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Guidance on Natural Organic Matter in Drinking Water



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Guidelines for Canadian Drinking Water Quality: Guidance on Natural Organic Matter in Drinking Water

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Guidance on Natural Organic Matter in Drinking Water

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Other Guideline Technical Documents for the Guidelines for Canadian Drinking Water Quality can be found on the following web page: www.canada.ca/en/health-canada/services/environmental-workplace-health/reports-publications/water-quality.html

Background on guidance documents

Health Canada works with the provinces, territories and federal agencies to establish the Guidelines for Canadian Drinking Water Quality. Over the years, new methodologies and approaches have led Health Canada, in collaboration with the Federal-Provincial-Territorial Committee on Drinking Water, to develop a new type of document, guidance documents, to provide advice and guidance on issues related to drinking water quality for parameters that do not require a formal Guideline for Canadian Drinking Water Quality.

Guidance documents are developed to provide operational or management guidance related to specific drinking water-related issues (e.g., boil water advisories), to make health risk assessment information available when a guideline is not deemed necessary.

Guidelines are established under the Guidelines for Canadian Drinking Water Quality specifically for contaminants that meet all of the following criteria:

1. exposure to the contaminant could lead to adverse health effects;
2. the contaminant is frequently detected or could be expected to be found in a large number of drinking water supplies throughout Canada; and
3. the contaminant is detected, or could be expected to be detected, at a level that is of possible health significance.

If a contaminant of interest does not meet all these criteria, Health Canada, in collaboration with the Federal-Provincial Territorial Committee on Drinking Water, may choose not to develop a Guideline Technical Document. In that case, a guidance document may be developed.

Guidance documents undergo a similar process as Guideline Technical Documents, including public consultations through the Health Canada Web site. They are offered as information for drinking water authorities and to help provide guidance in spill or other emergency situations.

Executive Summary

Natural organic matter (NOM) is an extremely complex mixture of organic compounds and is found in all groundwater and surface waters. Although NOM has no direct impact on health, it affects the efficacy of drinking water treatment processes and consequently the safety of drinking water. NOM may also affect consumer satisfaction because it can contribute to undesirable colour, tastes and odours in drinking water.

Health Canada completed its review of NOM in drinking water and the impact that it can have on drinking water treatment processes. This guidance document was prepared in collaboration with the Federal-Provincial-Territorial Committee on Drinking Water and reviews and assesses risks associated with the impact of NOM on drinking water treatment processes and the safety of drinking water.

Assessment

The health effects of NOM are due to its impact on drinking water treatment processes that are aimed to protect drinking water quality and public health. NOM can impact processes designed to remove or inactivate pathogens, contribute to the formation of disinfection by-products and favour the development of biofilms in the distribution system. Its presence may also create conditions that result in increased lead and/or copper concentrations in treated water due to its influence on corrosion.

The treatability and reactivity of NOM vary significantly in Canada, as each water source has unique features. Because NOM consists of numerous organic compounds, it cannot be measured directly. However, there are a number of other parameters that can be used to provide an indication of the concentration and character (i.e., chemical, physical and biodegradability properties) of NOM. It is important to understand variations in NOM concentrations and character in order to select, design and operate appropriate water treatment processes.

No practical health-based value can currently be derived for NOM in drinking water. The development of an effective NOM control strategy needs to be based on a good understanding of:

- variations in the concentration and character of NOM in the source water, including those due to climate change, landscape changes or source water protection programs;
- NOM's impact on water treatment processes and the impact of water treatment on NOM, for the full range of water quality conditions; and
- its potential impacts on water quality in the distribution system.

Source-specific treatability studies, including bench- and/or pilot-scale testing, are essential to determine the most effective treatment option(s) to remove NOM, decrease its reactivity to form disinfection by-products, reduce its potential to contribute to corrosion, and produce biologically stable water for distribution. The lack of a source-specific treatability study may result in the selection of inappropriate treatment, an increase in disinfection by-product concentrations following the implementation of treatment or other unintended consequences. As water sources or treatment processes can change over time, it is important to routinely monitor the concentration and character of NOM and to evaluate its impact on treatment, water quality and distribution system conditions.

The intent of this document is to provide provinces, territories, other government departments and stakeholders (such as water system owners, consultants, equipment suppliers

and laboratories) with guidance on the impacts of NOM on the overall quality of drinking water, including its potential effects on drinking water treatment processes and consequently on the safety of drinking water. It summarizes the factors that affect the concentration and character of NOM and discusses the points to consider when developing a NOM control strategy. It also provides specific guidance on treatment, monitoring, and water quality goals.

International considerations

Drinking water guidelines, standards and/or guidance from other national and international organizations may vary due to the date of the assessments as well as differing policies and approaches.

International organizations have not established numerical limits for NOM in drinking water. The United States Environmental Protection Agency's rule for disinfectants and disinfection by-products requires removal of total organic carbon (TOC) by surface water facilities using conventional or lime softening water treatment with levels of TOC above 2 mg/L in their source water. The World Health Organization suggests optimized NOM removal as a means to minimize biofilm growth in the distribution system. The European Union regulations include TOC as a general water quality indicator; in some jurisdictions, chemical oxygen demand (COD) can be used in place of TOC. In Australia, guidance has been developed for water utilities to help them understand and control the impact of NOM.

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Part A. Guidance on natural organic matter in drinking water

A.1 Introduction

Natural organic matter (NOM) is an extremely complex mixture of organic compounds that vary greatly in terms of their physical and chemical characteristics. NOM occurs naturally in the environment and may also be the result of human activities. NOM is found in particulate, colloidal and dissolved forms in all ground and surface waters, as well as in rainwater. While exposure to NOM in the environment is commonplace and is not associated with direct health effects, the presence and characteristics of NOM will have significant impacts on drinking water treatment processes aimed at protecting public health. NOM plays a critical role in drinking water treatment for a number of reasons. First and foremost, NOM can contribute indirectly to health impacts in many ways, including:

- it exerts a coagulant demand which can lead to suboptimal coagulation conditions and a deterioration of pathogen log removal capability;
- it exerts a chemical disinfectant demand or interferes with ultraviolet (UV) disinfection which can lead to a deterioration of pathogen log inactivation capability;
- it forms regulated and non-regulated disinfection by-products (DBPs) when it reacts with disinfectants;
- it favours the development of distribution system biofilms that can harbour pathogens; and
- it influences corrosion and may create conditions that result in increases in lead and/or copper concentrations as a result of corrosion of lead- and/or copper-bearing materials (e.g., piping, fittings).

Water utilities can also be significantly impacted by a number of NOM-caused operational issues, namely:

- increased coagulant dose;
- poor floc formation or settling;
- shorter filter run times;
- more frequent backwashes;
- increased sludge production;
- reduced hydraulic capacity;
- membrane fouling, higher transmembrane pressure and energy consumption, more frequent chemical cleaning and shorter membrane life; and
- reduced effectiveness of adsorption and ion exchange processes.

NOM can also lead to an increase in consumer complaints because it can contribute to undesirable colour, tastes and odours in drinking water. These and other problems are further discussed in subsequent sections of this document.

A.2 Application

All water utilities should implement a risk management approach, such as the source-to-tap or water safety plan approach, to ensure water safety. These approaches require a system assessment that involves characterizing the water source, describing the treatment barriers that prevent or reduce contamination, highlighting the conditions that can result in contamination, and identifying control measures. Operational monitoring is then established and operational/management protocols are instituted (e.g., standard operating procedures, corrective actions and

incident responses). Compliance monitoring is determined and other protocols to validate the water safety plan are implemented (e.g., record keeping, consumer satisfaction). Operator training is also required to ensure the effectiveness of the water safety plan at all times.

When developing and implementing a risk management approach, it is important to understand how NOM can indirectly result in health impacts. NOM can increase prior to changes in turbidity and flow and can remain elevated after turbidity and flow have returned to baseline conditions. Thus, changes in NOM may go undetected and a deterioration in pathogen log removal or inactivation may occur if appropriate monitoring is not in place. The goal of the NOM control strategy should be to ensure protection from microbial risks at all times, while minimizing DBP, lead and copper concentrations and controlling biofilm formation in the distribution system.

The water system owner should strive at all times to appropriately characterize NOM and adequately remove it to achieve water quality goals. Water utilities may require multiple treatment processes to effectively balance microbial and chemical risks throughout the year. Water system owners should contact the appropriate drinking water authority in the affected jurisdiction to confirm if specific requirements will apply to their source/system.

A.2.1 Source-specific treatability study

Source-specific treatability studies are recommended to determine the most effective treatment option(s) to adequately remove NOM and to meet water quality goals related to microbial risks, DBPs, biological stability and corrosion control. Developing a strong understanding of the source water is essential to ensure a reliable, robust and resilient treatment strategy is selected. Source-specific monitoring prior to facility design is necessary to assess seasonal variations in NOM and forecast extreme conditions due to changes in climate. The treatability study should include bench- and/or pilot-scale testing, as well as DBP formation potential tests that are representative of distribution system conditions.

A.2.2 Monitoring

The concentration and character of NOM should be monitored in raw, treated and distribution system water to ensure that:

- treatment is optimized for NOM and turbidity removal;
- DBP, lead and copper concentrations are as low as reasonably achievable; and
- biofilm formation is minimized.

A source-specific monitoring plan should be developed to ensure that water utilities are aware of:

- raw water quality changes with respect to NOM concentration and character;
- the impact that NOM has on water treatment processes through all water quality conditions;
- the impact that treatment has on NOM concentration and character; and
- the impacts on distribution water quality.

The monitoring plan should be comprehensive and include source characterization, operational and compliance monitoring; it should also demonstrate that water quality goals are consistently met for microbial risks, DBPs, biological stability and corrosion control. Ideally, continuous online monitoring should be used for highly variable sources (i.e., those that fluctuate with precipitation/snowmelt events) and critical processes (e.g., coagulation).

A.2.2.1 Source water assessments

Source water assessments should be part of routine system assessments. They should include an understanding of NOM sources in the watershed/aquifer, the conditions that lead to changes in the concentration and/or character of NOM (e.g., precipitation/snowmelt events, algal blooms, drought, fire), and the factors that enhance the reactivity of NOM to form DBPs (e.g., reaction conditions, water age, and inorganic compounds such as ammonia, bromide, iodide, and sulphur).

Surface and subsurface sources should be characterized with regard to NOM and inorganic compounds. The frequency of source water characterization monitoring will depend on the variability of the source; highly variable sources should be monitored more frequently.

A.2.2.2 Treatment and operational monitoring

The concentration and/or character of NOM can have a significant influence on the selection, design and operation of water treatment processes. In order to determine the most appropriate treatment processes, water utilities should have knowledge about:

- the origin, occurrence and fluctuations in NOM;
- interactions between NOM and other water constituents (e.g., enhanced reactivity due to bromide);
- interactions with chemicals used during treatment (e.g., NOM creates a disinfectant and coagulant demand that must be overcome to produce microbiologically safe drinking water);
- interactions between NOM and unit processes (e.g., NOM fouls adsorbents and membranes); and
- its impacts on distribution system water quality (e.g., DBPs and biological stability).

The appropriate type and level of treatment should take into account source-specific fluctuations in water quality, including seasonal and/or short-term degradation, variability in treatment performance and distribution system conditions. Ongoing operational monitoring and treatment optimization will help ensure that water utilities achieve water quality goals and maximize public health protection for the full range of water quality conditions. Maintaining current knowledge of best practices and remaining aware of advancements in the drinking water industry are important aspects of the source-to-tap or water safety plan approach to ensure water safety.

A.2.2.3 Distribution system

Biodegradable organic matter (BOM) encourages biofilm growth in the distribution system. Biofilms can provide a habitat for the survival of pathogens that may have passed through drinking water treatment barriers or entered the distribution system directly via an integrity breach. The most important elements for controlling the growth of bacteria in distribution systems are maintenance of a disinfectant residual, limitation of BOM, and corrosion control. Maintaining the physical/hydraulic integrity of the distribution system and minimizing negative- or low-pressure events are other key components of a source-to-tap or water safety plan approach.

Distribution system water quality should be regularly monitored, including DBPs and biological stability indicators (e.g., variability of disinfectant residual, biofilm formation rate, corrosion rate). Operations/maintenance programs should also be in place (e.g., water age control, water main cleaning, cross-connection control, asset management) and strict hygiene should be practiced during all water main construction (e.g., repair, maintenance, new installation) to ensure drinking water is transported to the consumer with minimal loss of quality.

A.2.2.4 Suggested parameters and frequencies

Table 1 outlines suggested parameters, sampling locations and frequencies that can form the basis of a comprehensive monitoring program. Many of the listed parameters (e.g., disinfectant residual, DBPs) are already being monitored in most treatment facilities as part of a source-to-tap approach to producing safe drinking water. Other parameters are relatively easy to use (e.g., UV absorbance or transmittance) and provide rapid results. Suggested water quality targets are outlined in Table 2. These are suggested as guidance only based on the literature review that was completed to develop this document. As some water sources can be extremely reactive (e.g., form more DBPs), more stringent water quality targets may be required.

Water utilities should employ the most appropriate methods and parameters to routinely monitor raw, treated and distribution system water quality, establish baseline conditions and detect changes that require process modifications. Systems that exhibit low DBP concentrations, have stable biological water quality (e.g., biostability) and baseline data indicating that NOM does not influence corrosion may consider reduced monitoring.

Table 1. Suggested parameters to monitor

Parameter	Location	Frequency		
		Variable source	Stable source	Ideal
Organic colour (true colour)	Raw and treated	Daily	Weekly	Online
UV absorbance (at 254 nm) or UV transmittance	Raw and filtered ^a	Daily	Weekly	Online
Chemical oxygen demand (COD)	Raw, treatment processes ^b and treated	Daily	Weekly	Online
Dissolved or total organic carbon (DOC or TOC)	Raw and treated ^a	Weekly	Monthly	Online
Specific UV absorbance (SUVA)—calculate from UV ₂₅₄ and DOC	Raw and treated ^a	Weekly	Monthly	Daily
Inorganic compounds that can enhance the reactivity of NOM to form DBPs: – Ammonia – Bromide – Iodide – Sulphur	Raw and treated	Quarterly Quarterly Quarterly Quarterly	Quarterly Quarterly Quarterly Quarterly	Quarterly Quarterly Quarterly Quarterly
Coagulant demand	Coagulation process ^c	Daily	Daily	Online
Zeta potential or streaming current—when NOM controls or influences coagulant dose	Coagulation process ^c	Online	Online	Online
Disinfection by-products (DBPs)	Distribution system	Quarterly (measure DOC and inorganic compounds on same day to calculate specific DBP yields to assess NOM reactivity)		

Parameter	Location	Frequency		
		Variable source	Stable source	Ideal
Biological stability: <ul style="list-style-type: none"> – Disinfectant residual – Biofilm formation rate—measured by adenosine triphosphate (ATP) accumulated on mild steel coupons – Corrosion rate—measured by linear polarization resistance using mild steel coupons 	Distribution system	Weekly Every two weeks Monthly	Weekly Monthly Monthly	Online
Influence of NOM on corrosion: <ul style="list-style-type: none"> – Lead – Copper 	In accordance with corrosion control program In accordance with corrosion control program			

- ^a Disinfection will reduce UV absorbance without an associated reduction in DOC. Thus to calculate the treated water SUVA, UV absorbance at 254 nm (UV254) should be measured in filtered water pre-disinfectant addition and divided by the treated water DOC, then multiplied by 100.
- ^b COD decreases across each treatment process. Monitoring locations will vary depending on the process trains in place (e.g., flocculation, clarification, filtration) and the water utility's continuous improvement program.
- ^c Strict pH control is critical for NOM removal. As alkalinity affects pH control, pH and alkalinity are other important coagulation process monitoring parameters.

Table 2. Suggested treated water quality targets

Parameter	Units	Source with high specific DBP yield or extensive distribution system	Source with low specific DBP yield
Organic colour	TCU	5–10	<15
UV absorbance (at 254 nm)	cm ⁻¹	0.02–0.04	0.02–0.07
UV transmittance	Percent	90–95	85–95
COD	mg/L O ₂	<5	<5
DOC—for DBP control	mg/L C	<2	<4
DOC—for biological stability	mg/L C	<1.8	<1.8

C = carbon

COD = chemical oxygen demand

DOC = dissolved organic carbon

O₂ = oxygen

TCU = true colour units

Part B. Supporting information

B.1 Description of natural organic matter

NOM is an extremely complex mixture of organic compounds varying in polarity, acidity, charge density, and molecular mass; NOM can also range from biodegradable (i.e., labile or semi-labile) to less biodegradable (i.e., recalcitrant or refractory). Because NOM comprises numerous organic compounds, it can be categorized based on its polarity (i.e., hydrophobic or hydrophilic) and acid/neutral/base properties. This approach results in six NOM fractions, as outlined in Table 3. Compound classes within these fractions have also been identified. Compound classes provide the highest level of specificity possible, due to the number of compounds that can be present (Minor et al., 2014).

The size and shape of NOM is influenced by the pH and ionic strength of the water; at low pH and high ionic strength, NOM can have a rigid, compact, coil shape whereas at high pH and low ionic strength, it can have a flexible linear filament shape (Ghosh and Schnitzer, 1980; Braghetta et al., 1997). Some compounds can exhibit both hydrophobic and hydrophilic properties (i.e., amphipathic) (Leenheer and Croué, 2003) and possess both negative- and positive-charged functional groups (i.e., amphoteric) (Ghosh and Schnitzer, 1980; Braghetta et al., 1997; Her et al., 2007; Amy, 2008). Fractions containing polysaccharides, proteins and amino sugars have the highest molecular weights (>10 kDa), whereas the molecular weights of humic and fulvic acids typically range from 2 kDa to 5 kDa and from 0.5 kDa to 2 kDa, respectively (Bond et al., 2012; Sillanpää et al., 2015a). Lignin and tannin derivatives are also abundant in the high to medium molecular weight fractions. The smallest NOM fractions (<0.5 kDa) tend to be hydrophilic compounds (Sillanpää et al., 2015a). The most biodegradable fractions include carbohydrates, amino acids and proteins while the most recalcitrant comprise lignins, tannins and terpenoids.

Table 3. NOM fractions and compound classes^a

Fraction	Compound classes
<i>Hydrophobic</i>	
Acids	<u>Strong Acids</u> Humic and fulvic acids, high molecular weight alkyl monocarboxylic and dicarboxylic acids, aromatic acids
	<u>Weak acids</u> Phenols (e.g., lignin), tannins, medium molecular weight alkyl monocarboxylic and dicarboxylic acids
Bases	Proteins, aromatic amines, high molecular weight alkyl amines
Neutrals	Hydrocarbons (e.g., terpenoids), aldehydes, high molecular weight methyl ketones and alkyl alcohols, ethers, furans, pyrrols
<i>Hydrophilic</i>	
Acids ^b	Hydroxyl acids, sugars, sulphonics, low molecular weight alkyl monocarboxylic and dicarboxylic acids
Bases	Amino acids, purines, pyrimidines, low molecular weight alkyl amines
Neutrals	Proteins, carbohydrates (e.g., polysaccharides, low molecular weight alkyl alcohols, aldehydes and ketones), cellulose and cellulose derivatives

^a Adapted from Stevenson, 1982; Thurman, 1985; Edzwald, 1993; Imai et al., 2001; Leenheer and Croué, 2003; Reckhow et al., 2007; Amy, 2008; Bond et al., 2011

^b Hydrophilic acids can also be reported as transphilic NOM (Martin-Mousset et al., 1997)

B.2 Sources and occurrence of natural organic matter

The concentration and character (i.e., chemical, physical and biodegradability properties) of NOM can be highly variable because of the numerous hydrological and biogeochemical processes that affect the sources of NOM (Aiken and Cotsaris, 1995). This is briefly described in the following sections, along with parameters that have historically been used to quantify organic matter, including: 1) organic colour, as a measure of humic and fulvic acids; and 2) organic carbon, the key constituent of NOM (Thurman, 1984). Other parameters that can be used to measure and characterize NOM are discussed later in this document.

B.2.1 Sources

There are two natural sources of NOM: allochthonous (i.e., derived from the terrestrial ecosystem) and autochthonous (i.e., derived from the plants and microorganisms growing in the water body) (Aiken and Cotsaris, 1995). Anthropogenic (human) activities can also contribute to NOM.

Allochthonous NOM is exported to aquatic environments as precipitation moves through the atmosphere and vegetative canopy, infiltrates organic soil layers and percolates downward through mineral soil layers (Aitkenhead-Peterson et al., 2003). Soil humus, plant litter, microbial biomass and root exudates contribute to allochthonous NOM (Kalbitz et al., 2000). Allochthonous NOM tends to be hydrophobic in nature. These and other factors that influence the concentration and character of allochthonous NOM are described in Table C-3.1 of this document.

Autochthonous NOM is derived from phytoplankton, algae, cyanobacteria and macrophytes (i.e., plants attached to or rooted in the substrata of lakes and streams) and can account for 5–100% of the DOC concentration, depending on certain conditions (Bertilsson and Jones, 2003; Wetzel, 2003; Bade et al., 2007; Tomlinson et al., 2016). When allochthonous inputs are high, such as in coloured water sources or during precipitation/snowmelt events (i.e., stormflow conditions), the proportion of autochthonous NOM tends to be low. Conversely, when allochthonous inputs are low, such as in clear water sources or during dry periods when there is little runoff, the proportion of autochthonous NOM tends to be high. Autochthonous NOM encompasses a wide range of compounds: mono- and polysaccharides, amino acids, peptides, proteins, nucleic acids, organic acids, lipids and fatty acids (Pivokonsky et al., 2006; Henderson et al., 2008). DOC is generated by the production and decomposition of the microbial and plant biomass within water sources (Nguyen et al., 2002; Zhou et al., 2014). Algal inputs tend to dominate in large lakes, whereas macrophytes tend to be the major contributor in small lakes (Wetzel, 1992; Bertilsson and Jones, 2003). Algal and cyanobacterial blooms, in particular, represent a source of DOC that can be periodic and intense. Cyanobacterial blooms may be associated with additional water quality issues due to the potential presence of cyanobacterial toxins. Autochthonous NOM tends to be hydrophilic in nature and nitrogen-rich.

Anthropogenic sources of NOM include septic systems, wastewater treatment and stormwater discharges, agricultural runoff and industrial discharges. Anthropogenic NOM is reported to be hydrophilic in nature (Imai et al., 2001) and nitrogen-rich (Dotson and Westerhoff, 2009; Mitch et al., 2009). Watersheds heavily impacted by anthropogenic sources may observe a decrease in TOC or DOC after the improvement of wastewater or stormwater treatment (Reckhow et al., 2007).

Raw water NOM concentrations represent the net effect of hydrological and biogeochemical processes in the watershed or aquifer (Eckhardt and Moore, 1990). The concentration and character of NOM, and therefore its treatability (i.e., potential to be removed)

and reactivity (i.e., potential to form DBPs), vary significantly from one source to another because each water source has unique features. For example, Kerekes et al. (1982) reported that two lakes in Nova Scotia only 1 km apart had TOC concentrations of 5.6 and 17.2 mg/L, respectively. In the low TOC lake, the retention time was 1.27 years and organic soils were absent, whereas in the high TOC lake, the retention time was 0.35 years and organic soils were present. Longer retention times tend to lower the DOC concentration, as noted in Table C-3.1. However, Curtis and Adams (1995) reported that the evapoconcentration of refractory NOM resulted in increased DOC concentrations with increasing retention time in the sub-humid and semi-arid zones of Alberta. Other researchers have reported similar findings regarding the variability and uniqueness of NOM for sources in close proximity to each other (Aiken and Cotsaris, 1995; Ågren et al., 2007; Reckhow et al., 2007; Goss and Gorczyca, 2013; Kent et al., 2014).

NOM concentrations are typically lower in groundwater sources because the organic matter is subjected to adsorption and microbial degradation processes as it is transported through the soil (Thurman, 1985; Aiken and Cotsaris, 1995; Aitkenhead-Peterson et al., 2003). However, these processes are limited by the amount of biodegradable NOM that is present. Conversely, some groundwater flows through aquifer materials that are rich in organic matter resulting in high organic carbon concentrations (Thorstenson et al., 1979; Hem, 1985; Aravena et al., 1995; Lemieux et al., 2019). According to published sources, organic carbon concentrations in some North American groundwaters range from <0.1–22 mg/L (see Table C-3.2). Seasonal variability can occur on a per well basis hence reliance on a single sample to represent groundwater quality may be misleading (Washington State Department of Health and University of Washington, 2017).

The NOM in groundwater tends to be more hydrophilic and recalcitrant in nature (Diem et al., 2013) and almost as reactive as surface water NOM on a mg/L DOC basis (Owen et al., 1995; Reckhow et al., 2007; Tubić et al., 2013). For example, the Washington State Department of Health and University of Washington (2017) reported specific DBP yields of 67 µg trihalomethanes (THMs)/mg DOC and 29 µg haloacetic acids (HAAs)/mg DOC for small groundwater supplies using chlorination. Groundwater sources can have higher concentrations of bromide (Chowdhury, 2018) or iodine (Lemieux et al., 2019) which can contribute to their reactivity (i.e., potential to form DBPs).

In summary, localized conditions play a very significant role in establishing the concentration and character of NOM (Bourbonniere, 1989; Mulholland, 2003; Reckhow et al., 2007; Sillanpää, 2015). Occurrence data presented below highlight the variability that can occur, with or without an associated change in DOC concentration. The data also show that although groundwater tends to have lower NOM concentrations, some sources can have elevated concentrations. Thus, both surface and subsurface sources should be characterized.

B.2.2 Occurrence

The concentration and character (i.e., chemical, physical and biodegradability properties) of NOM can be highly variable because of the numerous hydrological and biogeochemical processes that export, generate or degrade NOM, as described in Table C-3.1.

B.2.2.1 Concentration

The monitoring data that were available for this review are summarized below (see Tables 4 and 5). Non-detect data were excluded from the statistical analysis. Results are presented to show the variability in detectable concentrations that can occur spatially.

Monitoring data from the provinces and territories for TOC and colour in raw water are summarized in Table 4. These data demonstrate that TOC concentrations and organic colour vary spatially and tend to be lower in groundwater (although some groundwater sources can have elevated concentrations). Table 5 presents the DOC monitoring data collected in 2009 and 2010 from select drinking water sources in every region of Canada (Health Canada, 2016). These data also demonstrate lower DOC concentrations in groundwater with minimal change between raw and treated water concentrations. For the surveyed surface water supplies, average treated water DOC ranges from 3.2–3.4 mg/L in summer and 2.8–3.5 mg/L in winter. As some jurisdictions are not represented in Table 4, Table C-3.3 presents Environment Canada’s long-term DOC surface water monitoring data (2000–2015) for select regions or river basins across Canada (Environment Canada, 2017). These data also demonstrate that there is significant variability in NOM concentrations spatially.

Table 4. Total organic carbon and organic colour data for raw water from select jurisdictions in Canada

Jurisdiction ^a		TOC ^b (mg/L)		Colour ^c (TCU)	
		Ground	Surface	Ground	Surface
Newfoundland and Labrador ^b (MDL ^d = 0.5 mg/L) (MDL = 2 TCU)	No. detects/samples	322/350	833/833	204/350	832/833
	Median	1.2	6.5	6.0	43.0
	Mean	2.0	7.0	14.3	53.5
	90th percentile	4.3	11.4	35.0	107.0
Nova Scotia (MDL = 0.5 mg/L) (MDL = 5 TCU)	No. detects/samples	53/126	136/140	50/152	142/149
	Median	1.2	4.6	7.4	27.5
	Mean	2.3	5.8	11.7	43.5
	90th percentile	6.7	10.9	22.0	86.7
New Brunswick (MDL = 0.2–1.0 mg/L) (MDL = 1-5 TCU)	No. detects/samples	893/1,389	324/324	86/235	37/45
	Median	2.0	4.8	3.0	28.0
	Mean	2.1	4.8	10.2	31.1
	90th percentile	3.4	6.0	16.0	48.6
Quebec (MDL = 0.2 mg/L) (MDL = 1 TCU)	No. detects/samples	129/129	91/91	No data provided	5/5
	Median	2.8	6.0		52.0
	Mean	3.1	6.2		53.2
	90th percentile	5.1	9.7		66.0
Manitoba (MDL = 0.5–1.0 mg/L) (MDL = 5 TCU)	No. detects/samples	564/723	456/458	225/721	433/458
	Median	2.9	10.9	10.0	26.2
	Mean	4.0	11.6	14.0	31.5
	90th percentile	8.2	16.2	30.0	60.0

^a Data is summarized for jurisdictions that provided information.

^b For Newfoundland and Labrador, data is for dissolved organic carbon.

^c Colour is in true colour units (TCU); data represents filtered samples whereby suspended particles (e.g., clay, iron and manganese oxides) that can give the appearance of colour are removed.

^d MDL = method detection limit

Table 5. Dissolved organic carbon data from Health Canada National Survey (2009–2010)^b

Source Type	Sample Type	Summer DOC (mg/L) ^a				Winter DOC (mg/L) ^a			
		n	Median	Mean	90th Percentile	n	Median	Mean	90th Percentile
Well	Raw	18	0.7	1.8	4.0	16	0.7	2.0	4.4
	Treated	17	0.8	1.6	4.3	15	0.8	1.7	4.5
Lake	Raw	21	4.0	7.3	11.5	20	4.6	6.6	9.7
	Treated	21	2.8	3.2	5.2	20	2.4	3.5	5.6
River	Raw	26	5.9	7.2	14.2	21	4.7	5.8	10.0
	Treated	26	2.6	3.4	6.0	21	2.6	2.8	5.4

^a Method detection limit of 0.2 mg/L

^b Health Canada, 2016

B.2.2.2 Character

A number of studies have characterized the six NOM fractions (see Table 3) found in several Canadian source waters (Montreuil, 2011; Newfoundland and Labrador Department of Environment and Conservation, 2011; Lamsal et al., 2012; Goss and Gorczyca, 2013). These studies are summarized in Table C-3.4. Montreuil (2011) studied a lake source in Nova Scotia and observed significant temporal variability in the six NOM fractions while DOC concentrations remained low with minimal change. Goss and Gorczyca (2013) studied a river source in Manitoba and also observed significant temporal variability in the six NOM fractions but with fluctuations in DOC concentrations. The other two studies demonstrate that NOM character can vary significantly by location.

The results of these fractionation studies demonstrate the variability that can occur in NOM character—with or without an associated change in DOC concentration. In addition, the results indicate that the hydrophilic neutral fraction can, at times, comprise a significant portion of NOM. This fraction can be particularly problematic, as discussed in subsequent sections.

NOM can also be fractionated using more rapid methods or according to its size or fluorescence, as outlined in section B.5.2.4. The biodegradability properties of NOM are discussed in detail in subsequent sections.

B.3 Environmental considerations

Environmental factors may change the NOM concentration and/or the relative contribution of allochthonous, autochthonous or anthropogenic inputs and thereby change its character. These changes can impact water sources and water treatment processes, as discussed below.

B.3.1 Seasonal or weather-related effects

A number of researchers have reported an increase in NOM concentration and a change in its character following snowmelt, spring runoff or heavy rain (Gregory, 1998; Billica and Gertig, 2000; Tseng et al., 2000; Goslan et al., 2002; Volk et al., 2002; Eikebrokk et al., 2004; Fearing et al., 2004a, 2004b; Hurst et al., 2004; Chow et al., 2006; Sharp et al., 2006; Parsons et al., 2007; Reckhow et al., 2007; Kraus et al., 2010; Carpenter et al., 2013; Kundert et al., 2014; McVicar et al., 2015; James et al., 2016). DOC concentrations can rapidly increase four- to five-fold during precipitation/snowmelt events that flush terrestrial NOM into a water body (Thurman, 1985; Saraceno et al., 2009). The highest concentrations can occur in the summer and autumn when temperatures are warmer, biological activity is high and high-intensity/short-duration rainstorms

are frequent (Aitkenhead-Peterson et al., 2003).

Precipitation and snowmelt events can significantly impair the coagulation process for a number of reasons. First, water quality changes occur during stormflow conditions that create challenging coagulation conditions (e.g., pH, alkalinity, ionic strength, divalent ion concentration) (Gregory, 1998; Billica and Gertig, 2000; Tseng et al., 2000; Davis and Edwards, 2014). Second, NOM has been observed to increase prior to changes in turbidity or flow and can remain elevated after turbidity and flow have returned to baseline conditions (Soulsby, 1995; Hurst et al., 2004; McVicar et al., 2015; James et al., 2016). Thus, if the coagulant dose is controlled based on flow or turbidity, coagulant may be under-dosed, leading to suboptimal coagulation conditions (Hurst et al., 2004; Parsons et al., 2007; Kundert et al., 2014; McVicar et al., 2015; James et al., 2016). It is well known that suboptimal coagulation conditions lead to a significant deterioration in pathogen log removal credits (Ongerth and Pecoraro, 1995; Patania et al., 1995; Edzwald and Kelley, 1998; Coffey et al., 1999; Emelko et al., 1999, 2005; Dugan et al., 2001; Harrington et al., 2001; Huck et al., 2001, 2002; Dai and Hozalski, 2002; Betancourt and Rose, 2004; Hendricks et al., 2005; O'Melia, 2006; Hijnen and Medema, 2007). Rainstorms during winter or spring can be of particular concern, as low temperature can reduce the efficacy of the coagulation process (Hurst et al., 2004).

B.3.2 Other environmental influences

An increase in DOC concentrations over the past several decades has been reported in Canada (Eimers et al., 2008; Keller et al., 2008; Couture et al., 2012; Chowdhury, 2018), North America (Driscoll et al., 2003; SanClements et al., 2012), the United Kingdom (Evans et al., 2005; Sharp et al., 2006; Worrall and Burt, 2009), northern Europe (Eikebrokk et al., 2004) and Japan (Imai et al., 2001). At sites where DOC has increased, waters have also often become more coloured (Ekström et al., 2011; Kritzberg and Ekström, 2012; Weyhenmeyer et al., 2014).

Pagano et al. (2014) conducted a comprehensive review of increasing DOC trends and noted that researchers suggest decreased atmospheric acid deposition (i.e., sulphur emission controls) and climate change agents as two key considerations. Monteith et al. (2007) reviewed data for North America, the United Kingdom and Europe and reported that declining acid deposition explained >85% of the increasing DOC trends, except in the United Kingdom and Newfoundland. In these regions, increasing sea salt deposition explained DOC declines in some areas. The authors found no trends between DOC and increasing temperature or atmospheric carbon dioxide concentration. Ekström et al. (2011) conducted plot-scale acidification experiments and confirmed that reduced acid deposition results in increased DOC and colour concentrations, implying an increase in NOM mobility with sulphur emission controls. With regard to organic colour, this trend has been linked to iron complexing with DOC (Weyhenmeyer et al., 2014). However, the mechanisms are not completely understood. Black and Christman (1963) also found that iron was always present with organic colour but no relationship could be established between the iron content and colour.

Increasing DOC and/or colour concentrations can significantly impact water utilities using coagulation and filtration processes. Anderson et al. (2017) reported a four-fold increase in alum dose (i.e., 12.9 to 49.5 mg/L) and a 1.75-fold increase in lime use at a full-scale facility where true colour increased from approximately 20 in 1990 to approximately 50 in 2015. The authors also reported that the plant hydraulic capacity was reduced by 26%. Parsons et al. (2007) reported that the average coagulant dose at full-scale facilities in the United Kingdom increased from approximately 40 mg/L in 1992–1997 to 70–100 mg/L in 1998–2002 due to increased colour. Eikebrokk et al. (2004) conducted pilot-scale studies and reported that a 75% increase in

colour in low turbidity waters (<0.3 nephelometric turbidity unit [NTU]) increased the coagulant dose, sludge production, number of backwashes and residual TOC by 64%, 64%, 87%, and 26%, respectively. In addition, filter run times and hydraulic capacity were reduced by 47% and 10%, respectively. The authors also forecast increased chemical consumption for pH adjustment and increased biological growth in the distribution system due to higher residual organic carbon concentrations. Other researchers have noted that higher residual organic carbon concentrations contribute to increased DBP formation (Imai et al., 2001; Sharp et al., 2006). Chowdhury (2018) analyzed 15 years of water quality data (2000–2015) for 304 surface water and 137 groundwater sources in Newfoundland/Labrador and observed increasing trends for DOC, organic colour and trihalomethanes THMs.

Researchers have also found that wildfires can result in long term (>10 years) water quality degradation that substantially changes the concentration and character of NOM, and thereby significantly impacts water treatment processes (Emelko et al., 2011a; Geng et al., 2011; Emelko, 2019). Wildfires are forecast to increase in frequency as a result of changes in climate (van der Linden et al., 2018). Other water quality changes that are forecast to occur as a result of a changing climate and exacerbate NOM-related impacts include: increased water temperature; increased variability in runoff; and increased nutrient loading due to extreme runoff events (van der Linden et al., 2018). An increase in the frequency and severity of algal growth and cyanobacterial blooms is forecast to be associated with these changes (van der Linden et al., 2018).

A comprehensive review of the expected impacts of climate change on the treatability of NOM can be found elsewhere (Ritson et al., 2014). Arctic and sub-arctic regions are expected to respond differently from temperate regions (Pagano et al., 2014).

B.4 Impact of natural organic matter

Although NOM does not have direct health effects, it critically affects drinking water treatment and can contribute to indirect health impacts, as well as operational and aesthetic issues.

B.4.1 Indirect health impacts

B.4.1.1 Pathogen log reductions

Drinking water treatment typically comprises physical removal barriers (e.g., clarification, filtration) that are assigned pathogen “log removal” credits, and inactivation barriers (i.e., primary disinfection) that are assigned “log inactivation” credits. NOM critically impacts both.

For chemically assisted clarification/filtration processes, NOM exerts a coagulant demand that must be overcome before neutrally-charged floc particles can form. Neutrally-charged floc particles are essential for filters to perform properly and meet turbidity requirements for pathogen removal (Hall and Packham, 1965; Semmens and Field, 1980; Dempsey et al., 1984; Edwards and Amirtharajah, 1985; Amy et al., 1989; Edzwald and Van Benschoten, 1990; White et al., 1997; Shin et al., 2008). NOM concentrations can increase without a change in turbidity or flow and therefore may go undetected. Yet, increased NOM concentrations trigger the need to increase the coagulant dose to achieve neutrally-charged floc particles; otherwise, suboptimal coagulation conditions exist and a loss in pathogen log removal can occur.

James et al. (2016) reported *Cryptosporidium* breakthrough and an increase in particle counts (2–5 µm, 5–10 µm and 10–15 µm) at a full-scale direct filtration treatment plant as a result of an increase in colour in the source water. Other researchers have reported the breakthrough of particles >2 µm at pilot-scale during periods of elevated TOC (Billica and Gertig, 2000; Carlson

and Gregory, 2000). Several other studies (Ongerth and Pecoraro, 1995; Patania et al., 1995; Dugan et al., 2001; Huck et al., 2001) report that *Cryptosporidium* removal by clarification/filtration can significantly deteriorate during suboptimal coagulant conditions (e.g., treatment effectiveness decreased by 2.0 to 3.4 logs as compared with optimal conditions).

It is well known that NOM exerts a chemical oxidant demand (i.e., chlorine, chlorine dioxide, ozone) that must be overcome before pathogen log inactivation requirements can be met (AWWA, 2011a; MWH, 2012). Grunet et al., (2018) and Léziart et al. (2019) re-confirmed the critical impact that NOM can have on primary disinfection. Grunet et al., (2018) studied chlorine and chlorine dioxide. The authors observed that the disinfectant concentration decayed rapidly when DOC concentrations were ≥ 2 mg/L. Careful monitoring was recommended under these conditions. Léziart et al. (2019) found that an organic turbidity of 1 NTU, generated by 2 mg/L of humic acids, interfered with chlorine efficacy whereas 5 NTU of inorganic turbidity, generated by chalk carbonates (e.g., aquifer formation material), had no measurable impact. Disinfection is typically applied after treatment processes that remove NOM to ensure efficient inactivation of pathogens and to minimize the formation of DBPs (see section B.4.1.2). Additional information on how NOM affects chemical oxidant demand, decay and disinfection is published elsewhere (Health Canada, 2009a, 2018).

Several studies have examined the effect of particles on UV disinfection efficacy, and most have concluded that the UV dose–response of microorganisms is not affected by variations in turbidity up to 10 NTU (Christensen and Linden, 2002; Batch et al., 2004; Mamane-Gravetz and Linden, 2004; Passantino et al., 2004). However, the presence of humic acid particles and coagulants has been shown to significantly affect UV disinfection efficacy, with lower inactivation levels being achieved. Templeton et al. (2005, 2007) found that in unfiltered influent samples (range = 4.4–9.4 NTU), UV disinfection of bacteriophages in the presence of humic acid flocs was reduced by a statistically significant degree (≈ 0.5 log) as compared with particle-free water. Templeton et al. (2005) also found that UV-absorbing organic particles (i.e., NOM) shielded particle-associated bacteriophages from UV light, whereas inorganic kaolin clay particles (i.e., inorganic turbidity) did not. The extent of shielding is more likely to depend on the particle type (e.g., size, structure, chemical composition), the number of large particles (e.g., ≥ 25 μm), the level of pathogen aggregation with particulate matter and the desired inactivation level than on the turbidity level (Caron et al., 2007; Hargy and Landry, 2007; Templeton et al., 2008; Kollu and Örmeci, 2012). In addition, UV transmittance at a wavelength of 254 nm is affected by dissolved and particulate matter that inhibits the penetration of UV light through the water. In general, every 10% decrease in UV transmittance results in a 50% reduction in the UV dose (Hofmann, 2008). Thus, water with a UV transmittance of 85% will need more reactors to achieve pathogen log inactivation requirements than water with a UV transmittance of 95%. Manufacturers usually specify a minimum UV transmittance below which the system will not function properly. Pretreatment to remove NOM may be necessary to meet the manufacturer’s specification for minimum UV transmittance, in order to achieve pathogen log inactivation requirements and ensure safe operation of the equipment.

It is important that water utilities understand the pathogen log reductions that can be achieved when operating under optimal conditions and the impact of short- and long-term treatment upsets (Hurst et al., 2004). The application of the “robustness index” suggested by Huck et al. (2001) provides a simple and practical means of identifying events and periods when the coagulation and clarification processes become unstable. Information on how to use quantitative microbial risk assessment (QMRA) to evaluate the robustness of physical removal and/or inactivation barriers is also available (Health Canada, 2019a).

B.4.1.2 Formation of disinfection by-products

Chemical disinfectants react with NOM to form potentially harmful DBPs (Rook, 1974; Stevens et al., 1976). After many years of research, it is generally accepted that all NOM fractions contribute to DBP formation, although some fractions form more DBPs than others (Hoehn et al., 1980; Croué et al., 1993; Owen et al., 1993, 1995; Martin-Mousset et al., 1997; Goslan et al., 2002; Liang and Singer, 2003; Hua and Reckhow, 2007a; Bond et al., 2014; Hua et al., 2014). It is also recognized that some non-regulated DBPs are more cytotoxic and genotoxic than regulated DBPs such as THMs and HAAs (Stalter et al., 2016; Wagner and Plewa, 2017). Although THMs and HAAs can be used as indicators for the presence of other DBPs (Health Canada, 2006, 2008a), it is important to recognize that their formation pathways and reaction rates are different (Reckhow and Singer, 1984; Liang and Singer, 2003; Hua and Reckhow, 2007a; Bond et al., 2012). For example, Plourde-Lescelleur et al. (2015) reported THM:HAA formation potential ratios of 0.66–3.35 for six Canadian surface water sources (test conditions = pH 8.0, residual chlorine 1.0 mg/L at 22°C for 24 h). Archer and Singer (2006a) reported that, as the hydrophilic fraction increases, THMs are preferentially produced over HAAs. Thus, THMs and HAAs should be managed conjunctively.

Other factors that affect DBP formation include water temperature, pH, disinfection conditions (i.e., disinfectant, dose, contact time, residual) and the presence of reactive species such as bromide, iodide, ammonia and sulphur (Liang and Singer, 2003; Ates et al., 2007; Hua and Reckhow, 2007b; Reckhow et al., 2007; Fabris et al., 2008; Kristiana et al., 2009). Bromide, in particular, has been shown to increase DBP formation rates three- to ten-fold (Symons et al., 1993; Westerhoff et al., 2004; Heeb et al., 2014). As a result, similar DOC concentrations can produce a wide range of DBP concentrations, depending on the character and reactivity of the NOM (Fabris et al., 2008). For example, Hua and Reckhow (2007a) reported that the reactivity of the raw water for Winnipeg, Manitoba, (DOC = 7.9 mg/L; bromide <10 µg/L) was approximately half that of Repentigny, Quebec, (DOC = 7.1 mg/L; bromide = 46 µg/L). In some cases, NOM is nitrogen-rich and contributes to the formation of nitrogenous DBPs (N-DBPs) (Leenheer and Croué, 2003; Mitch et al., 2009). Rain events can also transport DOC that is rich in precursors, resulting in elevated DBP concentrations (Goslan et al., 2002; Fearing et al., 2004a, 2004b; Reckhow et al., 2007; Kraus et al., 2010; Delpla and Rodriguez, 2016; Wright et al., 2016).

Bond et al. (2011, 2012) completed a comprehensive review of the literature and summarized the potential for various NOM components to form THMs, HAAs and N-DBPs, as outlined in Table 6. Reckhow et al. (2007) reported that tannin and lignin were significant precursors, representing 60% and 25–30% of the THM formation potential, respectively. The authors also suggested that proteins could be important precursors during early stages of NOM formation, but these would biodegrade (see section B.4.1.3) and terpenoids would take their place as another important source of THM precursors.

Table 6. NOM precursors and their DBP formation potential^a

Precursor	Potential for removal by coagulation	Formation potential		
		THMs	HAAs	N-DBPs
Humic and fulvic acids	High	Primary source	Primary source	Possibly important for halonitromethanes
Carboxylic acids	Medium	β -dicarbonyl species important precursors	β -dicarbonyl species important precursors	Probably minor
Amino acids	Low	Low except for two compounds ^b	Important for some compounds ^c	Significant
Proteins	Low	Variable; important during algal blooms	Not known; may be significant	Uncertain
Carbohydrates	Low	Important at pH 8	Probably minor	Insignificant

^a Adapted from Bond et al., 2011, 2012

^b Tryptophan and tyrosine

^c Aspartic acid, histidine, asparagine, tryptophan

It is important that water utilities understand the source-specific reactivity of NOM when selecting a disinfectant, in order to mitigate the formation of potentially harmful DBPs (Hua and Reckhow, 2007a). Determining the specific DBP yield (i.e., $\mu\text{g DBP/mg DOC}$) can help, because DOC removal is generally easier to achieve than a decrease in reactivity, particularly when bromide is present (Croué et al., 1993; Goslan et al., 2002; Reckhow et al., 2007; Fabris et al., 2008; Tubić et al., 2013). Different mitigative measures may be necessary to minimize one group of DBPs compared with another, depending on NOM reactivity; technologies targeting the removal of specific NOM fractions may be necessary (Bond et al., 2011). In addition, Kastl et al. (2016) reported that NOM removal requirements should be linked to distribution system conditions. For example, a distribution system with a residence time of 7 days and temperature of $>15^\circ\text{C}$ will require a different level of NOM removal to meet DBP guidelines than one with a residence time of 3 days and temperature of $<15^\circ\text{C}$ (Rodriguez and Sérodes, 2001; Kastl et al., 2016).

Source-specific treatability studies, including DBP formation potential tests, should be conducted when evaluating different mitigative measures and/or alternative treatment options. To ensure that an effective NOM control strategy is implemented, the treatability study should be specifically designed to: 1) assess seasonal variations in NOM; and 2) be representative of distribution system conditions. It is important to note that formation potential test methods that use very high chlorine doses may not correctly determine differences in DBP yield when bromide is present (Bond et al., 2014). This is because chlorine can out-compete bromine when it is in excess relative to bromine. Under typical operating conditions, bromine is much more effective at forming DBPs than chlorine (Bond et al., 2014). Standard methods are available to assess DBP formation potential (APHA et al., 2017). The use of uniform formation conditions (Summers et al., 1996) enables the direct comparison of results to assess the effectiveness of various treatment options (AWWA, 2011a). Practical guidance to help water utilities conduct DBP formation potential tests is available (Alexander et al., 2019).

The guideline technical documents for THMs and HAAs specify that water utilities

should make every effort to maintain concentrations as low as reasonably achievable. Hence, the removal of NOM is a recommended best practice to minimize the formation of both regulated and non-regulated DBPs (Health Canada, 2006; 2008a). This may require specific monitoring to ensure adequate precursor removal (see Tables 1 and 2 in section A.2.3).

It is critical that efforts made to minimize DBP formation not compromise the effectiveness of disinfection. More information can be obtained from the appropriate guideline technical documents for THMs, HAAs, chlorite/chlorate, *N*-nitrosodimethylamine and bromate (Health Canada, 2006, 2008a, 2008b, 2011, 2018).

B.4.1.3 Biological stability

The biological stability of drinking water refers to the concept of maintaining microbiological water quality from the point of production to the point of consumption (Prest et al., 2016). Heterotrophic organisms make up the majority of bacteria in drinking water and draw their energy for growth, multiplication and production of biofilm matrix materials from the degradation of organic carbon compounds (Vu et al., 2009; Prest et al., 2016). BOM encourages bacterial growth and biofilm development in the distribution system and premise plumbing, which can lead to issues that have public health significance. Biofilms provide a habitat for the survival of fecal pathogens that may have passed through drinking water treatment barriers or entered the distribution system directly via an integrity breach (Leclerc, 2003). It has been shown that enteric viruses and protozoa can be detected in biofilms (Howe et al., 2002; LeChevallier, 2003; Chang and Jung, 2004); although these organisms cannot grow in this environment, they can accumulate and be released over an extended period of time (Howe et al., 2002; Warnecke, 2006; Wingender and Flemming, 2011). Additionally, opportunistic premise plumbing pathogens (OPPPs) such as *Legionella pneumophila* and non-tuberculous mycobacteria (e.g., *M. avium*, *M. intracellulare*) have adapted to grow and persist in distribution and plumbing system biofilms. Biofilms can also create difficulties with maintaining adequate disinfectant residuals and can be involved in nitrification in distribution systems where chloramine is used (Wilczak, 2006).

While biofilm microorganisms utilize the constituents with the shortest biodegradation half-lives first, they are adept at consuming all types of available NOM constituents leaving a treatment facility to support their growth in the distribution system (Fischer, 2003; Camper, 2014). Table 7 summarizes the biodegradation half-life for NOM constituents (Reckhow et al., 2007).

Treatment processes also significantly impact the composition and concentration of organic nutrients. For example, oxidants such as chlorine and ozone produce biodegradable products upon reaction with NOM (Alarcon-Herrera et al., 1993; Bursill, 2001; Reckhow et al., 2007). It is well known that ozone transforms NOM to BOM (Owen et al., 1993, 1995); thus biologically active filtration may be necessary to stabilize treated water (GLUMRB, 2012). Chlorine can also react with organic matter thereby increasing the amounts of assimilable organic carbon (AOC) and biodegradable DOC (BDOC) which can exacerbate the problem of biofilm growth in distribution systems (Drikas et al., 2003; Reckhow et al., 2007; Croft, 2012; Camper, 2014).

Table 7. Biodegradable constituents^a

Constituent	Biodegradation half-life (days)
Sugars and starches	2
Proteins	8
Hemicellulose	25
Cellulose	40
Lipids (e.g., fats and waxes)	60
Lignins	150
Tannins	200
Terpenoids (e.g., geosmin and 2-methylisoborneol)	400

^a Adapted from Reckhow et al., 2007; Satchwill et al., 2007

Limited publications have measured this aspect, and the numerical values reported have varied from site to site (Camper, 2014). LeChevallier et al. (1992) observed AOC increases of 20% and 44% when samples of granular activated carbon (GAC)-sand filter effluent were treated with 1 mg/L free chlorine or 2 mg/L monochloramine for 30 min, respectively. Zacheus et al. (2001) studied the microbiological quality of water and pipeline deposits in 16 full-scale distribution systems in Finland. The mean rate of bacterial biomass production was higher in water in the distribution system (1.0 ng carbon/L/h) than water leaving the treatment facilities (0.22 ng carbon/L/h). Liu et al. (2002) noted that variations of AOC in distribution systems were affected by chlorine oxidation (increase in AOC) and bacterial activity (decrease in AOC). Different patterns were observed in different distribution systems and by season. In one system, a 120% increase in AOC concentrations was observed from the treatment plant to the end of the distribution system during December, whereas a 35% decrease in AOC concentrations was observed across the same sites in the spring of the following year (Liu et al., 2002).

In drinking water supplies in North America, minimum disinfectant residuals are typically recommended to control biofilm growth (LeChevallier et al., 1996; LeChevallier and Au, 2004). In some European countries, the approach taken to achieve biological stability is through the reduction of growth-supporting nutrients in water (Lautenschlager et al., 2013). In the Netherlands, water supply companies aim at limiting regrowth in distributed water in the absence of a disinfectant residual by meeting stringent targets for AOC (10 µg/L) (Lautenschlager et al., 2013). Levels of BOM (e.g., AOC, BDOC) are an important factor in the biostability of potable water supplies but are only one component influencing changes in water quality (Prest et al., 2016). Other compounds have been identified as having roles in controlling microbial growth in the distribution system: phosphorus, ammonia, manganese, iron and humic substances (Camper, 2004; Prest et al., 2016).

Multiple factors affect the biostability of distributed water: type and concentration of organic and inorganic nutrients, type and concentration of residual disinfectant, biofilms and sediments, and distribution system conditions (e.g., disinfectant residual decay, water temperature, residence time, hydraulic conditions, pipe material and diameter, pH, corrosion rate) (LeChevallier et al., 2015a; Prest et al., 2016).

Comprehensive reviews of biological stability can be found elsewhere (Prévost et al., 2005; van der Kooij and van der Wielen, 2014; LeChevallier et al., 2015a, 2015b; Prest et al., 2016). In general, strategies to improve biostability and minimize biofilm development in the distribution system and premise plumbing include optimized NOM removal, maintaining an effective disinfectant residual, maintaining low levels of biostability indicators in treated water

(e.g., AOC, BDOC, biofilm formation rate), controlling corrosion and managing water temperatures (e.g., optimize storage facility turn-over rates; install mixers to prevent thermal stratification). Additional guidance on monitoring the biological stability of drinking water distribution systems is available (Health Canada, in preparation).

B.4.1.4 Corrosion impacts

Corrosion is the deterioration of a material that results from a reaction with its environment. Corrosion in drinking water distribution systems can be caused by several factors, including the type of materials used in pipes and fittings, the age of the piping and fittings, the stagnation time of the water and the water quality in the system (including its pH and alkalinity). Other drinking water quality parameters that can influence corrosion include temperature, calcium, free chlorine residual, chloramines, chloride, sulphate and NOM (Health Canada, 2009b).

NOM has been shown to affect lead and copper corrosion (Korshin et al., 1996, 2000, 2005; Edwards and Sprague, 2001; Dryer and Korshin, 2007; Liu et al., 2009; Valentine and Lin, 2009; Schock and Lytle, 2011; Arnold et al., 2012; Zhou et al., 2015; Masters et al., 2016). The effects of NOM on metal surfaces can be varied. NOM can provide a protective film, decreasing corrosion (especially over a long timeframe), or it can increase corrosion through a variety of mechanisms: 1) NOM could complex with calcium ion and prevent protective scale formation; or 2) NOM could act as a food source for microorganisms, which could in turn attack the pipe surface and increase corrosion (Schock and Lytle, 2011).

Schock et al. (1996) reported varied impacts of NOM on lead and copper solubility. In some instances, NOM was observed to form soluble organic complexes with lead, resulting in an increase in dissolved lead concentrations. NOM can also adsorb/adhere to the interior surface of lead pipes, decreasing lead solubility. Korshin et al. (2005) demonstrated that NOM can impact both the morphology (physical structure) and size distribution of lead particles and can prevent or impair the formation of the more stable scales of cerussite and hydrocerussite. NOM was observed to prevent the formation of cerussite and impair the formation of hydrocerussite (i.e., imperfect and dispersed crystals were observed). Specifically, in the absence of NOM, approximately 90% of lead particles were found to be >5 µm. The addition of NOM decreased particle size, as larger particles broke down due to the accumulation of surface charge. The sharpest increases in soluble lead concentrations were observed between 0 and 3.5 mg/L DOC (Korshin et al., 2005) and between 0 and 2 mg/L DOC (Korshin et al., 2000). In a factorial experiment, Zhou et al. (2015) observed that NOM increases (from 1 mg/L to 7 mg/L DOC) resulted in significant increases in lead release in simulated partial lead service line replacements. In bench-scale work by Trueman et al. (2017), the authors observed that the presence of humic acid increased lead release from lead coupons as a result of both uniform and galvanic corrosion. Although the addition of orthophosphate lowered the lead release, the addition of humic substances made the orthophosphate less effective. The authors suggested that the complexation of lead and humic substances inhibits lead precipitation with phosphate.

Similarly, NOM has also been observed to increase copper release even at low concentrations (0.1–0.2 mg/L DOC), although the relationship between DOC and copper release was not observed to be linear (Korshin et al., 1996). The authors suggested that NOM adsorbs to the inorganic crystal structures at high DOC concentrations, causing mobilization and dispersion, whereas at very low concentrations, NOM cannot cover the entire surface, which results in patchy crystalline product coverage and creates conditions for copper pitting. The apparent contradictory effects of NOM on copper release were further explored by Edwards and Sprague

(2001). The authors observed that NOM interferes with pipe ageing by forming the more soluble cupric-hydroxide, copper carbonate and copper-NOM complexes that prevent the formation of the more stable malachite. Copper pipe ageing is a significant factor in copper release control, with older copper pipes being associated with the more stable tenorite and malachite scales (Lagos et al., 2001; Edwards and McNeill, 2002). NOM can also be protective of copper, by acting as a food source for bacteria, consuming dissolved oxygen and triggering re-deposition when a suitable catalyst is present (Edwards and Sprague, 2001). The presence of NOM can lead to decreased copper release, as the NOM can sorb onto freshly formed copper pipe surfaces, which decreases soluble copper complexation capacity and causes re-deposition (Edwards and Sprague, 2001). In practice, Arnold et al. (2012) demonstrated that removing NOM was an effective method to decrease blue-water issues in a school with new copper plumbing. The authors suggested that NOM removal accelerated the natural ageing process.

Peng et al. (2013) observed that iron release increased in the presence of NOM (DOC = 1 mg/L) and that other inorganics (lead, vanadium, chromium, copper and arsenic) could be released from iron at various levels of chloride (0–250 mg/L).

The interactions of NOM with metal surfaces are complex, with multiple factors influencing the interactions such as exposure time and pH (Korshin et al., 2005; Liu et al., 2009). NOM characteristics have also been observed to be important for lead and copper release (Willison and Boyer, 2012) and inconsequential for lead (Dryer and Korshin, 2007). Further research is needed to explore the significant impact and mechanisms of NOM and metal release in both distribution system and premise plumbing materials. Researchers currently recommend that NOM be removed to minimize lead and copper concentrations (Valentine and Lin, 2009; Arnold et al., 2012; Zhou et al., 2015). More information on corrosion control, lead and copper is available elsewhere (Health Canada, 2009b, 2019b, 2019c).

B.4.2 Operational issues

B.4.2.1 Coagulation process

The goal of coagulation is to destabilize (i.e., neutralize the charge of) colloidal particles (including pathogens) so that they effectively aggregate during flocculation and are subsequently removed by clarification and/or filtration. Coagulation also deals with removing NOM by a phase change that converts dissolved organic matter into particles: either directly by precipitation or by adsorption onto particles created by the coagulant (Edzwald and Haarhoff, 2012). When metal coagulants are added to the water, chemical reactions occur with both particles and NOM. Therefore, when a coagulant is added, the NOM acts as a ligand that complexes the positively charged metal ions, exerting a coagulant demand that must be overcome before flocculation can occur (Edzwald and Haarhoff, 2012). The character of NOM also has a significant impact on the coagulation process. For example, a two-fold increase in coagulant dose is needed to coagulate equal mass concentrations of fulvic acids as compared with humic acids (Edzwald, 1993; Rigobello et al., 2011).

Thus, coagulation should be viewed as an “integrated” process that considers both NOM and particles (i.e., turbidity) while having regard to their different coagulation characteristics (Edzwald and Haarhoff, 2012). For example, for the pH conditions of most water sources (pH 6–8), NOM and particles carry a negative charge that becomes more negative with increasing pH. However, the negative charge of NOM is typically between 5–15 µeq/mg carbon, while that of particles is between 0.05–0.5 µeq/mg particle, depending on the particle type (Edzwald, 1993). Pernitsky and Edzwald (2006) estimated the charge for both the NOM and particle components for a variety of water sources to demonstrate that, in most cases, coagulant dosing is controlled

by NOM, not by turbidity. Turbidity must increase significantly, in the absence of an associated NOM increase, for turbidity to control the coagulant dose. As NOM concentrations can rapidly increase four- to five-fold during storm events, it is important that water utilities have a good understanding of NOM's impact on coagulant dosing (Edzwald, 1993; Pernitsky, 2003; Hurst et al., 2004; McVicar et al., 2015; James et al., 2016). Failure to adjust the coagulant dose in accordance with a change in NOM may contribute to suboptimal coagulation conditions and a decrease in pathogen log removal capability (Edzwald, 2017).

Given the importance of coagulation chemistry to ensure pathogen log removals, water utilities should consider both NOM and turbidity when defining optimum pH and coagulant dose conditions (Edzwald, 1993; Edzwald and Tobiason, 1999; Edzwald and Haarhoff, 2012). Jar testing is one of the most commonly used techniques to simulate coagulation treatment and to determine the coagulation potential for a water source (Black and Willems, 1961; Chow et al., 2004). It should be noted, however, that some NOM fractions cannot be removed by coagulation at any pH or dose (Kavanaugh, 1978; Babcock and Singer, 1979; Owen et al., 1993, 1995; Volk et al., 2002; Chow et al., 2004, 2006; Carpenter et al., 2013). More detailed discussions on the principles of coagulation and process optimization are presented elsewhere (Edzwald and Van Benschoten, 1990; Edzwald, 1993; Gregor et al., 1997; Edzwald and Tobiason, 1999; Pernitsky, 2003; Eikebrokk et al., 2006; Dempsey, 2006; Pernitsky and Edzwald, 2006; Edzwald and Kaminski, 2009; AWWA, 2011a, 2011b; Edzwald and Haarhoff, 2012; Davis and Edwards, 2014).

B.4.2.2 Membrane treatment

NOM has been identified in numerous studies as being responsible for membrane fouling, which can significantly impair water treatment operations. It is generally accepted that the hydrophilic neutral fraction of NOM, comprising polysaccharides and proteins in macromolecular and/or colloidal form (i.e., biopolymers), is responsible for membrane fouling (Amy and Cho, 1999; Carroll et al., 2000; Cho et al., 2000; Fan et al., 2001; Kimura et al., 2004; Lee et al., 2006; Her et al., 2007; Amy, 2008; Kennedy et al., 2008; Hallé et al., 2009; Peldszus et al., 2011; Croft, 2012; Chen et al., 2014; Kimura et al., 2014; Rahman et al., 2014; Siembida-Lösch et al., 2014, 2015; Yamamura et al., 2014; Chon and Cho, 2016). It is hypothesized that once fouling is initiated by biopolymers, a decrease in electrostatic forces allows hydrophobic NOM to adsorb to the membranes, resulting in further fouling (Peldszus et al., 2011; Croft, 2012; Chon and Cho, 2016). Rahman et al. (2014) reported that biopolymer concentrations as low as 0.1 mg/L resulted in reversible (i.e., removable by backwashing/air scour) and irreversible (i.e., removable by chemical cleaning) fouling. Her et al. (2007) reported fouling by protein-like substances that were not detected in the feed water due to low concentrations (detection limit not given).

Other factors that influence membrane fouling include membrane characteristics (e.g., type of membrane, pore size distribution, material, surface charge, hydrophobicity), operating conditions (e.g., flux, recovery, pretreatment, backwashing, chemical cleaning), and water quality (e.g., pH, ionic strength, concentration and character of the foulants) (Amy, 2008; Huck and Sozański, 2011).

Water utilities should have a good understanding of how the NOM in their source water will interact with membranes to avoid configurations that incur significant fouling. Pretreatment may be necessary to reduce biopolymer concentrations (Carroll et al., 2000; Peldszus et al., 2011; Siembida-Lösch et al., 2014; Chon and Cho, 2016). Pretreatment should be customized to each individual source, as effectiveness is source-specific (Fabris et al., 2007; Gao et al., 2011;

Siembida-Lösch et al., 2015). A program of regular backwashing and periodic chemical cleaning, using proper foulant-based cleaning chemicals, should also be in place to remove accumulated foulants (Alspach et al., 2014).

B.4.3 Aesthetic

It is well established that NOM is responsible for such aesthetic concerns as colour, taste and odour (Hassler, 1947).

Colour caused by the presence of organic substances can occur in both surface and ground waters (Black and Christman, 1963; Thurman, 1984; Tan and Sudak, 1992). Organic colour tends to be caused by the presence of humic and fulvic acids, which are black- to yellow-coloured substances (Stevenson, 1982). Black and Christman (1963) reported that 87% of the compounds responsible for colour in 10 U.S. sources were colloidal and 3.5–10 nm in size. By contrast, Ratnaweera et al. (1999) reported that 40% of the compounds responsible for colour in seven Finnish sources were <10 kDa (approximately 1 nm). Highly coloured sources tend to have a higher concentration of high molecular weight humic acids, which may account for these differences in size distribution (Edwards and Amirtharajah, 1985; Aitkenhead-Peterson et al., 2003). Fulvic acids represent a more complex mixture of low molecular weight compounds that are more hydrophilic than humic acids, and they have a significant impact on the required coagulant dose (see section B.4.2.1). Also, a higher proportion of fulvic acids are non-coagulable at any pH or coagulant dose (Hall and Packham, 1965; Kavanaugh, 1978; Babcock and Singer, 1979). As humic and fulvic acids are important DBP precursors, adequate colour removal may be necessary to meet DBP guidelines (Chaulk, 2015). For example, Tan and Sudak (1992) reported THM formation potentials of 250–262 µg/L (7-day formation potential test at 20°C and at around pH 8) for a highly coloured groundwater supply with naturally occurring humic and fulvic acids (TOC = 3.93–4.70 mg/L; UV absorbance = 0.1829–0.1907).

Tastes and odours can be caused by volatile compounds produced by the microbial biomass (e.g., actinomycetes, cyanobacteria, fungi) that is washed in from the terrestrial environment or is naturally present in the aquatic system/aquifer (Hrudey et al., 1992; Zaitlin and Watson, 2006; AWWA, 2011a). Watson (2003) identified approximately 200 volatile organic compounds that produce undesirable tastes and odours. Terpenoids (e.g., geosmin and 2-methylisoborneol), sulphides and polyunsaturated fatty acids were identified as the most odorous. Geosmin and 2-methylisoborneol are environmentally stable compounds (i.e., not easily biodegraded) that can be transported significant distances from where the compounds are produced (Satchwill et al., 2007). Other researchers identified pyrimidines as problematic (Chorus et al., 1992; Zaitlin and Watson, 2006; Peter et al., 2009). Zacheus et al. (2001) found that actinomycetes and fungi can survive in the soft deposits (i.e., accumulated deposits containing organic and inorganic matter) of water distribution systems. As a result, the distribution system may constitute a source of taste and odour problems.

Chlorine reactions with NOM may also contribute to tastes and odours (AWWA, 2011a). In particular, nitrogen-rich NOM can form odorous aldehydes (Hrudey et al., 1988), N-chloraldimines (Freuze et al., 2004, 2005) or nitriles (Freuze et al., 2004, 2005; Brosillon et al., 2009) when appropriate conditions exist with either chlorine or chloramines. Reaction pathways depend on: the choice of disinfectant, disinfectant to amino acid molar ratio, pH, temperature and reaction time (Froese et al., 1999; Brosillon et al., 2009). Amino acids have been identified as the primary odour-causing precursor and they can be released by the lysis of bacterial or algal cells or when proteins are oxidized (How et al., 2018). Table 8 provides the odour threshold concentrations that have been reported in the literature for these compounds. As nitriles have

much higher odour threshold concentrations (see Table 8), they are not typically implicated in taste and odour events (Freuze et al., 2005). Odour-causing compounds (e.g., terpenoids) and precursors (e.g., amino acids, proteins) are not effectively removed by conventional treatment. Thus, other processes may be necessary to minimize tastes and odours (Rice and Gomez-Taylor, 1987; Bruchet et al., 1992; Froese et al., 1999). Once odorous compounds are formed, they can persist in the distribution system for up to 500 hours (≈ 21 days) at 15°C (Freuze et al., 2004, 2005). Concentrations can also increase in the distribution system due to the release of amino acids or peptides from the biofilm (Brosillon et al., 2009). Guidance material to assist water utilities assess and minimize objectionable tastes and odours is available elsewhere (AWWA, 2011c).

Table 8. Odour threshold concentrations reported for NOM-related compounds^a

Odorous by-product	Odour threshold concentration	Reported odour
Aldehydes	0.15–30 µg/L	Swampy swimming pool ^b
N-chloraldimines	0.20–3 µg/L	Floral swimming pool ^c
Nitriles	210–430 µg/L	

^a Adapted from How et al., 2018

^b Froese et al., 1999

^c Freuze et al., 2004, 2005

B.5 Measurement and characterization

An effective NOM control strategy requires a good understanding of the origin, occurrence and variation that occurs in the source water (Volk et al., 2002). Water utilities should have a good understanding of:

- their water source and the nature and generation of NOM;
- whether NOM changes seasonally or with precipitation/snowmelt events; and
- how NOM interacts with treatment processes.

B.5.1 Considerations for quantifying natural organic matter

Although the numerous organic compounds that contribute to NOM cannot be measured directly, there are a number of surrogates that can be used to provide an indication of the NOM concentration. The most commonly used surrogates include TOC, DOC, UV absorbance and chemical oxygen demand (Sillanpää et al., 2015b). In addition, UV absorbance and UV transmittance are mathematically related; hence the latter can also provide an indication of NOM concentration. As NOM is a major contributor to organic colour, this parameter may also be relevant (Matilainen et al., 2011).

TOC quantifies all organic carbon in a water sample and is the sum of particulate and dissolved organic carbon. DOC is operationally defined as the organic carbon that has passed through a 0.45 µm filter (APHA et al., 2017). As the filter can leach some organic carbon to the sample, it is recommended that at least 50 mL of organic-free water be passed through the filter and filter assembly before filtering the DOC sample (Karanfil et al., 2002, 2005). TOC and DOC are measured indirectly from the carbon dioxide that is produced by UV-catalyzed chemical oxidation or by high-temperature combustion.

UV-visible light absorbance at 254, 350 and 440 nm can be linearly correlated to DOC concentration in some freshwater systems. However, linear correlations are less likely to be found in sources with strong autochthonous or anthropogenic inputs or where DOC has been

extensively degraded by natural UV light (e.g., long retention time in lake) (Minor et al., 2014). The measurement of UV254 has historically been used in the water industry (Edzwald et al., 1985). Samples should be filtered to remove particle-related variations in UV absorbance (APHA et al., 2017). It is generally accepted that a change in UV absorbance provides a good indication of changes in NOM (Pernitsky, 2003; Wright et al., 2016). Online or daily monitoring of UV absorbance provides valuable information to operators about pending impacts to the coagulant dose, as NOM concentrations can change without any observed fluctuation in flow or turbidity (see section B.3.1). Otherwise, operators are not aware of coagulant under-dosing until turbidity spikes are observed in clarified water or filter effluent. It is important that correlations be developed on a source-specific basis, because the relationship between NOM and UV absorbance is unique to each source (Pernitsky, 2003). In some cases, it is not possible to establish a correlation between UV254 and DOC (Cho et al., 2010; Sadrnourmohamadi et al., 2013; Minor et al., 2014). Monitoring UV-visible light absorbance over a broader range of wavelengths may be more appropriate in some cases (Wright et al., 2016). Alternatively, the lack of a correlation may be due to the presence of NOM that has low UV absorbance (e.g., proteins, sugars) or a high nitrate content, which may interfere with this measurement (Leenheer and Croué, 2003). Monitoring UV-visible light absorbance over a broader range of wavelengths may also provide a more advanced characterization.

UV transmittance is a relative measure of how much light passes through a water sample (at a wavelength of 254 nm typically through a 1 cm path length) compared with how much light passes through pure deionized water (which has a UV transmittance of 100%). Since UV absorbance and UV transmittance are mathematically related as per the formula below (Bolton, 2013), no information is lost by choosing one parameter over the other.

$$\text{UV absorbance (in cm}^{-1}\text{)} = 2 - \log_{10} \text{UV transmittance (in \%)}$$

A chart to convert UV transmittance to UV absorbance (and vice versa) is provided in Table C-3.5 of this document.

Chemical oxygen demand serves to give some indication of the concentration of oxidizable organic matter in a water sample (Frisch and Kunin, 1960; Stoddart and Gagnon, 2014). Historically, the chemical oxygen demand test method (using potassium dichromate) was not sensitive enough for drinking water (Rittman and Huck, 1989). More sensitive methods have since been developed. One involves using potassium permanganate as the oxidant (ISO, 1993); the other is a photoelectrochemical oxygen demand (peCOD) method using UV activated titanium dioxide as the oxidant (Zhao et al., 2004; ASTM, 2017). Information on the mean oxidation state of organic carbon during water treatment can also be gleaned using a ratio of the molar concentrations of COD and TOC/DOC (Li et al., 2018).

Colour has historically been measured using colorimetric methods. The presence of suspended particles (e.g., clay, iron and manganese oxides) can give waters the appearance of colour and should be removed by filtering the sample through a 0.45 µm filter before measurement of NOM-related organic colour (e.g., colour due primarily to the presence of humic and fulvic acids – see section B.4.3); a filtered sample is operationally defined as “true colour” (APHA et al., 2017). Researchers have also used visible light absorbance at 420 nm as a measure for organic colour (Ekström et al., 2011; Weyhenmeyer et al., 2014). However, a wavelength between 450 nm and 465 nm is proposed as a standard spectrophotometric method (APHA et al., 2017). The spectrophotometric method requires that samples be filtered through a 0.45 µm filter (Hongve and Åkesson, 1996; APHA, 2017). The comparison of true and apparent colour results

can help water utilities determine if colour complaints are NOM-related. Apparent colour applies to unfiltered samples and is a useful measure to assess the presence of iron and manganese oxides in the distribution system (Reiber and Dostal, 2000; Imran et al., 2005).

Measurement of the above-noted parameters is simple and fast to perform, and some tests can be automated. They can indicate a change in water quality is occurring; however, they do not offer information about the character of the NOM. Edzwald et al. (1985) found that UV₂₅₄ divided by the mg/L of DOC was a helpful indicator of NOM character. This concept later became known as specific UV absorbance (SUVA), which is discussed below. The calculation of specific colour (i.e., true colour divided by mg/L DOC) may also provide useful information (Chow et al., 2005).

B.5.2 Natural organic matter characterization

B.5.2.1 Specific ultraviolet absorbance

The concept of SUVA has been developed as an operational indicator of NOM character and coagulation effectiveness for NOM removal (Edzwald and Van Benschoten, 1990; Edzwald and Tobiasson, 1999). Table 9 presents the generally accepted relationships between SUVA, NOM composition, UV absorbance, coagulation and potential TOC removal. The calculation of SUVA is widely used to assess NOM character because it is easy and inexpensive to determine and is a good indicator of changes in source water quality (Westerhoff et al., 1999; Imai et al., 2001; Weishaar et al., 2003; Reckhow et al., 2007). For example, Volk et al. (2002) monitored DOC and UV₂₅₄ for the White River (Muncie, Indiana) on a daily basis for 22 months. During this period, SUVA ranged from 1.40 L/mg·m to 10.51 L/mg·m. Lower values were associated with periods of low runoff and high algal activity (i.e., hydrophilic, autochthonous NOM), whereas high values were associated with snowmelt and storm runoff. During a typical rainfall event, SUVA increased from 2.6 L/mg·m to 4.5 L/mg·m within 12 h, indicating that hydrophobic, allochthonous NOM was being flushed into the source from the terrestrial watershed. Archer and Singer (2006b) analyzed 18 months of surface water data from the U.S. Environmental Protection Agency (EPA) Information Collection Rule and found a clear relationship between SUVA, source water characteristics and the effectiveness of coagulation for the removal of organic carbon.

In general, high SUVA sources (>4 L/mg·m) have NOM that is amenable to coagulation. However, the hydrophilic neutral fraction can have a high SUVA, which can be misleading with respect to the potential for organic carbon removal using coagulation (Edzwald, 1993). Also, achieving DBP guideline limits will depend on the raw water NOM concentration and whether a sufficient amount of reactive NOM can be removed. If the post-coagulation DOC residual remains reactive with respect to DBP formation, other technologies targeting the removal of specific NOM fractions may be necessary (Bond et al., 2011). As humic and fulvic acids are important DBP precursors, adequate colour removal may be necessary to meet DBP guidelines (Chaulk, 2015). Low SUVA sources tend to have NOM that is not amenable to coagulation (Pernitsky, 2003).

Table 9. Relationship between SUVA and potential TOC removal^a

SUVA (L/mg·m)	NOM composition	UV absorbance	Coagulation	Potential TOC removal
<2	Mostly hydrophilic ^b and low molecular weight compounds	Low	NOM has little influence on coagulant dose (i.e., mainly non-coagulable NOM)	0 ^c –40%; higher end for waters with high TOC
2–4	Mixture of hydrophilic and hydrophobic compounds; mixture of molecular weights	Medium	NOM influences coagulant dose	40–60%; higher end for waters with high TOC
>4	Mostly hydrophobic and high molecular weight compounds	High	NOM controls coagulant dose	60–80%; higher end for waters with high TOC

^a Adapted from Edzwald and Tobiason, 1999; AWWA, 2011a (Table 3-22)

^b The hydrophilic neutral fraction can have a high SUVA, which can mislead water treatment designers regarding the potential for organic carbon removal using coagulation (Edzwald, 1993).

^c Owen et al., 1993; Hargeheimer et al., 1994; Carpenter et al., 2013.

B.5.2.2 Chemical usage

Tracking chemical usage (e.g., coagulant dose, chlorine demand) and calculating the specific dose or demand (i.e., mg/L per mg/L DOC) can help water utilities assess changes in NOM character. For example, Chow et al. (2005) reported that the specific coagulant dose decreased when allochthonous NOM inputs increased. Also, Hwang et al. (2001) reported that the hydrophilic base fraction of NOM produces significant chlorine demand, as outlined in Table 10. This fraction comprises compounds that are biodegradable (e.g., amino acids). Thus, the removal of these compounds using biological filtration techniques (see section B.6.2.5) may decrease chlorine demand and DBPs (Prévost et al., 1998). Summers et al. (2013) cautioned that DOC and chlorine demand do not correlate to THMs and HAA5 (i.e., monochloroacetic acid, monobromoacetic acid, dichloroacetic acid, dibromoacetic acid and trichloroacetic acid) due to the presence of some compounds that exhibit strong chlorine demand but do not produce DBPs. By contrast, Roccaro et al. (2008) reported a linear correlation between THMs and chlorine consumption ($R^2 = 0.94$). The authors noted that chlorine consumption could be used to predict THM concentrations in the distribution system when NOM oxidation and halogenation processes dominate, compared with other reactions that consume chlorine (e.g., oxidation of inorganic species, photolytical and corrosion processes). Caution is recommended when assessing trends related to chlorine demand or specific chlorine demand, as reactions are likely to vary based on seasonal and weather-related effects, the treatment processes in place and where chlorine is added.

Table 10. Impact on chlorine demand by NOM fractions^a

Fraction	Chlorine demand
Hydrophobic	+
Hydrophilic – acids and neutrals	++
Hydrophilic – bases	++++
Colloidals	+

+ Lowest demand
++++ Highest demand

^a Adapted from Hwang et al., 2001

B.5.2.3 Disinfection by-products

Actual DBP concentrations measured in the distribution system provide a good indication of the reactivity of NOM. It is recommended that parameters used to characterize NOM be measured in conjunction with DBP samples to estimate the specific DBP yield (e.g., µg DBP/mg DOC). Also, inorganic compounds that enhance the reactivity of NOM to form DBPs should be characterized (i.e., ammonia, bromide, iodide and sulphur).

B.5.2.4 Other methods

NOM compounds can be fractionated using commercially available solid-phase extraction sorbents. However, measurement of the six NOM fractions (see Table C-3.4) is time- and labour-intensive (Minor et al., 2014; Goss et al., 2015) and may not correlate well with DBP formation potential results (Wright et al., 2016). Several researchers have investigated more rapid assessment methods (Martin-Mousset et al., 1997; Chow et al., 2004, 2006; Rosario-Ortiz et al., 2007; Dittmar et al., 2008; Ratpukdi et al., 2009).

Goss et al. (2017) compared three prepackaged solid phase extraction cartridges for the isolation of hydrophobic and hydrophilic fractions from three surface waters in Manitoba. The authors reported that the method could be used at water treatment plants to rapidly assess raw water quality, adapt treatment processes and verify treatment performance. By contrast, Wright et al. (2016) reported that solid phase extraction cartridges leaked variable amounts of organic carbon, skewed TOC results and correlated poorly with DBP formation potential results. The authors did not recommend using this method as a monitoring tool.

NOM compounds can also be physically fractionated based on differences in molecular size using membrane fractionation or size exclusion chromatography (Koudjonou et al., 2005). NOM is typically fractionated into four size ranges: <1, 1–10, 10–30 and >30 kDa. Size exclusion chromatography can be supplemented with organic carbon (LC-OCD) and/or organic nitrogen (LC-OND) detection (Huber et al., 2011). These methods are commonly reported in peer reviewed literature but are not yet used routinely by water utilities.

Fluorescence is another method that shows promise (McKnight et al., 2001; Wright et al., 2016). The advantages of fluorescence include rapid analysis with minimal sample preparation (Fellman et al., 2010; Bridgeman et al., 2011; Markechová et al., 2013; Sanchez et al., 2013). The fluorescent fractions of NOM exhibit intensity peaks at specific wavelengths; this allows their classification as terrestrial, microbial or anthropogenic organic matter, as well as humic-, fulvic- or protein-like compounds. Researchers have developed mathematical tools or algorithms to evaluate the large datasets that are generated and to compare differences between samples. Field units are commercially available and research continues as to how this method can be integrated as a routine monitoring tool (Bridgeman et al., 2011; Murphy et al., 2013; Wright et al., 2016; Peleato et al., 2017; Frank et al., 2018; Li et al., 2020).

A comprehensive review of these and other methods for the isolation and analysis of NOM is presented elsewhere (Minor et al., 2014).

B.5.3 Biological stability

BDOC and AOC are the two most widely used parameters for measuring the biological stability of water.

BDOC refers to the portion of DOC available to be utilized by heterotrophic bacteria (Escobar and Randall, 2001). Testing consists of measuring the DOC in the water before and after incubation with an inoculum of a natural bacterial population. The BDOC value is considered a measure of the hydrolyzable pool of carbon available for bacterial regrowth.

AOC represents the most readily degradable portion of the BDOC that can be taken up by bacteria and converted into organic biomass (Escobar and Randall, 2001). The test for AOC determines the growth potential of the water by measuring the growth yield of two pure strains of bacteria (*Pseudomonas fluorescens* strain P17, *Spirillum* strain NOX) over several days and comparing these observations against a calibration curve for the growth produced using solutions of organic carbon standards (e.g., acetate or oxalate) (LeChevallier et al., 2015a). The AOC concentration is considered one measure of the biostability of water for heterotrophic bacterial growth (Escobar and Randall, 2001). AOC is often used as the method to predict bacterial regrowth, as it returns a value that corresponds to a bacterial count (Escobar and Randall, 2001).

It has been suggested that the AOC:BDOC ratio can be used as an indication of the relative biological stability of the biodegradable organic compounds present in drinking water (Escobar and Randall, 2001). Both methods are time-consuming and require a high level of analytical expertise. Camper (2004) and van der Kooij et al. (2015) reported that carbon compounds not measured by AOC and BDOC may also influence biofilm growth and that these measurements alone may not be sufficient for estimating regrowth potential.

The water industry has been investigating the benefits of more rapid methods that use adenosine triphosphate (ATP) measurements or flow cytometry technology (Hammes et al., 2012; Besmer et al., 2014; Nescerecka et al., 2014; Pharand et al., 2014; Gilmore and Summers, 2015; Liu et al., 2015; van der Kooij et al., 2015; Besmer and Hammes, 2016; Elhadidy et al., 2016).

ATP measurements are gaining popularity as an indicator of microbiological biomass (Siebel et al., 2008). ATP methods are low cost, rapid and require a modest amount of training (LeChevallier et al., 2015a). Still, an understanding of the meaning of the measurements as they relate to other water quality dimensions such as viable and culturable cell counts is necessary when considering the inclusion of ATP analysis in a monitoring program (Siebel et al., 2008; Hammes et al., 2010).

Flow cytometry has also emerged as a potential tool for rapid online monitoring of general microbial water quality (Prest et al., 2013, 2016). Because of its ability to measure changes in bacterial cell counts, flow cytometry has been proposed as one of several methods for assessing biological stability (Lautenschlager et al., 2013; Prest et al., 2013, 2016; Nescerecka et al., 2014). The technology is advanced and has considerable requirements for equipment, user training and data processing (Hammes and Egli, 2010). Several studies have investigated online biostability monitoring using flow cytometry, but standardized methods have not yet been developed for drinking water applications (Hammes and Egli, 2010; Lautenschlager et al., 2013; Prest et al., 2013). In an investigation of a full-scale chlorinated drinking water system, Nescerecka et al. (2014) found that flow cytometry, in combination with ATP measurements, provided more meaningful information than heterotrophic plate counts for assessing and understanding biological stability at various points in the distribution system. In van der Kooij et al. (2015) it is reported that the use of ATP methods to assess biofilm formation potential and

biofilm accumulation rates provided an improved understanding of biological instability in distributed water (without a disinfectant residual) that would not have been revealed based on the assessment of AOC only.

LeChevallier et al. (2015a) completed a statistical analysis of full-scale data for six water utilities and concluded that the most useful measures to assess biological stability were variability in disinfectant residual (measured by the coefficient of variation), biofilm formation rate (measured by ATP accumulated on mild steel coupons) and changes in corrosion rates (measured by linear polarization resistance using mild steel coupons). LeChevallier et al. (2015b) provide guidance to help water utilities produce biologically stable water and establish an appropriate system-specific monitoring program. Guidance on monitoring the biological stability of drinking water distribution systems is also available in Health Canada (in preparation).

B.6 Treatment and distribution system considerations

The source-to-tap or water safety plan approach, which includes careful selection of the highest quality water source and source water protection, is an accepted approach to manage risks to drinking water safety (O'Connor, 2002; CCME, 2004; WHO, 2012). Source-specific treatability studies, including bench- and/or pilot-scale testing, should be conducted to determine the most suitable treatment alternatives for the full range of water quality conditions (Valade et al., 2009; Huck and Sozański, 2011).

Temporal variations in the concentration and character of NOM can have a significant influence on the selection, design and operation of water treatment processes (Sillanpää, 2015). More variable weather patterns associated with climate change will place increased importance on proper process selection (Huck and Coffey, 2004) and day-to-day process control (Wright et al., 2016). Water utilities should integrate risks related to changes in climate (e.g., algal blooms, drought, fire, flood) into the process to maximize the reliability, robustness and resilience of their systems (Emelko et al., 2011b; Irias, 2019).

B.6.1 Choice of appropriate treatment

To appropriately select, design and operate water treatment facilities, an understanding of the variations in the concentration and character of NOM is necessary—for the full range of conditions encountered over the year, for both surface and groundwater sources (AWWA, 2011a; Sillanpää, 2015). To determine the most appropriate treatment processes, water utilities should have knowledge regarding the following (Ivančev-Tumbas, 2014):

- the origin, occurrence and fluctuations in NOM;
- interactions with other water constituents (e.g., enhanced reactivity due to bromide);
- interactions with chemicals used during treatment (e.g., NOM creates a disinfectant and coagulant demand that must be overcome to produce microbiologically safe drinking water);
- interactions with unit processes (e.g., NOM fouls adsorbents and membranes); and
- impacts on distribution system water quality.

The appropriate type and level of treatment should take into account source-specific fluctuations in water quality, including short-term degradation, variability in treatment performance and distribution system conditions (Kastl et al., 2016).

A source-specific treatability study should be conducted to assess and compare treatment options for the removal of NOM (Goss and Gorczyca, 2013; Plourde-Lescelleur et al., 2015; Kastl et al., 2016). The treatability study should include bench- and/or pilot-scale testing and

consider concomitant water quality goals related to microbial risks, DBPs, biological stability and corrosion control. Parameters to be considered as part of a treatability study include chemical doses and residuals, turbidity, organic content (e.g., DOC, UV254, COD, colour), organic character (e.g., hydrophobicity, size, specific UV absorbance), pH and alkalinity, anions (e.g., bromide, chloride, fluoride, nitrate/nitrite, orthophosphate, sulphate), DBP formation potential that is representative of the distribution system, biostability, and corrosion characteristics (Gregor et al., 1997; Karanfil et al., 2007; Shin et al., 2008; Cho et al., 2010; Brown and Cornwell, 2011). The optimum solution will be source-specific, and multiple treatment processes may be needed to adequately remove NOM at all times of the year (Collins et al., 1986; Chang et al., 2001; Hua and Reckhow, 2007a; Karanfil et al., 2007; Fabris et al., 2008; Kristiana et al., 2009; Carpenter et al., 2013; Hua et al., 2015; Sillanpää, 2015). The lack of a source-specific treatability study may result in the selection of inappropriate treatment, an increase in disinfection by-product concentrations following the implementation of treatment or other unintended consequences.

B.6.2 Treatment options

A number of treatment options are available to remove NOM. Optimized coagulation is the most commonly used method, as it is effective in most applications. However, its applicability should be carefully analyzed on a source-specific basis because coagulation can only remove some NOM fractions; the remaining fractions (i.e., those not removed by coagulation) may react with disinfectants such that DBP guidelines are not achieved. For example, allochthonous NOM tends to be hydrophobic in nature and is generally amenable to coagulation, whereas hydrophilic NOM tends to be more difficult to treat (Volk et al., 2002; Chow et al., 2004, 2006). In fact, for sources high in hydrophilic neutral NOM, coagulation will be ineffective (Chow et al., 2006). As a result, it is very important that jar testing and DBP formation potential testing be performed to determine the feasibility of optimized coagulation for NOM removal. Additional or alternative treatment options include nanofiltration, ion exchange, GAC or powdered activated carbon (PAC), biological filtration and oxidation processes.

The literature cautions that the specific DBP yield (i.e., $\mu\text{g DBP/mg DOC}$) can sometimes be greater in the treated water than in the disinfected raw water (Jacangelo et al., 1995; Singer et al., 2007; de la Rubia et al., 2008; Newfoundland and Labrador Department of Environment and Conservation, 2011). This is attributed to a higher bromide:DOC ratio following treatment. Because bromide is not removed by most treatment processes, more brominated DBPs may form following treatment if NOM removal is inadequate. As a result, it is very important that a source-specific treatability study be performed to assess and compare treatment options; this study should include bench- and/or pilot-scale testing to determine the DBP formation potential.

Treatment options and their reported effectiveness are briefly discussed below. Results are presented to demonstrate that NOM removal can be highly variable. More detailed information regarding treatment is available in other sources (Parsons et al., 2007; AWWA, 2011a; Bond et al., 2011; Huck and Sozański, 2011; Sillanpää, 2015).

It is important that water treatment operators understand the NOM removal mechanisms, since changes in treatment practices can significantly impact water quality (Ivančev-Tumbas, 2014). Thus, operator training is also needed to ensure the effective operation of treatment barriers at all times (Smeets et al., 2009). Maintaining current knowledge of best practices and remaining aware of advancements in the drinking water industry are important to ensure water safety.

B.6.2.1 Optimized coagulation

Coagulation is a complex chemical process that can be optimized for both NOM and turbidity removal (Edzwald and Harhoff, 2012). Coagulation involves two primary mechanisms: one consists of charge neutralization and the formation of insoluble precipitates; the other involves adsorption onto aluminum or ferric hydroxide floc (i.e., sweep coagulation) (Dempsey et al., 1984). Each mechanism is favoured by a particular set of operating conditions related to pH and coagulant dose. As pH increases, NOM becomes increasingly negatively charged, but coagulant hydrolysis products with lower positive charge dominate. Thus, at pH >7 a four-fold increase in the coagulant dose is necessary to overcome NOM's negative charge compared with that required at pH 5.5. Above pH 7, NOM removal is poor (Semmens and Field, 1980; Edzwald and Van Benschoten, 1990; Edzwald and Tobiasson, 1999). Physical factors (such as mixing of the coagulant and mixing conditions in the flocculator) can affect floc formation; in most cases, however, the coagulation chemistry controls the process (Kavanaugh, 1978; Vadasarukkai and Gagnon, 2015; Vadasarukkai, 2016).

The choice of coagulant will depend on the characteristics of the water to be treated. Available coagulant choices (e.g., aluminum- and ferric-based coagulants, inorganic polymer flocculants, organic polyelectrolytes, composite coagulants and novel coagulants) are discussed elsewhere (Sillanpää and Matilainen, 2015; Sillanpää et al., 2018). While some coagulants provide a wider operational window with respect to pH, it is noteworthy that for all metal coagulants the pH of minimum solubility increases as temperature decreases (Pernitsky, 2003). For alum, optimum performance generally occurs at pH values close to the pH of minimum solubility (i.e., 6.5–6.7 at 4°C and 6.0–6.2 at 20°C) (Edzwald and Kaminski, 2009). As the pH of minimum solubility is higher at lower temperatures, a higher coagulant dose may be needed to overcome the more negative charge on NOM with the lower positive charge on coagulant hydrolysis products, as noted above. Strict pH control is necessary for optimum coagulation; pH should be kept constant from coagulant addition to after filtration to effectively remove floc particles. Even a small pH change can release NOM that was previously incorporated into flocs (Slavik et al., 2012). Jar testing is recommended to optimize coagulant selection.

NOM also determines the size, structure, and strength of the flocs, controlling both the extent and the rate of the clarification or filtration processes (Eikebrokk and Saltnes, 2001; Newcombe and Dixon, 2006; Parsons et al., 2007). Several studies have demonstrated that low-density NOM flocs are more amenable to flotation than to sedimentation (Plummer et al., 1995; Edzwald and Kelley, 1998; Edzwald et al., 1999, 2000, 2003; Harrington et al., 2001; Edzwald, 2010; Gregory and Edzwald, 2011). Alternatively, the addition of a coagulant aid (e.g., activated silica, bentonite, lime, polymer) may be needed to form settleable flocs (Semmens and Field, 1980; Edwards and Amirtharajah, 1985; Gregor et al., 1997).

The charge-driven nature of NOM coagulation means that electrophoretic monitoring is appropriate (Bond et al., 2011). Otherwise, operators are not aware of coagulant under-dosing until spikes in settled water or filter effluent turbidity are observed (Pernitsky, 2003). Ideally, the raw water should be continuously monitored to optimize the coagulant dose (Pernitsky, 2003; Newcombe and Dixon, 2006; Sharp et al., 2006; Shin et al., 2008). Online monitoring tools for NOM include TOC, DOC, UV absorbance/transmittance and COD; for particle destabilization they include zeta potential or streaming current (Conio et al., 2002; Newcombe and Dixon, 2006; AWWA, 2011b). Maximum NOM removals have been reported when the coagulant particle charge is near neutral as measured by zeta potential (Sharp et al., 2006; Sharp, 2015) or streaming current (McVicar et al., 2015). Failure to adjust the coagulant dose in accordance with a change in NOM may contribute to suboptimal coagulation conditions and a decrease in pathogen log

removal capability (Edzwald, 2017).

Table 11 summarizes the variability in DOC removal achieved at several full-scale chemically assisted filtration plants. Low to zero organic carbon removals are reported for certain periods. Hargesheimer et al. (1994) reported 0% removal at various times during 1993, namely early March, May, August and December. It is possible that these timeframes represent snow cover (March, December) or base flow conditions with low allochthonous NOM inputs (May and August). Carpenter et al. (2013) reported 0% removal in early September 2011 for Plant 1 and in August 2011 for Plant 2. NOM is expected to be generated by autochthonous sources at this time (i.e., late summer) and would likely be hydrophilic in nature.

Table 11. Range of TOC/DOC removals reported in full-scale studies

Reference	Source water quality				Treatment processes	TOC/DOC % removal (mean)
	Parameter ^a	Min	Max	Mean		
Hargesheimer et al., 1994	TOC	0.9	4.5	2.4	Conventional filtration (alum)	0–28 (8.7)
Jacangelo et al., 1995	DOC	1.2	7.8	2.1–3.5 ^b	Conventional with GAC (coagulant not specified)	8–48 ^c
Volk et al., 2002 ^d	DOC	2.15	11.90	4.00	Coagulation (ferric chloride and cationic polymer), flocculation, clarification	7.1–66 (34.7)
	UV254	0.037	0.830	0.118		
	SUVA	1.40	10.51	2.81		
Volk et al., 2002 ^d	DOC	2.15	11.90	4.00	Conventional (ferric chloride and cationic polymer) with GAC	16.9–72.9 (41.9)
	UV254	0.037	0.830	0.118		
	SUVA	1.40	10.51	2.81		
Chow et al., 2005 ^e	DOC	8.2 ^f	11.8	Not given	Conventional filtration (alum and cationic polymer)	36–57 (47)
Chow et al., 2005 ^e	DOC	11.6 ^f	15.8	Not given	DAF filtration (alum and cationic polymer)	56–65 (62)
Carpenter et al., 2013 ^g	DOC	0.9	2.2	1.3	Direct filtration (Plant 1) (coagulant not specified)	0–50 (28.2)
	UV254	0.01	0.10	0.03		
	SUVA	2.00	4.41	2.73		
Carpenter et al., 2013 ^g	DOC	0.9	2.2	1.3	Direct filtration (Plant 2) (coagulant not specified)	0–45 (27.9)
	UV254	0.01	0.10	0.03		
	SUVA	2.00	4.41	2.73		
Nova Scotia Environment, 2016	TOC	4.3	8.3	6.2	Conventional filtration (Plant 1) (alum)	27–78 (66)
	TOC	10.4	22.7	15.6	Conventional filtration (Plant 2) (alum)	71–89 (80)

^a TOC/DOC = mg/L; UV254 = cm⁻¹; SUVA = L/mg·m

^b Range in annual means from 1978 to 1992

^c Range in annual means not provided

^d Daily monitoring from August 28, 1998 to June 13, 2000

- ^e Monthly monitoring from January 2001 to July 2002
- ^f Interpretation from a graph
- ^g Monthly monitoring from April 14, 2010 to September 22, 2011

Table 12 summarizes the TOC compliance monitoring data published by the U.S. EPA (2016) as part of its third Six-Year Review. The data represent the TOC removal (in percent) achieved at conventional surface water treatment plants as a function of the influent water quality matrix established by the Disinfectants/DBP Rule. In general, the U.S. EPA concluded that regulated facilities are achieving higher removals than mandated (see section B.8), although some facilities have not been able to achieve removal requirements. The report cautioned that the data analysis could not determine which facilities are permitted to determine alternative performance criteria or which may have treated water TOC less than 2 mg/L (U.S. EPA, 2016).

Table 12. TOC removals reported from U.S. EPA compliance monitoring data (2006–2011)^a

Influent TOC (mg/L)	Influent alkalinity (mg/L CaCO ₃)		
	0–60 mg/L	>60–120 mg/L	>120 mg/L
>2–4	Mean ^b 41.7%	Mean 35.2%	Mean 30.4%
	Median ^c 41.6%	Median 35.1%	Median 30.1%
>4–8	Mean 54.7%	Mean 46.8%	Mean 44.1%
	Median 54.3%	Median 46.3%	Median 43.9%
>8	Mean 66.2%	Mean 46.3%	Mean 46.9%
	Median 66.4%	Median 44.2%	Median 47.8%

^a Adapted from U.S. EPA (2016)

^b Mean TOC removal

^c Median TOC removal

At pilot-scale, Braun et al. (2014) assessed conventional treatment (in parallel with membrane filtration, ion exchange and GAC) for a three-year period that included an extended drought and two distinct flood periods. The authors reported variable DOC removal with conventional treatment (range = 32–61%; alum dose = 20–160 mg/L; coagulation pH = 6.0–6.5). The authors also noted that water quality was the best during the drought period, but conventional treatment achieved its lowest DOC removal and highest variability during this period. During drought periods, NOM tends to be generated by autochthonous sources and be hydrophilic in nature. This supports the full-scale study results indicating that hydrophilic NOM can be challenging to treat.

The results from numerous bench-scale studies were reviewed and summarized by Bond et al. (2011). In these studies, removal percentages were determined for DOC, UV254, THM precursors and HAA precursors for numerous treatment processes, including coagulation and coagulation combined with other processes (see Table 13). Bench-scale results published by Plourde-Lescelleur et al. (2015) are also summarized in Table 13; in this study, two coagulants were compared for conventional treatment and alum coagulation combined with ion exchange, ozonation or PAC was assessed for six Canadian surface water sources. The results summarized in Table 13 confirm the variable performance that can be achieved using coagulation alone and that enhanced removals can be achieved by integrating coagulation with other processes.

Collectively, the full-, pilot- and bench-scale results indicate that coagulation can be effective, however poor results can also be observed. These findings reiterate the need to conduct jar testing to confirm the applicability of coagulation to adequately remove NOM and DBP precursors for the full range of water quality conditions. The other processes noted in Table 13 are further discussed in subsequent sections.

Table 13. Range of removals reported in bench-scale studies

Treatment process	Percent removal (mean)				
	DOC	UV254	THM precursors	HAA precursors	
Coagulation-based processes					
Coagulation					
Bond et al., 2011 ^a					
Alum	17–33 (25)	3–80 (46)	7–71 (36)	15–78 (38)	
Plourde-Lescelleur et al., 2015 ^b					
Alum	26–70 (54)	34–85 (69)	48–83 (70)	48–93 (79)	
Ferric	13–74 (53)	30–88 (68)	44–90 (72)	69–97 (81)	
Ion exchange–coagulation					
Bond et al., 2011 ^a	42–76 (59)	47–96 (79)	27–88 (70)	52–80 (67)	
Plourde-Lescelleur et al., 2015 ^b	39–75 (63)	47–90 (77)	50–92 (76)	61–97 (84)	
Alum coagulation–PAC					
Plourde-Lescelleur et al., 2015 ^b	58–86 (77)	57–96 (88)	73–93 (85)	91–99 (96)	
Alum coagulation–ozonation					
Bond et al., 2011 ^a	16–34 (23)	49–69 (61)	47–58 (51)	60–81 (71)	
Plourde-Lescelleur et al., 2015 ^b	21–69 (54)	55–93 (82)	59–90 (78)	48–97 (80)	
Pre-ozonation–alum coagulation					
Bond et al., 2011 ^a	0–30 (15)	42–69 (60)	51–66 (57)	48–76 (66)	
Nanofiltration^c (section B.6.2.2)					
Bond et al., 2011 ^a	86–93 (90)	89–99 (96)	66–98 (87)	67–99 (87)	
Plourde-Lescelleur et al., 2015 ^b	77–89 (84)	79–93 (87)	75–98 (89)	88–100 (97)	
Oxidation processes (sections B.6.2.5 and B.6.2.6)					
Ozonation					
Bond et al., 2011 ^a	8–16 (12)	28–77 (58)	0–43 (14)	-50 to 20 (4)	
Ozonation–biological sand					
Bond et al., 2011 ^a	Not available	Not available	-5 to 54 (42)	-4 to 68 (51)	
Ozonation–UV					
Bond et al., 2011 ^a	17–56 (33)	90–94 (92)	48–89 (67)	Not available	
UV–H₂O₂					
Bond et al., 2011 ^a	-11 to 20 (-1)	20–59 (34)	8–73 (43)	DCAA -197 to -11 (-79)	TCAA 6–69 (24)
UV–H₂O₂–biological sand					
Bond et al., 2011 ^a	38–80 (59)	45–81 (64)	42–85 (60)	DCAA 3–63 (36)	TCAA 42–85 (62)

^a Range of data from numerous bench-scale studies presented in Bond et al., 2011

^b Range of data for six Canadian surface water sources

^c Molecular weight and cutoff = 100–400 kDa

B.6.2.2 Membrane filtration

Four types of pressure-driven membranes are currently used in drinking water treatment: microfiltration (MF), ultrafiltration (UF), nanofiltration (NF) and reverse osmosis (RO). Membranes are generally classified by the type of substances they remove, operating pressure and pore size or molecular weight and cutoff (MWCO). MF and UF are referred to as low-pressure membranes and are used for particle/pathogen removal. The predominant removal mechanism is straining or size exclusion. NF and RO are referred to as high-pressure membranes and are used for the removal of NOM and inorganics (e.g., sodium, chloride, calcium,

magnesium). The predominant removal mechanism is differences in solubility or diffusivity.

The size distribution of NOM varies between sources, but generally over 50% of NOM molecules have a molecular weight of <1kDa and 80% have a molecular weight of <10kDa (Sillanpää et al., 2015a). As a result, a tight NF membrane is required to remove the majority of DBP precursors, as shown in Figure 1. Studies indicate that the optimum MWCO for NOM removal is 0.2–0.3 kDa (Jacangelo et al., 1995, 1997; Bond et al., 2011; Sillanpää et al., 2015a).

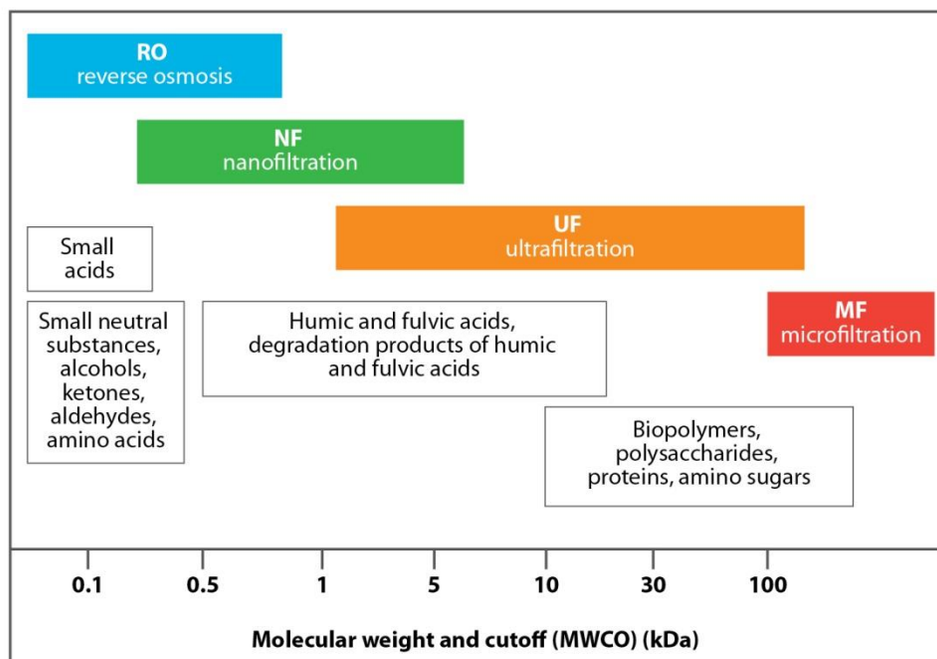


Figure 1. Natural organic matter fractions removed by membrane processes (adapted from Sillanpää et al., 2015a)

Figure 1 illustrates that MF membranes cannot remove any NOM fractions other than biopolymers. UF membranes may remove some NOM, as shown in Figure 1, but DBP formation potential may not decrease adequately. For example, Lamsal et al. (2012) reported an overall DOC removal of 66% for full-scale UF membranes (absolute pore size = 0.01 µm). THM and HAA formation potentials decreased by 54% and 77%, respectively but remained high at 200 µg/L and 80 µg/L, respectively (test conditions = chlorine 1.0±0.4 mg/L at room temperature for 24 h). Ødegaard et al. (2000; 2010) reported colour removals of >85% for 27 full-scale NF membrane plants in Norway (raw water colour = 35–50 mg Pt/L; MWCO = 1–2 kDa).

At pilot-scale, Braun et al. (2014) assessed an integrated membrane system (MF nominal size = 0.2 µm; NF MWCO = 270 Da) for the same three-year period described above for conventional treatment (see section B.6.2.1). DOC removal ranged from 89–97% (data interpreted from a graph) and represented the highest percent removals when compared to conventional treatment, ion exchange and GAC which were tested in parallel to the integrated membrane system. As with conventional treatment (see section B.6.2.1), the MF/NF system achieved the lowest DOC removal during the drought period. Bench-scale results published by Bond et al. (2011) and Plourde-Lescelleur et al. (2015) for NF membranes are summarized in Table 13. Reported removals were high, similar to the pilot-scale results.

Collectively, the full-, pilot- and bench-scale results indicate that membrane treatment can be effective when the membrane pore size is optimized for NOM removal ($\approx 200\text{--}300$ Da). However, it is important to consider fouling potential as NOM is known to foul membranes. Indicators of fouling potential include low SUVA, high hydrophilic fraction, high dissolved nitrogen or high biopolymer concentration (Lee et al., 2006; Amy, 2008; Croft, 2012; Kimura et al., 2014; Siembida-Lösch et al., 2014). Also, UF membranes cannot remove AOC (usually <1 kDa) unless combined with a tight NF or RO membrane or biological treatment (Sillanpää et al., 2015a). The use of a coagulant or absorbent may improve DBP precursor removal by MF/UF processes (Jacangelo et al., 1997). Pretreatment requirements for NOM removal should be considered as part of a source-specific treatability study whenever the optimum MWCO for NOM removal (i.e., $0.2\text{--}0.3$ kDa) is not used. Pilot testing is recommended to assess fouling potential and the need for pre-treatment (Huehmer and Voutchkov, 2007).

B.6.2.3 Ion exchange

Ion exchange is a process in which ions from the raw water are exchanged with ions within the solid phase of a resin. It was recognized in the early 1960s that ion exchange processes could remove NOM, mainly because NOM was found to foul ion exchange resins used to remove other contaminants (Frisch and Kunin, 1960; Ungar, 1962; AWWA, 2011a). The dominant removal mechanism involves ion exchange (i.e., electrostatic), with hydrophobic adsorption and hydrogen bonding also playing a role (Fu and Symons, 1990; Bolto et al., 2002).

Ion exchange can effectively remove charged NOM compounds of small and large molecular weights, making it generally more effective than coagulation; however, similar to coagulation, ion exchange is less effective at removing neutral (e.g., uncharged) hydrophilic compounds (Kim and Symons, 1991; Bolto et al., 2002; 2004; Parsons et al., 2007; Cornelissen et al., 2008; Humbert et al., 2008). Fettig (1999) reported that the non-removable fraction of NOM by full-scale ion exchange treatment plants could vary from $<10\%$ to around 40% . Hongve et al. (1999) found that full-scale ion exchange was more effective for DOC removal (8.1 mg/L decreased to 1.7 mg/L) compared to colour removal (75 mg/L Pt decreased to 30 mg/L Pt). Longer contact times, or another treatment process, may be necessary when improved colour removal is necessary (Hongve et al., 1999; Ødegaard et al., 1999).

Ion exchange has received significant attention since 2000 as a result of the development of the magnetic ion exchange resin, which was specially designed for NOM removal (Slunjski et al., 2000; Drikas et al., 2003; Fearing et al., 2004c; Budd et al., 2005; Singer et al., 2007, 2009; Bond et al., 2010; Brown and Cornwell, 2011). Levchuk et al. (2018) summarized the results of 22 studies (1 full-scale; 21 bench-scale) conducted between 1997–2016, 50% of which assessed the effectiveness of magnetic ion exchange.

Factors influencing the performance of ion exchange include NOM concentration and character, water quality (particularly the concentration of competing anions such as bicarbonate and sulphate), resin properties (polymer composition, porosity and charged functional groups) and operational variables (resin dose, contact time, regeneration frequency). However, ion exchange processes do not remove turbidity; hence they are typically applied with a turbidity removal process (Drikas et al., 2003).

Several researchers reported that ion exchange was more effective than coagulation for NOM removal, either on its own (Drikas et al., 2003; Singer et al., 2007, 2009) or in combination with coagulation (Drikas et al., 2003; Brown and Cornwell, 2011; Braun et al., 2014).

Drikas et al. (2011) reported average DOC removal of 54% for a full-scale ion exchange-alum coagulation facility (resin dose $\approx 8\text{--}16$ mL/L for 10 minutes; alum dose $\approx 6\text{--}10$ mg/L). The

authors attributed the majority of DOC removal to the use of magnetic ion exchange resin. Singer et al. (2009) completed a comprehensive review of numerous full- and pilot-scale studies involving 21 sources in Australia and the United States and reported DOC removals of 36–80% (resin dose \approx 0.2–2.8 mL/L; contact time not given). The authors attributed the wide range in DOC removal to the presence of hydrophilic NOM with base and neutral charge.

Table 14 summarizes the results from several other pilot-scale studies. Researchers studying magnetic ion exchange (alone or with coagulation) reported a wide variability in DOC removal; removal was reported to be affected by the NOM character (Fearing et al., 2004c; Singer et al., 2007; Braun et al., 2014), with higher DOC removals being observed with increasing SUVA (Singer et al., 2009). Braun et al. (2014) demonstrated that integrating GAC as an additional process can improve NOM removal. The authors also noted that magnetic ion exchange achieved its lowest DOC removal and highest variability during a drought period, similar to conventional and membrane treatment discussed above. Fearing et al. (2004c) highlighted the variability that occurs due to resin dose and contact time.

Table 14. Range of DOC removals reported in pilot-scale studies

Treatment process	DOC percent removal	Process details	Reference
Magnetic ion exchange	35–67	Resin dose = 15–20 mL/L Contact time = 15–20 min	Singer et al. 2007
	64–74	Resin dose = 6–8 mL/L Contact time = 15 min	Drikas et al., 2003
Magnetic ion exchange with coagulation	64–76	Resin dose = 6 mL/L Contact time = 10–15 min	Drikas et al., 2003
	10–20	Resin dose = 2 mL/L Contact time = 10–20 min	Fearing et al., 2004c
	\approx 50	Resin dose = 20 mL/L contact time = 5 min	Fearing et al., 2004c
	66.1–82.1	Resin dose = 30 mL/L Contact time = 60 min	Fearing et al., 2004c
	52–81 ^a	Resin dose = 15 mL/L Contact time = 10 min	Braun et al., 2014
Magnetic ion exchange with coagulation and GAC	74–91 ^a	Resin dose = 15 mL/L Contact time = 10 min	Braun et al., 2014

^a Interpreted from a graph.

Table 15 summarizes the results from several bench-scale studies that assessed other resins. Results demonstrate the variability in performance that can occur by resin and system, similar to the magnetic ion exchange resin results presented above.

Table 15. Range of DOC removals reported in bench-scale studies

Reference	Resin	DOC percent removal
Afcharian et al., 1997 ^a (France)	Lewatit MP 500	59
	Lewatit S6328A	73
Hwang et al., 2001 (United States)	Dowex MSA (Strong base)	Source A – 77
		Source B – 71
		Source C – 57
		Source D – 59
	Imac HP661 (Weak base)	Source A – 56
		Source B – 52
Bolto et al., 2002 ^b (Australia)	IRA 458	Source A – 69
		Source B – 90
	IRA 958	Source A – 72
		Source B – 90
	SIR 22P	Source A – 84
		Source B – 85
Humbert et al., 2008 ^c (France)	IRA 938	75
	Dowex 11	68
	Dowex MSA	61
	IRA 958	58
	Ambersorb	46
Brezinski et al., 2019 ^d (Rainy River, Ontario)	Amberlite PWA 9	71.7
	Dow TAN	69.4
	Purolite A502P	68.7
	Purolite A860	67.7

^a The authors also assessed percent removal of UV254, fluorescence, BDOC and chlorine demand.

^b The authors also assessed other resins for source A.

^c Results were interpreted from a graph for 30 minutes contact time; the authors also assessed 3 and 120 minutes.

^d The authors also assessed removal of THM formation potential.

The use of ion exchange in combination with other processes can have some operational benefits and impacts that should be considered in a source-specific treatability study. For example, ion exchange prior to coagulation can reduce the coagulant dose and associated sludge production, lower the settled water turbidity, reduce the use of pH adjustment chemicals, reduce the disinfectant dose, and stabilize distribution system chlorine residuals (Budd et al., 2005; Brown and Cornwell, 2011). Ion exchange processes may also remove some bromide from sources that have low alkalinity and sulphate concentrations, due to minimal competition for ion exchange sites (Singer et al., 2007). Humbert et al. (2008) reported improved performance of activated carbon for pesticide removal when ion exchange was used to enhance NOM removal. However, Brown and Cornwell (2011) noted that ion exchange treatment (e.g., chloride addition) and lower doses of sulphate-based coagulants could increase the potential for corrosion, due to changes in the chloride:sulphate mass ratio. More information on the chloride:sulphate mass ratio and its impacts is available in Health Canada (2019b)

Collectively, the full-, pilot- and bench-scale results indicate that ion exchange can be effective when the resin dose and contact time is optimized for NOM removal, however poor results have also been reported. These findings reiterate the need to conduct a treatability study, including bench- or pilot-scale testing, to determine the optimum configuration and to assess unintended consequences for the full range of water quality conditions (Fearing et al., 2004c;

Brown and Cornwell, 2011). The handling and disposal of residuals generated by ion exchange processes should also be considered.

Water utilities that use ion exchange for the removal of other anions (e.g., arsenic, chromium, nitrate, uranium) should be aware that NOM competes for ion exchange sites and can decrease process efficacy (Frisch and Kunin, 1960; Ungar, 1962). Pretreatment for NOM removal may be required to ensure that the process remains economical for its intended purpose (Bursill, 2001).

B.6.2.4 Activated carbon

Activated carbon is an absorbent material that provides a surface on which ions or molecules in the raw water can concentrate. It can be applied in two ways: slurry applications using powdered activated carbon (PAC) or fixed bed reactors with granular activated carbon (GAC) (AWWA, 2011a). The removal mechanisms involve adsorption of dissolved organic matter onto PAC or GAC, as well as biodegradation of BOM in GAC fixed bed reactors if an active biofilm forms. Chowdhury et al. (2010) found that biofilms can form in GAC macropores even in the presence of chlorine.

The primary use of PAC and GAC in water treatment is to remove micropollutants as well as taste- and odour-causing compounds. The use of PAC offers the advantage of providing virgin carbon when required (e.g., during the taste and odour season). GAC fixed bed reactors are operated similarly to conventional rapid rate filters; hence the GAC characteristics (e.g., type, particle size, reactivation method) and operating conditions (e.g., filter velocity, empty bed contact time, backwashing regime, filter run time) influence their performance.

The large specific surface area and well-developed porous structure of GAC can provide high sorption capacity for organic molecules (Simpson, 2008). However, GAC has not been widely used as a primary NOM control strategy because the adsorption capacity of GAC tends to be quickly exhausted (i.e., in the order of months) and regeneration can be costly (Prévost et al., 1998; Huck and Sozański, 2011). For example, the U.S. EPA (2016) reports a 120-day reactivation frequency for systems with TOC of <6 mg/L using an empty bed contact time of 10 min. The removal of high molecular weight hydrophobic NOM by conventional treatment processes can significantly increase the operational life of GAC (Karanfil et al., 2007). In addition, once the adsorption capacity is exhausted, GAC can continue to remove NOM through the biodegradation mechanism, albeit at lower efficiencies (Bond et al., 2011; Gibert et al., 2013a).

Thus, integrating PAC or GAC as an additional process can improve NOM removal (AWWA, 2011a). For PAC, the dose and contact time are important factors. Results for two full-scale conventional water treatment plants (both using alum coagulation) are reported in the literature. In one case, a PAC dose of 150 mg/L improved DOC removals by 20% and THM precursors by 80% (Kristiana et al., 2011). In the other case, a PAC dose of 11 mg/L improved DOC removals by 7% but achieved no improvement in the removal of THM precursors (Carrière et al., 2009). Jar testing is recommended to optimize the PAC type, dose and contact time. For GAC, studies indicate that the pore volume should be in a size range that matches the source-specific NOM for GAC to be effective (Karanfil et al., 2007; Gibert et al., 2013b). As noted by Karanfil et al. (2007), the surface area and total pore volume are not sufficient criteria for selecting a GAC for NOM removal, because these parameters do not provide information about the accessible pore region. The authors suggest that water utilities request detailed information about the pore size distribution and the pH of the point of zero charge for candidate GACs. Rapid small-scale column tests should be conducted to compare the performance of alternative GACs,

particularly for low SUVA sources (Ates et al., 2007; Karanfil et al., 2007). Abrasion of GAC particles should also be considered, as abrasion can lead to the loss of GAC material and stratification within the bed, both of which are undesirable (Gibert et al., 2013b).

Water utilities that use activated carbon for the removal of pesticides or other trace contaminants should be aware that NOM competes for adsorption sites and can decrease process efficacy (Haist-Gulde and Happel, 2012). Pretreatment may be required for NOM removal, to ensure that the process remains economical for its intended purpose (Bursill, 2001).

B.6.2.5 Biological treatment

Biological treatment involves targeting the removal of the BOM fraction that encourages biofilm growth in the distribution system (section B.4.1.3) and increases chlorine demand (section B.5.2.2) (Prévost et al., 1998). The effectiveness of biological treatment therefore depends on the amount of BOM that is present in the water to be treated, the microbial community consuming the BOM and the temperature (Drewes et al., 2009; Diem et al., 2013). Recalcitrant or refractory NOM is unlikely to be removed by biological processes unless it is oxidized to transform it into BOM. Biological treatment generally improves the biological stability of the water and decreases DBP concentrations, as well as tastes and odours (Servais et al., 2005).

The main biological treatment processes for drinking water include riverbank filtration, rapid granular media filtration without the maintenance of a disinfectant residual across the bed and slow sand filtration.

Riverbank filtration

Riverbank filtration (RBF) involves locating vertical or horizontal water supply wells near a river to use the riverbank and adjacent aquifer as a natural filter to remove contaminants, including BOM. As water proceeds to the groundwater table, concentrations are lowered through adsorption, biodegradation and dilution with groundwater (Piet and Zoeteman, 1980; Bize et al., 1981; Kuehn and Mueller, 2000; Ray et al., 2002).

Kuehn and Mueller (2000) reported that RBF decreased DOC and AOC concentrations by 27% and 63%, respectively. Weiss et al. (2003) monitored three full-scale RBF sites and reported that TOC/DOC concentrations and THM/HAA formation potentials were lower in the RBF wells by approximately 35–70% and 50–80%, respectively. Wang et al. (2002) found that TOC concentrations decreased by approximately 50% through the RBF process while BDOC was completely removed. Drewes et al. (2009) reported that TOC concentrations at three full-scale RBF sites were consistently decreased from 3–10 mg/L to 1–3.5 mg/L.

Water utilities considering RBF should be aware that the oxygen demand created by the biodegradable NOM and other contaminants such as ammonia can change redox conditions and cause the dissolution of manganese, which may require treatment (Appelo and Postma, 1996). More information on manganese in drinking water is available elsewhere (Health Canada, 2019d).

Engineered biological filtration

Engineered biological filtration involves the use of granular media filters (i.e., anthracite/sand or GAC) without the maintenance of a disinfectant residual across the bed. Biological activity within the filters can be influenced by a number of factors: water quality, temperature, oxidant dose and type, and backwashing procedures (Huck et al., 2001). The process is typically preceded by an oxidation step (e.g., ozonation) that transforms NOM into BOM to

make it more readily biodegradable (Evans et al., 2013a); when biologically active carbon (BAC) filters are used after ozonation, the process is referred to as ozone-BAC. If a biological treatment step is not used after ozonation, increased biofilm growth in the distribution system is highly likely (Juhna and Melin, 2006).

Emelko et al. (2006) reported TOC removals of 13–23% and BOM removals of 72–93% (measured as oxalate) for a full-scale plant at warm (21–25°C) and cold (1–3°C) temperatures. Evans et al. (2013b) reported average AOC removals of 31–42% for 14 full-scale biological filters over a one-year timeframe. Stoddart and Gagnon (2015) reported a decrease in THMs and HAAs of 10–20 µg/L and 6–10 µg/L, respectively, following the conversion of anthracite-sand filters to biological filters in a full-scale direct filtration plant.

Water utilities considering biological filtration for an existing facility should be aware that some utilities have reported unwanted algae or biogrowth, shorter filter run times and problems maintaining a chlorine residual (Brown et al., 2016). There is extensive guidance available to help water utilities understand the mechanisms associated with biological filtration, as well as to identify and implement appropriate monitoring (Prévost et al., 2005; Juhna and Melin, 2006; Evans et al., 2013a, 2013b; Brown et al., 2016; Nyfennegger et al., 2016).

Slow sand filtration

Slow sand filtration (SSF) generally consists of untreated water flowing by gravity at a slow rate through a bed of submerged porous sand. During operation, biological growth occurs within the sand bed and gravel support. In addition, bacteria and other materials in the source water accumulate on the surface to form a “schmutzdecke”, the layer of solids and biological growth that forms on top of a slow sand filter. The biological growth within the filter and the schmutzdecke both contribute to the effectiveness of SSF. Depending on the source water quality, it may take weeks or months for this biological growth to develop (Bellamy et al., 1985a, 1985b; Logsdon et al., 2002).

Amy et al. (2006) reported that conventional SSF can decrease BDOC and AOC concentrations by <80% and <65%, respectively, whereas DOC and THM precursor removal was limited to between <15–30% and <20–35%, respectively. This level of removal, however, is generally not sufficient to comply with DBP drinking water guidelines (Pyper, 1985; Collins et al., 1991; Graham, 1999), particularly in winter when low temperatures reduce biological activity (Collins et al., 1992). The addition of ozone or GAC has been reported to achieve colour reduction and improve DBP precursor removal (Graham, 1999; Di Bernardo and Pereira Tangerino, 2006; Ødegaard et al., 2006; Steele et al., 2006).

In a review of the literature, Graham (1999) reported that with pre-ozonation, DOC and THM formation potential removals ranged from 18% to 55% and from 20% to 64%, respectively. Overall, the author suggested that the addition of ozone increased DOC removal by 10% while THM formation potential was halved. DiBernardo and Pereira Tangerino (2006) used bench-scale experiments and observed that colour removal increased from 33% to 63% when GAC was added (after SSF) and ranged from 21.5% to 53% when the water was oxidized (before SSF) with ozone or ozone/hydrogen peroxide (H₂O₂). When both GAC and oxidation with ozone or ozone/H₂O₂ were used in combination with SSF, colour removal ranged from 44% to 68%.

The enhancement of SSF with ozone and/or GAC can create a number of operational issues. Ozone can increase filter headloss and thereby shorten filter runs (Graham, 1999; Logsdon et al., 2002; Di Bernardo and Pereira Tangerino, 2006). The ozone residual should also be quenched before it reaches the schmutzdecke; otherwise the biomass becomes inactive, and biologically unstable water will be produced (Melin et al., 2006; Ødegaard et al., 2006). The

filtered effluent may also contain high concentrations of heterotrophic bacteria that should be removed/inactivated (Ødegaard et al., 2006). Steele et al. (2006) cautioned that the dissolved oxygen demand associated with the inclusion of a GAC layer must be considered. In addition, water temperature is an important design factor when considering SSF and the selection of any ancillary processes (Jabur, 2006).

Although Gottinger et al. (2011) conclude that enhanced SSF can provide significant reductions in colour and organic matter, pilot testing is recommended to ensure that the source water can be successfully treated (Logsdon et al., 2002). It should also be noted that the ozonation of water containing naturally occurring bromide can result in the formation of bromate. Water utilities using ozone should characterize their source water to assess water quality parameters (i.e., bromide, temperature, pH, alkalinity, NOM, ammonia) and how these change on a seasonal basis. Quarterly monitoring of raw water bromide is recommended to characterize the source water and allow correlation to bromate (and brominated DBPs). More information on bromate can be obtained from Health Canada (2018).

B.6.2.6 Oxidation processes

Oxidation processes include ozone, chlorine dioxide and advanced oxidation processes such as ozone/UV, ozone/H₂O₂, UV/H₂O₂, and Fenton's reaction. Under typical water treatment conditions, oxidation processes transform the nature of the organics rather than remove bulk NOM (Owen et al., 1993; Świetlik et al., 2004). As a result, oxidation processes are generally used for disinfection, taste and odour control and degradation of target organic contaminants. Ozone and chlorine dioxide tend to make NOM less reactive with chlorine, which generally results in decreases in THMs and tri-HAAs; however, some DBPs may increase, such as halonitromethanes and haloketones (Reckhow, 2017). Advanced oxidation processes can, in principle, remove a variety of NOM, but they can also increase the formation of DBPs and dichloroacetic acid (DCAA) in particular (Bond et al., 2011). In a few studies reviewed by the authors, differences between DCAA and trichloroacetic acid (TCAA) formation were reported (see Table 13). In cases denoted by negative values, an increase in DBPs occurred as a result of treatment. The authors recommended careful assessment of oxidation processes when they are used for DBP control. As oxidative processes can result in the reduction of some DBPs while increasing others, mitigative measures tend to focus first on minimizing DBP formation by maximizing NOM removal (AWWA, 2011a). The use of alternative disinfectants to reduce DBP formation should therefore be considered with caution (Reid Crowther & Partners, 2000).

Water utilities should be aware that all oxidants, including chlorine, produce biodegradable products upon reaction with NOM (see section B.4.1.3). As a result, biologically active filtration may be necessary to stabilize treated water (see section B.6.2.5). Water utilities should also be aware that all oxidants reduce UV absorbance, which affects SUVA without an associated reduction in NOM concentration. Thus, it is important to select appropriate sampling sites when measuring UV absorbance to calculate SUVA.

B.6.3 Distribution system

The biodegradable portion of NOM (i.e., BOM) impacts distribution system water quality by providing a source of nutrients that contributes to bacterial regrowth and biofilm development. Biofilms can provide a habitat for the survival of pathogens of fecal origin that may have passed through drinking water treatment barriers. OPPPs such as *Legionella* and non-tuberculous mycobacteria (e.g., *M. avium*, *M. intracellulare*) are also commonly found in biofilms of piped drinking water supplies (Fricker, 2003; Falkinham, 2015). The potential for the multiplication of

OPPPs in distribution system and plumbing system biofilms is of increasing concern to the water industry. In the United States, the most frequently reported cause of outbreaks associated with drinking water is *Legionella* associated with building plumbing systems (largely in hospitals or health care facilities that fall outside the jurisdiction of water utilities) (Beer et al., 2015).

The impact of organic carbon levels on the growth and survival of OPPPs after drinking water treatment has been investigated. Falkinham et al. (2001) observed higher mycobacterial numbers in distribution system samples than in those collected immediately downstream from treatment facilities, and the increase was correlated with AOC and BDOC levels ($r^2 = 0.65$). *M. avium* and *M. intracellulare* were not detected in any water samples collected immediately after treatment; however, they could be recovered in the distribution system and in biofilm samples from water meters on these same systems (Falkinham et al., 2001).

Studies on the effects of organic carbon on OPPP numbers in drinking water distribution systems in the absence of a disinfectant residual have also been conducted in order to provide specific information on the impact of nutrient levels on their growth in biofilms. Norton et al. (2004) reported that *M. avium* could be recovered from biofilms at nutrient levels as low as 50 µg/L AOC in model distribution systems where no disinfection was applied. Van der Wielen and van der Kooij (2013) observed that gene copies of *L. pneumophila* were sporadically found in unchlorinated distributed water from surface water and groundwater treatment plants with AOC levels above 10 µg/L and were not observed in systems with AOC levels below 5 µg/L. Wullings et al. (2011) observed that *L. pneumophila* DNA was detected more frequently in biofilm samples in a distribution system fed with drinking water with a high NOM concentration (8 ppm carbon) than in biofilm samples from a distribution system fed with drinking water having a low NOM concentration (<0.5 ppm carbon).

Collectively, these studies highlight the importance of organic carbon removal and the maintenance of an effective disinfectant residual in order to minimize biofilm development in the distribution system and premise plumbing. Guidance material to assist water utilities to develop control programs for treated drinking water is available elsewhere (LeChevallier and Au, 2004). In general, the most important elements for controlling the growth of bacteria in distribution systems are maintenance of a disinfectant residual, limitation of BOM and corrosion control.

Water utilities should be aware that, when applied as secondary disinfectants, free chlorine and chloramines possess different capabilities in terms of disinfectant power, reactivity with organic and inorganic material, biofilm penetration, potential for DBP formation and potential for nitrification. There is extensive guidance available in other publications to assist water utilities in selecting chemical disinfectants (LeChevallier and Au, 2004; AWWA, 2011a; Health Canada, 2013).

A well-maintained distribution system is a critical component of the source-to-tap or water safety plan approach to provide safe drinking water (Fisher et al., 2000). Distribution system optimization is a complex process involving numerous concomitant goals (e.g., microbial, DBPs, corrosion, physical integrity). Distribution system water quality should be regularly monitored, including indicators of biological stability (see section B.5.3). Operations/maintenance programs should be in place (e.g., water age control, watermain cleaning, cross-connection control, asset management) and strict hygiene should be practiced during watermain repairs to ensure drinking water is transported to the consumer with minimum loss of quality (Kirmeyer et al., 2001, 2014).

B.7 Monitoring and treated water quality targets

Water system owners should collect water quality information to optimize their water treatment processes, meet regulatory requirements related to DBPs, lead and copper, as well as minimize biofilm formation. Site-specific conditions and treatment objectives influence monitoring requirements, including, but not limited to parameter selection, analysis method and frequency. The monitoring frequency is typically based on source variability and/or the critical nature of a treatment process. Highly variable water sources and critical processes should therefore be monitored on a more frequent basis.

Raw water monitoring should be conducted to characterize the source and better understand the conditions that lead to changes in the concentrations and/or character of NOM (e.g., precipitation/snowmelt events, algal blooms, drought, fire), and the factors that enhance the reactivity of NOM to form DBPs (e.g., reaction conditions, water age, inorganic compounds such as ammonia, bromide, iodide and sulphur). Ongoing operational monitoring and treatment optimization will help to ensure that water utilities adequately remove NOM to meet concomitant water quality goals related to microbial risks, DBPs, biological stability and corrosion control. Table 1 (see section A.2.3) suggests parameters and recommends sampling frequencies. Additional guidance is available elsewhere (Kornegay et al., 2000; WHO, 2014).

Once data is collected, it should be analyzed to assess the following:

- if, and how, source water quality is changing (e.g., true colour, UV absorbance/transmittance, DOC, SUVA);
- if a correlation exists between raw water DOC and other surrogates used to measure NOM concentration (e.g., true colour, UV absorbance, UV transmittance);
- how NOM is impacting water treatment processes (e.g., chemical usage and specific chemical dose/demand) and if control limits should be established;
- how treatment is impacting NOM (e.g., residual NOM concentration, change in SUVA, specific DBP yields, specific colour) and if control limits should be established;
- distribution system impacts (e.g., DBP concentrations, lead/copper concentrations);
- biological stability (e.g., variability in disinfectant residual, biofilm formation rate, changes in corrosion rates); and
- if a correlation exists between treated water NOM surrogates (e.g., DOC, true colour, UV absorbance, UV transmittance, COD) and distribution system water quality (e.g., DBPs, specific DBP yields, lead, copper, biological stability).

A continuous improvement process should be in place to ensure water treatment is optimized to achieve water quality goals and maximize public health protection for the full range of water quality conditions. Treated water quality targets are suggested in Table 2 (see section A.2.3) for the surrogate parameters most commonly used to provide an indication of NOM concentrations. Treated water quality targets will be source- and system-specific for the following reasons.

- Some sources can have a higher specific DBP yield (e.g., $\mu\text{g DBP/mg DOC}$) than other sources, as discussed in section B.4.1.2. This may be due to source-specific differences in NOM character (e.g., some NOM fractions form more DBPs than others) or the presence of inorganic compounds that increase DBP formation rates (e.g., ammonia, bromide, iodide and sulphur). Source-specific DBP yields can be determined as discussed in section B.5.2.3. Sources with higher specific DBP yields are considered more “reactive.”
- Some systems have extensive distribution systems. As noted in section B.4.1.2, a distribution system with a residence time of 7 days and a temperature of $>15^{\circ}\text{C}$ will

require a different level of NOM removal to meet DBP guidelines than one with a residence time of 3 days and a temperature of <15°C.

For more reactive sources and extensive distribution systems, water should be treated to more stringent requirements, as there is a greater potential for DBP formation. Less reactive sources have more flexibility with respect to upper control limits for most of the parameters listed in Table 2 (see section A.2.3), with the following exceptions.

- COD: because COD measures oxidizable organic matter, a highly reactive source with 2 mg/L TOC and a less reactive source with 4 mg/L TOC can have comparable COD concentrations (Stoddart and Gagnon, 2014; Dabrowska, 2016).
- DOC for biological stability: a DOC of less than 1.8 mg/L is suggested to minimize the biofilm formation rate and disinfectant variability regardless of source water quality or secondary disinfectant used for residual control (free chlorine or chloramine) (LeChevallier et al., 2015a, 2015b).

B.8 International considerations

NOM has a fundamental impact on drinking water treatment processes aimed at protecting public health. As a result, some jurisdictions have established regulatory requirements or voluntary targets to minimize its impacts on drinking water quality.

The U.S. EPA (1998) mandates a treatment technique for removal of TOC to reduce the formation of DBPs. It applies to surface water facilities using conventional or lime softening water treatment when the TOC in the source water exceeds 2 mg/L. Performance criteria for the treatment technique are based on the raw water TOC and alkalinity. Utilities with raw water sources containing NOM that is poorly removed by coagulation are permitted to conduct jar testing to determine alternative performance criteria for avoiding the use of excessive alum dosages that result in limited additional TOC removal. The rule requires monitoring of DBPs, disinfectant residuals, TOC and alkalinity. Facilities with alternative performance criteria must also monitor magnesium hardness removal, DOC, UV₂₅₄ and SUVA. A monitoring plan must be developed and implemented that includes monthly sampling for TOC in the raw water and filter effluent, as well as total THM (i.e., chloroform, bromoform, bromodichloromethane, chlorodibromomethane) and HAA5 monitoring that is representative of the entire distribution system. TOC removal is calculated as a running annual average computed quarterly from monthly samples.

The World Health Organization suggests optimized NOM removal as a means to minimize biofilm growth in the distribution system (WHO, 2011). Organic carbon is also suggested as an operational parameter in water safety plans to monitor control measures.

The European Union drinking water regulations include TOC as a general water quality indicator parameter for supplies $\geq 10,000$ m³/d (EU, 2014). The regulations specify “no abnormal change” as the parametric value. In some jurisdictions, oxidizability (measured as chemical oxygen demand) can be used in place of TOC. A parametric guideline value of 5 mg/L O₂ is specified (EU, 2014). French regulations specify guideline limits for treated water intended for human consumption for several chemical and organoleptic parameters, including TOC (i.e., 2 mg/L and no abnormal change) and oxidizability (i.e., 5 mg/L O₂) (Government of France, 2007).

The Dutch approach to safe drinking water includes measures to control or limit microbial activity in the distribution system in the absence of a disinfectant residual (Smeets et al., 2009). This requires the production of biologically stable drinking water with an AOC target of below 10 µg/L (van der Kooij, 2000; Smeets et al., 2009; Lautenschlager et al., 2013). Investment in

both advanced treatment and distribution system infrastructure is necessary to achieve the AOC target. Groundwater supplies typically use aeration and filtration with GAC in some cases to remove chemical contaminants, followed by UV disinfection to reduce the colony counts after GAC. Locations with high methane concentrations require more aeration, whereas locations with high ammonia concentrations use “dry rapid sand filtration” (e.g., the sand bed is not saturated) to allow more oxygen transfer to the water. Surface water and riverbank filtration systems have unique combinations of multiple treatment processes that may include coagulation–sedimentation, rapid sand filtration, GAC, dune filtration, softening, advanced oxidation or ozonation, membrane filtration (UF and/or RO) and slow sand filtration (Smeets, 2017).

In Australia, guidance has been developed to help water utilities understand and control the impact of NOM within the context of the Australian Drinking Water Guidelines Framework (Cooperative Research Centre for Water Quality and Treatment, 2005).

Part C. References, acronyms and tables

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C.2 Acronyms

ACU	apparent colour units
AOC	assimilable organic carbon
ATP	adenosine triphosphate
BDOC	biodegradable organic carbon
BOM	biodegradable organic matter
COD	chemical oxygen demand
CU	colour units
DAF	dissolved air flotation
DCAA	dichloroacetic acid
DBP	disinfection by-product
DOC	dissolved organic carbon
EPA	Environmental Protection Agency (United States)
EU	European Union
GAC	granular activated carbon
GLUMRB	Great Lakes – Upper Mississippi River Board
H ₂ O ₂	hydrogen peroxide
HAA	haloacetic acid
HAA5	haloacetic acid 5
LC-OCD	liquid chromatography-organic carbon detection
LC-OND	liquid chromatography-organic nitrogen detection
MDL	method detection limit
MF	microfiltration
MWCO	molecular weight and cutoff
N-DBPs	nitrogenous-DBPs
NF	nanofiltration
NOM	natural organic matter
NTU	nephelometric turbidity unit
OPPP	opportunistic premise plumbing pathogen
PAC	powdered activated carbon
peCOD	photoelectrochemical oxygen demand
POC	particulate organic carbon
RBF	riverbank filtration
RO	reverse osmosis
SSF	slow sand filtration
SUVA	specific UV absorbance
THM	trihalomethane
TCAA	trichloroacetic acid
TCU	true colour units
TOC	total organic carbon
UF	ultrafiltration
UV	ultraviolet
UV254	ultraviolet absorbance at 254 nm wavelength
WHO	World Health Organization

C.3 Tables

Table C-3.1: Factors influencing allochthonous natural organic matter concentration and character

Factor	Comment	Reference(s)
Percent wetlands in the watershed	Wetlands have a high DOC production rate and the DOC tends to be high in organic acidity. Even 1% wetlands cover can influence DOC concentrations and character.	Eckhardt and Moore, 1990; Grieve, 1994; Dalva and Moore, 1991; Cool et al., 2014
Soil composition	Soils with the highest organic content (i.e., humus) export more DOC to aquatic environments. Mineral soils, particularly those rich in iron, aluminum or clay, tend to adsorb DOC.	Dalva and Moore, 1991; Kalbitz et al., 2000; Aitkenhead-Peterson et al., 2003
Forest cover	Coniferous-dominated watersheds can produce approximately 50% more DOC than hardwood-dominated watersheds.	Cronan and Aiken, 1985; Dalva and Moore, 1991; Kalbitz et al., 2000, 2006
Retention time	In general, the longer the retention time the lower the DOC concentration, due to biogeochemical processes that degrade and adsorb DOC. The residual DOC tends to be recalcitrant (i.e., not easily biodegraded).	Aiken and Cotsaris, 1995; Hanson et al., 2011; Reckhow et al., 2007
Watershed hydrology	In the absence of wetlands, DOC concentrations tend to increase relative to streamflow. However, not all storms elicit the same response, due to variations in soil texture, antecedent soil moisture and precipitation/ snowmelt conditions.	Eckhardt and Moore, 1990; Grieve, 1994; Soulsby, 1995; Carpenter et al., 2013
Flow pathways	Water passes through different soil layers depending on soil texture and antecedent moisture conditions. Variations in flow pathways can result in five-fold increases in DOC in short periods of time (i.e., hours to days).	Thurman, 1985; Aitkenhead-Peterson et al., 2003; Saraceno et al., 2009; Pellerin et al., 2012
Channel slope	Mildly sloped watersheds tend to have higher DOC concentrations than steeply sloped watersheds.	Eckhardt and Moore, 1990; Aitkenhead-Peterson et al., 2003; Cool et al., 2014
Watershed size	Small watersheds tend to have highly variable DOC concentrations, whereas large watersheds tend to be less variable. Watersheds with a high land-to-water ratio tend to have higher DOC concentrations.	Sinsabaugh and Findlay, 2003; Eikebrokk et al., 2004; Ågren et al., 2007

Table C-3.2: Groundwater organic carbon concentrations reported in the literature

Reference	Study details	TOC/DOC concentration ^a (mg/L)
Robinson et al., 1967	Groundwater from four small towns in Illinois	TOC range = 1.5–7.8 mg/L
Leenheer et al., 1974	60 public water supply wells in 19 U.S. states	DOC range = <0.1–15.0 mg/L Median = 0.8 mg/L Average = 1.4 mg/L
Thorstenson et al., 1979	9 wells in the Fox Hills-Basal Hell Creek aquifer ^b	TOC range = 1.9–20 mg/L Median = 3.1 mg/L Average = 5.1 mg/L
Leenheer and Bagby, 1982	~40 production wells in Idaho	DOC range = 1.5–11 mg/L Median = 4.8 mg/L Average = 5.4 mg/L
Aravena et al., 1995	26 domestic or commercial wells in the Alliston aquifer (Ontario)	DOC range = 0.9–18.0 mg/L Median = 7.1 mg/L Average = 7.0 mg/L
Bradner et al., 2005	30 public supply wells in the Biscayne aquifer (Florida)	DOC range = 0.6–22 mg/L Median = 9.4 mg/L
Washington State Department of Health and University of Washington, 2017	18 public supply wells in Island and San Juan Counties (Washington State)	DOC range = 0.54–7.86 mg/L

^a Median and average concentrations were calculated when the complete dataset was published; otherwise the data is as reported by the author(s).

^b An extensive aquifer supplying domestic and municipal wells for communities in Montana, North Dakota and South Dakota that are not located on the Missouri River; the aquifer extends into Saskatchewan.

Table C-3.3: Dissolved organic carbon data from Environment Canada (2000–2015)

Region	River Basin	Number of Samples	Number of Detects ^a	10th Percentile (mg/L)	Median (mg/L)	Mean (mg/L)	90th Percentile (mg/L)
East	Maritime Coast	94	92	1.00	2.25	3.43	8.65
	Newfoundland–Labrador	1,111	1,111	2.60	4.70	5.02	7.60
	North Shore–Gaspé	42	42	5.01	6.25	6.34	7.99
	Saint John–St. Croix	89	89	3.48	4.40	5.79	9.92
Central	Winnipeg	136	136	9.10	9.75	9.81	10.55
Prairie	Assiniboine–Red	1,153	1,153	8.14	10.90	12.46	18.80
	Churchill	292	292	6.64	11.80	11.12	16.60
	Lower Saskatchewan–Nelson	507	507	4.77	14.00	12.89	20.80
	Missouri	188	188	1.62	4.02	4.56	8.33
	North Saskatchewan	594	594	0.24	2.01	4.98	14.40
	South Saskatchewan	818	818	0.49	1.24	2.15	4.79
Pacific	Columbia	4,308	3,175	0.73	1.34	1.77	3.00
	Fraser	3,503	3,374	1.20	2.95	3.58	6.70
	Okanagan–Similkameen	1,118	1,079	1.10	2.86	3.15	5.59
	Pacific Coastal	2,510	2,217	0.90	2.00	2.67	4.70
	Peace–Athabasca	443	442	0.34	2.00	2.34	4.89
Arctic	Arctic Coast	148	136	0.60	2.45	3.09	6.50
	Keewatin–Southern Baffin Island	40	40	1.98	3.45	3.26	4.20
	Lower Mackenzie	697	690	1.10	3.60	6.26	16.44
	Yukon	619	518	0.70	1.90	3.51	9.23

^a Method detection limit = 0.5 mg/L

Table C-3.4: Spatial and temporal variation in NOM fractions for select Canadian water sources

Study/ Source	Sample Date	DOC (mg/L)	Hydrophobic (%)			Hydrophilic (%)		
			Acid	Base	Neutral	Acid	Base	Neutral
Montreuil, 2011								
Pockwock Lake, Nova Scotia	Jun 2009	2.6	8.3	17.6	20.5	0.0	19.2	34.5
	Aug 2009	2.5	3.6	0.4	36.8	6.8	6.4	46.0
	Oct 2009	2.9	0.0	1.3	31.7	19.1	35.5	12.4
	Dec 2009	2.8	19.1	29.6	7.6	2.5	8.3	32.9
	Jan 2010	2.9	16.1	2.1	22.4	3.5	11.2	44.8
	Jul 2010	2.5	14.0	1.9	3.2	7.0	2.5	71.3
	Aug 2010	2.1	28.0	4.9	0.9	12.0	0.9	53.3
	Range (Mean)	2.1-2.9 (2.6)	0.0-28.0 (12.7)	0.4-29.6 (8.3)	0.9-36.8 (17.6)	0.0-19.1 (7.3)	0.9-35.5 (12.0)	12.4-71.3 (42.2)
Newfoundland and Labrador Department of Environment and Conservation, 2011								
Community A	Feb 2011	3.6	29.7	2.9	2.9	8.8	1.8	53.8
Community B	Sep 2010	7.7	22.0	1.1	3.4	50.7	1.6	21.2
Community C	Sep 2010	10.8	62.4	0.9	0.0	5.5	0.2	31.0
Community D	Jan 2011	5.2	60.5	1.6	2.0	9.2	2.0	24.9
Community E	Sep 2010	8.3	57.7	1.0	3.0	3.5	1.0	33.9
Community F	Dec 2010	9.0	63.5	6.8	2.1	11.6	0.8	15.2
	Range (Mean)	3.6-10.8 (7.4)	22.0-63.5 (49.3)	0.9-6.8 (2.4)	0.0-3.4 (2.2)	3.5-50.7 (14.9)	0.2-2.0 (1.2)	15.2-53.8 (30.0)
Lamsal et al., 2012								
French River, Nova Scotia	Not given	5.3	35.3	2.2	4.5	6.1	1.5	50.4
Goss and Gorczyca, 2013								
Red River, Manitoba	Sep 2010	11.3	21.8	1.9	21.8	12.9	1.9	39.7
	Nov 2010	12.0	36.0	3.6	18.2	11.6	5.6	25.1
	Feb 2011	8.0	37.1	7.7	13.2	17.1	7.0	17.9
	Jun 2011	8.7	27.1	2.4	11.5	2.3	5.3	51.3
	Range (Mean)	8.0-12.0 (10.0)	21.8-37.1 (30.5)	1.9-7.7 (3.9)	11.5-21.8 (16.2)	2.3-17.1 (11.0)	1.9-7.0 (4.9)	17.9-51.3 (33.5)

Table C-3.5: Conversion chart for UV transmittance and UV absorbance

Conversion formula: $UV\ absorbance\ (cm^{-1}) = 2 - \log_{10}\ UV\ transmittance\ (\%)$

UVT (%)	UVA (cm ⁻¹)						
1	2.0000	26	0.5850	51	0.2924	76	0.1192
2	1.6990	27	0.5686	52	0.2840	77	0.1135
3	1.5229	28	0.5528	53	0.2757	78	0.1079
4	1.3979	29	0.5376	54	0.2676	79	0.1024
5	1.3010	30	0.5229	55	0.2596	80	0.0969
6	1.2218	31	0.5086	56	0.2518	81	0.0915
7	1.1549	32	0.4949	57	0.2441	82	0.0862
8	1.0969	33	0.4815	58	0.2366	83	0.0809
9	1.0458	34	0.4685	59	0.2291	84	0.0757
10	1.0000	35	0.4559	60	0.2218	85	0.0706
11	0.9586	36	0.4437	61	0.2147	86	0.0655
12	0.9208	37	0.4318	62	0.2076	87	0.0605
13	0.8861	38	0.4202	63	0.2007	88	0.0555
14	0.8539	39	0.4089	64	0.1938	89	0.0506
15	0.8239	40	0.3979	65	0.1871	90	0.0458
16	0.7959	41	0.3872	66	0.1805	91	0.0410
17	0.7696	42	0.3768	67	0.1739	92	0.0362
18	0.7447	43	0.3665	68	0.1675	93	0.0315
19	0.7212	44	0.3565	69	0.1612	94	0.0269
20	0.6990	45	0.3468	70	0.1549	95	0.0223
21	0.6778	46	0.3372	71	0.1487	96	0.0177
22	0.6576	47	0.3279	72	0.1427	97	0.0132
23	0.6383	48	0.3188	73	0.1367	98	0.0088
24	0.6198	49	0.3098	74	0.1308	99	0.0044
25	0.6021	50	0.3010	75	0.1249	100	0.0000