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RELATION BETWEEN CONDUCTIVITY AND ION CONTENT IN URBAN WASTEWATER

Relation entre conductivité et composition ionique dan les eaux usées urbaines

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ABSTRACT

Wastewater conductivity has been monitored for extended periods of time by in situ probes and on grabbed samples in four communities (from 1,000 to 350,000 PE). In parallel, the concentrations of the main ionic contributors, such as calcium, sodium, potassium, magnesium, ammonium, orthophosphate, chloride and sulphate have been measured and their variations with respect to time compared to human activity patterns. It appears that sodium, potassium, ammonium and ortho-phosphate, which contribute to about 34% to wastewater conductivity, exhibit diurnal variations in phase with human activity evaluated by absorbance at 254 nm. However calcium ($\approx 22\%$ of wastewater conductivity) is out-of-phase. This release, ahead of the one of other cations and anions, could be related to sewer concrete corrosion or to groundwater infiltration. The combination of these different ionic contributions creates a conductivity pattern which cannot be easily related to human activity. It makes difficult to integrate conductivity in a monitoring system able to detect ion-related abnormalities in wastewater quality.

Keywords: Conductivity; corrosion; urine; calcium; potassium, infiltration.

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RÉSUMÉ

Les variations de la conductivité d'eaux usées urbaines ont été suivies sur de longues durées à l'aide de sondes placées in situ en entrée d'installations de traitement et sur des échantillons prélevés automatiquement. Quatre communautés (entre 1 000 et 350 000 habitants) ont été choisies pour cette étude. On a également dosé sur les échantillons les principaux ions (calcium, sodium, potassium, magnésium, ammonium, orthophosphates, chlorures et sulfates). Il apparaît que le sodium, le potassium, l'ammonium et les ortho-phosphates contribuent pour 34 % à la conductivité des eaux usées et présentent des variations journalières en phase avec la pollution carbonée soluble, estimée à partir de l'absorbance à 254 nm. Par contre, le calcium, qui contribue pour 22 % à la conductivité, présente un déphasage qui peut être dû à son transport dans le réseau d'assainissement du fait de la corrosion des conduites en béton ou à des infiltrations d'eaux claires. Finalement, la combinaison de ces différentes contributions ioniques conduit à une variabilité de la conductivité qu'il n'est pas facile de lier à l'activité humaine, et donc à des rejets accidentels dans le cadre d'un système de détection de variation anormale de la qualité des eaux usées.

Mots clés: Conductivité; corrosion; urine; calcium; potassium, infiltration.

1. INTRODUCTION

Domestic wastewater treatment systems of any kind (continuous or batch activated sludge reactors, anaerobic digesters, lagoons, etc.) are subjected to variations in flow and load due mainly to predictable changes in urban activity (normal diurnal cycle, week-ends and holidays). But accidental releases of industrial pollution and weather-related events (storm, long rain periods) may also occur. They can be considered as critical situations, which require steps to be taken to protect especially the biological stage: their rapid detection can be used in the control strategy of the system to trigger the opening of gates and to direct part or totality of the flow towards a temporary storage tank. As regulations require the treatment of all incoming wastewater in suitable facilities, the protection of the biological stages by detection of critical events is more and more necessary.

Wastewater characteristics are generally difficult to obtain rapidly. Automated chemical and biochemical analysis is proposed for COD, ammonium, nitrate, etc. and devices based on spectrophotometry are also available (HOCHEDLINGER et al., 2005; LANGERGRABER et al., 2003). For toxics it appears unrealistic to organize a specialized detection at the plant inlet as well as in the sewer network. The spectrum of possible components is very large: heavy metals, fuels, solvents, pesticides, herbicides, detergents, road de-icing products, etc. Respirometry-based systems are available for wastewater treatment plants (LE BONTE et al., 2005) but they are difficult to use along the sewer system, especially as many require the use of activated sludge. Conductivity and pH probes belong to a class of sensors that can be easily placed in situ either at the inlet or outlet of a wastewater treatment plant (KIM OANH and BENGTSSON, 1995) or further upstream in the sewer system to provide information on wastewater ionic content. Such a project has been initiated by the Greater Nancy (France) to implement early-warning systems based mainly on electrochemical probes in various locations of the sewer network (GRAPIN, 2004). But conductivity variations have been found difficult to understand. More recently, similar projects have been started in Austria and the Netherlands (SCHILPEROORT et al., 2006). The variations of the conductivity/calcium ratio have been proposed to detect easily pollution in rivers by untreated sewage (NIREL and REVACLIER, 2003). The conductivity c of a liquid sample is a linear function of the concentrations (C_{i}) of the n ions present:

$$\mathbf{c} = \sum_{i=1}^{n} \lambda_i \cdot \mathbf{z}_i \cdot \mathbf{C}_i \tag{1}$$

where λ_i is the mobility of ion i and z_i its electrical charge. Its variation will be related to cations and anions (mostly inorganic) present in the wastewater. In the case of release of high concentrations of ions (bases, acids, salts) the conductivity variations could be used to warn the plant operators and help to locate the discharge point in the sewer network. In order to evaluate whether conductivity could be used to detect changes in (waste)water composition, reference patterns should be obtained under dry weather conditions. For that purpose, the relation between the wastewater concentration of main ions and conductivity has been examined over extended periods of time in communities of different sizes.

2. MATERIALS AND METHODS

Wastewater grabbed samples have been collected at four different locations in the north-eastern part of France with auto-samplers (Sigma 900P, Hach Company, Loveland, Colorado for Nancy and Pont-à-Mousson, ASP-Station 2000, Endress+Hauser for Colombey-les-Belles and Villey-Saint-Etienne).

In Nancy-Maxéville (WWTP capacity = 350,000 PE), the device was installed on a loop providing primary settled wastewater, as shown in figure 1. On the same loop was installed a well-mixed tank, continuously fed with the same wastewater, in which were located a conductivity probe (CDC 565, Radiometer Analytical, Villeurbanne, France) and a pH (Type 06 242 040, Prolabo, Paris, France) probe. These probes were connected respectively to a PHM220 pHmeter and a CDM210 conductimeter (Radiometer Analytical, Villeurbanne, France). The pHmeter and the conductimeter were themselves connected via a RS232 multiplexer (Black Box Network Products n.v., Zaventem, Belgium) to a PC. A procedure developed under HPView (Hewlett-Packard) enabled the reading and storage of pH and conductivity values every 5 min. Data were collected over two periods, in winter and in summer. Calibration was checked weekly.

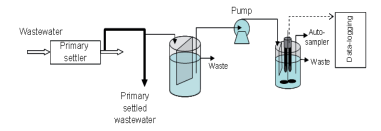


Figure 1. Experimental set-up at the Nancy-Maxéville wastewater treatment plant. Site expérimental de la station d'épuration de Nancy-Maxévill.

In Pont-à-Mousson (WWTP capacity = 16,200 PE), Colombey-les-Belles (1,200 PE) and Villey-Saint-Etienne (1,050 PE), raw wastewater samples were collected at the inlet of the wastewater works. The pH and conductivity of these grabbed samples were measured as above. Data were collected in spring for Pont-à-Mousson and in summer for Colombeyles-Belles and Villey-Saint-Etienne.

After filtration of all grabbed samples (on paper filter of pore diameter $\approx 7 \,\mu$ m), calcium, sodium and magnesium were determined by atomic absorption and potassium by atomic emission with a Varian AA20 spectrophotometer, according to the manufacturer's guidelines. Adequate dilution with deionised water was performed with a Hamilton auto-diluter in order to be in the linear response range of the device for each element. For the analysis of anions (sulphate, chloride, nitrate, ortho-phosphate), the samples were filtrated again (pore diameter = $0.45 \,\mu\text{m}$) and analyzed with a Dionex HPLC, according to the manufacturer's guidelines. The overall precision is estimated to less than 5% for these cations and anions. Ammonium was determined by the Nessler method modified to accommodate small samples (Hach method 380 on DR/2000). Ortho-phosphate was also determined spectrophotochemically (Hach method 490 on DR/2000). The overall precision is estimated to less than 5% for ammonium and ortho-phosphates. Absorbance at 254 nm (A_{254}) was measured on filtrated samples with an Anthelie Light spectrophotometer (Secomam, Domont, France). It is used as a surrogate parameter for COD (THOMAS et al., 1993), encompassing, due to the paper filter pore size, dissolved COD (< 0.45 µm) and colloidal COD (MRKVA, 1983). A coefficient of correlation larger than 0.84 was obtained between A_{254} and COD (Hach method 435) (Figure 2) confirming the results found by WU et al. (2006).

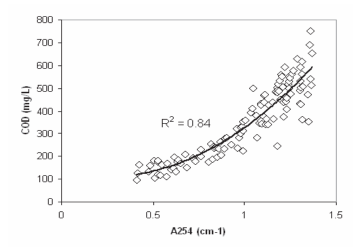


Figure 2. Correlation between absorbance at 254 nm (A₂₅₄) and COD. Corrélation entre l'absorbance à 254 nm (A₂₅₄) et la DCO.

3. RESULTS AND DISCUSSION

Figure 3 presents three examples of one-week monitoring of the variations of conductivity and pH in Nancy, Colombeyles-Belles and Villey-Saint-Etienne in summer. For Colombey and Villey the weather was dry but some rainfall had occurred two days before sampling started in Villey, which may have caused some disturbances on the first samples. Some small rainfalls occurred in Nancy. Nancy is a large community, with many small (but no large) industries of various natures, such as laundries, house-painting businesses, hardware stores, carrepairs, restaurants, hospitals, and a complex sewer system.

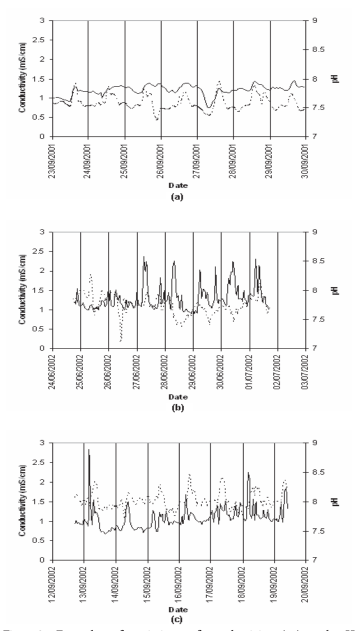


Figure 3. Examples of variations of conductivity (--) and pH (---) for three different communities in summer: (a) Nancy; (b) Colombey-les-Belles and (c) Villey-Saint-Etienne. Exemples de variations de la conductivité (--) et du pH (---) pour trois sites expérimentaux en été; (a) Nancy; (b) Colombey-les-Belles and (c) Villey-Saint-Etienne.

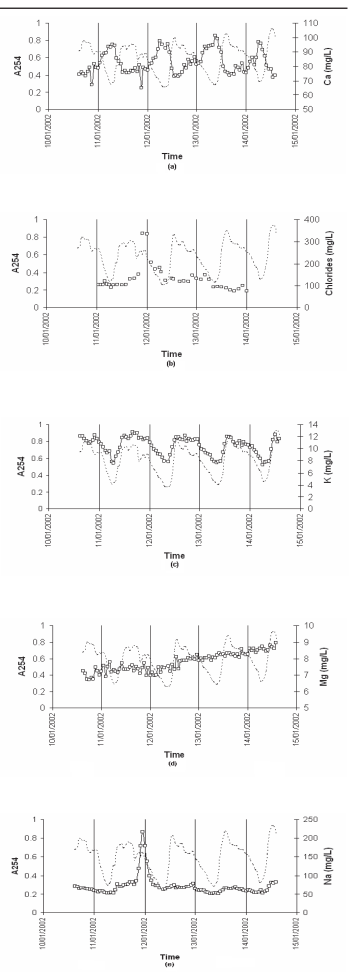
Diurnal patterns of organic and nitrogen pollutions are generally reported at the inlet of municipal wastewater treatment plants, reflecting the diurnal activity of inhabitants (HENZE *et al.*, 2002; METCALF and EDDY, 1991; PETERSEN *et al.*, 2002). In Nancy a diurnal pattern of variation of conductivity and pH is clearly visible. It is not so for the villages: it should be mentioned that Villey does not have any industry and Colombey has only small ironmongery and a small industrial bakery. To explain such variations, it is necessary to look more closely to the variations of cations and anions. It can be seen also that the variations are largely dampened for Nancy due to the size of the sewer network (750 km), much larger that the ones of the villages (between 10 and 20 km).

It can be seen in figure 4, which presents data collected in Nancy in winter, that the variations can be very different from one ion to another. Calcium and potassium exhibit diurnal variations. Potassium concentration is maximal when the soluble organic pollution (monitored by the surrogate parameter A_{254}) is maximal. It is the contrary for calcium, which increases at the end of the night. In this series magnesium kept increasing. A diurnal variation can be observed for sodium, but it is masked here by the large peak at 220 mg/L, which is due to rainfall and washout of road de-icing salt (which is generously spread on the streets in the North-East of France) in the sewer network. A peak of chloride can be observed at the same time. The concentration of nitrates is low, although slightly higher than the values reported in literature (0.5 mg N-NO₂/L in HENZE et al., 2002). This could be due to some nitrification occurring in the sewer system that could be favoured by the steep slopes in many parts of Greater Nancy. Diurnal variations can also be observed.

Table 1 summarizes the drinking water composition which can be used as a reference to analyze the wastewater data. In table 2, data on urine composition have been collected from literature (UDERT *et al.*, 2003a, 2003b). Urine is the main source of ammonia in domestic wastewater as urea is easily hydrolyzed according to the following reaction (UDERT *et al.*, 2003b):

$$\mathrm{NH}_{2}(\mathrm{CO})\mathrm{NH}_{2} + 2\mathrm{H}_{2}\mathrm{O} \rightarrow \mathrm{NH}_{3} + \mathrm{NH}_{4}^{+} + \mathrm{HCO}_{3}^{-} (2)$$

The ratio of each species with respect to total potential ammonia $\left(NH_{4\,hyd}^{+}\right)(\eta)$ has been calculated for urine and average Nancy wastewater. Urine contains also large amounts of potassium, sodium and chloride. Due to high time-correlation between ammonia and potassium, it is likely that a large part of the potassium released in the sewer is related to urine. Furthermore $\eta_{K,urine}$ is of the same order of magnitude than $\eta_{K,wastewater}$. The average sodium content in wastewater is higher than in the drinking water but $\eta_{Na,urine}$ is lower than $\eta N_{a,wastewater}$. As stated in ERIKSSON *et al.* (2002), sodium is often included in powder laundry detergents as a counterion



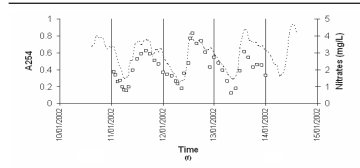


Figure 4. Variations of the concentrations (open symbols) of some anions (chlorides and nitrates) and cations (sodium, potassium, magnesium and calcium) in Nancy (winter season). (- - -) A₂₅₄: (a) Ca; (b) Chlorides; (c) K; (d) Mg; (e) Na; (f) Nitrates. Variations des concentrations (symboles ouverts) de quelques anions (chlorures ry nitrates) et cations (sodium, potassium, magnésium et calcium) à Nancy (en hiver). (- -) A₂₅₄: (a) Ca; (b) Chlorides; (c) K; (d) Mg; (e) Na; (f) Nitrates.

for sulphate (diluting agent), phosphate, silicate, carbonate or borates (builders). Grey-water from washing activities is therefore a source of sodium. During winter season, de-icing salt is also the source of sudden peaks of sodium. In Nancy, the calcium content in wastewater is much higher than in drinking water. As urine contains almost no calcium, its origin should be found elsewhere.

Table 3 summarizes the range of variations of conductivity as well as calcium, potassium, magnesium and sodium concentrations for the four sampling sites. In table 4, the contributions of the different ions to the total conductivity have been roughly estimated based on the average concentrations measured in the four sites in winter and summer. More accurate evaluation will require extended sampling sessions and the present work should assess the real need for. In Nancy, which is the only location investigated in two different seasons, calcium

Table 1.Drinking water characteristics (average concentrations).Tableau 1.Caractéristiques des eaux potables (concentrations moyennes).

Location	Na (mg/L)	K (mg/L)	Mg (mg/L)	Ca (mg/L)	Cl ⁻ (mg/L)	SO ₄ ²⁻ (mg/L)
Nancy	12	2.5	11	21	20	97
Pont-à- Mousson	5.6	4.9	15	83	-	-
Villey-Saint- Etienne	5.7	3.6	12	108	-	-
Colombey- les-Belles	5	1.5	4	130	-	-

Table 2.Comparison of urine and wastewater composition.Tableau 2.Comparaison des compositions de l'urine et des eaux usées

	Concentration (mg/L)	CV%	Range (mg/L)	$\eta_{\rm urine}$	$\eta_{\rm wastewater}$	
$\mathrm{NH_4^+}(\mathrm{mgN/L})$	480	29				
Urea (mgN/L)	7700	20				
$\mathrm{NH_{4\ hyd}^{+}}(\mathrm{mgN/L})$	8180					
Phosphate (mgP/L)	740	14		0.09	0.33	
Calcium (mg/L)	190			0.02	3.3	
Sodium (mg/L)	2800		1800-5800	0.34	2.8	
Potassium (mg/L)	2200		1300-3100	0.27	0.4	
Magnesium (mg/L)	100	21		0.01	0.3	
Chloride (mg/L) 3800			2300-7700	0.46	3.8	
Sulphate (mg/L)	1500	29		0.18	3.3	

Table 3.	Range of variations of conductivity and concentrations in calcium, potassium, sodium and magnesium at
	the four sampling sites.

 Tableau 3.
 Domaine de variations de la conductivité et des concentrations en calcium, potassium, sodium et magnésium

 pour les quatre sites expérimentaux.

	Conductivity (μS/cm)		Calcium (mg/L)		Sodium (mg/L)		Magnesium (mg/L)		Potassium (mg/L)	
	Min	Max	Min	Max	Min	Max	Min	Max	Min	Max
Nancy	750	1450	65	101	51	217	7	9	7	13
Pont-à- Mousson	190	1860	33	136	19	194	2	23	4.7	67
Villey-Saint- Etienne	680	2840	56	306	41	457	9.2	30	12	59
Colombey-les- Belles	900	2380	23	173	45	261	4	9	10	30

Table 4.	Contributions of the various ions to total conductivity.
Tahleau 4	Contributions des différents ions à la conductivité totale

Ion	λ (mS.m ² /mol)	z	Average concentration (mol/m ³)	Average conductivity by element (µS/cm)	Contribution to total conductivity (%)	
Cl-	7.63	1	3.78	28.8	25.9	
Ca ²⁺	5.95	2	2.07	24.6	22.1	
Na ⁺	5.01	1	3.09	15.5	13.9	
SO4 2-	8	2	8.54 10-1	13.7	12.3	
NH_{4}^{+}	7.34	1	1.8	13.2	11.9	
PO ₄ ³⁻	9.28	3	2.68 10-1	7.5	6.7	
Mg^{2+}	5.31	2	3.29 10-1	3.5	3.1	
K^+	7.35	1	0.26	1.9	1.7	
Total					97.6	

varies between 70 and 100 mg/L in winter and between 50 and 70 mg/L in summer (out of rain events which decrease significantly the concentrations by dilution). The major contributors are calcium and chloride. Potassium contributes less than 2%. Figure 5 compares the measured conductivity and the conductivity estimated from the ion concentrations for Villey-Saint-Etienne. In that village, calcium and sodium are the main cations. The large peaks observed at the beginning of the campaign are difficult to explain with respect to the population normal activity (Figure 6). They might be related to the rain event which occurred previously. A clear periodicity can be observed for potassium. A first maximum is observed between 9:00 am and 11:00 am every morning. A second maximum is visible in the afternoon, when inhabitants are coming back from work. For calcium, sodium and magnesium, it is more difficult to figure out the relationship between their variations and the activity of the inhabitants. Obviously the early morning peaks for calcium and magnesium are occurring before the potassium peak. This explains the erratic variations of conductivity seen in figure 3. In Colombey (Figure 6), diurnal variations could be observed for potassium and magnesium,

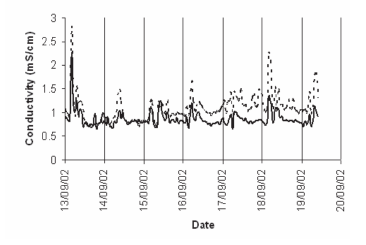
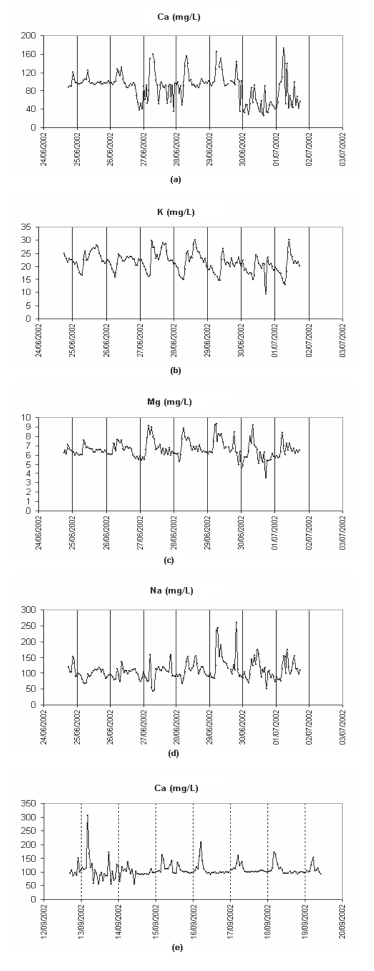


Figure 5. Comparison of measured (- - -) and estimated (----) conductivities for Villey-Saint-Etienne. *Comparaison des conductivités mesurées (---) et estimées (---) pour Villey-Saint-Etienne.*



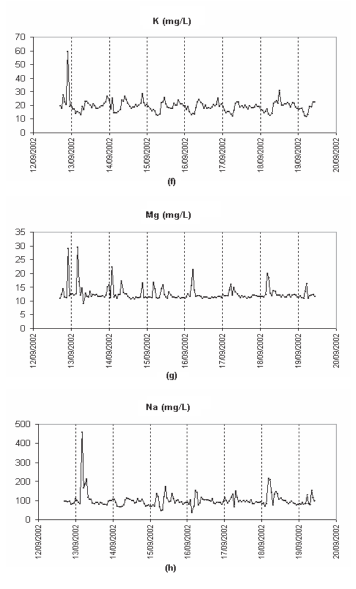
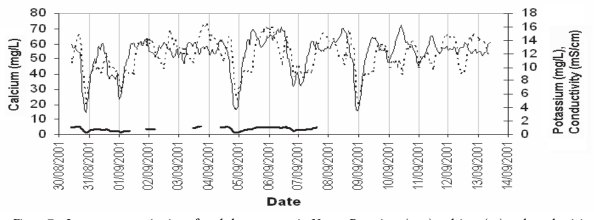
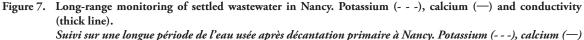


Figure 6. Calcium, potassium, magnesium and sodium: (a), (b), (c), (d) in Colombey-les-Belles; (e), (f), (g), (h) in Villey-Saint-Etienne.
Calcium, potassium, magnésium et sodium : (a), (b), (c), (d) in Colombey-les-Belles; (e), (f), (g), (b) in Villey-Saint-Etienne.

and to some extent for sodium (in spite of two high peaks on the Saturday morning which could be related to the intensive use of laundry machines). The variations of calcium are difficult to explain by anthropogenic reasons. But again, it appears that the early morning peaks for calcium and magnesium occur before the potassium peak.

In order to confirm the out-of-phase variations of potassium and calcium, repeated long-term monitoring sessions of calcium and potassium were performed on the Nancy settled wastewater in summer and winter. One of them is shown in figure 7. Except during rain events when both concentrations drop sharply as well as conductivity, the calcium rises before potassium in the morning. It is not so much potassium which counter-balances calcium in conductivity than ammonia (which





Suivi sur une longue période de l'eau usée après décantation primaire à Nancy. Potassium (- - -), calcium (—, et conductivité (trait épais).

is highly correlated to potassium). As calculated in table 4, sodium, ammonia and ortho-phosphate represent about 32% of the total conductivity. One reason of the presence of calcium at the end of nights could be the corrosion of the mains, made of concrete, in which wastewater is flowing. Concrete contains mainly calcium aluminate, silicate and hydroxide. Depending upon the physico-chemical conditions, calcium can be released into the sewer system. pH is much lower at night, when ammonia, the major contributor to its balance, is minimum. The sewage flow is also minimal and anaerobic conditions favour the reduction by sulphate-reducing bacteria of S-containing substances into H₂S. In the sewer gas phase, H₂S is transformed into sulphuric acid which reacts with the calcium components of concrete (DE BELIE *et al.*, 2004; ROBERTS *et al.*, 2002).

Part of calcium is incorporated in sewer sediment particles such as anapaite, whitlokite (containing also magnesium) and apatite. Apatite has been detected in Nancy sewer system (EL SEMRANI *et al.*, 2004). The soluble fraction of calcium is transported to the wastewater treatment plant. The arrival of the calcium just ahead of the ammonia and potassium in the morning would result from the first-flush effect described by KREBS *et al.* (1999) but at a lower scale as it is governed by the rapid transition between night and day flows. Another hypothesis is related to the possible infiltration into the sewer system of groundwater rich in calcium. For comparison, groundwater quality data have been extracted from the Rhin-Meuse Basin Waterboard database (http://www.eau-rhinmeuse.fr) (Table 5) for piezometers the closest to the sampling

Table 5.Groundwater characteristics near the four sampling sites. Data from the Rhin-Meuse Basin Water Board.
Coefficient of Variation (CV) calculated on 35 data points for Francheville and Colombey-les-Belles (except
for Mg: 33 data points), 6 data points for Tomblaine and 18 data points for Loisy over five years (2000 to
2005).

Tableau 5.Caractéristiques des eaux souterraines près des quatre sites expérimentaux Données de l'Agence de l'Eau
Rhin-Meuse. Coefficient de variation (CV) calculés sur 35 données pour Francheville et Colombey-les-
Belles (sauf Mg: 33 données), 6 données pour Tomblaine et 18 données pour Loisy sur cinq ans (2000 to
2005).

	Franch (5 km fron Saint-En (Dogger li	m Villey- tienne)	Colombey- (Dogger li		Tombla (Greater N (Lower 2 sandsto	Jancy) Frias	Loisy (5 km from Pont-à-Mousson) (Moselle alluvium)		
Depth	65 m		0 1	0 m		708 m		8 m	
	Average	CV (%)	Average	CV (%)	Average	CV (%)	Average	CV (%)	
Conductivity (μ S/cm)	638	15	468	20	1733	10	946	20	
Potassium (mg/L)	4.6	16	0.92	34	14.7	6.4	2	100	
Magnesium (mg/L)	26	12	4.8	100	21	10	10	17	
Calcium (mg/L)	68	8	102	16	84	10	135	12	
Sodium (mg/L)	56	9	6	50	246	18	44	37	

sites. Tomblaine data are probably obtained at a too large depth to be really representative of the situation in the Greater Nancy sewer system. The high conductivity and sodium content at this location are related to the underground salt layer. Sparse data obtained closer to the ground layer give calcium values between 100 and 150 mg/L (personal communication from B. Lartiges, LEM) which could indeed explain the higher values observed in the wastewater samples in the night. The potential calcium flux, estimated from the wastewater flowrate at the inlet of the plant and the calcium data, doubles between night and day (150 (min.) to 300 (max.) kg/h), indicating anyway an anthropogenic source of calcium during the day. For Pont-à-Mousson, the maximal calcium value is close to the groundwater average value, which is in favour of the infiltration hypothesis. In the case of Colombey-les-Belles and Villey-Saint-Etienne, night calcium values are higher than the groundwater values. This leaves open the possible effect of pipe corrosion for these two sites. In conclusion it is difficult to select one hypothesis or the other based on the available data.

4. CONCLUSIONS

Wastewater conductivity has been monitored for extended periods of time in four communities of different size range (from 1,000 to 350,000 PE) and socio-economical profile. In parallel, the concentrations of the main ionic contributors, such as calcium, sodium, potassium, magnesium, ammonium and ortho-phosphate have been measured and their variations with respect to time compared to human activity patterns. It appears that, if sodium, potassium, ammonium and orthophosphate, which contribute to about 34% to wastewater conductivity, exhibit diurnal variations in phase with human activity evaluated by absorbance at 254 nm, calcium (≈ 22% of wastewater conductivity) is out-of-phase. Its release might be related to sewer concrete corrosion or to groundwater infiltration but more experiments should be run to select the proper hypothesis, especially in locations with different groundwater characteristics. The combination of these different ionic patterns creates a conductivity pattern which does not facilitate its relation to a reference human activity that could be used in an automated system to detect abnormalities in wastewater quality. If events such as large rainfalls, which can cause in winter an increase of the conductivity by washout of de-icing salt and during the rest of the year a decrease by dilution, toxic spills linked to the release of ionic species might be difficult to detect.

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