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TOXIC METALS IN URBAN RUNOFF: PHYSICO-CHEMICAL MOBILITY ASSESSMENT USING SPECIATION SCHEMES

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ABSTRACT

Physico-chemical characterization of lead, zinc and cadmium has been carried out on eight samples from both separate and combined sewers. Dissolved and particulate total metal levels have been determined. A speciation scheme has been used to further divide these phases in two dissolved fractions, bioavailable and stable, and five particulate fractions, ion-exchangeable, acid-soluble, reducible, oxidizable and residual. Total dissolved lead concentrations were found to be largely below European Community (EC) directives for drinking water production. Lead was represented by stable forms while zinc and cadmium were mainly bioavailable. Particulate concentrations were higher than natural levels found in the Seine-Normandie basin. Zinc was characteristic for municipal effluents, and lead and cadmium were more concentrated in separate sewers. Correlations have been observed between particulate heavy metals and volatile matter on the ring highway, or with iron at the Savigny site. The particulate metal speciation scheme has demonstrated the important dependence of heavy metal distribution on the sampling site, as well as the potentially more mobile nature of zinc and cadmium, compared to that of lead. The structural modifications undergone by solids during their transport along the sewer networks had a significant impact on particulate metal mobility. This has been clearly noted for lead, whose mobility decreased from upstream to downstream in the studied system.

KEYWORDS

Heavy metals; pollution; impact; storm sewer overflow; combined sewer overflow; detention pond; sediment; speciation.

INTRODUCTION

Originally, only hydraulic impacts of urban runoff were considered and assessed for waste water management. During the last decade, it has been demonstrated that roof runoff (Förster, 1990), as well as highway runoff, contains high pollutant concentrations (Hémain, 1984; Torno et al., 1985). In fact, runoff appears to be the main problem in improving surface water quality in highly urbanised areas (Marsalek, 1991). Sewer suspended solids and sediments are especially interesting since a very large fraction of pollutants is fixed on them (Chebbo, 1992). In most cases, the particulate matter directly reaches surface
waters by combined or separate sewer overflows. There, through transport and settling, they may pose a threat to the aquatic environment. In particular, solids attract toxic metals, i.e. zinc, lead, copper and cadmium, which may be released into the dissolved phase. Mobility and toxicity of these metals must be determined for assessing urban runoff impact on receiving waters and for developing suitable treatment procedures.

Different kinds of experimental approaches have been developed to study metal transport and biogeochemical cycling processes, as well as to predict their biological impact, with acute or long term effect studies (Allen et al., 1980; Nelson and Donkin, 1985). Field measurements of river suspended solids and sediments surrounding sewer overflows give direct indications of the total amount of heavy metal pollution introduced into the receiving system and its accumulation by sedimentation in river beds (Estèbe, 1993). Caged or free bioindicators have also been utilized to determine sewer overflow toxicity (Shutes et al., 1990; Fraboulet et al., 1993). Laboratory assays, like sediment re-suspension experiments in a reactor, can also yield interesting data on the release capacities of polluted particles (Lebreton et al., 1993, Bussy, 1993).

Finally, numerous physico-chemical speciation schemes have been developed to give indications of the heavy metal distribution in sediments and waters (Florence, 1982; Welté and Montiel, 1983). It is well known that exact species determination in aqueous and solid environmental samples is excessively difficult because of the complexity of natural systems, and that heavy metal physico-chemical speciation schemes cannot really extract well defined fractions, because of sample handling mistakes (Kersten and Förstner, 1987), non-specific reactions to particular mineral or organic fractions (Megellati, 1982), or metal redistribution during the fractionation procedure (Ajayi and Vanloon, 1989). However, the data obtained from these schemes are considered as operational (Martin et al., 1987) and provide useful information on the relative mobility and bioavailability of toxic metals (Morisson et al., 1989; Bourg et al., 1992).

This paper presents some of the results obtained during a three year study of storm sewer discharges. The heavy metal speciation scheme included both dissolved and particulate elements (Flores-Rodríguez et al., 1990). The particulate part, slightly modified from Tessier et al. (1979), consisted of five fractions. The dissolved part was simpler, with a unique organic resin extraction (Figura and McDuffie, 1977) as close as possible to the biological membrane uptake. A wide range of situations was reviewed, with pure road runoff, as well as combined sewer overflows, separate sewers, and wet and dry detention ponds.

**SAMPLING SITES**

<table>
<thead>
<tr>
<th>Sample Reference</th>
<th>Sampling Site</th>
<th>Sampling Date</th>
<th>Sewer Type</th>
<th>Catchment Type</th>
<th>Catchment Area (ha)</th>
<th>Rain Intensity (mm/h)</th>
<th>Number of Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Ring highway</td>
<td>May 89</td>
<td>road</td>
<td></td>
<td>3</td>
<td>n.a.</td>
<td>0,9</td>
</tr>
<tr>
<td>2p, 2S</td>
<td>Savigny pond, sewer</td>
<td>Mar 89</td>
<td>S</td>
<td>urban</td>
<td>500</td>
<td>1.3</td>
<td>8,8</td>
</tr>
<tr>
<td>3</td>
<td>Créteil Lake SSO</td>
<td>Sep 89</td>
<td>S</td>
<td>urb+shop</td>
<td>75</td>
<td>n.a.</td>
<td>1,1</td>
</tr>
<tr>
<td>4</td>
<td>La Morée sewer</td>
<td>Mar 89</td>
<td>S</td>
<td>very urban</td>
<td>1,300</td>
<td>1.3</td>
<td>1,1</td>
</tr>
<tr>
<td>5</td>
<td>Béguigneaux pond</td>
<td>Oct 89</td>
<td>S</td>
<td>urb+indust</td>
<td>606</td>
<td>4.4</td>
<td>6,13</td>
</tr>
<tr>
<td>6</td>
<td>La Molette pond</td>
<td>Aug 89</td>
<td>C</td>
<td>urban</td>
<td>8,000</td>
<td>5-13</td>
<td>7,11</td>
</tr>
<tr>
<td>7</td>
<td>La Molette pond</td>
<td>Aug 91</td>
<td>C</td>
<td>urban</td>
<td>8,000</td>
<td>1.2</td>
<td>4,9</td>
</tr>
<tr>
<td>8</td>
<td>Clichy CSO</td>
<td>Jul 91</td>
<td>C</td>
<td>very urb</td>
<td>12,000</td>
<td>n.a.</td>
<td>0,1</td>
</tr>
</tbody>
</table>

n.a. = not available; S = separate; C = combined; Part. = particulate.

Eight sewer flows, following storm events, were studied at seven different sites, most of them near Paris. Table 1 shows the principal site characteristics. The storm outlet of a portion of the Paris ring highway was chosen as a road runoff sample. La Morée separate sewer carries waters from a part of Seine-Saint-Denis
Toxic metals in urban runoff

Laboratory Analyses

Laboratory samples were split into dissolved and particulate phases using a 30 min 6000 g centrifugation (Jouan, model Plasma R 1000), followed by a tangential membrane filtration (Millipore, model Minitan S) using 0.45 μm Millipore Durapore membranes. Total dissolved lead, zinc and cadmium were determined directly by furnace atomic absorption spectrophotometry (AAS) with a HGA 700 Perkin Elmer spectrophotometer and an AS 70 autosampler. The bioavailable fraction was extracted by a chelating resin 24 hour contact. The chosen resin, Chelex 100, had previously been transformed into its calcium form by means of a 1 M CaCl₂ solution as indicated by Fugita and McDuffie (1977). It was estimated to be one of the closest resins to biological membrane uptake, except for S-compounds (Monison et al., 1989). Stable dissolved metals were measured by AAS in the resin-extracted sub-sample supernatants. Bioavailable metals were determined by subtracting stable fractions from total dissolved levels (Flores-Rodríguez et al., 1990).

Total particulate metals were determined by flame or furnace AAS, after an acid digestion by a mixture of concentrated HNO₃ and HClO₄. The sediment samples also underwent a chemical speciation, using a five successive extraction speciation scheme adapted from Tessier et al. (1979) by Flores-Rodríguez (1992). Following the extraction order, these fractions were:

- ion exchangeable, obtained after a 1 M CH₃COONH₄ extraction at pH 7 and room temperature, during 2 hours,
- acid-soluble, obtained after a 1 M CH₃COOH/CH₃COONa extraction, at pH 5 and room temperature, during 2 hours,
- reducible, obtained after a 0.04 M NH₂OH·HCl in 25% CH₃COOH extraction, at pH 2 and 85°C during 5 hours,
- oxidizable, obtained after a 3 hour 30% H₂O₂ in 0.02 M HNO₃ extraction at 96°C, in a water bath,
- residual, obtained after the addition of a HNO₃/HClO₄ concentrated mixture, evaporation to dryness on a hot plate, and a second extraction with 15% HNO₃ under a slight heating.

Results and Discussion

Dissolved Metals

Table 2 displays mean lead, zinc and cadmium concentrations in seven dissolved samples collected during, or just after, storm events. Levels stayed usually below or near surface water EC directives for drinking water production (EC Dir. 80/778). The total dissolved metals appeared to be much smaller than values previously measured in Paris urban area (STU, 1985). However, it may be important to notice that these latter data were obtained with bulk waters firstly acidified and then filtered, allowing possible metal release from suspended solids into the final dissolved samples.

The extraction carried out with the chelating resin revealed that cadmium and zinc were mostly found in bioavailable form in separate and combined water samples, whereas the majority of lead was often stable (Fig.1). The different pattern observed for Créteil lake may be due to its wet detention pond status, as well
as its large surface (40 ha versus 4 ha for Savigny pond) which tends to favour the influence of the groundwater. By comparing sewer metal concentrations to pond levels at Savigny site (samples 2$_S$ and 2$_P$), a purification capacity of this pond for dissolved metals was observed, especially for zinc and lead. For the same storm event as the studied one on Savigny site (but for a larger catchment area), La Monée sewer (sample 4) displayed much higher zinc levels. If total lead and cadmium concentrations were slightly lower, their relative bioavailable fractions were increased. The storm intensity did not modify the total levels of lead and cadmium at La Molette site. However, their bioavailable fractions were more important for a small rain (sample 7) than for a big storm (sample 6). Among the three metals, zinc was the most affected by rainfall intensity, with an observed doubled bioavailable fraction. Finally, Béquigeaux dry detention pond (sample 5) displayed similar patterns to sites in Paris, except for the cadmium level of its dissolved samples, which was strongly increased by industrial inputs.

**TABLE 2 Values of Total Dissolved Metals in Sewer Water Samples**

<table>
<thead>
<tr>
<th>Sample Reference</th>
<th>Dissolved Metal ($\mu$g/l)$^*$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$2_p$</td>
<td>Pb 3.0  Zn 3.5  Cd 0.7</td>
</tr>
<tr>
<td>$2_s$</td>
<td>Pb 17.8  Zn 11.9  Cd 0.9</td>
</tr>
<tr>
<td>3</td>
<td>Pb 3.6  Zn 29.0  Cd 0.4</td>
</tr>
<tr>
<td>4</td>
<td>Pb 12.7  Zn 53.0  Cd 0.8</td>
</tr>
<tr>
<td>5</td>
<td>Pb 9.6±0.6  Zn 14.0±8.0  Cd 1.4±0.4</td>
</tr>
<tr>
<td>6</td>
<td>Pb 6.1±1.9  Zn 28.7±1.1  Cd 0.4±0.1</td>
</tr>
<tr>
<td>7</td>
<td>Pb 6.6±1.6  Zn 56.0±1.4  Cd 0.3±0.3</td>
</tr>
</tbody>
</table>

S.T.U. Measurements$^5$

<table>
<thead>
<tr>
<th>Drinking Water Production (EC dir.)$^5$</th>
</tr>
</thead>
<tbody>
<tr>
<td>200-1 100 500-1 900 5-50</td>
</tr>
</tbody>
</table>

$^5$ from S.T.U. (1985); $^*$ Surface Water Quality EC Directives for Drinking Water Production (80/778)

$^a$ for multiple sample sites, mean ± standard deviation (see text for sample identification)

**Particulate Metals**

The automobile's impact on heavy metal pollution (high lead and cadmium concentrations of runoff suspended solids - sample 1) is clearly demonstrated in Table 3. Another study of the same portion of Paris ring highway, focused on road tunnel aerosols, supports this observation (Lebreton et al., 1992). It showed
higher particulate metal levels than our analysis, as well a significant tendency to leach out, which could be attributed to its lack of water contact. Particulate lead and cadmium are also carried down to Créteil Lake SSO (sample 3) from a catchment area which principally consists of an important shopping centre and its extensive car park areas, the latter exerting an important influence on sewer effluent quality. A battery production plant was identified in the Béquigneaux detention pond catchment area (sample 5) which explains the high levels of cadmium and zinc found in this sample. Savigny and La Moëve separate sewers (samples 2 and 4) showed lower heavy metal concentrations than combined sewer samples taken from the same area. The combined sewer sample levels did not vary significantly with catchment area (Clichy - sample 8), with a concentration very similar to La Moëve, but for a much larger drained surface. Particulate lead and cadmium concentrations were similar for separate and combined samples. By contrast, zinc levels were notably more important on combined sewer sites, as well as less variable among the storm events. A study of French combined sewage metal composition (Colin, 1986) has shown that zinc is the heavy metal which is the most important in flow coming into a sewage plant, by at least a factor 4. Thus, when a notable storm event occurred, runoff solids, drained by La Moëve combined sewer, produced a dilution of the particulate zinc concentration normally found in the municipal sewage effluent.

### TABLE 3 Total Particulate Metal Concentrations in Samples and Associated Pollution Factors

<table>
<thead>
<tr>
<th>Sample</th>
<th>Particulate Metals (g/kg d.w.)</th>
<th>Pollution Factor</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Pb</td>
<td>Zn</td>
</tr>
<tr>
<td>1</td>
<td>2.36</td>
<td>1.11</td>
</tr>
<tr>
<td>2</td>
<td>0.23</td>
<td>0.38</td>
</tr>
<tr>
<td>2p</td>
<td>0.05</td>
<td>0.11</td>
</tr>
<tr>
<td>3</td>
<td>3.18</td>
<td>1.76</td>
</tr>
<tr>
<td>4</td>
<td>0.96</td>
<td>0.77</td>
</tr>
<tr>
<td>5</td>
<td>0.46</td>
<td>3.85</td>
</tr>
<tr>
<td>6</td>
<td>0.95</td>
<td>2.03</td>
</tr>
<tr>
<td>7</td>
<td>0.36</td>
<td>2.67</td>
</tr>
<tr>
<td>8</td>
<td>1.40</td>
<td>3.08</td>
</tr>
</tbody>
</table>

AFNOR regulation\(^{o}\) | 0.30 | 3.00 | 0.015 | n.a. |
Natural levels\(^{x}\) | 0.03 | 0.10 | <0.001 | 13  |
Circ. highway aerosol\(^{+}\) | 4.8  | 2.33 | 0.027 | 50  |
Down. Paris Seine sed.\(^{*}\) | 0.28 | 0.68 | 0.009 | 19  |

\(^{o}\) French maximum limits for sewage plant sludge agriculture recycling (AFNOR U44-041/1975),
\(^{x}\) Pereira-Ramos (1989), \(^{+}\) Lebrelon and Thévenot (1992) and \(^{*}\) Estèbe et al. (1993).
(see text for sample identification)

Finally, most of our samples reach the river Seine. Considering the natural levels of Seine basin sediments, which have been statistically calculated (Pereira-Ramos, 1989 - Table 3), our results demonstrated that urban runoff and CSO outfalls may have a very negative influence on receiving water bodies. Thus, it should be useful to have suspended solid separation treatment before the sewage reaches the river Seine. A recent study of the Seine sediment first layer, on a site located downstream from Paris and its main combined sewer outfalls supported this, showing great similarity to our samples and a significant increase of natural heavy metal levels (Estèbe et al., 1993). They imply a significant sewage impact on river sediment quality. Measured metal concentrations, particularly lead, often exceed allowed limits for agricultural use of solid waste (AFNOR U44-041/1975). If such solids were removed, using for instance dry detention ponds, the resulting sludge could not always be recycled. However, Savigny wet detention pond showed real purifying capacity, if effluent levels are compared with levels found in its connected sewer (samples 2p and 2s). Local water authorities have defined a pollution factor for sediments, by dividing measured metal concentrations by calculated natural levels (Pereira-Ramos, 1989). This factor allowed us to more easily evaluate sediment heavy metal pollution, since natural soil concentration may vary significantly, depending on the site (Robbe, 1982). The right part of Table 3 displays pollution factors for lead, zinc, cadmium and iron for all our
"Parasite" samples. One principal characteristic of separate sewer waters should be noted: the significant variability of their quality, which has been the argument for not treating these waters. Pollution factors calculated for combined sewers are more homogenous since they only vary from 12 to 47, while the separate samples vary from 4 to 106 (except Savigny pond, which can be considered as unpolluted). Another remarkable fact is that iron levels were roughly constant among the samples, with a maximum pollution factor of 2. We can conclude that natural levels of this metal were found in our samples. This major soil characteristic can be considered as typical in the Seine basin, independent from the general heavy metal pollution level.

This property has allowed us to use iron as a natural mineral fraction tracer. At the same time, we chose the volatile matter ratio as the organic matter tracer, since it is generally utilized to measure the organic matter content of sewage effluents (AFNOR T90-0291/1970). By plotting correlations between these two parameters and the heavy metal concentrations, we wanted to roughly distinguish the origin of metallic pollutants or, at least, their transport mode. These correlations are illustrated for zinc in Figure 2. The relations between sites are clearly displayed on the graphs. Similar conclusions were made for lead and cadmium. Créteil sediment (sample 3) characteristics were consistent with the ring highway (sample 1) results, probably because of the numerous parking areas included in the catchment area. La Morée (sample 4) was closer to Savigny (sample 2), which is located in the same county. The small rainfall we studied at La Molette pond (sample 7) presented some analogies with Clichy outfall (sample 8), but during an important storm event La Molette samples were more similar to the separate sewer samples collected in the same county at the Savigny and La Morée sites. The non-homogeneous reparation of solids in a dry detention pond was clearly notable within La Molette pond. The weak intensity of the small rainfall increased the phenomenon of selective settling which induces sampling representativity problems.

![Figure 2. Correlations between total particulate zinc and either total particulate iron, or volatile matter](image)

Sites: O: 1, x: 2, *: 3, #: 4, +: 5, O: 6, f: 7, X: 8
(see text for sample identification)

Linear regressions were done on sample sites when possible. Correlation coefficients are listed in Table 4. Significant correlations were only found on two sites. Solids sampled in the runoff detention chamber of Paris ring highway (sample 1) indicated a strong link between heavy metal concentrations and volatile matter content. By contrast, Savigny pond (sample 2) metal levels showed a strong correlation with the level of particulate iron. The first site, typically representative of automobile pollution, may concentrate pollutants from organic materials like tyres, oils, etc. These products have also been listed by Morrison (1985) as important heavy metal sources. On the second site, sediment samples showed a very strong correlation between iron and the three studied elements (correlation coefficients ranged from 0.83 to 0.95). This may be explained by the fact that location of the catchment area, outside Paris and not very urban, favours the presence of soil particles rather than pure road runoff solids. Tessier et al. (1982) have also observed important correlations between iron and heavy metals for river sediments, as well as Breilhante (1990), for different types of solids. The relationship to volatile matter, although not as good as for the ring highway,
was notable. This last revealed the impact of road runoff on the quality of these samples. However, the mineral nature of the source, or at least of the transport mode of heavy metals on this site, is more important. Although Béquigneaux (sample 5) is a separate sewer site like the two former, its sediments did not show good correlation with the two studied parameters. Particularly, lead did not show a relationship with iron, even though it is strongly bound to inorganic particles (Audlitia and Pickering, 1987). The industrial outfall may create this difference by modifying deeply the sample characteristics.

La Molette large storm (sample 6) showed some links between heavy metals and iron, as well as volatile matter, like it was observed for Savigny pond. The small rainfall (sample 7), which occurred on the same site, presented the lowest correlation coefficient for both parameters. Thus, the nature and transport of the metallic pollutants in combined sewers is strongly dependent on storm event duration. A heavy rainfall gave results closer to separate sewer ones (like Savigny samples) than did small events. This difference between the storms at La Molette was verified by calculating the regression coefficients for all the samples which were collected on this site. We generally obtained lower values, except for lead (which links with iron), which was greater for both events.

<table>
<thead>
<tr>
<th>Site Reference</th>
<th>Parameter</th>
<th>Pb</th>
<th>Zn</th>
<th>Cd</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 (n=9)</td>
<td>VM</td>
<td>0.77</td>
<td>0.86</td>
<td>0.63</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.49</td>
<td>0.34</td>
<td>0.01</td>
</tr>
<tr>
<td>2 (n=8)</td>
<td>VM</td>
<td>0.55</td>
<td>0.65</td>
<td>0.57</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.92</td>
<td>0.95</td>
<td>0.83</td>
</tr>
<tr>
<td>5 (n=13)</td>
<td>VM</td>
<td>0.12</td>
<td>0.06</td>
<td>0.06</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.09</td>
<td>0.63</td>
<td>0.80</td>
</tr>
<tr>
<td>6 (n=11)</td>
<td>VM</td>
<td>0.48</td>
<td>0.26</td>
<td>0.44</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.49</td>
<td>0.49</td>
<td>0.62</td>
</tr>
<tr>
<td>7 (n=9)</td>
<td>VM</td>
<td>0.0003</td>
<td>0.38</td>
<td>0.89</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.54</td>
<td>0.19</td>
<td>0.00002</td>
</tr>
<tr>
<td>6+7 (n=20)</td>
<td>VM</td>
<td>0.008</td>
<td>0.18</td>
<td>0.26</td>
</tr>
<tr>
<td></td>
<td>Fe</td>
<td>0.70</td>
<td>0.02</td>
<td>0.43</td>
</tr>
</tbody>
</table>

(see text for sample identification)

As these solids are discharged directly to the rivers, it was interesting to evaluate the portion of the particulate heavy metals which could be released to pollute the natural environment. Total metal cannot provide this kind of indication. With these sequential chemical extractions, physico-chemical sediment speciation may offer more information. We used this process upon some particulate samples. Figure 3 displays the distribution of heavy metals among the five fractions of the chosen scheme. Regardless of the sites or the metals, the most important fractions were the acid-soluble and reducible fractions, if the residual fraction was not taken into account. This latter introduced greater differences for lead and iron than for zinc and cadmium. Thus, these two latter heavy metals showed similar distributions, while the lead pattern appeared to be closer to iron. This may be linked to the good correlations usually found between the total particulate concentrations of these two elements. Moreover, the large particulate lead residual fraction implied that its stable nature may render it essentially non-toxic for the environment. In 1979, Wilber et al. also found this metal mostly in residual forms in runoff waters. The ring highway (sample 1) was the only site where samples containing important ion-exchangeable fractions for all the metals were found. This agrees with previous work carried out on the same site, but with tunnel road aerosols (Lebreton and Thévenot, 1992), and with the study of Revitt et al. (1987), which found that particulate lead in road runoff
and gully pot outflow was mainly present in the ion-exchangeable fraction. In addition to this fraction, we found heavy metals in acid-soluble and reducible forms in our road runoff sample. However, the correlations previously plotted for this site displayed notable links between heavy metals and volatile matter, which is related to organic matter, destroyed with the oxidizable fraction.

This discrepancy may be due to the reagent used for the extraction, based on the breaking down of humic acids/metal bonds, while heavy metals were more probably adsorbed on different organic matter with weaker chemical bonds. The Créteil Lake SSO (sample 3) was not as close to the ring highway sample as the correlations indicated. Particulate heavy metals were extracted primarily in more stable fractions. This could be explained by the sewer sample, which allowed an in-pipe evolution of sediments. The result differed from those obtained with longer separate sewers (samples 2 and 4), probably because of the short travel time.

A comparison of metal distributions in these samples, particularly with lead, indicates their progressive stabilization from upstream (road runoff) to downstream (the sewer system). The significant runoff processes in the larger catchment areas implied a greater suspended solids transport. These solids tend to be rich in
humic organic matter, which may form more stable chemical bonds than with runoff from small catchment areas like Créteil. Finally, it is important to remark that for substantial metal concentrations, the largest fractions were most easily released (i.e. Créteil and Béquigneaux), while less concentrated samples had a large residual fraction.

CONCLUSION

Concentrations of dissolved lead, zinc and cadmium in separate and combined samples were generally below EEC water quality directives for sources of drinking water. In some cases, zinc and cadmium exceeded these values. Moreover, they were mostly found in bioavailable form, whereas lead was more stable. Particulate heavy metal concentrations varied over a wide range. All were much higher than natural levels calculated for the Seine-Normandie basin, and for measured Seine sediment levels, and higher than French regulations for residual metals in sludge used for agricultural recycling. Lead and cadmium were mainly found in runoff, but zinc was concentrated in municipal effluents diluted at the final outfall. Few correlations could be found between iron and volatile matter, except for very specific sites. Dry detention ponds presented very different results because of a kind of selective settlement which occurred during and after storms. It appeared that the structural modifications undergone by solids during their transport along the sewer networks had a significant effect on particulate heavy metal distributions (and their related mobilities). These were also very site-dependent. Runoff from roads and car parks showed significant metal mobility. Particulate pollutants were transformed during travel through the sewage network to more stable forms found in the detention ponds. This was particularly notable for lead, whose exchangeable forms disappeared, and whose residual fraction increased.

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