

Measurement of pollution levels of N-nitroso compounds of health concern in water using ultra-performance liquid chromatography–tandem mass spectrometry

Y. Kadmi, L. Favier, A.I. Simion, L. Rusu, M.L. Pacala, D. Wolbert

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1 **Highlights:**

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3 - A UHPLC/MS/MS method for N-nitrosamines quantification in water samples was
4 developed.

5 - A solid phase extraction procedure was proposed for analytes extraction.

6

7 - The proposed extraction procedure allows the extraction and purification of surface water
8 samples.

9

10 - Proposed method successfully applied to the detection of pollution in real water samples.

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12

1 **MEASUREMENT OF POLLUTION LEVELS OF N-NITROSO**
2 **COMPOUNDS OF HEALTH CONCERN IN WATER USING ULTRA-**
3 **PERFORMANCE LIQUID CHROMATOGRAPHY-TANDEM**
4 **MASS SPECTROMETRY**

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7 **Yassine Kadmi^{a,b*}, Lidia Favier^{a,b*}, Andrei Ionut Simion^c, Lacramioara Rusu^c,**
8 **Mariana Liliana Pacala^d, Dominique Wolbert^{a,b}**

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11
12 ^a*Ecole Nationale Supérieure de Chimie de Rennes, CNRS, UMR 6226,*
13 ⁱ*11 Allée de Beaulieu, CS 50837, 35708 Rennes Cedex 7, France*

14 ^b*Université Européenne de Bretagne, France*

15 ^c*“Vasile Alecsandri” University of Bacau, Faculty of Engineering, Department of Chemical and Food*
16 ^e*Engineering, 157 Calea Marasesti, 600115 Bacau, Romania*

17 ^d*“Lucian Blaga” University of Sibiu, Faculty of Agricultural Science, Food Industry and Environmental*
18 ^l*Protection, 7-9 Dr. I. Ratiu St., 550012 Sibiu, Romania*

19
20 **lidia.favier@ensc-rennes.fr ; yassine.kadmi@gmail.com*

21
22
23
24
25
26
27 *Corresponding Authors:

28
29 Dr. Lidia Favier, Associate Professor

30 Ecole Nationale Supérieure de Chimie de Rennes, CNRS, UMR 6226, Université Européenne
31 de Bretagne, 11 Allée de Beaulieu CS 50837, 35708 Rennes Cedex 7, France.

32 Tel.: +33 233238135; Fax: +33 223238120

33 E-mail: lidia.favier@ensc-rennes.fr

34
35 Dr. Yassine Kadmi

36
37 Ecole Nationale Supérieure de Chimie de Rennes, CNRS, UMR 6226, Université Européenne
38 de Bretagne, 11 Allée de Beaulieu CS 50837, 35708 Rennes Cedex 7, France.

39 E-mail: yassine.kadmi@gmail.com

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Abstract

This paper reports the development of a highly sensitive analytical method combining solid-phase extraction (SPE) with ultra-high performance liquid chromatography coupled with tandem mass spectrometry (UHPLC-MS/MS), for the monitoring of ultra-trace levels of N-nitrosamines in water samples.

Under optimized analytical conditions, chromatographic separation was performed in three minutes, in isocratic mode, using an Acquity UHPLC C18 column and a mobile phase consisting of acetonitrile, water, and formic acid (60:40:0.1, v/v/v) at a flow rate of 0.4 mL min⁻¹. Electrospray ionization tandem interface was employed prior to mass spectrometric detection. Good linearity ($R^2 \geq 0.9987$) and low limits of detection (0.04 to 0.4 ng L⁻¹) and quantification (0.1-1.2 ng L⁻¹) were obtained. The extraction recoveries ranged from 98 ± 1% to 100 ± 1% and the relative standard deviations were less than 1.53%. The matrix effect was between 98 ± 2 and 100 ± 1%.

The obtained results clearly demonstrate that the developed method is accurate and highly sensitive for the simultaneous determination of N-nitroso-n-propylamine, N-nitrosomorpholine, N-nitrosomethylethylamine and N-nitrosodimethylamine at ultra-trace levels (ng L⁻¹) in different types of water samples. Therefore, this method can be a useful analytical tool for future toxicological, water quality surveillance studies and for the investigation of drinking water quality.

Keywords: water quality; N-nitrosamines; solid phase extraction; ultra-high liquid chromatography-tandem mass spectrometry

1 1. Introduction

2
3 The occurrence of N-nitrosamines (NAms), in water is considered as an emerging
4 issue, due to their mutagenic and carcinogenic effects at ultra trace levels (ng L^{-1}). As a
5 consequence, in recent years, this class of molecules has received an increased interest from
6 environmental and analytical chemists. They are generally produced during different
7 industrial processes such as cosmetics, metal casting, leather tanning, food (beverages and
8 meats) or during the rubber or dyes manufacturing. Thus, these kinds of applications
9 commonly lead to wastewater and groundwater contamination. For example, high
10 concentrations (2 mg L^{-1}) of N-nitrosodimethylamine (NDMA) were found in Ontario, in
11 downstream industrial water of a tyre factory (Mitch et al., 2003). More recently, another
12 study carried out in Switzerland reported the presence of N-nitrosamines in the influents of an
13 urban wastewater treatment plant at a concentration level ranging from 5 to 25 ng L^{-1} (Krauss
14 et al., 2009; Sedlak et al., 2005).

15 Moreover, it was found that drinking water disinfection processes with chlorine,
16 monochloramine, chlorine dioxide, and ozone generally lead to the formation of these
17 compounds in treated water, by the reaction between disinfectant and the nitrogen containing
18 organic matter (Andrzejewski et al., 2008; Kadmi et al., 2015a, 2015b; Schreiber and Mitch,
19 2005).

20 Several studies reported the occurrence of NAms in treated drinking water from
21 different sites situated in North America and Canada. Other studies confirmed their presence
22 in drinking water (Richardson, 2009; Zhao et al., 2006) and of NDMA, NMOR in surface
23 water (Kosaka et al., 2010; Zhao et al., 2008).

24 N-nitrosamines are a class of non-halogenated emerging disinfection by-products
25 (DBPs) which have been recently identified in drinking water. NAms are alkylating agents
26 characterized by the presence of the N-nitroso group and may be aliphatic or ring structures.
27 Different studies reported that these molecules are significantly more toxic than the regulated
28 DBPs (Oya et al., 2008).

29 In response to their suspected adverse risks on human health, different guidelines have
30 been implemented in United States and Canada for these molecules and more particularly, for
31 NDMA, which is one of the most detected. However, in the European Community these
32 molecules are not yet regulated. Indeed, NAms are not listed in the Drinking Water Directive
33 (Council Directive 98/93/EC), but a few European Union (EU) member states have regulated
34 their presence in drinking water. Provisional standard values were proposed only in

1 Netherlands and in Germany for NDMA and N-nitrosomorpholine (NMOR) (Kadmi et al.,
2 2014; Planas et al., 2008). In addition, the French government proposed recently the addition
3 of NDMA in the EU Directive for drinking water, and a guideline value of 100 ng L⁻¹,
4 according with the guideline of the World Health Organization (WHO, 2011).

5 Therefore, in the light of new regulations at European level and for water quality
6 monitoring purposes it is of great interest to develop fast, sensitive and environmentally
7 friendly analytical methods for the monitoring of trace and ultra-trace levels of N-
8 nitrosamines in water samples. The major challenge in the determination of these molecules is
9 to attain the high sensitivity required for the quantification of trace levels in environmental
10 samples.

11 Presently there are no standard analytical methods for the quantification of NAmS in
12 environmental samples in the range of nanogram NAmS per liter. Due to the low
13 concentration level of these compounds in the aquatic environment and in drinking water an
14 extraction and pre-concentration step of analytes is generally required.

15 N-nitrosamines are polar compounds with low molecular weights (< 200 g mol⁻¹),
16 usually water soluble and have low octanol/water (K_{ow}) partition coefficients. Consequently,
17 they are difficult to extract with organic solvents or to remove by adsorption.

18 Several selective analytical techniques have been reported for the quantification of
19 NAmS. The analytical strategies currently used mainly consists of two steps, i.e., analysis by
20 gas chromatography (GC) or liquid chromatography (LC) and an extraction/concentration
21 procedure for the concentration of analytes. The NAmS have been determined in water
22 samples by using GC coupled with different types of detectors, such as gas chromatography–
23 coupled with mass spectrometry (GC/MS) (Huang et al., 2013; Reyes-Contreras et al., 2012;
24 Ventana and Ruiz, 2006) and gas chromatography–tandem mass spectrometry (GC/MS/MS)
25 (Llop et al., 2012; McDonald et al., 2012). However, these methods are limited to the analysis
26 of volatile and thermally stable compounds. Other studies focused on the determination and
27 quantification of NAmS by liquid chromatography using a fluorescence detector (Cha et al.,
28 2006) and high pressure liquid chromatography-tandem mass spectrometry (HPLC/MS/MS)
29 (Cheng et al., 2011; Zhao et al., 2006).

30 The aim of the present work was to develop a rapid and robust analytical SPE-
31 UHPLC-MS/MS protocol for the simultaneous quantification of N-nitrosamines in water
32 samples. The developed analytical procedure has been selected in order to attain the
33 selectivity, sensitivity and sample throughput which is needed for the quantification of these
34 molecules in water samples. Based on occurrence and toxicity data, the N-nitrosamines

1 selected for this work were: N-nitroso-n-propylamine (NDPA), N-nitrosomorpholine
2 (NMOR), N-nitrosomethylethylamine (NMEA) and N-nitrosodimethylamine (NDMA).

3 The study of the performance of the developed method was carried out in terms of
4 method detection limits (MDL), method of quantification limits (MQL), linearity, extraction
5 recovery and matrix effect.

6 The developed analytical method was then applied to real water samples (surface
7 water and treated water samples from public water system) collected from different locations
8 in Brittany region (France) in order to measure the pollution levels. Since these molecules are
9 usually not considered in routine monitoring programs in Europe, and especially in France,
10 information about the contamination level with these emerging DBPs is very limited.

12 **2. Materials and methods**

14 **2.1. Chemicals, standards and preparation of stock solutions**

15
16 Individual standard solutions (2000 mg L⁻¹ in methanol) of N-nitrosamines were
17 purchased from LGC Standards (Wesel, Germany). The main physico-chemical
18 characteristics of the studied molecules as well as their toxicity are shown in Table 1.

20 **Table 1**

21
22 Acetonitrile (LC-MS grade) and formic acid (> 95%) were obtained from J.T. Baker
23 (Deventer, Netherlands). Methanol and GC-grade dichloromethane were purchased from
24 Fischer Scientific-Bioblock (Illkirch, France). Acetic acid (100%) was supplied by Acros
25 Organics (Noisy-le-Grand, France). All reagents used in this work were of the highest
26 analytical grade (suitable for trace analysis). The ultrapure water used for the preparation of
27 the samples was obtained from an Elga Option-Q DV-25 system (Antony, France). Nitrogen
28 for drying (99.99% purity) was from Air Liquid (France).

29 A standard mix stock solution of 100 mg L⁻¹ containing the target analytes (NDPA,
30 NMOR, NMEA and NDMA) in methanol was prepared and stored in a freezer at -20°C for up
31 to three months. Working solutions were freshly prepared prior to UHPLC/MS/MS analysis
32 with 60: 40 (v/v) acetonitrile/ultrapure water at the desired concentration by appropriate
33 dilution of the stock standard solution.

1 Cartridges used for off-line SPE analytes extraction were Sep-Pak Plus[®]AC-2 cartridges
2 (400 mg, 85 µm) purchased from (Waters, Guyancourt, France).

3 Safety precautions were taken when handling N-nitrosamines because of their
4 carcinogenic risks for humans and animals. The disposal of the resulted waste followed a
5 proper safety procedure.

6 7 **2.2. Samples collection**

8
9 Water samples were directly collected from different surface and drinking water from
10 different locations situated in Brittany region (France) in June-July and October-November
11 2014 and in June-July 2015. Sample sites can be showed in Fig. 1.

12 Water samples were collected in pre-cleaned amber glass bottles (4-L) with Teflon lined
13 caps to ensure sample integrity. Before sampling bottles were rinsed several times with the
14 same water samples in order to avoid internal contamination. In order to remove suspended
15 particles water samples were vacuum filtered through cellulose membrane (0.45µm; Sartorius,
16 Germany) and stored at 4°C under light protection until analysis (within one week of
17 collection). No additives were placed in the samples to prevent their contamination.

18 19 **Figure 1**

20 21 **2.3. Methods**

22 23 **2.3.1. Sample concentration and clean-up**

24
25 An SPE-off line method was performed and developed in order to extract, purify and
26 concentrate the water samples. A 12-port position Visiprep SPE vacuum manifold obtained
27 from Supelco (Bellefonte, PA, USA) was used. Several sorbents were evaluated for the
28 extraction of analytes. Sep-Pak Plus[®]AC-2 cartridges (400 mg, 85 µm; Waters, Guyancourt,
29 France) showed the best extraction recoveries for all considered molecules, even for
30 hydrophilic molecules such as NDMA and NMOR.

31 Briefly, cartridges were initially pre-conditioned with 8 mL of methanol followed by 8
32 mL of dichloromethane, 8 mL of acetonitrile and 8 mL of ultrapure water. A sample volume
33 of 250 mL was loaded with a light vacuum on the cartridge and a flow rate of
34 approximately 3 mL min⁻¹. The sorbent was then rinsed with 5 mL of ultrapure water

1 acidified at pH 2 with acetic acid. Analytes adsorbed on SPE cartridges were eluted
2 successively with 6 mL of dichloromethane, 4 mL of acetonitrile, and 2 mL of methanol at a
3 flow rate ranging from 2 to 3 mL min⁻¹. Cartridges were air-dried for few minutes to remove
4 the water drops. After the extraction step, the elution solvent was collected in conical
5 graduated glass tube Pyrex® (VWR, Fontenay-sous-Bois, France).

6 Eluates were then concentrated by evaporation under a high-purity and moderate
7 nitrogen flow in an N-Evap evaporation system (Organomation, Berlin, MA, USA) to a final
8 volume of 0.1 mL (concentration factor of 2.500). Extracts were reconstituted with
9 acetonitrile/ultrapure water (60:40, v/v), transferred to an injection vial and stored at 4°C until
10 further analysis.

11 12 2.3.2 Instrumentation

13
14 The target compounds were separated by a Waters Acquity UHPLC H-Class system,
15 containing a binary pump, an auto-sampler and a thermostated column compartment (Waters,
16 Saint-Quentin en Yvelines, France). Chromatographic separation of N-nitrosamines was
17 carried out on Ethylene Bridged Hybrid (BEH) C18 column (100 mm × 2.1 mm, 1.7 μm)
18 from Waters. The used column is packed with a C18 reverse phase bounded to an ethylene-
19 bridged hybrid (BEH) substrate. In the chromatographic system, column was protected by an
20 in-line filter unit purchased from Waters (Saint-Quentin en Yvelines, France). The analytical
21 column compartment was maintained at 45°C. The auto-sampler was conditioned at 5°C. For
22 the optimization of the chromatographic analysis and MS/MS characterization, standard
23 solutions of each N-nitrosamine in mobile phase were used.

24 Isocratic elution was carried out with LC-MS grade acetonitrile/ultrapure water mix
25 (60:40, v/v) with added 0.1% (v/v) formic acid. The flow rate was 0.4 mL min⁻¹, and the
26 injection volume was 5 μL.

27 Tandem mass spectrometry (MS/MS) determination was performed on a Quattro
28 Premier Triple Quadrupole Mass Spectrometer (Waters, France) equipped with an
29 electrospray ionization source (ESI). ESI experiments were designed in positive and negative
30 ionization mode (ESI) to determine the optimal MS/MS detection conditions. Quantitative
31 analysis was carried out in Multiple-Reaction Monitoring (MRM) mode. In addition, the cone
32 voltage and the collision energy were optimized in order to achieve the best sensitivity.

33 The optimal MS/MS conditions selected for the analysis of target compounds were:
34 capillary voltage 3 kV; cone voltage 40 V; source temperature 120°C and desolvation

1 temperature 350°C. The considered cone and desolvation gas flows were 75 and 750 L h⁻¹,
2 respectively. High purity argon (99.99% purity, Air Liquid, Paris, France) was employed as
3 collision gas at a flow rate of 0.12 mL min⁻¹. Dry nitrogen used as desolvation, nebulization
4 and cone gas was produced by a nitrogen generator (Peak Scientific, Inchinann, UK). The
5 argon pressure used in the collision cell was 3.52×10⁻³ mbar. The analytical system control
6 and data acquisition were processed using Masslynx software, version 4.1 (Waters, Saint-
7 Quentin en Yvelines, France).

9 **2.4. Quality parameters**

10
11 The ultra-high performance liquid chromatography tandem mass spectrometry method
12 was evaluated in terms of linearity, instrumental limits of detection (IDL) and quantification
13 (IQL), method detection (MDL) and quantification limits (MQL), precision; but also through
14 the extraction recovery on the whole extraction protocol and matrix effects. To evaluate the
15 practical applicability of the developed method, analytical quality parameters were
16 investigated using spiked water samples.

17 Linearity of the instrumental method was investigated for all analytes, from the
18 calibration curves, at seven calibration levels ranging from 0.1 to 100 µg L⁻¹. Standard
19 solutions were prepared by spiking calculated volumes of ultrapure water. The standard
20 calibration curves were generated by least squares linear regression. This method was used to
21 determine the slope, intercept, and correlation coefficient (R²) of the calibration equation. The
22 applied calibration model for all curves was $y = ax+b$ (weighting 1/x), where y was the peak
23 area, x was the concentration of the compound, a = the slope of the curve, and b = the
24 intercept.

25 The sensitivity of the developed instrumental method was determined in terms of
26 instrumental detection and quantification limits. IDL and IQL were defined as the
27 concentration which gave a signal-to noise ratio (S/N) above 3 and 10, respectively. The noise
28 was measured from six independent analyses.

29 The linearity of the analytical method was determined by passing the entire extraction
30 and clean-up method (SPE) on ultrapure and surface water samples spiked with the selected
31 compounds at concentrations ranging from 0.04 to 40 ng L⁻¹ considering a pre-concentration
32 factor of 2500. These samples were used to build the sample calibration curves (SCC).
33 Moreover, they are used to determine the MDL and MQL defined as the lowest concentration
34 which gave an S/N of 3 and an S/N above 10, respectively. These samples were also used to

1 calculate the repeatability (intra-day precision) of the method. It was determined for the
2 lowest level of concentration (0.4 ng L^{-1}) by analyzing the same spiked sample in six
3 replicates within a single day and results were expressed in terms of relative standard
4 deviation (RSD).

5 For the SPE extraction methodology of N-nitrosamines, SPE recoveries (R, %) were
6 determined quantitatively at different concentration levels. These recovery studies permitted
7 the evaluation of the efficiency of the proposed SPE method. They were investigated by
8 analyzing water samples at three concentration levels (0.4 , 4 and 40 ng L^{-1} , respectively)
9 spiked before and after extraction and clean-up procedure with appropriate amounts of the
10 mix N-nitrosamines standard solution. SPE recoveries were calculated as the ratio between
11 the resulting peak areas of the both extracted and non-extracted samples.

12 Matrix effect ($ME = C/D$) was determined for each analyte and sample as previously
13 described by Matuszewski et al. (2003). It was calculated as the ratio between responses (peak
14 areas) obtained in the presence of the matrix (C: samples spiked after extraction) to the
15 responses (peak areas) obtained in absence of the matrix (D: pure standard solution). This
16 method of calculation was used in many studies to evaluate the matrix effect in environmental
17 and biological samples. In this work, the matrix effect was evaluated by using real water
18 samples and was expressed as percentage. It will be noticed that, the used sample was first
19 checked to verify that no analyte was detected in the matrix.

20

21 **3. Results and discussions**

22

23 The aim of this study was the development of a rapid and sensitive method for the
24 simultaneous extraction and analysis of N-nitrosamines in water samples. In the light of the
25 lack of rapid, sensitive and robust methods for the analysis of these molecules in water
26 samples, the methodology proposed in this work has been focused especially on the
27 determination of these molecules at concentration levels relevant for environmental water
28 samples. As indicated above, the selection on the target molecules was mainly based on their
29 toxicity and occurrence in environmental waters.

30

31

32

33

3.1. Optimization of the MS/MS detection conditions

Mass spectrometer settings were first investigated in order to determine the optimal MS/MS detection conditions. Parameters of the mass spectrometer were obtained by direct infusion of a standard solution of each analyte (1 mg L^{-1}) into the source of the mass spectrometer. For analyte infusion a fixed flow rate of $10 \text{ }\mu\text{L min}^{-1}$ was used.

For the ionization of the target analytes, electrospray ionization (ESI) and atmospheric pressure ionization (APCI) modes with positive or negative ionization were investigated. Obtained data clearly showed that the optimal analytes responses were observed in positive ionization mode due to its high sensitivity (data not shown). Thus, a positive ionization mode (ESI^+) combined with multiple reaction monitoring (MRM) was considered in this study for further quantification purposes.

The influence of capillary voltage, source temperature and desolvation temperature were also studied and the optimum operating conditions are presented in Table 2. The effects of desolvation gas flow (rate $600\text{-}900 \text{ L h}^{-1}$) and cone gas flow rate ($0\text{-}100 \text{ L h}^{-1}$) were analyzed. No significant influence on the signal of precursor ion was observed. These results are similar with the ones previously reported by Mourao Rodrigues et al. (2006) for the analysis of pesticides. For further analytical purposes the considered values for desolvation gas flow and cone gas flow were 750 L h^{-1} and 75 L h^{-1} , respectively as recommended by the manufacturer. In addition, the mobile phase composition was evaluated because this parameter is crucial for the separation and detection of target molecules. It was found that formic acid increases the ionization of all considered analytes.

Table 2

The selection of MRM transitions and associated acquisition parameters (collision energy and cone voltage) were evaluated for best response under positive mode conditions (ESI^+) by direct infusion in the source of a standard solution of each compound (1 mg L^{-1}) into the mass spectrometer. For these tests the cone voltage in the mass spectrometer scan mode was varied from 10 to 35 eV and the collision energy from 9 to 20 eV.

In this study, two sensitive MRM transitions were considered for each N-nitrosamine according to the requirements regarding mass spectrometric confirmation defined by the EU Commission Decision 2002/657/EC. Indeed, two transitions have to be recorded for each analyte in order to have a sufficient number of identification points for a suitable

1 confirmation. Hence, in this work, the peak area of the most intense transition was used for
2 quantitative purposes and the less intense one for confirmation. The cone voltage and the
3 collision energy were also investigated. The optimized MS/MS transitions as well as specific
4 cone voltage, collision energy, are presented in Table 3.

5
6 **Table 3**

7
8 **3.2. Optimization of separation conditions**

9
10 Separation and ionization of analytes is generally affected by the composition of the
11 mobile phase. Thus, in this work, the influence of mobile phase composition and mobile
12 phase additives on the separation of the target molecules was studied. Different compositions
13 of the mobile phase (i.e., acetonitrile/water and methanol/water) modified with acetic acid, or
14 formic acid (0.05, 0.1, 0.2, and 0.3%) were investigated in order to obtain an efficient
15 separation of N-nitrosamines using the BEH C18 column. An important increase in the
16 measured signal intensity was observed for the four analytes using acetonitrile/water (60:40,
17 v/v), modified with formic acid. In addition, the measured responses were higher than those
18 obtained in both mobile phases containing acetic acid (data not shown). Indeed, the use of
19 formic acid improves the ionization efficiency. Obtained data showed that a very low
20 concentration of formic acid lead to a lack of protons while a high concentration conduct to a
21 ion suppression. Thus, both conditions would reduce the analytical sensitivity (data not
22 shown). Therefore, 0.1% of formic acid was chosen as additive for the mobile phase in this
23 work.

24 The influence of column temperature and flow rate was also investigated. Column
25 temperatures from 35°C to 50°C were tested, and 45°C was selected as the working
26 temperature. Flow rates from 0.2 to 0.5 mL min⁻¹ were assayed, and the obtained data
27 indicated that a flow rate of 0.4 mL min⁻¹ significantly improves the resolution, peak shape,
28 intensity of the response, and retention times. Under optimized analytical conditions all the
29 analytes were separated with high sensitivity and selectivity within a run time of three
30 minutes.

31
32 **3.3. Method performance**

1 The performance of the developed analytical method was investigated in terms of
2 recovery, linearity, limits of detection and quantification. The main evaluated quality
3 parameters were indicated in Table 4 and Table 5.

4 5 3.3.1. Recovery

6
7 The sensitivity of the analytical system is not sufficient to directly analyze the N-
8 nitrosamines in the range of concentration found in real water samples (ng L^{-1}). For this
9 reason, prior to their instrumental analysis an enrichment step is required for the analyte
10 extraction and pre-concentration but also to remove the interfering components from the
11 matrix. Current extraction methods for NDMA or other molecules of the class of N-
12 nitrosamines from aqueous samples were developed and they are typically based of liquid-
13 liquid extraction (LLE) or solid phase microextraction (SPME) (Perez et al., 2008; Huang et
14 al., 2010; Taguchi et al., 1994). However, the LLE generally requires large amounts of
15 organic solvents, potentially harmful for environment, and the analytical process is time
16 consuming and labor-intensive (Hung et al., 2010). Indeed the time required for analyte
17 extraction and quantification with such technique may range from 3 to 16 h.

18
19 To overcome the drawback of extraction methods based on LLE some studies focused
20 their attention on the extraction of these molecules by solid phase extraction because, this
21 technique generally needs shorter processing times and less volumes of organic solvents. SPE
22 methods use different types of sorbents for the extraction of specific N-nitrosamines from
23 aqueous samples (Charrois et al., 2004; Cheng et al., 2006). However, several studies in
24 literature reported extraction recoveries below 50% and noticed the improvement in SPE
25 procedure in order to attain lower quantification limits.

26 In the light of these considerations, a pre-concentration methodology based on solid
27 phase extraction was developed in this work. SPE experiments were carried out after the
28 optimization of UHPLC/MS/MS conditions. The optimization of the extraction process was
29 performed in order to attain excellent recoveries for all target molecules in a single extraction
30 step. The optimized conditions used in this work were previously described (Materials and
31 methods section).

32 The SPE extraction recoveries were determined by extracting ultrapure water samples
33 spiked with each target molecule at three quality control concentration levels (0.4 , 4 and 40 ng L^{-1} ,

1 respectively). Six different sets of extractions ($n = 6$) for each sample were carried out.
2 Extracted samples were then analyzed by using the developed UHPLC/MS/MS method.
3 Extraction recoveries were obtained by comparing peak areas of the analytes obtained from
4 water fortified before extraction to those fortified after extraction.

5 As shown in Table 4, high mean recoveries were obtained for the considered analytes
6 (between $98 \pm 1\%$ and $100 \pm 1\%$). Among the target molecules, at the spiked level of 40 ng L^{-1} ,
7 NMOR showed the highest recovery ($100 \pm 1\%$). This value is higher than the one reported by
8 Zhao et al. (2006) for NMOR in spiked water samples for the same level of concentration. For
9 their SPE methodology a mean recovery of 65% was obtained. For NDMA, MNEA and
10 NDPA low extraction recoveries were reported (between 75 and 82%).

11

12

Table 4

13

14 For NMOR, recoveries in the same range of magnitude were obtained by Jurado-Sanchez et
15 al. (2009) using LiChronut and Oasis HLB cartridges.

16

17

18 In this study for NDMA, the calculated recoveries were higher than 98% (Table 4).
19 Similar recoveries were obtained by other authors (Charois et al., 2004; Plumlee et al., 2008).
20 Moreover, the values of the calculated relative standard deviations (RSD) were below 1.53%
21 for all analytes and all control concentrations tested (Table 4).

22

23 3.3.2. Linearity

24

25 For N-nitrosamines studied in this work the calibration curves were linear over the
26 considered concentration range. It will be noticed that, for linearity studies all samples were
27 analyzed in triplicate. For instrumental calibration curves the tests were carried out without
28 organic interfering species (ultrapure water). They were determined by using serial dilutions
29 of standard solutions containing the selected analytes and were established by plotting peak
30 areas against the analyte concentration. Obtained data showed good correlation mean
31 coefficients for all N-nitrosamines. The calculated values are between 0.9987 and 0.9992
32 (Table 5). The lower correlation coefficient was obtained for NDMA and the higher for
33 NDPA.

34

Table 5

1 As previously stated, sample calibration curves were also studied. They were determined from
2 the sample analysis in a given matrix, surface water samples spiked with selected analytes at
3 the same concentrations with the ones considered for instrumental calibration curves. For
4 these assays blanks were periodically run to confirm the absence of any contamination. The
5 calculated mean correlation coefficients were lower (between 0.9962 and 0.9983) than the
6 ones obtained in spiked ultrapure water. All the obtained results for ICC and SCC were
7 considered as satisfactory. Indeed, the linearity was considered as satisfactory when the R^2
8 was > 0.99 .

10 3.3.3. Matrix effect

12 One of the major drawbacks of electrospray mass spectrometry is that the ionization
13 source is highly sensitive to co-extracted matrix components. The mechanism and the origin
14 of the matrix effect is not fully understood but it may originate from the competition between
15 an analyte and the co-eluting, undetected matrix components which reacts with primary ions
16 formed in the LC-MS/MS interface (Matuszewski et al., 2003; Kadmi et al., 2016).

17 Matrix effects can be highly variable, difficult to control or predict, and analyte
18 specific (Chambers et al., 2007). Indeed, the presence of matrix components may lead to
19 suppression (decrease in analyte ionization) or enhancement of the analyte response (ion
20 enhancement) due to co-eluting matrix constituents affecting the detection capability,
21 repeatability and accuracy of the assay (Bijlsma et al., 2009). Hence, such phenomenon
22 severely affects the quantification of the analyte by electrospray mass spectrometry (Antignac
23 et al., 2005; Caban et al., 2012). Therefore, the elimination of such effect is critical in the
24 development of reliable analytical methods. Ignoring such phenomena may adversely affect
25 the reliability of determination of analyte concentrations.

26 As previously stated, if unseen, undetected, endogenous compounds present in real
27 water samples co-elute with the target molecules, they may affect the ionization efficiency of
28 analytes leading to the increase or decrease in their MS response. In most cases it is
29 impossible to completely eliminate the matrix effect.

30 Several action levels are proposed in literature to minimize their consequences in the
31 final determination allowing obtaining accurate and reliable LC/MS/MS data. They include
32 optimization of sample preparation to remove interfering compounds, changing
33 chromatographic parameters to avoid the co-elution phenomenon, and changing MS
34 conditions to reduce the occurrence of the matrix effect in the ion source (Chambers et al.,

1 2007; Van Eckhaut et al., 2009). However, the most direct means to obtain maximum
2 sensitivity and signal reproducibility is through reduction of matrix components prior to the
3 LC-MS/MS analysis by applying a selective extraction and improved sample clean-up
4 methodology (Gomez et al., 2006). Such approach will limit the presence of interfering
5 compounds in the final extract and will definitively overcome the problem of ion suppression.
6 Numerous authors demonstrated the evidence of this approach. Other studies indicated that it
7 is not a universal strategy but only solutions case by case for each analyte/matrix combination
8 (Antignac et al., 2005).

9 Thus, in this work the matrix effect of the optimized SPE-UHPLC/MS/MS analytical
10 method was also investigated. It was evaluated by performing matrix effect experiments with
11 different river water samples (collected in summer or in winter 2014) spiked with the target
12 molecules at 0.4 ng L^{-1} . They were performed in six replicates ($n=6$) for each analyte to assess
13 the variability of instrumental response. Chromatographic peak areas (responses) of each
14 analyte from the spiked after extraction samples were compared to those obtained from the
15 standard solution at the same level of concentration (0.4 ng L^{-1}) and the matrix effect was
16 calculated as described above (Section 2.4). The obtained data in spiked river water samples
17 collected in summer are shown in Table 5. For the selected analytes the determined matrix
18 effects and the corresponding relative standard deviations were between 98 ± 2 and $100 \pm 1\%$.
19 These data demonstrated the absence of a detectable matrix effect for the considered analytes
20 determined in ground water extracts. The obtained data were very similar with the ones
21 measured in ultrapure water samples spiked for the same level of concentration. They are
22 consistent to the ones obtained using spiked surface water samples collected in winter, for
23 which, higher concentration in natural organic matter are expected (data not shown). In
24 addition, it will be noticed that preliminary tests were carried out with river water samples
25 spiked with the target molecules and their corresponding blank samples and the obtained data
26 showed that the blank samples were free from interfering compounds (data not shown).

27 The satisfactory results (ME close to 100%) obtained in different aqueous matrices
28 clearly indicate no significant effects from the matrix composition of the environmental water
29 samples. The sample pretreatment procedure (clean-up and pre-concentration) developed in
30 this work efficiently reduced the amount of the co-extracted substances. The obtained results
31 indicated that the analytical procedure (SPE-UHPLC/MS/MS) developed in this study allows
32 to a reliable quantification of the target molecules in real water samples. They suggest that,
33 the matrix effect have a minimal influence on the results of the proposed method. These
34 results are quite similar to the ones reported previously, by Reyes-Contreras et al. (2012),

1 Kadmi et al. (2014) and Kadmi et al. (2015c). In their work, Hung et al. (2010) also reported
2 that the complicated natural water matrix did not affect the performance of the SPME-
3 GC/MS/MS method developed for the trace analysis of N-nitrosamines. At contrary, the
4 matrix effect appears as significant in the case of the analysis of NDMA in wastewater
5 samples because of the high organic matter concentrations. For this kind of matrix, it was
6 suggested to use an internal standard to eliminate analytical errors (instrumental and
7 extraction recovery) caused by the matrix effect for the quantification of NDMA at low
8 concentration levels (Topuz et al., 2012). However, the use of isotopically labeled internal
9 standards, is expensive and they are not always commercially available for the analysis of
10 molecules of interest.

11 For the analysis of drinking water samples, supplementary recovery tests may be
12 necessary to be conducted because, chlorinated water includes residual chlorine that could
13 affect the extraction recovery of analytes. Under these conditions, sodium thiosulfate must be
14 added to reduce residual chlorine in chlorinated water samples (Pepich et al., 2004). Indeed, it
15 is well-known that sodium thiosulphate acts as a reducing agent and it is suitable for the
16 reduction of residual chlorine in water, preventing the chlorine interference in the analysis and
17 extraction of analytes (Guerra Simões et al., 2007).

18 19 3.3.4. Limits of detection and quantification

20
21 As stated previously, the instrumental limits of detection and quantification (IDL and
22 IQL) of the proposed method were calculated based on a signal to noise ratio (S/N) of 3 and
23 10, respectively. The determined IDLs and IQLs ranged from $0.1 \mu\text{g L}^{-1}$ (for NMOR) to $1 \mu\text{g}$
24 L^{-1} (for NDPA) and from $0.25 \mu\text{g L}^{-1}$ (NMOR) to $3 \mu\text{g L}^{-1}$ (for NDPA), respectively. For
25 these parameters the lowest values were obtained for NDMA and NMOR (Table 6).

26
27 **Table 6**

28
29 The calculated MDLs of the selected N-nitrosamines for the SPE-UHPLC/MS/MS
30 method were in the 0.1 to 0.4 ng L^{-1} range (except 0.04 ng L^{-1} for NMOR).
31 It should be pointed out that, the excellent method detection and quantification limits obtained
32 in this study make possible the analysis of N-nitrosamines at ultra-trace levels. Similar results
33 were reported by Asami et al. (2009) for NDMA by using an SPE-UHPLC/MS/MS and by

1 Kadmi et al. (2014) for NMEA. The obtained results, are better than those reported by Grebel
2 et al. (2006) using a SPME-GC/NCD methodology which are in the range of 57-193 ng L⁻¹.
3 A low method detection limit was also reported using a GC/LRMS technique. Templeton and
4 Chen (2010) developed an SPE with Ambersorb 484 and GC/LRMS methodology for the
5 analysis of N-nitrosamines with a MDL ranging from 0.9 to 4.4 ng L⁻¹. Charois et al. (2004)
6 reported similar results by using GC/LRMS-PCI (ammonia positive chemical ionization) and
7 SPE with Ambersorb 572 and LiChrosorb EN as sorbents.

8 On the other hand, the California Department of Public Health set a notification level
9 for NDMA at 10 ng L⁻¹ for drinking water. The obtained MDL for this molecule using the
10 SPE-UHPLC/MS/MS method developed in this work is below this notification level. This
11 suggests that the proposed analytical methodology can be considered as an interesting tool for
12 the monitoring NDMA in drinking water samples. For NMOR and NDMA the results
13 revealed that the MDL are much lower than the provisional standard values proposed in
14 Netherlands and Germany (Planas et al., 2008). NMOR is considered to be less toxic
15 compared to NDMA. Consequently, until now, any notification has been set for this N-
16 nitrosamine in drinking water.

17 The overall results presented in this study demonstrate the analytical performance and
18 sensitivity of the developed method. Indeed, the proposed SPE-UHPLC/MS/MS analytical
19 strategy allow quantification limits in the ultra-trace range (ng L⁻¹) and an enrichment factor
20 of 2500 for all the target compounds (sample volume, 250 mL to 0.1 mL). The obtained MDL
21 and MQL fulfill with all N-nitrosamines guideline regulations (stringent or less stringent) and
22 with the provisional standard values considered in some European Countries.

23 Moreover, further research is needed to verify the applicability of the proposed
24 analytical strategy for the analysis of wastewater samples. For such samples the main
25 analytical limitation is related to the relatively high organic matter concentrations and as
26 consequence, to the matrix effect. Thus, in this case it is necessary to evaluate if there is
27 alteration in the extraction recoveries, elution profile or symmetry peak loss by matrix effect,
28 in order to prove if the developed method is appropriate for the analysis N-nitrosamines in
29 this kind of water samples. Furthermore, future research should focus to confirm the use of
30 the proposed analytical method for the detection of target molecules during the disinfection
31 water process and in swimming pool waters.

32
33

3.4. Application of the proposed method for the determination of pollution level in real water samples

The applicability of the developed analytical method was assessed through the analysis of several surface water and tap water samples collected from different rivers and locations, in Brittany region (France), during different periods (summer and winter 2014 and summer 2015).

Samples were all extracted and analyzed under the optimized analytical (SPE-UHPLC/MS/MS) conditions in 6 replicates ($n = 6$) according to the procedure described above. Quality standards were used as controls. It should be noticed that, for the sampling periods trip blanks are also carried out. The obtained data showed that the trip blanks are free of detectable target molecules.

The concentration of each N-nitrosamines determined in the collected samples are listed in Table 7.

Table 7

Data analysis showed that all target molecules were detected at least one time in collected water samples (surface and tap water). For all of them, the measured concentrations were in the range of nanogram per liter. As expected, the NDMA was most frequently detected. For the samples collected in summer the detected concentrations did not exceed 0.32 ng L^{-1} . In winter, the highest detected concentration was 0.67 ng L^{-1} . According to literature data, NDMA is the one of the N-nitrosamines which is most frequently detected in raw waters (Planas et al., 2008; Zhao et al., 2008). MNEA was detected in one river water sample (sample 5) collected in June-July and in October-November 2014, and the measured concentrations were 0.43 and 0.59 ng L^{-1} , respectively. However, in this river samples the other nitrosamines were not detected, suggesting that the nitrosamine contamination depends on the location and source of water. These concentrations are much lower than the ones reported by Kim et al. (2013) for NMEA, in Korea, in Nakdong river (in the range 6.2 to 17.7 ng L^{-1}).

NDPA was only found in two of the six analyzed samples collected in October-November 2014. However, in June-July 2014 this molecule was found only in sample 3. For NDPA, the measured concentrations were in the range of 0.35 to 0.81 ng L^{-1} . The study

1 conducted by Kim et al. (2013) reported also the presence of NDPA in Nakdong river (Korea)
2 at concentrations up to 455.4 ng L⁻¹.

3 NMOR was only found in sample 6 with a concentration of 0.17 ng L⁻¹. Templeton and
4 Chen (2010), Zhao et al. (2008) also detected this N-nitrosamine in raw water samples.

5 Among the all considered N-nitrosamines, in the analyzed tap water samples only,
6 NDMA was detected in one of the six samples, at a very low concentration (0.4 ng L⁻¹).
7 Although the molecule was found in tap water, the detected concentration is lower than the
8 California's notification level (10 ng L⁻¹) or the Ontario's acceptable limit (9 ng L⁻¹) (Kadmi
9 et al., 2015a). Li et al. (2015), in a survey study of distribution of N-nitrosamines in drinking
10 waters of east of China found NDMA with the highest detection frequency (31%). Zhao et al.
11 (2006), in a monitoring study reported for NDMA higher concentrations between 51.7 and
12 108.2 ng L⁻¹. More recently, Rusell et al. (2012), regarded this molecule as the most prevalent
13 N-nitrosamine in drinking water.

14 NMEA and NDPA were only found in one of the six analyzed tap water samples
15 (sample 5 and sample 3, respectively). The found values are quite similar to the one measured
16 for the NDMA.

17 The analysis of the presented data clearly showed that, in most of the analyzed water
18 samples these molecules are not detected or present at concentrations levels below the
19 detection limits of the developed method. The selected N-nitrosamines were found only in
20 few of the collected samples at concentration levels much lower than those fixed by different
21 international organizations and regulatory authorities.

22 While the NDMA was the N-nitrosamine most frequently found in this work, in near
23 future, it is clear that this family of nitrogenous pollutants of health concern may come under
24 increasing scrutiny as water contaminants and disinfection by-products.

26 4. Conclusion

27
28 In this paper, a suitable SPE-UHPLC/MS/MS analytical methodology for the analysis
29 of N-nitroso compounds of health concern was developed. The proposed procedure enables
30 the simultaneous extraction, pre-concentration and quantification of N-nitroso-n-propylamine,
31 N-nitrosomorpholine, N-nitrosomethylethylamine and N-nitrosodimethylamine in various
32 water samples (drinking and surface waters). The presented data clearly demonstrated that,
33 the developed SPE-UHPLC/MS/MS method is highly sensitive and selective for their analysis

1 at ultra-trace levels (few ng L⁻¹). The analysis of the considered N-nitrosamines was
2 performed in 3 min.

3 Good linearity, precision, accuracy, lower limits of detection, and quantification were
4 obtained for all target molecules. The Sep-Pak[®] AC-2 cartridges used for analytes extraction
5 from water samples led to satisfactory extraction recoveries and to high pre-concentration
6 factors of 2500. In addition, no significant matrix effect for the considered N-nitrosamines
7 was observed in surface water samples. Moreover, the developed analytical technique
8 provides low MDLs allowing the quantification of N-nitrosamines at concentration levels
9 below the ones determined by many monitoring programs or below the notification levels
10 established by different legislations. The excellent detection limits of the developed method
11 make ultra-trace N-nitrosamine analysis possible.

12 In addition, the proposed method was successfully applied for the analysis of these
13 molecules in real water samples and was shown to be convenient and reliable for their
14 analysis in surface and tap water samples. The developed procedure is certainly, a powerful
15 analytical tool for future toxicological, epidemiological and screening studies for the
16 investigation of water pollution. It permits to rapidly initiate specific actions to minimize their
17 environmental release or impact.

18 The application of the proposed method to the analysis of surface and drinking water
19 samples from Brittany region (France) revealed that the selected N-nitrosamines were
20 determined only in few of the collected samples, at concentration levels much lower than
21 those fixed by different international organizations and regulatory authorities. Research is
22 ongoing in order to have a more detailed evaluation of the pollution levels of N-nitrosamines
23 associated with surface and drinking water in Brittany region in order to provide a more
24 adequate spatial and temporal coverage.

25

26

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14 **Figure captions**

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16 **Figure 1.** Water sampling locations on the Brittany territory considered in this study.

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Table 1. Physico-chemical properties and toxicity of the considered N-nitrosamines.

Nitrosamines (Abbr.)	Molecular formula	Molecular weight (g·mol⁻¹)	log K_{o/w}	Water solubility (mg·L⁻¹)	Standard U.S. EPA* cancer classification group
NDMA	C ₂ H ₆ N ₂ O	74.082	-0.57	1,000,000	B2
NMEA	C ₃ H ₈ N ₂ O	88.108	0.04	300,000	B2
NDPA	C ₆ H ₁₄ N ₂ O	130.188	1.36	13,000	B2
NMOR	C ₄ H ₈ N ₂ O ₂	116.059	-0.44	861,527.5	2B (IARC)**

*United States Environmental Protection Agency

**IARC: International Agency for Research on Cancer

Table 2. Optimized MS/MS parameters.

Parameter	Value
Source temperature (°C)	120
Capillary voltage (kV)	3.0
Desolvation temperature (°C)	350
Desolvation gas flow (L h ⁻¹)	750
Cone gas flow (L h ⁻¹)	75

Table 3. Ionization mode, MRM transitions used for quantification and confirmation purposes and optimized values used for cone voltage and collision energy for the analysis of N-nitrosamines.

Analyte	Ionization mode	Transition of quantification (m/z)	Transition of confirmation (m/z)	Cone voltage (V)	Collision energy (eV)
NDMA	ESI ⁺	74.7 > 42.8	75.0 > 58.5	25	10
NMEA		88.7 > 60.7	88.7 > 42.8	25	10
NDPA		131.5 > 43.8	130.8 > 88.6	22	11
NMOR		116.9 > 86.2	116.9 > 41.7	28	10

Table 4. Extraction recoveries (%) and relative standard deviation (RSD, %) obtained with the proposed SPE methodology for each analyte in ultrapure water for different spiking levels.

Analyte	Conc. spiked (ng L ⁻¹)	Conc. measured (ng L ⁻¹)	Recovery (%) ± RSD (%; n = 6)
NDMA	0.40	0.39	98±1
	4.00	3.92	98±1
	40.0	39.33	98±1
NMEA	0.40	0.39	99±1
	4.00	3.97	99±1
	40.0	39.75	99±1
NDPA	0.40	0.39	98±2
	4.00	3.94	99±1
	40.0	39.46	99±1
NMOR	0.40	0.40	100±1
	4.00	4.00	100±1
	40.0	40.0	100±1

Table 5. Regression coefficients (R^2) for the instrumental and sample calibration curves, matrix effects and relative standard deviation (RSD) of the selected N-nitrosamines spiked at 0.4 ng L^{-1} .

Analyte	ICC ^a (R^2)	SCC ^b (R^2)	Matrix effect (%) \pm RSD (%; n = 6)
NDMA	0.9987	0.9962	98 \pm 2
NMEA	0.9991	0.9983	99 \pm 2
NDPA	0.9995	0.9978	98 \pm 2
NMOR	0.9992	0.9961	100 \pm 1

^a Instrument calibration curve

^b Sample calibration curve

Table 6. Instrumental detection limit (IDL), instrumental quantification limit (IQL), concentration factor (FC) of the extraction method and method limit of detection (MDL) and method quantification limit of quantification (MQL).

Analyte	IDL ($\mu\text{g L}^{-1}$)	IQL ($\mu\text{g L}^{-1}$)	FC	MDL (ng L^{-1})	MQL (ng L^{-1})
NDMA	0.25	1.50	2500	0.10	0.60
NMEA	0.50	2.00		0.20	0.80
NDPA	1.00	3.00		0.40	1.20
NMOR	0.10	0.25		0.04	0.10

Table 7. Maximal concentrations (ng L^{-1}) measured in surface and tap water samples.

Sampling point	N-nitrosamines detected (ng L^{-1}) \pm RSD (%), (n = 6) ^b			
	NDMA	NMEA	NDPA	NMOR
Summer				
(June-July 2014)				
River water 1	0.2 \pm 0.1	ND ^a	< LD	ND
River water 2	ND ^a	ND	ND	ND
River water 3	< LD ^b	ND	0.6 \pm 0.3	< LD
River water 4	ND	ND	ND	ND
River water 5	ND	0.4 \pm 0.3	ND	ND
River water 6	0.3 \pm 0.3	ND	ND	< LD
Winter				
(October-November 2014)				
River water 1	ND ^a	ND	0.4 \pm 0.2	ND
River water 2	< LD	ND	ND	< LD
River water 3	0.4 \pm 0.2	ND	0.8 \pm 0.4	ND
River water 4	0.2 \pm 0.2	ND	ND	ND
River water 5	ND	0.6 \pm 0.2	< LD	ND
River water 6	0.7 \pm 0.5	ND	ND	0.2 \pm 0.1
Summer				
(June-July 2015)				
Tap water 1	< LD	ND ^a	< LD	ND
Tap water 2	ND	ND	ND	ND
Tap water 3	< LD	ND	< LD	ND
Tap water 4	ND	ND	ND	< LD
Tap water 5	ND	< LD	ND	ND
Tap water 6	0.4 \pm 0.2	ND	ND	< LD

^a NF: not found^b LD: limit of detection

