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# 1 Sources and behavior of perchlorate in a shallow Chalk aquifer under

# 2 military (World War I) and agricultural influences

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### 11 Abstract

12 Perchlorate ( $ClO_4$ ) has been detected at concentrations of concern for human health on a large scale in groundwater used for drinking water supplies in NE France. Two sources are suspected: a military 13 14 source related to World War I (WWI) and an agricultural source related to past use of Chilean nitrate fertilizers. The sources and behavior of ClO<sub>4</sub><sup>-</sup> have been studied in groundwater and rivers near the 15 Reims city, by monitoring monthly the major ions and ClO<sub>4</sub> concentrations for two years (2017 – 16 17 2019), and by measuring the isotopic composition of ClO<sub>4</sub><sup>-</sup> and NO<sub>3</sub><sup>-</sup> in water samples. ClO<sub>4</sub><sup>-</sup> was detected throughout the study area with high concentrations (>  $4 \mu g \cdot L^{-1}$ ) detected mainly downgradient 18 19 of the Champagne Mounts, where large quantities of ammunition were used, stored and destroyed 20 during and after WWI. A WWI military origin of ClO<sub>4</sub><sup>-</sup> is inferred from isotopic analysis and 21 groundwater ages. Different tendencies of ClO4- variation are observed and interpreted by a combination of ClO<sub>4</sub> concentrations, aquifer functioning and historical investigations, revealing major 22 sources of ClO<sub>4</sub><sup>-</sup> (e.g., unexploded ordnance, ammunition destruction sites) and its transfer 23 24 mechanisms in the aquifer. Finally, we show that concentrations of ClO<sub>4</sub><sup>-</sup> in groundwater seems 25 unlikely to decrease in the short- to medium-term.

# 26 Keywords

27 Perchlorate; Isotope; Groundwater; World War I; Chilean nitrate fertilizer

# 28 Highlights

29	•	High ClO <sub>4</sub> <sup>-</sup> concentrations are detected mainly downgradient of the Champagne mounts.
30	•	ClO <sub>4</sub> <sup>-</sup> contamination of water comes mostly from WWI military sources.
31	•	Factors governing ClO <sub>4</sub> <sup>-</sup> transfer in the Champagne Chalk aquifer are evidenced.
32	•	ClO <sub>4</sub> <sup>-</sup> contamination seems unlikely to decline in the short- to medium-term.

2

### 33 1 Introduction

34 Perchlorate ( $ClO_4^{-}$ ) is an environmental pollutant of growing concern due to its widespread occurrence 35 in water and its adverse health effects by interfering with thyroid uptake of iodine and production of 36 hormones (e.g., Brabant et al., 1992; Braverman et al., 2005; Greer et al., 2002). The high solubility 37 and nonreactivity of ClO<sub>4</sub><sup>-</sup> under typical geochemical conditions make it highly mobile in the aquatic 38 environment (Brown and Gu, 2006; Urbansky, 1998). The origin of  $ClO_4$  in the environment can be 39 synthetic or natural (Cao et al., 2019). ClO<sub>4</sub><sup>-</sup> is widely used as an oxidant in solid rocket fuels and 40 explosives (Urbansky, 1998). Therefore, the use and manufacture of these products may constitute an 41 important source of  $ClO_4$  contamination. Other synthetic sources of  $ClO_4$  include fireworks, air bags, 42 matches, road flares, chlorate herbicides and bleach products (Aziz et al., 2006; Trumpolt et al., 2005). 43 In natural environments, ClO<sub>4</sub> can form atmospherically and accumulate by dry and wet deposition in 44 vadose soils in arid and semi-arid environments such as the Atacama Desert (Chile), the southwestern 45 United States, northwestern China, and Antarctica (Jackson et al., 2015, 2016; Lybrand et al., 2016; 46 Rao et al., 2007). The natural NO<sub>3</sub>-rich salt deposits in the Atacama Desert (Chilean nitrate), with 47 ClO<sub>4</sub> as a minor component (0.1-0.5 wt.%), have been refined and distributed worldwide for use as  $NO_3^{-1}$  fertilizer, especially during the first half of the 19<sup>th</sup> century, thus representing a potential 48 49 widespread source of ClO<sub>4</sub><sup>-</sup> contamination in waters (Ericksen, 1983; Rajagopalan et al., 2006). Over 50 the last two decades, ClO<sub>4</sub> contamination has been reported from many countries including the USA, Canada, Chile, China, India, UK and France, related to various origins (e.g., Cao et al., 2019, 2018; 51 Furdui et al., 2018; Jackson et al., 2005; Kannan et al., 2009; McLaughlin et al., 2011; Qin et al., 2014; 52 53 Sturchio et al., 2014; Urbansky, 2002; Vega et al., 2018).

In France, recommended levels of  $ClO_4^-$  in drinking water were first issued in 2011 by the French Agency for Food, Environmental and Occupational Health & Safety (ANSES): 15  $\mu$ g·L<sup>-1</sup> for adults and 4  $\mu$ g·L<sup>-1</sup> for children under 6 months. The level was subsequently reduced from 15 to 5  $\mu$ g·L<sup>-1</sup> for adults (ANSES, 2018). According to the national measurement campaign conducted by ANSES from 58 2011 to 2012, most of the sites having high concentrations of  $ClO_4^-$  (> 4 µg·L<sup>-1</sup>) were located in NE 59 France (ANSES, 2013).

The large extent of ClO<sub>4</sub><sup>-</sup> contamination in NE France is unlikely to have been caused by point sources 60 61 related to industrial activities (Figure 1). However, a potential link between the spatial distribution of 62 high ClO<sub>4</sub><sup>-</sup> concentrations and the position of the trench areas of WWI (1914 – 1918) was observed. 63 Thus, a military source of ClO<sub>4</sub><sup>-</sup> related to WWI seems likely (Hubé, 2016; Hubé and Bausinger, 2013; Ricour, 2013) but has never been clearly demonstrated. Indeed, synthetic ClO<sub>4</sub> produced industrially 64 65 by electrolysis was largely used in explosives manufacturing during WWI. According to Hubé (2014), ~131,000 T of (per)chlorate explosives (primarily composed of NH<sub>4</sub>ClO<sub>4</sub>, KClO<sub>4</sub>, or NaClO<sub>3</sub>) were 66 67 used during WWI, mainly in grenade and trench artillery. In addition, ClO<sub>4</sub> could also be present in other explosives including black powder and nitro group explosives (e.g., TNT, nitroglycerine and 68 nitrocellulose), as Chilean nitrate (with ClO<sub>4</sub> impurity) was intensively used in the manufacturing of 69 70 these explosives.



71

Figure 1 : Distribution of ClO<sub>4</sub><sup>-</sup> contamination in groundwater and the positions of the WWI trench areas in NE France
 (Jaunat et al., 2018)

74 High levels of  $ClO_4^-$  have also been detected in some regions outside the trench areas (e.g., Oise, Pas-75 de-Calais; Figure 1). An agricultural source was suspected, as large quantities of Chilean nitrate was 76 used as fertilizer in France between 1880 and 1950 (Lopez et al., 2015), especially for sugar beet and 77 wheat cultivation (Zimmermann, 1917). Around the year 1928, the estimated amount of Chilean nitrate applied annually for wheat cultivation in France was between 150 and 400 kg·ha<sup>-1</sup> and could 78 exceed 800 kg·ha<sup>-1</sup> for beet cultivation (Lopez et al., 2014). For a better management of water 79 80 resources in NE France, it is now necessary to clarify the source of ClO<sub>4</sub><sup>-</sup> contamination (military 81 and/or agricultural). In addition to hydrogeological and historical investigations, isotopic analysis of 82  $ClO_4$  can provide a direct approach for  $ClO_4$  source apportionment.

Measurements of stable isotope ratios of chlorine ( ${}^{37}$ Cl/ ${}^{35}$ Cl) and oxygen ( ${}^{18}$ O/ ${}^{16}$ O,  ${}^{17}$ O/ ${}^{16}$ O) and the fractional abundance of the radioactive isotope  ${}^{36}$ Cl in ClO<sub>4</sub><sup>-</sup> ions have shown that three primary source types of ClO<sub>4</sub><sup>-</sup> (synthetic; "Atacama" from nitrate salts mined in the Atacama Desert of Chile; and indigenous natural atmospheric deposition) can be clearly distinguished isotopically (Bao and Gu, 2004; Böhlke et al., 2017, 2009, 2005; Hatzinger et al., 2011; Jackson et al., 2010; Poghosyan et al., 2014; Sturchio et al., 2014, 2011, 2009, 2006). Additional background information about the ranges of isotopic composition of ClO<sub>4</sub><sup>-</sup> is detailed in the references above and summarized in Cao et al. (2019).

90 In this study, a representative study area with two potential sources of ClO<sub>4</sub><sup>-</sup> contamination (military 91 and/or agricultural) was selected east of the Reims city in NE France (Figure 1). In this agricultural 92 and military (WWI) context, the primary objectives of this study are to 1) assess the extent of  $ClO_4^{-1}$ 93 contamination and its spatio-temporal evolution in the Chalk aquifer, 2) clarify the sources of ClO<sub>4</sub><sup>-</sup> 94 contamination (military and/or agricultural) and 3) understand the mechanism of transport and predict 95 the evolution of ClO<sub>4</sub> in groundwater in the short- to medium- term. To achieve these goals, we used 96 an approach combining continuous monitoring of ClO<sub>4</sub><sup>-</sup> concentrations, isotopic composition 97 measurements, historical and hydrogeological investigations, which could be further applied in other 98 ClO<sub>4</sub><sup>-</sup> contaminated areas.

### 99 2 Materials and methods

#### 100 **2.1 Description of the study area**

#### 101 2.1.1 Location and land use

The study area is situated east of the city of Reims, in the Champagne region in NE France (Figure 1 and Figure 2). It covers about 500 km<sup>2</sup> between the Vesle River (as the southern boundary) and the Suippe River (as the northern and eastern boundaries; Figure 2). Land use in this area is largely agricultural (> 80%) with wheat, barley, sugar beet, and alfalfa as major crops. Forest covers about 15% of the study area, mainly on the Berru and Champagne Mounts and the riparian areas. Urban lands including towns, villages, and industrial sites represent only about 3% of the study area (CORINE Land Cover geographic database).

#### 109 2.1.2 Geological and hydrogeological context

110 A detailed geological and hydrogeological description of the study area is given by Cao et al. (2020). 111 The Chalk formation (Upper Cretaceous, 66 - 100 Ma) constitutes almost the entire surface of the study area, with only a limited exposure of Tertiary formation at Berru Mount (Figure 2). The Tertiary 112 113 formation includes a succession of permeable (sand and coarse limestone) and impermeable deposits 114 (clay and marl) where only a few thin aquifers of limited extent are developed with some springs 115 flowing from the sand layers (Laurain et al., 1981). These aquifers are insufficiently productive to be 116 used as drinking water supplies. The Chalk formation of the study area includes Coniacian (C3), 117 Santonian (C4) and Campanian (C5) Chalks (Figure 2), which have similar lithology and 118 hydrodynamic properties despite their different ages of formation. It is a pure, fine-grained carbonate 119 rock, characterized by dual porosity (matrix and fracture) providing both slow and rapid flowpaths for 120 groundwater (e.g., Foster, 1975; Headworth et al., 1982; Price, 1987). The total porosity of the Chalk 121 is about 40% (Crampon et al., 1993) with only 1% effective porosity (Vachier et al., 1987). The first 10-20 m of the Chalk is significantly fractured; the density of fractures decreases with depth and 122 distance to river valleys (Allen et al., 1997; Mangeret et al., 2012; Vachier et al., 1987). In addition, 123

- the Chalk formation is partially covered by superficial (Quaternary) formations including graveluche
  (a periglacial formation up to 10 m thick), colluvium (1-3 m thick) and alluvium (up to 10 m thick)
- 126 with relatively high content of clay and silt (Allouc et al., 2000; Vernhet, 2007) (Figure 2).



Figure 2 : Geological and hydrogeological map of the study area recorded at high water level with the location of sampling points (modified from Cao et al., 2020). Source of geological map: Laurain et al. (1981), Allouc and Le Roux. (1995); source of water level data: Rouxel-David et al. (2002a).

The chalk aquifer is an important groundwater resource of the region. Precipitation is the only recharge source of the aquifer, mainly from November to March, due to the excess of rainfall compared to evapotranspiration (Cao et al., 2020; Chiesi, 1993). The surface runoff is usually considered as very low or absent for the Champagne Chalk aquifer (Chiesi, 1993; Foster, 1975;

Mathias et al., 2006). The study area is divided into two parts by the groundwater divide line across
the summit of Berru Mount and the Champagne Mounts, which delimits the Vesle River watershed in
the south and the Suippe River watershed in the north (Figure 2).

#### 138 2.1.3 Suspected sources of ClO<sub>4</sub><sup>-</sup> contamination

139 Groundwater contamination by ClO<sub>4</sub> in the study area is potentially caused by both the WWI-related 140 military source and the agricultural source related to the past use of Chilean nitrate fertilizer. The 141 study area was intensively marked by events of WWI (Facon, 2018; Laurent, 1988). As shown in 142 Figure 3, almost the entire study area was crossed by trenches, particularly in the center of the area (Taborelli, 2018). From 1914 to 1918, these trench areas were the scenes of several intensive battles of 143 144 WWI (e.g., the Battles of Champagne), especially on the Champagne Mounts. During WWI, military 145 tunnels were dug by the German army, such as the tunnels of Mont Cornillet (N1), Mont Perthois (N2) 146 and Mont sans Nom (N3) (Figure 3). These tunnels, usually equipped with quantities of ammunition, connected the German front positions with the rear and allowed the German army to fire until the last 147 148 moment. It is highly probable that quantities of ammunition still remain underground at these sites, 149 representing potential sources of groundwater contamination of  $ClO_4^-$  and other pyrotechnic 150 compounds. Furthermore, it is likely that the precise locations of some of these underground sites are 151 unknown.

152 After WWI, large quantities of unused ammunition were still present on the French land and it was necessary to clear the battlefield in order to make these lands habitable and cultivable on a large scale. 153 154 Thus, unused ammunitions were either dismantled to recycle valuable materials or destroyed by 155 explosion. The ammunition explosion sites were recognizable in historical aerial photographs by anthropogenic forms such as alignments of shell-holes, an access road, and a storage area. Several 156 157 explosion sites were identified in the study area from inspection of historical aerial photographs (IGN 158 Remonter le temps database; Figure 3). While traces of N5 and N6 could no longer be recognized, the 159 destruction activities on site N1 lasted until the 2000s and some military wastes are still present at this 160 site (BASOL database). The sites and activities listed are not exhaustive since the compilation is based 161 on existing data from historical archives and/or former aerial photographs that did not cover the entire 162 study area. In addition, traces of military activities could have been quickly erased, making it difficult 163 to identify all related sites. The ammunitions exploded on battlefields during WWI or destroyed after 164 the war could have released  $ClO_4^-$  into the environment. In addition, large quantities of unexploded 165 ammunitions (about 30% of the total quantity of ammunition used) could still persist in the subsoil 166 (Desailloud and Wemeau, 2016), representing a continuous and diffuse source of  $ClO_4^-$  contamination.



<sup>167</sup> 

- 172 consumption of nitrogen in Marne (the department where the study area is located) was between 7,000
- 173 and 10,000 T in the 1950s (Lopez et al., 2014). The use by spreading and the storage of these Chilean
- 174 nitrates could represent diffuse or point sources of ClO<sub>4</sub><sup>-</sup> contamination in groundwater.

Figure 3 : Trenches, military tunnels and ammunition destruction sites related to WWI on the study area (position of trenches from Taborelli, 2018)

<sup>170</sup> As a traditional agricultural area for wheat and sugar beet, the study area was extensively cultivated

<sup>171</sup> with the use of Chilean nitrate fertilizer from 1880 to 1950. It was estimated that the average annual

#### 175 **2.2** Sampling and analysis methods

176 An intensive sampling network was established in the study area with a total of 36 sampling points 177 including 26 groundwater (19 boreholes, 3 pumping stations and 4 springs) and 10 surface water (2 in 178 the Aisne-Marne Canal, 2 in the Vesle River, 3 in the Suppe River and 3 in their tributaries) (Figure 179 2), as surface water is here closely related to groundwater of the Chalk aquifer. The boreholes and 180 pumping stations have long screen depths which cover at least half of the well depths (Table 1), thus providing a weighted average of groundwater. A first screening campaign was carried out in June 181 182 2017, which yielded a chemical map of the study area and the extent of  $ClO_4^-$  contamination. Fourteen sampling points (primarily points with high concentrations of ClO<sub>4</sub>, well distributed in the study area; 183 184 Figure 2) were then selected for monthly monitoring for 2 years to observe the spatio-temporal 185 evolution of groundwater geochemistry and ClO<sub>4</sub><sup>-</sup> concentrations. In addition, groundwater ages, water 186 flow rate in rivers, explosive concentrations, and isotopic compositions of ClO<sub>4</sub><sup>-</sup> and NO<sub>3</sub><sup>-</sup> were 187 measured.

188 The measurement methods and results of groundwater level, physico-chemical parameters, major ions 189 and groundwater dating using CFCs and  $SF_6$  have been fully described and interpreted previously in 190 Cao et al. (2020). These results are directly used in this study.

Water discharge of rivers was measured by the cross-section method using an OTT C2 Small Current Meter.  $ClO_4^-$  was analyzed by ion chromatography at BRGM (Orléans, France) with a quantification limit of 0.5  $\mu$ g·L<sup>-1</sup>. Thirty-nine explosives (see SI.1) were analyzed in waters with an RPHPLC-DAD system by the company Envilytix GmbH (Wiesbaden, Germany), as described by Bausinger et al. (2007).

196 Isotopic analyses of  $ClO_4^-$  were carried out in water samples collected from 3 sites in the study area: 197 FBN4 (at high and low water levels), FVDV (at high water level) and PY (at high water level; Figure 198 2). In order to obtain a pure military isotopic signature of  $ClO_4^-$ , a water sample was also collected at 199 the limit of the Suippes Military camp (outside the study area; Figure 1), as the only source of  $ClO_4^-$ 200 here that is related to WWI without any potential agricultural influence. Water samples were collected 201 using PVC columns containing ClO<sub>4</sub><sup>-</sup> specific ion exchange resin (IX resin) and then purified and 202 analyzed as described previously by Gu et al. (2011), Hatzinger et al. (2011, 2018), and Böhlke et al. 203 (2017). The extraction and purification of ClO<sub>4</sub><sup>-</sup> was done in the Department of Civil and 204 Environmental Engineering of Texas Tech University (USA) and the key steps can be summarized as 205 follows: 1) the resin was washed by deionized water and flushed with 4M HCl to remove  $NO_3^{-}$ ,  $SO_4^{2^-}$ , 206 HCO<sub>3</sub><sup>-</sup> and organics; 2) the absorbed ClO<sub>4</sub><sup>-</sup> was eluted from the IX resin using a solution of 1M FeCl<sub>3</sub> 207 and 4M HCl (Gu et al., 2007, 2001; Gu and Brown, 2006); 3) eluted ClO<sub>4</sub> was purified by a series of precipitation, liquid-liquid extraction, evaporation, and cation exchange processes, then crystallized as 208 209 a ClO<sub>4</sub> salt for isotopic analysis. The relative abundances of stable isotopes of chlorine (<sup>37</sup>Cl and <sup>35</sup>Cl) 210 and oxygen (<sup>18</sup>O, <sup>17</sup>O and <sup>16</sup>O) in ClO<sub>4</sub>- were measured using isotope-ratio mass spectrometry (IRMS) 211 at the Environmental Isotope Geochemistry Laboratory of the University of Delaware (USA).

For isotopic analysis of  $NO_3^-$ , water samples were collected in 100-mL polyethylene bottles after filtration through 0.45 µm membranes. Nitrogen and oxygen isotope ratios were measured using an automated denitrifier method as described in Morin et al. (2009) and Savarino et al. (2013). This technique uses *Pseudomonas aureofaciens* bacteria to convert  $NO_3^-$  to  $N_2O$ , which is then analyzed for its isotopic composition after thermal decomposition to  $O_2$  and  $N_2$ . Isotopic analysis was performed on a Thermo Finnigan MAT253 equipped with a gas-bench interface at the Laboratoire de Glaciologie et Géophysique de l'Environnement at the University of Joseph Fourier Grenoble (France).

#### 219 2.3 Statistical analysis methods

Interactions between ClO<sub>4</sub><sup>-</sup> concentrations and groundwater level as well as major ions were calculated by using an approach based on a semi-parametric regression model. More precisely, a generalized additive model (GAM) with cubic splines accounting for autocorrelation of data through a first order autoregressive model (AR1) was used. Each GAM was carried out using the Akaike information criterion (AIC) score and the likelihood ratio. The corresponding adjustment coefficient of determination (Adj.  $R^2$ ) associated with a significance test at a P-value < 0.05 was used to characterize correlations between response variable and the explicative one. 227 The GAM was also adopted to decipher time-dependent changes in perchlorate concentrations. A model was built for each of the 14 sampling points, which were monitored monthly. Each of these 228 229 GAM was built considering a Gamma distribution of perchlorate concentration and the log link 230 function was used as it was the most appropriate for this kind of distribution. Two time-dependent 231 smooth functions were used to build each model. Seasonal changes in perchlorate concentrations are represented by the first term taking into account days of year. The penalized cubic regression spline 232 233 was used for this smooth function to allow a nonlinear response of perchlorate to time during a 234 calendar year. The second term was constructed to filter out the long-term trends (inter-annual 235 variations). For convenient output, we have redefined the time-scale for this term as a continuous 236 variable. For this second term, the thin plate regression spline was used.

237 For the two smooth functions, the smoothness was controlled by the number of knots and the 238 associated effective number of degrees of freedom. The optimal number of knots was estimated 239 through cross-validation. In consequence, the basis dimension for the two smooth functions was 240 adapted to have enough degrees of freedom to fit the data while these values remained small enough to 241 maintain reasonable computational efficiency and avoided over-fitting the data. For the two smooth 242 functions, the method of the finite difference was used to calculate the first derivative of the function 243 and the associated confidence interval. This first derivative was used to estimate the rate of change to 244 identify periods of statistically significant change (either increase or decrease in perchlorate 245 concentration).

The underlying assumption of homogeneity for model residuals has been checked by plotting deviance residuals against fitted values. QQ plots (sample quantiles against theoretical quantiles) and Shapiro tests were used to assess normality of the model residuals. We have examined autocorrelation of the model residuals; the autocorrelation function values showed that the model residuals were not correlated as they dropped to small values within a couple of days. Validity of the model was also assessed through AIC score and Adj.R<sup>2</sup>. Analysis of deviance was used to assess the significance of

- the null hypothesis for the two smooth functions. All statistical tests were carried out using R-software
- 253 (R core Team, 2018).

## 254 **3 Results and discussion**

### 255 **3.1 Occurrence of ClO<sub>4</sub><sup>-</sup> and explosives**

256 Perchlorate was detected at almost all sampling sites (33 out of 36) (Table 1). Mean ClO<sub>4</sub><sup>-</sup> 257 concentrations measured during the two years monitoring were > 4  $\mu$ g·L<sup>-1</sup> at 17 sites, including two 258 sites with mean concentrations > 15  $\mu$ g·L<sup>-1</sup>, representing 49% and 6% of the sampling sites, 259 respectively. Low concentrations (< 4  $\mu$ g·L<sup>-1</sup>) were measured at 19 sites, representing 51% of the 260 sampling points.

261 Table 1 : Properties of sampling points, water table depth, concentrations of  $Cl^{-}$ ,  $NO_{3}^{-}$  and  $ClO_{4}^{-}$  and isotopic compositions 262 of  $NO_{3}^{-}$  in ground- and surface water samples (N: number of sampling; NA: not available. -: not applicable)

			Depth	Screen	Water table	Cl	NO <sub>3</sub> -	ClO <sub>4</sub> -	NO3 <sup>-</sup>	NO <sub>3</sub> -
Name	Туре	Ν	(m)	depth (m)	depth (m)	$(mg \cdot L^{-1})$	$(mg \cdot L^{-1})$	$(\mu g \cdot L^{-1})$	δ <sup>18</sup> O (% <i>o</i> )	δ <sup>15</sup> N (%)
FA	Borehole	21	35	3.5 - 30	2 - 6	$35.4 \pm 6.3$	$39.7 \pm 6.0$	$3.1 \pm 0.9$	-1.4	-0.7
FAP	Borehole	21	15	6.5 - 15	3 - 12	$30.7 \pm 9.5$	$28.4 \pm 6.7$	$5.4 \pm 2.2$	-1.9	1.7
FBN1	Borehole	21	48	4 - 48	26 - 40.5	$25 \pm 8.8$	$28.9 \pm 7.6$	$3.6 \pm 1.7$	2.1	3.9
FBN4	Borehole	21	28	16 - 28	2.5 - 18	$52.9 \pm 2.6$	$54.3 \pm 2.6$	$20.8 \pm 3.2$	-0.6	1.0
FBN5	Borehole	21	47	24 - 43	22.5 - 35	$15 \pm 9.2$	$31.6 \pm 11.3$	$7.9 \pm 3.7$	-0.6	1.1
FEP1	Borehole	21	25	7 - 25	7.5 - 17	$35.2 \pm 1.0$	$36.7 \pm 1.4$	$12.9 \pm 2.4$	-0.3	0.5
FP1	Borehole	21	19	9 - 15	4.5 - 13	$28 \pm 2.9$	$29.9 \pm 3.9$	$6.6 \pm 1.9$	-1.3	1.1
FPM1	Borehole	21	24	NA	NA	$7.6 \pm 0.7$	$19 \pm 0.8$	$14.1 \pm 2.0$	-1.4	0.4
FPM3	Borehole	21	21	7 - 21	9 - 18	$18.6 \pm 5.0$	$24 \pm 5.7$	$3.5 \pm 0.8$	-0.5	0.3
FVDV	Borehole	21	22	12.5 - 21	7 - 15	$39 \pm 1.6$	$40 \pm 2.3$	$44.4 \pm 6.9$	-1.7	0.8
PDO	Pumping station	21	25	7 - 25	NA	$33.9 \pm 1.2$	$42.1 \pm 2.0$	$11.5 \pm 1.7$	-1.8	0.6
PP	Pumping station	21	80	23 - 80	NA	$21.8 \pm 1.1$	$25.8\pm0.9$	$9.2 \pm 4.0$	-1.5	0.0
PS	Pumping station	4	16	7 - 16	NA	$23.6 \pm 1.4$	$32.4 \pm 2.7$	$4.1 \pm 1.4$	-0.2	1.6
FCE	Borehole	1	85	NA	NA	7.2	3.1	0.5	NA	NA
FNA	Borehole	1	47	NA	NA	20.0	24.3	1.3	NA	NA
FBN3	Borehole	3	56	12 - 56	16 - 22	$13.4 \pm 2.4$	$15.2 \pm 2.2$	$1.3 \pm 0.4$	5.4	7.5
FBN2	Borehole	4	32	10 - 32	17 - 26	$31.6 \pm 4.1$	$18.2 \pm 12.5$	$2 \pm 1.7$	6.6	10.3
FEP2	Borehole	6	23	NA	15 - 18.5	$8.8 \pm 0.6$	$9.5 \pm 0.8$	$2 \pm 1.3$	-0.3	-0.5
FPM2	Borehole	5	35	NA	30 - 31	$16.2 \pm 5.1$	$22.1 \pm 5.2$	$6.1 \pm 0.6$	-1.3	0.6
FP2	Borehole	4	21	NA	8 - 13	165.8 ± 18.9	$38 \pm 3.3$	$1.2 \pm 0.7$	-0.9	3.6
FP	Borehole	4	23	7 – 22.5	12 - 17	$39.0 \pm 3.3$	$45 \pm 5.0$	$5.9 \pm 0.9$	-1.7	0.6
FB	Borehole	1	33	NA	12.8	28,0	36.6	2.2	NA	NA
SEP	Spring	12	-	-	-	$23.5 \pm 1.9$	$33.1 \pm 2.6$	$5.5 \pm 3.0$	-1.5	1.8
SCG	Spring	2	-	-	-	$28.5\pm0.2$	$40.7 \pm 0.1$	$4.7 \pm 2.1$	NA	NA
SB1	Spring	1	-	-	-	16.5	7.8	< 0.5	NA	NA
SB2	Spring	1	-	-	-	18.4	2.3	2.1	NA	NA
RS1	Suippe River	21	-	-	-	$25.8 \pm 1.2$	$29.3 \pm 1.9$	$3.4 \pm 0.8$	1.0	3.7
RPY	Py River	21	-	-	-	$20.7 \pm 1.5$	$28.5 \pm 1.6$	$11.6 \pm 3.1$	NA	NA
RS2	Suippe River	21	-	-	-	$24.3 \pm 1.3$	$29.6 \pm 1.3$	$6.5 \pm 2.1$	0.8	2.6
RA	Arnes River	4	-	-	-	$22.1 \pm 1.0$	$26.1 \pm 1.2$	$1.5 \pm 1.0$	0.4	3.1
RS3	Suippe River	4	-	-	-	$25.2 \pm 0.4$	$28.8\pm0.8$	$5.1 \pm 1.5$	0.4	3.1
RV1	Vesle River	3	-	-	-	$29.3 \pm 1.1$	$25.9 \pm 3.7$	$1.2 \pm 0.5$	0.7	5.9
RV2	Vesle River	3	-	-	-	$25 \pm 1.9$	$28.5 \pm 2.8$	$0.9 \pm 0.4$	0.4	4.7
RP	Prosnes River	1	-	-	-	21.3	25.6	3.6	NA	NA
C1	Canal	1	-	-	-	18.3	12.7	< 0.5	NA	NA
C2	Canal	1	-	-	-	15.5	13.1	< 0.5	NA	NA

263 An analysis of the geographic distribution of ClO<sub>4</sub> is presented in Figure 4, revealing some major trends and potential sources of  $ClO_4^-$ . Lower concentrations of  $ClO_4^-$  (< 4 µg·L<sup>-1</sup>) were mainly found 264 on the Berru Mount, in the Vesle River and in the Aisne-Marne Canal. As mentioned above, the 265 Tertiary formation on the Berru Mount is represented by a succession of permeable and impermeable 266 267 layers, which contains several small aquifers in which water could be renewed quickly by precipitation. 268 As a result, low levels of  $ClO_4^-$  were detected in this area. In the Vesle River,  $ClO_4^-$  concentrations ranged from 0.9  $\pm$  0.4 µg·L<sup>-1</sup> to 1.2  $\pm$  0.5 µg·L<sup>-1</sup>. The Vesle River originates far away upstream, 269 receiving groundwater discharge from outside the study area that is little affected by ClO<sub>4</sub>-270 contamination. In the Aisne-Marne Canal,  $ClO_4^-$  was not detected (< 0.5 µg·L<sup>-1</sup>), indicating that the 271 canal has little or no input from the contaminated groundwater or river water (Vesle River) of the 272 study area. 273

274 Most of the sampling sites with  $ClO_4^-$  concentrations exceeding 4  $\mu$ g·L<sup>-1</sup> were located downgradient of 275 the Champagne Mounts (Figure 4), where large quantities of ammunitions were used, stored, and 276 destroyed during and after WWI. The highest concentrations of ClO<sub>4</sub> were found at borehole FVDV, with a maximum of 62.5  $\mu$ g·L<sup>-1</sup> (in September 2018) and an average of 44.4 ± 4.9  $\mu$ g·L<sup>-1</sup>. Plume-like 277 patterns of  $ClO_4^-$  were observed along the sections A – A' and B – B' (the same direction as the 278 groundwater flow line; Figure 4). At FBN4, FEP1 and SEP (A – A'), mean ClO<sub>4</sub> concentrations over 279 the two years monitoring were  $20.8 \pm 3.2 \ \mu g \cdot L^{-1}$ ,  $12.9 \pm 2.4 \ \mu g \cdot L^{-1}$  and  $5.5 \pm 3.0 \ \mu g \cdot L^{-1}$  respectively, 280 281 indicating a progressive decrease with distance downgradient. A similar pattern was observed at FPM1, FPM2 and FPM3 (B – B') with concentrations of 14.1  $\pm$  2.0  $\mu$ g·L<sup>-1</sup>, 6.1  $\pm$  0.6  $\mu$ g·L<sup>-1</sup> and 3.5  $\pm$  0.8 282 283  $\mu$ g·L<sup>-1</sup> respectively. The high ClO<sub>4</sub><sup>-</sup> levels and the observed plume-like patterns indicated that potential point-sources of ClO<sub>4</sub><sup>-</sup> could be present upstream at FVDV, FBN4 and FPM1. Specifically, the 284 military tunnel N1 and the ammunition destruction sites (N4 and N5) were likely responsible for the 285 286 high concentrations of ClO<sub>4</sub><sup>-</sup> measured at these sites (Figure 4).

287 Although ClO<sub>4</sub><sup>-</sup> concentrations in river waters were generally lower than those of groundwater, some

river sites had relatively high ClO<sub>4</sub><sup>-</sup> concentrations (>10  $\mu$ g·L<sup>-1</sup>), such as RPY in the Py River (Figure 4

and Table 1). The Py River is downstream from the Suippes military camp that represents a potential source of  $ClO_4^-$  contamination to the Py river watershed (Figure 1). In the Suippe River upstream of the confluence with the Py River (RS1), low  $ClO_4^-$  concentrations ( $3.4 \pm 0.5 \ \mu g \cdot L^{-1}$ ) were measured while, downstream of this confluence (RS2), higher concentrations ( $6.5 \pm 2.1 \ \mu g \cdot L^{-1}$ ) were found.



293

294 Figure 4 : Spatial distribution of ClO<sub>4</sub> contamination on the study area and military sites related to the WWI

Unlike the widespread contamination of  $ClO_4^-$  on the study area, organic explosives have not been detected in surface and groundwater samples, which could be explained by their low persistence and mobility in soil and water (Clausen et al., 2006). During WWI, nitro group explosives such as TNT, nitroglycerine and nitrocellulose were largely used. TNT can be rapidly degraded in most soil and aquifer systems; therefore, its presence is typically restricted to areas near its introduction to the environment. At most sites, TNT can be completely attenuated in the surface soil, thereby preventing 301 contamination of the unsaturated zone (UZ) or groundwater (Clausen et al., 2006). Nitroglycerin is
302 soluble when present alone and is subject to rapid biodegradation, but when present with nitrocellulose
303 it is insoluble. Nitrocellulose is also insoluble, resulting in its low mobility in the environment (Quinn,
304 2015).

#### 305 **3.2** Sources and fate of ClO<sub>4</sub><sup>-</sup> in the Chalk aquifer

306 3.2.1 Isotopic composition of  $ClO_4^-$  and  $NO_3^-$ 

The results of Cl and O stable isotope analysis for  $ClO_4^-$  in water samples are presented by dual isotope plots in comparison to published data for synthetic, Atacama and selected US indigenous natural  $ClO_4^-$  occurrences (Table 3 and Figure 5). There is no evidence of  $ClO_4^-$  biodegradation, which is consistent with the typical oxic condition of the unconfined Chalk aquifer (Barhoum et al., 2014; Edmunds et al., 1987); the isotopic composition of  $ClO_4^-$  could therefore reflect initial values of the sources.

313 Table 3 : Isotopic compositions of ClO<sub>4</sub> in ground- and surface water samples (HW: high water; LW: low water)

Name	Sample date	δ <sup>18</sup> O (‰)	$\Delta^{17}O$ (%o)	δ <sup>37</sup> Cl (‰)
FVDV	04/05/18	-20,7	0,2	0,3
FBN4 (HW)	04/06/18	-21,8	0,2	0,2
FBN4 (LW)	01/18/19	-29,6	-0,3	-3,3
RPY	04/10/19	-22,9	1,3	-6.0
RM	02/22/19	-18,2	0,1	-0,2

314 The  $\delta^{37}$ Cl,  $\delta^{18}$ O and  $\Delta^{17}$ O values of ClO<sub>4</sub><sup>-</sup> in water samples collected at the Suppe military camp (RM; 315 pure military source), FVDV and FBN4 (at high water level) plotted exactly within the synthetic ClO<sub>4</sub><sup>-</sup> range, proving similar military sources of ClO<sub>4</sub><sup>-</sup> at these sites (Figure 5). However, the results for 316 FBN4 (at low water level) was different, with a lower  $\delta^{18}$ O value (-29.6%) falling outside the 317 published synthetic ClO<sub>4</sub><sup>-</sup> ranges. Nevertheless, the  $\delta^{37}$ Cl and  $\Delta^{17}$ O values (-3.3% and -0.3%), 318 319 respectively) were typical of synthetic ClO<sub>4</sub><sup>-</sup>. Therefore, ClO<sub>4</sub><sup>-</sup> at FBN4 (at low water level) was 320 interpreted as of synthetic origin, but different from other samples and currently reported synthetic 321 ClO<sub>4</sub><sup>-</sup> products. Indeed, the manufacturing processes of synthetic ClO<sub>4</sub><sup>-</sup> and the materials used more than 100 years ago during WWI may be different from those of today. Even during WWI, different 322 ClO<sub>4</sub> salts (NH<sub>4</sub>ClO<sub>4</sub> and/or KClO<sub>4</sub>) were used in explosives; they were produced in different facilities 323 17

324 and the method could also have evolved during the WWI conflict. This could possibly explain why 325 this "unusual" synthetic end-member was observed at FBN4 (at low water level). The different ClO<sub>4</sub>-326 isotopic compositions observed at FBN4 at high and low water levels suggest that there could be 327 different sources of ClO<sub>4</sub> at different depths. The estimated groundwater ages in the study area were less than 50 years (Cao et al., 2020). Moreover, because the thickness of UZ is less than 30 m (Table 1) 328 329 and considering that the flow rate is about 1 m year<sup>-1</sup> through the Chalk matrix, the time of transfer in 330 the UZ were less than 30 years (e.g., Barraclough et al., 1994; Brouyère et al., 2004; Chen et al., 2019; Wellings, 1984). Therefore, military sources of ClO<sub>4</sub><sup>-</sup> contamination are most likely related to military 331 activities after WWI (destruction of ammunitions) rather than during the conflict, or to the release of 332 333 ClO<sub>4</sub><sup>-</sup> from unexploded ordnance persisting in the subsoil (or unused ammunitions stored on the 334 surface).

335 In the RPY sample, a substantial fraction of  $ClO_4^-$  with an Atacama-type isotopic composition is evident. The lower value of  $\delta^{37}$ Cl (-6.0%) and the higher value of  $\Delta^{17}$ O (+1.3%) are both consistent 336 337 with a mixture of a synthetic end-member with typical Atacama ClO<sub>4</sub>. This could probably be explained by the nitrogen explosives of WWI (black powder and nitro group explosives) made with 338 339 Chilean nitrate and/or the past use of Chilean nitrate as fertilizer. Indeed, the PY River water consists 340 of aquifer discharge from the entire watershed. Although synthetic (military)  $ClO_4$  is indicated by 341 isotopic analysis at two sites with the highest  $ClO_4$  concentrations (FVDV and FBN4), Atacama  $ClO_4$ 342 from Chilean nitrate fertilizer might be present, especially at sites with low ClO<sub>4</sub><sup>-</sup> concentrations related to diffuse sources. 343

The  $NO_3^-$  isotope data at all the sampling points did not show any evidence of an Atacama source for the  $NO_3^-$  (Figure 5). However, this cannot rule out the possibility of the existence of Atacama  $NO_3^-$  in water samples, as Atacama  $NO_3^-$  could have been replaced and/or assimilated with the biogenic  $NO_3^$ in the soil (Böhlke et al., 2009). Indeed, the distinctive isotopic composition of oxygen in atmospheric  $NO_3^-$  is preserved only in hyper-arid environments and is lost in moist soils where higher biological activity occurs (Böhlke et al., 1997; Michalski et al., 2015). Therefore, more information is needed to

- 350 better evaluate the regional extent of ClO<sub>4</sub> sources related to the past use of Chilean nitrate in the
- 351 study area.



Figure 5: Summary of isotope data for ClO4 and NO3 in water samples of the study area, compared with major known ClO4 sources including synthetic, Atacama and indigenous natural ClO4 from the Southern High Plains (Texas) (SHP) (Ader et al., 2001; Bao and Gu, 2004; Böhlke et al., 2005, 2009; Jackson et al., 2005b, 2010; Parker et al., 2008; Plummer et al., 2006; Rajagopalan et al., 2006; Rao et al., 2007; Sturchio et al., 2007, 2011). Arrow in the  $\delta^{37}$ Cl vs  $\delta^{18}O$  graph represents the slope of biodegradation ( $\delta^{37}$ Cl = 0.4  $\delta^{18}O$ ) and arrow in the  $\Delta^{17}O$  vs  $\delta^{18}O$  graph represents the direction of mass dependent fractionation.

#### 359 3.2.2 Temporal variability of $ClO_4^-$ compared with groundwater levels and major ions

The temporal variation of  $ClO_{4^{-}}$  concentrations measured from June 2017 to June 2019 was compared with groundwater level fluctuation and the temporal variation of  $NO_{3^{-}}$  and  $Cl^{-}$  concentrations (two major agriculture-derived ions, Cao et al., 2020), in order to explore the potential sources of  $ClO_{4^{-}}$ (point and/or diffuse source), and the possible future evolution of  $ClO_{4^{-}}$  in the Chalk aquifer.

364 At most sites, inter-annual variations of  $ClO_4^-$  concentrations were observed and also revealed by the

- 365 statistical trend analysis (Figure 6 and Table 4). The periods when ClO<sub>4</sub> concentrations changed
- 366 significantly under the seasonal and annual effects were estimated with the statistical methods

- 367 presented in section 2.3 (results detailed in SI.2). Higher ClO<sub>4</sub><sup>-</sup> concentrations were observed in 2018
- 368 (Figure 6 and SI.2), corresponding with the higher groundwater level in 2018.

	Seasonal trend	Annual trend
FVDV	0,079	> 0.001
FBN4	0,097	0,209
FEP1	0,052	0,291
FBN5	0,003	0,007
FA	0,176	0,015
FBN1	0.028	0,374
FP1	0,161	0,165
FPM3	0,067	0,033
RS1	0,053	0,767
RPY	0,021	0,022
RS2	0,011	0,240
PDO	0,624	0,135
FPM1	0,097	> 0.001
PP	0,012	0,326

 $\begin{array}{l} 369\\ 370 \end{array} Table 4: Seasonal and annual effects on ClO4 concentrations (the effect is considered as statistically significant with P$  $value < 0.05, related values < 0.05 are marked in bold) \end{array}$ 

371 At sites having mean ClO<sub>4</sub><sup>-</sup> concentrations > 10  $\mu$ g·L<sup>-1</sup> (FVDV, FBN4, FEP1 and FBN5; Figure 6), 372 several peaks of  $ClO_4$  concentration were observed and the temporal variation of  $ClO_4$  was poorly correlated with the groundwater level fluctuation (Table 5). The peaks could possibly be explained by 373 374 localized flushing of ClO<sub>4</sub><sup>-</sup> from the UZ by natural (rainfall) or artificial (irrigation) recharge processes, 375 indicating the presence of point sources of ClO<sub>4</sub><sup>-</sup> contamination upstream of these sites. At FVDV and FBN4, the two most contaminated sites, the correlation coefficients between  $ClO_4$  and groundwater 376 377 level were the lowest (Table 4 and Figure 6), as ClO<sub>4</sub> concentrations here were mainly controlled by 378 the ClO<sub>4</sub>- transfer waves following flushing rather than the groundwater level fluctuation.

In contrast, at sites showing ClO<sub>4</sub><sup>-</sup> concentrations < 10  $\mu$ g·L<sup>-1</sup> (FA, FBN1, FP1 and FPM3; Figure 6), the temporal variation of ClO<sub>4</sub><sup>-</sup> was significantly correlated to the groundwater level fluctuation (Table 5). The correlation relationship was stronger at FA and FBN1 (Adj. R<sup>2</sup> = 0.42 and 0.54, respectively; Table 5) with larger groundwater level fluctuation; at FP1 and FPM3 where the groundwater level fluctuated less, the correlation relationship was weaker (Adj. R<sup>2</sup> = 0.35 and 0.20, respectively; Table 5). Diffuse sources of ClO<sub>4</sub><sup>-</sup> were presumed to be present at these sites. During low water level periods, most of the sources were apparently disconnected from the saturated zone. As water level rose, the 386 contamination source was re-activated and more contaminants were released in water and flushed into

387 the saturated zone, resulting in the increase of  $ClO_4$  concentrations.

388Table 5 : Correlation of ClO4 with groundwater levels and major ion concentrations. Values correspond to Adj.  $R^2$  issued389from the generalized additive model (statistically significant as the P-value < 0.05, related Adj.  $R^2$  values when P-value <</td>3900.05 are marked in bold; NA: not available)

Name	Water level	Cl.	NO <sub>3</sub> -	SO4 <sup>2-</sup>	Na <sup>+</sup>	<b>K</b> +	Mg <sup>2+</sup>	Ca <sup>2+</sup>
FVDV	0,07	-0,04	0,21	0,37	0,15	0,53	0,41	0,08
FBN4	0,07	0,03	0,32	0,24	0,03	0,18	0,31	0,01
FEP1	0,08	0,00	0,08	0,05	0,32	-0,02	-0,03	0,03
FBN5	0,22	0,71	0,64	0,02	0,28	0,67	0,41	0,61
FA	0,42	0,54	0,56	0,68	0,04	0,29	0,24	0,64
FBN1	0,54	0,53	0,67	0,55	-0,02	0,62	0,35	-0,02
FP1	0,35	0,17	0,25	0,23	0,28	0,48	-0,05	0,05
FPM3	0,20	-0,03	-0,04	-0,05	-0,04	0,17	0,46	0,18
RS1	NA	0,61	0,13	0,53	0,49	0,37	-0,06	0,05
RPY	NA	0,10	0,06	-0,01	-0,03	-0,06	-0,06	0,13
RS2	NA	0,51	0,11	0,36	0,02	0,17	0,53	0,13
PDO	NA	-0,05	0,06	0,01	-0,03	0,16	-0,03	0,19
FPM1	NA	0,28	0,51	-0,02	0,10	0,33	0,22	0,11
PP	NA	0,29	0,02	-0,05	0,16	0,03	0,08	-0,06

391 Generally, a poor correlation was observed between  $ClO_4^-$  and major ions (Table 5). The temporal 392 evolution of  $ClO_4^-$  was compared with the chronicles of  $NO_3^-$  and  $Cl^-$ , the two major agriculture-393 derived ions in groundwater of the Champagne Chalk aquifer (Cao et al., 2020). At most sites, the 394 temporal evolution of  $ClO_4^-$  was different from that of  $NO_3^-$  and  $Cl^-$  (Figure 6), indicating different 395 origins of ClO<sub>4</sub><sup>-</sup> versus NO<sub>3</sub><sup>-</sup> and Cl<sup>-</sup>. At FEP1, FPM1 and PP, despite the temporal heterogeneity of 396  $ClO_4$  levels, the concentrations of NO<sub>3</sub> and Cl were stable over time (Figure 6). As described in Cao 397 et al. (2020), estimated groundwater ages at these points were > 30 years in a piston flow model, which is related to the superficial formations limiting rapid transport of water and solutes ( $NO_3^-$  and 398 399 Cl<sup>-</sup>) from surface (agriculture-derived ions are distributed mainly in the soil area) to the saturated zone. 400 Consequently, the aquifer receives recharge mainly from upstream of the superficial formation 401 covered area. Water traveled laterally in the saturated zone during > 30 years and solute concentrations 402 were greatly buffered, resulting in stable NO3<sup>-</sup> and Cl<sup>-</sup> levels independent of water level fluctuation (Cao et al., 2020). The large variation of ClO<sub>4</sub> concentrations, in contrast to the temporal stability of 403 404  $NO_3^-$  and  $Cl^-$ , could be interpreted as an indication that the location of  $ClO_4^-$  sources is much deeper than those of agriculture-derived ions. As groundwater levels rose, the contamination front of ClO<sub>4</sub>-405

406 was reached, generating changes in  $ClO_4^-$  levels, despite the fact that flushing from the soil was largely 407 limited by superficial formations. At FPM3 (downstream from the borehole FPM1; Figure 2), a time 408 lag was observed between groundwater levels and concentrations of  $NO_3^-$  and  $Cl^-$ , likely as a result of 409 slow transport through the discontinuous graveluche formations. However,  $ClO_4^-$  levels varied with 410 groundwater levels without time lag, implying again that deeper  $ClO_4^-$  sources were more quickly 411 activated by the rise in water table.

412 At FA, unlike the positively correlated relationship between ClO<sub>4</sub><sup>-</sup> and groundwater level, NO<sub>3</sub><sup>-</sup> and Cl<sup>-</sup> 413 were negatively correlated with groundwater level (Figure 6 and Table 5), indicating a dilution effect. 414 At this site, groundwater dating showed 75-80% of modern water by the binary mixing model, 415 indicating a water table constituted mainly by freshly percolated rainwater that favors the process of 416 dilution (Cao et al., 2020). During the rapid flow process following precipitation, the low mineralized 417 rainwater entered the aquifer resulting in decrease of  $NO_3^-$  and Cl<sup>-</sup> concentrations. For ClO<sub>4</sub><sup>-</sup>, it seemed 418 that the inputs by the potential sources of ClO<sub>4</sub><sup>-</sup> in the UZ following the rise of groundwater level was 419 more important than the dilution process, which led to an increase in  $ClO_4$  with the groundwater level.

At FBN1 and FBN5, the temporal variabilities of  $ClO_4^-$ ,  $NO_3^-$  and  $Cl^-$  were well correlated (Figure 6 and Table 5). Indeed, located near the groundwater divide line (Figure 2), deep water table levels were observed (> 22 m, Table 1) and low permeability was suggested at these two sites (Cao et al., 2020). According to groundwater dating, the groundwater flow was well described by the exponential mixing model (mean residence time of < 20 years), indicating a spatially uniform recharge (Cao et al., 2020). ClO<sub>4</sub><sup>-</sup>, NO<sub>3</sub><sup>-</sup> and Cl<sup>-</sup> were thus flushed from the potential sources located in the UZ following recharge, showing a similar temporal variation.

427 At RPY (Py River) and RS2 (Suippe River after the confluence of Py River), a decrease in  $ClO_4^-$ 428 concentration was observed during high flow period (May 2018 and May 2019; Figure 6), implying a 429 dilution effect on  $ClO_4^-$  concentration. Indeed, the rivers are recharged by the Chalk aquifer with little 430 surface runoff produced. In river valleys, the Chalk is usually highly fractured with shallow 431 groundwater levels and preferential flow of rainwater is favored especially at high water level, resulting in dilution of the aquifer and the river water. A similar tendency was observed at PDO
(pumping station near RPY), which is consistent with a mixture of groundwater and surface water as
implied by groundwater dating (30% to 60% of modern water and an end-member of ~40-year old
water; Cao et al., 2020).



Figure 6 : Temporal variation of perchlorate concentrations compared with groundwater level fluctuations from June 2017
 to June 2019

The potential sources of ClO<sub>4</sub> located deeper than the sources of agriculture-derived ions in the UZ 439 440 refer most probably to unexploded ordnance still present underground after WWI. Indeed, some shells 441 fired during the war could reach > 10 meters underground without exploding (UNMAS, 2015). 442 Unused ammunitions could still be present underground in military tunnels of WWI. In addition, in the Champagne Mounts area, some unused ammunitions could have been cleaned up by burying in 443 444 specified boreholes (Debant, 2019). Over time, the release of the explosive charge occurs as a result of 445 the general corrosion of the envelope and/or its perforation. The time required for perforation was estimated between 250 and 450 years, at a rate of 1 mm/year on average (Parker et al., 2004). These 446 447 unexploded ordnances can be difficult to locate and clean up due to their number and their deep 448 underground location. Contamination plumes observed at some sites indicate that the ammunition 449 destruction sites may be the main point sources of ClO<sub>4</sub>, as repeated detonation causes an 450 accumulation of residual ClO<sub>4</sub><sup>-</sup> at these sites. The residue of ammunition destroyed by detonation can persist for more than 100 years in soils (Hubé, 2014). Despite the huge quantities of fully exploded 451 452 ordnance, their residue is estimated at around 0.01% and can only persist for a short time underground, 453 representing a relatively small contribution to the ClO<sub>4</sub><sup>-</sup> contamination of groundwater. Considering 454 that most of the potential sources of  $ClO_4$  have long persistence times, the  $ClO_4$  contamination in 455 groundwater of the study area may not decline in the short to medium term.

456

#### **3.3** Mass flow rate of perchlorate

457 In the Suippe River and its tributaries (Figure 4), the mass flow rate of  $ClO_4^-(M)$  has been estimated 458 according to the measured water flow rate (Q) and  $ClO_4^-$  concentrations (C):

459

$$M = C \times Q \tag{1}$$

460 The data of water flow rate and  $ClO_4^-$  concentration used for calculation was measured on October 461 2017. The estimated mass flow rate of  $ClO_4^-$  was 4.6 kg·month<sup>-1</sup> upstream the Suippe River (RS1) and 462 13.5 kg·month<sup>-1</sup> downstream of its confluence with the Py River (RS2) (Table 2). The increase of mass 463 flow rate from RS1 to RS2 (8.9 kg·month<sup>-1</sup>) was approximately equal to the contribution from Py

River (9.6 kg·month<sup>-1</sup>), implying that the contribution from the Chalk aquifer between RS1 and RS2 is 464 negligible with low groundwater discharge and low ClO<sub>4</sub> concentrations in the riparian aquifer. 465 The estimated mass flow rate of ClO<sub>4</sub><sup>-</sup> downstream along the Suippe River (RS3) was 14.6 kg·month<sup>-1</sup>. 466 467 The Arnes River, which joins the Suppe River between RS2 and RS3, has a  $ClO_4$  flux less than 0.1 kg·month<sup>-1</sup>. The increase of ClO<sub>4</sub><sup>-</sup> mass flow rate from RS2 to RS3 (1.1 kg·month<sup>-1</sup>) was due to the 468 469 contribution of Chalk aquifer discharge (zero runoff is assumed on the Champagne Chalk). The groundwater discharge from RS2 to RS3 was 322 L·s<sup>-1</sup>, estimated from the water flow rate at RS2, 470 RS3 and RA. According to equation (1), a  $ClO_4^-$  concentration of 1.3 µg·L<sup>-1</sup> in aquifer discharge on 471 472 October 2017 was estimated, which represents the average concentration from the Chalk aquifer of left 473 and right bank of the Suippe River.

474 *Table 2 : Mass flow rate of ClO4<sup>-</sup> in the Suippe River and its tributaries (data measured on October 2017)* 

1	Name	Location	Water flow rate $(L \cdot s^{-1})$	$ClO_4^-$ concentration ( $\mu g \cdot L^{-1}$ )	ClO <sub>4</sub> <sup>-</sup> mass flow rate (kg·month <sup>-1</sup> )
	RS1	Suippe River	401	4.3	4.6
	RS2	Suippe River	752	6.7	13.5
	RS3	Suippe River	1184	4.6	14.6
	RPY	Py River	276	12.3	9.1
	RA	Arnes River	110	< 0.5	< 0.1

The estimated monthly  $ClO_4^-$  flow rate on October 2017 represents approximately the minimum level in the Suippe River and its tributaries, as the water flow rate was measured during low discharge period, whereas  $ClO_4^-$  concentrations in rivers were relatively stable over time according to the low values of standard deviations presented in Table 1.

479 **4** Conclusions and perspectives

This study examined sources and evolution of  $ClO_4^-$  contamination in groundwater of NE France. The NE region of France is suspected to have multiple sources of  $ClO_4^-$  related to military activities of WWI and/or the past use of Chilean nitrate as fertilizer in agriculture. An intensive sampling network was established on a study area of a representative watershed, where  $ClO_4^-$  concentrations were monitored monthly for two years (2017-2019). The measured concentrations and isotopic contents of  $ClO_4^-$  and the historical investigations have been combined with previously published data on 486 groundwater dating as well as hydrologic and geochemical characteristics of the Chalk aquifer (Cao et 487 al. 2020), which allowed to clarify the sources of ClO<sub>4</sub><sup>-</sup> and to understand its evolution in groundwater. 488 This work produced the first precise  $ClO_4$  contamination mapping in the study area east of Reims city with ClO<sub>4</sub> concentrations in ground- and surface water ranging from < 0.5 to 62.5 µg·L<sup>-1</sup>. About half 489 of the sampling sites showed  $ClO_4^-$  concentrations > 4  $\mu g \cdot L^{-1}$  and most of these sites were located 490 491 downgradient of the Champagne Mounts area, where huge quantities of  $ClO_4^-$  were used, stored, or 492 destroyed during and after WWI. Point sources of ClO<sub>4</sub><sup>-</sup> were presumed to exist in the study area, as 493 indicated by the plume-like patterns of contamination observed at some sites. The isotopic signature of 494 ClO<sub>4</sub><sup>-</sup> at the two most contaminated sites showed a synthetic origin, proving the military source of 495 ClO<sub>4</sub><sup>-</sup> contamination in the study area. In addition, the estimated groundwater ages in the study area 496 were < 50 years, implying that ClO<sub>4</sub><sup>-</sup> contamination is related to sources that may still persist in the 497 subsoil long after the end of WWI (e.g., unexploded ammunition) or post-WWI military activities (e.g., 498 destruction of ammunition). The isotopic analysis of ClO<sub>4</sub> in river water showed a minor but distinct 499 component of Atacama ClO<sub>4</sub>, indicating the presence of some Chilean nitrate in the watershed.

500 Annual variations of ClO<sub>4</sub><sup>-</sup> concentration were observed, indicating the influence of recharge processes and groundwater levels on ClO4<sup>-</sup> contamination. Two major temporal trends of ClO4<sup>-</sup> concentration 501 502 were observed: 1) ClO<sub>4</sub> concentrations poorly correlated to groundwater level with peaks of 503 contamination due to flushing at sites having  $ClO_4$  concentrations > 10 µg·L<sup>-1</sup>, implying the presence 504 of point sources; 2)  $ClO_4^-$  concentrations highly correlated to groundwater level at sites showing  $ClO_4^-$ 505 concentrations  $< 10 \,\mu g \cdot L^{-1}$ , where diffuse sources were suggested. In addition, the rapid response of ClO<sub>4</sub><sup>-</sup> concentration following the rise of groundwater level compared with relatively stable 506 507 concentrations of agricultural ions at some sites indicated that the location of ClO<sub>4</sub> sources could be 508 much deeper than those of agricultural ions (mainly in the soil area). Considering the long persistence 509 time of the explosive residues related to unexploded ammunitions and ammunition destruction 510 activities, the ClO4<sup>-</sup> contamination in groundwater of the study area seems unlikely to decrease in the 511 short- to medium-term.

512 The multi-tool methodology developed in the study area could furtherly be applied to other ClO<sub>4</sub><sup>-</sup> 513 contaminated sites in NE France with suspected military and agricultural sources, with the aim of 514 making appropriate recommendations for a long-term management of groundwater resources. 515 Moreover, this characterization methodology could more widely applicable elsewhere in Chalk 516 aquifers or other multi-porosity mediums for the prediction of solute transport (natural or 517 anthropogenic) and for the evaluation of aquifer vulnerability. However, this research can be improved 518 and developed in several different aspects. The continuous monitoring of ClO4<sup>-</sup> concentrations and 519 geochemistry can be continued to obtain longer time series, with the aim to confirm and specify our 520 conclusions and also to study the evolution of ClO<sub>4</sub> in the Chalk aquifer under different climate 521 conditions. More measurements of water flow rate in rivers can be realized (during low, medium and 522 high discharge period) in order to better estimate the mass flow rate of ClO<sub>4</sub>. In addition, groundwater 523 sampling and analysis should also be realized at different depths of boreholes to obtain vertical 524 profiles of  $ClO_4$  concentrations as well as other chemical characteristics, which allows to explore their 525 relationships and to further confirm the position of potential ClO<sub>4</sub> sources underground. In this study, 526 isotopic analysis of ClO<sub>4</sub><sup>-</sup> has only been realized at sites with high levels of ClO<sub>4</sub><sup>-</sup>. This analysis should also be performed at sites having low concentrations of ClO<sub>4</sub><sup>-</sup> to further confirm whether traces of 527 Atacama  $ClO_4$  exist on these low contaminated sites related to diffuse sources. Finally, 528 529 hydrogeochemical numerical modeling tools could be a relevant complement of this study to better 530 understand the transfer mechanism of ClO<sub>4</sub> in the Chalk groundwater.

## 531 **Declaration of Competing Interest**

532 The authors declare that they have no known competing financial interests or personal relationships533 that could have appeared to influence the work reported in this paper.

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