Outliers	N	NDL	Min	Percentile			Max	Mean	%RSD	
						25th	75th	95th				
Organics	Ethylene Glycol	mg/L	-	-	-	-	-	-	-	-	-	
	Propylene Glycol	mg/L	-	-	-	-	-	-	-	-	-	
	F1 (C6-C10)	µg/L	0	24	24	25	25	25	25	25	0	
	F2 (C10-C16 Hydrocarbons)	µg/L	0	24	24	100	100	100	100	100	0	
	F3 (C16-C34 Hydrocarbons)	µg/L	0	24	24	200	200	200	200	200	0	
	F4 (C34-C50 Hydrocarbons)	µg/L	0	24	24	200	200	200	200	200	0	
	AEO C8-C9	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C10-C11	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C12-C13	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C14-C15	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C16-C18	µg/L	-	-	-	-	-	-	-	-	-	
	TOTAL AEOs	µg/L	-	-	-	-	-	-	-	-	-	
	LAS C10	µg/L	-	-	-	-	-	-	-	-	-	
	LAS C12	µg/L	-	-	-	-	-	-	-	-	-	
	TOTAL LAS	µg/L	-	-	-	-	-	-	-	-	-	
NP1EC	µg/L	-	-	-	-	-	-	-	-	-		
Metals	Mercury (Hg)	µg/L	-	-	-	-	-	-	-	-	-	
	Total Aluminum (Al)	µg/L	-	-	-	-	-	-	-	-	-	
	Total Cadmium (Cd)	µg/L	1	23	6	0.01	0.01	0.03	0.04	0.05	0.02	45
	Total Chromium (Cr)	µg/L	0	24	0	1.98	3.25	5.80	8.10	9.91	4.57	44
	Total Copper (Cu)	µg/L	1	23	0	2.73	3.39	4.55	5.34	5.6	4.00	20
	Total Extractable Gadolinium (Gd)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Iron (Fe)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Lead (Pb)	µg/L	0	24	1	0.03	0.07	0.10	0.18	0.20	0.09	45
	Total Lithium (Li)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Molybdenum (Mo)	µg/L	0	24	0	34	40	50	59	66	46.00	18
	Total Nickel (Ni)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Selenium (Se)	µg/L	0	24	0	0.65	1.13	1.46	1.85	2.05	1.30	24
Total Zinc (Zn)	µg/L	0	24	0	1.19	1.67	4.19	7.57	8.55	3.25	67	
Nutrients	Nitrate	mg/L	0	24	0	1.91	2.41	3.92	4.59	4.88	3.17	26
	Total Phosphorus	mg/L	-	-	-	-	-	-	-	-	-	-
Other	Morpholine	µg/L	-	-	-	-	-	-	-	-	-	

Note:

AEO= alcohol ethoxylates

LAS= linear alkylbenzene sulphonates

NP1EC= nonylphenol ethoxycarboxylate

NDL= number of values below detection limit (included in statistics at value of detection limit).

%RSD= percent relative standard deviation

Source: Table B3 from NK38-REP-03480-10033 [R-64]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>128 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table H-9 Inactive Drainage COPC Summary Statistics with Outliers Removed

Inactive Drainage (IAD)	Units	No. of Outliers	N	NDL	Min	Percentile			Max	Mean	%RSD	
						25th	75th	95th				
Organics	Ethylene Glycol	mg/L	0	24	24	5	5	5	5	5.0	0	
	Propylene Glycol	mg/L	0	24	24	5	5	5	5	5	0	
	F1 (C6-C10)	µg/L	-	-	-	-	-	-	-	-	-	
	F2 (C10-C16)	µg/L	-	-	-	-	-	-	-	-	-	
	Hydrocarbons)											
	F3 (C16-C34)	µg/L	-	-	-	-	-	-	-	-	-	
	Hydrocarbons)											
	F4 (C34-C50)	µg/L	-	-	-	-	-	-	-	-	-	
	Hydrocarbons)											
	AEO C8-C9	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C10-C11	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C12-C13	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C14-C15	µg/L	-	-	-	-	-	-	-	-	-	
	AEO C16-C18	µg/L	-	-	-	-	-	-	-	-	-	
	TOTAL AEOs	µg/L	-	-	-	-	-	-	-	-	-	
	LAS C10	µg/L	-	-	-	-	-	-	-	-	-	
LAS C12	µg/L	-	-	-	-	-	-	-	-	-		
TOTAL LAS	µg/L	-	-	-	-	-	-	-	-	-		
NP1EC	µg/L	-	-	-	-	-	-	-	-	-		
Metals	Mercury (Hg)	µg/L	-	-	-	-	-	-	-	-	-	
	Total Aluminum (Al)	µg/L	0	24	0	5.15	8.57	12.00	17.00	19.00	11.00	31
	Total Cadmium (Cd)	µg/L	0	24	6	0.01	0.02	0.03	0.09	4.76	0.22	441
	Total Chromium (Cr)	µg/L	1	23	0	1.98	0.10	0.12	0.17	0.17	0.11	19
	Total Copper (Cu)	µg/L	0	24	0	2.19	2.69	2.83	3.07	3.25	2.73	9
	Total Extractable	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Iron (Fe)	µg/L	0	24	0	14.0	17.0	25.0	29.0	38.0	22.0	26
	Total Lead (Pb)	µg/L	0	24	0	0.03	0.04	0.07	0.08	0.09	0.05	39
	Total Lithium (Li)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Molybdenum (Mo)	µg/L	-	-	-	-	-	-	-	-	-	-
	Total Nickel (Ni)	µg/L	0	24	0	0.41	0.48	0.59	0.73	0.75	0.55	17
	Total Selenium (Se)	µg/L	0	24	0	0.09	0.10	0.12	0.13	0.14	0.11	11
Total Zinc (Zn)	µg/L	-	-	-	-	-	-	-	-	-	-	
Nutrients	Nitrate	mg/L	-	-	-	-	-	-	-	-	-	
	Total Phosphorus	mg/L	-	-	-	-	-	-	-	-	-	
Other	Morpholine	µg/L	-	-	-	-	-	-	-	-	-	

Note:

AEO= alcohol ethoxylates

LAS= linear alkylbenzene sulphonates

NP1EC= nonylphenol ethoxycarboxylate

NDL= number of values below detection limit (included in statistics at value of detection limit).

%RSD= percent relative standard deviation

Source: Table B4 from NK38-REP-03480-10033 [R-64]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>129 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table H-10: Boiler Blowdown COPC Summary Statistics with Outliers Removed

Boiler Blowdown (BB)	Units	No. of Outliers	N	NDL	Min	Percentile			Max	Mean	%RSD
						25th	75th	95th			
Organics	Ethylene Glycol	mg/L	-	-	-	-	-	-	-	-	-
	Propylene Glycol	mg/L	-	-	-	-	-	-	-	-	-
	F1 (C6-C10)	µg/L	-	-	-	-	-	-	-	-	-
	F2 (C10-C16 Hydrocarbons)	µg/L	-	-	-	-	-	-	-	-	-
	F3 (C16-C34 Hydrocarbons)	µg/L	-	-	-	-	-	-	-	-	-
	F4 (C34-C50 Hydrocarbons)	µg/L	-	-	-	-	-	-	-	-	-
	AEO C8-C9	µg/L	-	-	-	-	-	-	-	-	-
	AEO C10-C11	µg/L	-	-	-	-	-	-	-	-	-
	AEO C12-C13	µg/L	-	-	-	-	-	-	-	-	-
	AEO C14-C15	µg/L	-	-	-	-	-	-	-	-	-
	AEO C16-C18	µg/L	-	-	-	-	-	-	-	-	-
	TOTAL AEOs	µg/L	-	-	-	-	-	-	-	-	-
	LAS C10	µg/L	-	-	-	-	-	-	-	-	-
	LAS C12	µg/L	-	-	-	-	-	-	-	-	-
	TOTAL LAS	µg/L	-	-	-	-	-	-	-	-	-
	NP1EC	µg/L	-	-	-	-	-	-	-	-	-
	Metals	Mercury (Hg)	µg/L	-	-	-	-	-	-	-	-
Total Aluminum (Al)		µg/L	-	-	-	-	-	-	-	-	-
Total Cadmium (Cd)		µg/L	0	24	0	0.01	0.01	0.01	0.01	0.01	22
Total Chromium (Cr)		µg/L	0	24	23	0.10	0.10	0.10	0.10	0.44	63
Total Copper (Cu)		µg/L	1	23	1	0.05	0.12	0.25	0.43	0.47	55
Total Extractable Gadolinium (Gd)		µg/L	0	-	-	-	-	-	-	-	-
Total Iron (Fe)		µg/L	1	23	5	1.0	1.0	2.7	6.1	7.0	2.2
Total Lead (Pb)		µg/L	0	24	0	0.06	0.09	0.15	0.17	0.21	0.11
Total Lithium (Li)		µg/L	0	-	-	-	-	-	-	-	-
Total Molybdenum (Mo)		µg/L	0	-	-	-	-	-	-	-	-
Total Nickel (Ni)		µg/L	0	24	17	0.02	0.02	0.02	0.05	0.09	0.03
Total Selenium (Se)		µg/L	0	24	24	0.04	0.04	0.04	0.04	0.04	0.04
Total Zinc (Zn)	µg/L	0	24	0	0.16	0.49	0.84	1.08	1.22	0.68	
Nutrients	Nitrate	mg/L	-	-	-	-	-	-	-	-	-
	Total Phosphorus	mg/L	-	-	-	-	-	-	-	-	-
Other	Morpholine	µg/L	-	-	-	-	-	-	-	-	-

Note:

AEO= alcohol ethoxylates

LAS= linear alkylbenzene sulphonates

NP1EC= nonylphenol ethoxycarboxylate

NDL= number of values below detection limit (included in statistics at value of detection limit).

%RSD= percent relative standard deviation

Source: Table B5 from NK38-REP-03480-10033 [R-64]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>130 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
--

Table H-11: Condenser Cooling Water COPC Summary Statistics with Outliers Removed

Condenser Cooling Water (CCW)	Units	No. of Outliers	N	NDL	Min	Percentile			Max	Mean	%RSD	
						25th	75th	95th				
Organics	Ethylene Glycol	mg/L	0	24	24	5	5	5	5	5	5.0	0
	Propylene Glycol	mg/L	0	24	24	5	5	5	5	5	5	0
	F1 (C6-C10)	µg/L	0	24	24	25	25	25	25	25	25	0
	F2 (C10-C16 Hydrocarbons)	µg/L	0	24	24	100	100	100	100	100	100	0
	F3 (C16-C34 Hydrocarbons)	µg/L	0	24	24	200	200	200	200	200	200	0
	F4 (C34-C50 Hydrocarbons)	µg/L	0	24	24	200	200	200	200	200	200	0
	AEO C8-C9	µg/L	0	24	24	0.03	0.03	0.03	0.03	0.03	0.03	0
	AEO C10-C11	µg/L	0	24	24	0.03	0.03	0.03	0.03	0.03	0.03	0
	AEO C12-C13	µg/L	3	21	7	0.03	0.03	5.52	18.0	21.0	4.01	151
	AEO C14-C15	µg/L	0	24	0	1.04	2.42	8.08	15	16	6.09	76
	AEO C16-C18	µg/L	2	22	14	0.03	0.03	0.12	0.40	0.45	0.10	128
	TOTAL AEOs	µg/L	2	22	0	1.04	3.53	17.25	24.00	47.00	11.77	93
	LAS C10	µg/L	1	23	13	0.06	0.06	0.20	0.20	0.20	0.11	64
	LAS C12	µg/L	2	22	6	0.06	0.07	0.42	0.60	0.92	0.28	82
	TOTAL LAS	µg/L	3	21	6	0.06	0.06	0.40	0.50	0.60	0.25	73
NP1EC	µg/L	0	24	24	0.01	0.01	0.01	0.01	0.01	0.01	0	
Metals	Mercury (Hg)	µg/L	0	24	21	0.01	0.01	0.01	0.01	0.10	0.01	129
	Total Aluminum (Al)	µg/L	1	23	0	2.76	7.03	30.00	69.30	111.00	21.82	121
	Total Cadmium (Cd)	µg/L	2	22	14	0.01	0.01	0.01	0.01	0.01	0.01	31
	Total Chromium (Cr)	µg/L	3	21	3	0.10	0.11	0.15	0.28	0.41	0.15	51
	Total Copper (Cu)	µg/L	1	23	0	0.66	0.85	1.00	1.36	1.48	0.95	21
	Total Extractable	µg/L	0	24	23	2.00	2.00	2.00	2.00	9.60	2.32	67
	Total Iron (Fe)	µg/L	1	23	0	2.0	9.0	56.5	118.9	178.0	35.5	128
	Total Lead (Pb)	µg/L	1	23	0	0.01	0.01	0.07	0.15	0.16	0.05	107
	Total Lithium (Li)	µg/L	1	23	2	0.50	1.76	2.06	2.34	2.49	1.83	27
	Total Molybdenum (Mo)	µg/L	1	23	0	1.13	1.16	1.26	1.32	1.34	1.21	5
	Total Nickel (Ni)	µg/L	0	24	0	0.48	0.53	0.64	0.90	0.93	0.61	21
	Total Selenium (Se)	µg/L	0	24	0	0.13	0.13	0.17	0.17	0.20	0.15	13
	Total Zinc (Zn)	µg/L	1	23	0	0.43	0.77	1.45	2.55	3.75	1.29	60
Nutrients	Nitrate	mg/L	0	24	0	0.15	0.25	0.39	0.41	0.44	0.32	27
	Total Phosphorus	mg/L	0	24	19	0.02	0.02	0.02	0.03	0.03	0.02	10
Other	Morpholine	µg/L	-	-	-	-	-	-	-	-	-	-

Note:

AEO= alcohol ethoxylates

LAS= linear alkylbenzene sulphonates

NP1EC= nonylphenol ethoxycarboxylate

NDL= number of values below detection limit (included in statistics at value of detection limit).

%RSD= percent relative standard deviation

Source: Table B6 from NK38-REP-03480-10033 [R-64]

Report

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>131 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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**Table H-12: Summary of Benthic Invertebrate Community Endpoints at Each Sampling Depth (5, 10, 15 M) and at Each Sampling Area (BH, DN, TP), May, June and August 2016**

	ENDPOINT	BH5	BH10	BH15	DN5	DN10	DN15	TP5	TP10	TP15
May	ABUNDANCE	2,898	2,125	2,007	3,989	4,176	2,518	4,487	4,039	2,373
	RICHNESS	8.6	8.9	8.9	7.6	7.3	8.1	7.5	7.9	7.6
	EVENESS	0.49	0.51	0.55	0.52	0.57	0.58	0.24	0.42	0.46
	SIMPSON'S DIVERSITY	0.79	0.80	0.83	0.79	0.80	0.83	0.56	0.74	0.77
	Bray-Curtis Dissimilarity Index (TP)	0.68	0.55	0.34	0.38	0.29	0.30	0.25	0.23	0.16
	Bray-Curtis Dissimilarity Index (BH)	0.31	0.34	0.22	0.39	0.55	0.28	0.61	0.58	0.31
	Hilsenhoff Biotic Index	5.9	6.0	5.6	6.2	6.0	5.2	6.0	5.9	5.7
June	ABUNDANCE	1,853	1,317	739	2,602	1,862	721	1,324	2,279	1,605
	RICHNESS	7.3	8.2	9.3	7.3	7.8	9.2	6.8	7.2	8.7
	EVENESS	0.33	0.43	0.38	0.36	0.44	0.44	0.33	0.36	0.36
	SIMPSON'S DIVERSITY	0.66	0.73	0.75	0.67	0.76	0.78	0.64	0.68	0.72
	Bray-Curtis Dissimilarity Index (TP)	0.38	0.51	0.49	0.44	0.38	0.56	0.46	0.53	0.46
	Bray-Curtis Dissimilarity Index (BH)	0.24	0.31	0.23	0.32	0.31	0.41	0.46	0.60	0.59
	Hilsenhoff Biotic Index	5.83	6.85	5.97	6.38	7.13	6.56	6.11	5.98	6.22
August	ABUNDANCE	9,120	3,538	2,106	1,484	3,067	2,183	5,587	3,701	2,016
	RICHNESS	8.3	8.3	8.4	8.2	7.9	8.6	7.6	7.3	7.9
	EVENESS	0.32	0.37	0.41	0.36	0.32	0.25	0.36	0.31	0.29
	SIMPSON'S DIVERSITY	0.69	0.70	0.75	0.70	0.66	0.58	0.69	0.59	0.64
	Bray-Curtis Dissimilarity Index (TP)	0.47	0.36	0.37	0.51	0.32	0.40	0.31	0.25	0.22
	Bray-Curtis Dissimilarity Index (BH)	0.33	0.25	0.21	0.70	0.30	0.37	0.46	0.37	0.38
	Hilsenhoff Biotic Index	5.85	6.46	6.42	6.08	6.42	7.08	5.98	6.11	6.28

Note:

TP = Thickson Point

BH = Bond Head

DN = Darlington Nuclear

Endpoints are given as arithmetic mean values

Source: Tables 3-10, 3-11 and 3-12 from NK38-REP-07260-00006 [R-67]

**Report**

Public Information		
Document Number: <b>N-REP-03443-1001</b>	Usage Classification: <b>Information</b>	
Sheet Number: <b>N/A</b>	Revision Number: <b>R000</b>	Page: <b>132 of 132</b>

Title: <b>2017 RESULTS OF ENVIRONMENTAL MONITORING PROGRAMS</b>
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**Appendix I: Compliance with Regulatory Document REGDOC-3.1.1**

The OPG annual EMP report was structured to comply with CNSC regulatory document S-99 *Reporting Requirements for Operating Nuclear Power Plants*. In May 2014, CNSC Regulatory Document REGDOC-3.1.1 *Reporting Requirements for Nuclear Power Plants* was published to replace S-99 [R-3]. It provides revised requirements for an annual report on environmental protection. OPG is required to comply with REGDOC-3.1.1 for the 2015 reporting year and has modified the annual EMP report such that the requirements in section 3.5 of REGDOC-3.1.1 are met. Corresponding sections are summarized in the table below.

**Table I-1: OPG EMP Report Compliance with Regulatory Document-3.1.1, Reporting Requirements for Nuclear Power Plants**

<b>REGDOC-3.1.1, Section 3.5 Requirement</b>	<b>Corresponding Section in OPG's Annual EMP Report</b>
1. A summary of the results of the environmental protection program and an analysis of the significance with respect to health and safety or persons and the protection of the environment, of the results of the environmental protection program	Executive Summary
2. The amount of nuclear substances (i.e. activity concentrations, flow rates and loadings) in SI units, released to the environment and monitored as part of the licensee's effluent/emission monitoring program, presented on an appropriate basis (weekly or monthly), along with a comparison to regulatory release limits for the nuclear substance	Section 2.1
3. The amount of nuclear substances measured in the environment, in SI units, as part of the licensee's radiological environmental monitoring program	Section 3.3 Section 3.4 (if any conducted within that year)
4. The results and calculations of the annual radiation doses to the representative persons and/or critical groups in comparison to the regulatory public dose limit with a description of the environmental transfer/exposure pathways associated with the operation of the nuclear power plant including the dispersion and dosimetric models used	Section 4.0
5. The amount of hazardous substances (i.e. concentrations, flow rates and loadings), in SI units released to the environment and monitored as part of the licensee's effluent/emission monitoring program, and measured in the environment as part of the licensee's environmental monitoring program	Section 2.2 Section 3.4 (if any conducted within that year)
6. For each parameter reported as part of the effluent/emission monitoring and environmental monitoring program, a description of the characteristics of the monitoring results, including but not limited to the sample frequency (e.g. daily, monthly, semi-annually), sample type (e.g. grab, composite, activity counts over time), statistical quantity reported (e.g. weekly/ monthly mean, annual average, annual total)	Section 3.0 Appendix D Section 2.0
7. A description of any significant events, findings or results in respect to the conduct of the environmental monitoring program	Section 5.0
8. A summary of any proposed changes to the environmental monitoring program	Section 6.0