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Removal of emerging micropollutants from wastewater by activated carbon adsorption:
experimental study of different activated carbons and factors influencing the adsorption of
micropollutants in wastewater

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HIGHLIGHTS

- BET surface and bulk density of activated carbons are correlated to micropollutants adsorption,
- The activated carbon dose and the contact time have a great influence on adsorption,
- FeCl₃ has a positive influence on micropollutants adsorption by activated carbon (+10-15%),
- Both the quantity and composition of organic matter impact the adsorption,
- UV-254 removals are correlated to several micropollutants removals.

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ABSTRACT

Activated carbon processes, initially designed for drinking water production, are tested for wastewater application in order to characterize their efficiency to remove micropollutants from wastewater treatment plants (WWTPs) discharges. In that purpose, a pilot was set up at the Seine Centre WWTP and is studied by the Paris sanitation service (SIAAP) and the Water Environment and Urban Systems laboratory (LEESU). The in-situ study raised several additional questions related to the structural and morphological properties of activated carbons, in order to select the proper material, the influence of operational parameters such as the activated carbon dose and the contact time, the role of organic matter concentration and composition, the presence of a residual concentration of methanol or the impact of ferric chloride addition. Thus, various complementary experiments were carried out at laboratory scale to improve the understanding of the micropollutants adsorption process on activated carbon, in particular on powdered activated carbon (PAC).

The results have highlighted a strong link between the efficiency of PACs and their specific surface (BET), which can be easily estimated by their bulk density. The study of the sorption process has also confirmed the strong influence of the PAC dose and the rapidity of the sorption kinetic. From an operational point of view, the ferric chloride injection seems to slightly improve most of the detected compounds adsorption, probably thanks to the coagulation of the dissolved organic matter colloidal fraction. In contrary, the presence in the water of a residual concentration of methanol seems to have no impact on the pharmaceuticals fate. The influence of the wastewater matrix where adsorption is performed is strong, with notably lower adsorption in settled water compared to various WWTP discharges. However, the dissolved organic carbon concentration is not always sufficient to explain sorption competitions in wastewater, and the composition and the distribution of the organic matter should be considered too. In particular, the carbon removal from biological treatments is the step that clearly modifies both the quantity and the composition of the organic matter. It has been observed that discharges from WWTPs operating with different biological processes (activated sludge, membrane bioreactor or biofiltration) have similar organic matter concentrations and compositions, and allows comparable removals of organic matter and micropollutants by adsorption. The lower performances on micropollutants observed in the settled water can be explained by the higher quantity of small protein-like molecules (fluorophores δ and γ), which compose the most competitive organic matter fraction for adsorption on activated carbon, compared to the other waters.

KEYWORDS
Adsorption; organic matter; pharmaceuticals; activated carbon; wastewater

INTRODUCTION

The presence of a large range of emerging micropollutants, particularly pharmaceuticals and hormones, personal care products or pesticides, has been highlighted in wastewater treatment plant (WWTP) discharges (Miège et al. 2009; Verlicchi et al. 2012; Loos et al. 2013; Mailler et al. 2015). Even if several hydrophobic, volatile or biodegradable compounds are strongly removed by conventional wastewater treatments (Ruel et al. 2012; Mailler et al. 2014b), most of micropollutants are not removed. Thus, various strategies of contamination reduction are assessed by the scientific community and water treatment engineers. Among them, the implementation of tertiary treatments dedicated to micropollutants elimination represents a relevant solution. In particular, adsorption on activated carbon appears to be efficient, very flexible and relatively cheap (Abegglen & Siegrist 2012), together with not producing oxidation by-products.

In this context, the Parisian public sanitation service (SIAAP) and the Water Environment and Urban Systems laboratory (LEESU) study, in collaboration with SAUR teams, the CarboPlus® process. This process is based on a fluidized bed of activated carbon (powder - PAC - or micro-grain - μGAC) to remove micropollutants by adsorption. A large-scale pilot based on this technology was set up at the Seine Centre WWTP (240 000 m³/days) to characterize the efficiency of activated carbon to remove a wide range of pollutants from WWTP discharges. The in-situ results are presented in (Mailler et al. 2015) for PAC.

In parallel to the in situ study, complementary laboratory scale experiments were conducted to i) better understand the relationships between activated carbon properties and the adsorption in wastewater and ii) to improve the understanding of the micropollutant adsorption mechanisms in wastewater application, in particular with PAC.

Activated carbon is characterized by different structural and morphological properties that can affect adsorption (Baudu et al. 2001; Li et al. 2002; Yu et al. 2008; Delgado et al. 2012). Thus, the first axis consists in studying the relationships between activated carbon properties and their efficiency to remove micropollutants. In particular, a focus is performed on the specific BET surface and the bulk density.

The second axis consists in studying the adsorption mechanism in wastewater representing a complex matrix. Indeed, the
literature highlighted that the matrix where adsorption is performed plays a crucial role in the fate of molecules (Ebrie et al. 2001; Ternes et al. 2002; de Ridder et al. 2010; de Ridder et al. 2011), particularly the organic matter which competes with pollutants for adsorption through direct sites competition or pore blocking. In addition, most of the studies were conducted for drinking water production or in surface waters, and the adsorption mechanisms were poorly assessed in wastewater. Thus, the influences of both the activated carbon dose and the contact time on the performances were assessed, as well as the influence of the organic matter and chemicals that can be injected within the process for operational purposes.

This article summarizes the results from the laboratory scale experiments conducted within this project. First, the relationships between micropollutants removals and activated carbon properties are assessed. Then, the influence of the dose of carbon, the contact time, the organic matter quantity and quality, the presence of a residual concentration of methanol and the injection of FeCl₃ were studied.

MATERIALS AND METHODS

1. Activated carbon characterization

11 adsorbents, including micro and mesoporous PACs and μGAC, have been selected based on their technical datasheets provided by the producers. Among them, 3 are commercialized by DaCarb® (PB 170®, PB 170-400® and PC 1000® - France), 3 by Chemviron® (WP 235®, Carbsorb 28® and Cyclecarb 305® - Belgium), 2 by Norit® (W 35® and SA Super® - Netherlands) and 3 by Jacobi® (LP 39®, MP 25® and Hydro XP 17® - Sweden). These activated carbons are recognized for their high organic pollutants affinity, but their use in wastewater was poorly studied in the literature.

Several structural properties such as the specific BET surface (m²/g), the porous volume (mL/g), the pore size distribution and the bulk density (g/cm³) have been measured on the 11 activated carbons. In addition, the particle size distribution and the micropollutant elimination have been determined for 4 of them: PB 170®, WP 235®, W 35® and PC 1000®.

These structural properties have been analyzed at the University of Technology of Compiègne (UTC, France). The specific BET surface and the pore size distribution were measured with an ASAP 2010 Micromeritics analyzer equipped
with a degasing station and a gas isotherm adsorption analyses station (nitrogen), according to the conventional methods used to determine these parameters. Fresh activated carbon samples (100 mg) were degased at least 12 h before to be analyzed. Results correspond to the specific surface in m²/g obtained with the BET (Brunauer, Emett and Teller) method.

The pore size distribution is determined with the BJH (Barrett, Joyner and Halenda) method, using the desorption curve of the same gas on the same analyzer. This method allows also calculating the microporous and mesoporous pore volumes. The bulk density was measured by weighting, with a high precision balance (0.01 mg), 50 to 100 mL of activated carbon, measured with a 100 mL graduated cylinder. The activated carbon is introduced by small doses (10 mg) and is compacted every 10 mL to minimize the vacuum between particles.

The particles size distribution of the 4 PACs was measured with a Mastersizer 2000 Malvern laser particle sizer. Every analysis corresponds to 15 000 light diffraction measures. 3 scans were performed at least per sample.

2. Pollutants and analytical procedures

For every sample, several parameters were measured in the dissolved phase: UV absorbance at 254 nm (UV-254), dissolved organic carbon (DOC) and concentrations of 16 pharmaceuticals and 2 pesticides (list and limit of quantification in supporting material - Table S1). Organic micropollutants were analyzed by the Institute of Analytical Sciences (ISA - Villeurbanne, France), while UV-254 and DOC were measured by the SIAAP laboratory (French accreditation - Colombes, France). Analytical protocols are validated and are given in (Mailler et al. 2015).

3D fluorescence spectrometry analyses were also performed by the LEESU. They allow generating a tridimensional matrix and to plot it on a 3D spectrum containing all excitation and emission spectra of the dissolved organic matter (DOM) present in a sample. This method gives the distribution of the different components of the DOM as they don’t fluoresce in the same zones of the spectrum. The detailed description of 3D fluorescence spectrometry is given in (Hudson et al. 2007). Analyses have been performed with a Jasco FP-8300 spectrofluorometer equipped with a 1 cm quartz cell.

3. Laboratory experiments protocols

The laboratory experiments have been performed in the Seine Centre WWTP (Colombes, France) between April 2013 and May 2015. Wastewater was punctually sampled from the WWTP discharges with 10 L glass bottles, properly rinsed and grilled, and used the same day for experiments. Then, samples were filtered on glass fibers filters (GF/F 0.7 µm) after
experiments, before to be sent to laboratories for analyses.

3.1. Efficiency comparison of 4 PACs

The efficiency of 4 PACs (PB 170®, WP 235®, W 35® and PC 1000® - Table 1) to remove micropollutants was assessed, by contacting individually 10 mg/L of each PAC with 1 L of Seine Centre WWTP (SEC) discharges during 45 min under strong mixing. After filtering, samples were conditioned and sent to ISA laboratory to analyze pharmaceuticals and pesticides.

3.2. PAC dose and adsorption kinetic

Regarding its performances, PB 170® was selected to characterize the relationship between PAC dose, contact time and micropollutant removal. First, 10 mg/L of this PAC were contacted with 1 L of SEC discharges under strong mixing during different contact times (5 - 10 - 30 - 45 - 60 min), which are close from the pilot contact time range. Then, 3 PAC doses (5 - 10 - 20 mg/L) were tested with 1 L of SEC discharges and during 45 min.

3.3. DOM and adsorption

The DOM has been identified to play a crucial role in the adsorption process because it induces a competition with the organic micropollutants for adsorption (Matsui et al. 2003; de Ridder et al. 2011; Delgado et al. 2012). These phenomena have been principally observed with surface water and natural organic matter, but some recent studies also studied it in wastewater (Margot et al. 2013; Altmann et al. 2014). In this context, it is necessary to determine the role of both the quantity and the quality of the DOM in the adsorption process. To this end, 7 types of wastewater featuring by different levels of organic matter (Table 1) have been contacted under strong mixing with PAC (10 mg/L of PB 170® during 45 min). Among the tested waters, 4 were WWTP discharges, from Seine Centre (SEC), Seine Aval (SAV), Seine Morée (SEM) and Seine Amont (SAM) WWTPs. The 3 other tested waters were outlet waters from the physico-chemical lamellar settling unit (SEC settled water) and the carbon biofiltration effluent (carbon removal) of SEC, as well as SEC settled water diluted with distilled water to reach a DOC level comparable to the WWTP discharges. The layouts of the different studied WWTPs are given in supporting material - Table S1, as well as the sampling points (red circles). The organic matter characteristics of these waters are different and given in Table 1. Among the 7 wastewaters, the
micropollutants adsorption has been assessed with SEC settled water, SEC carbon biofiltration effluent, SEC and SAV discharges.

A 3D fluorescence spectrometry characterization has also been performed on these 7 waters, before and after contact with PAC. The spectra interpretation follows the fluorophores defined in (Parlanti et al. 2002), as indicated for SEC settled water and discharges in supporting material - Figure S2. \( \alpha \), \( \alpha' \) and \( \beta \) correspond to humic-like substances, \( \gamma \) and \( \delta \) respectively represents tyrosine-like and tryptophan-like proteins. Then, \( I_1 \), \( I_2 \), \( I_3 \) and \( I_4 \) indexes, respectively corresponding to the ratios \( \alpha'/\alpha \), \( \beta/\alpha \), \( \gamma/\alpha \) and \( \delta/\alpha \), were calculated to evaluate the distribution. Finally, the humification index (HIX) and the biological activity index (BIX) were also evaluated (Zsolnay et al. 1999; Parlanti et al. 2002). The different fluorophores, HIX and BIX are given in Table 1 for the 7 tested wastewaters.

### Table 1 - The 7 studied wastewaters and their main DOM characteristics

<table>
<thead>
<tr>
<th>Biological treatment</th>
<th>SEC settled water</th>
<th>SEC settled water - diluted</th>
<th>SEC carbon biofiltration effluent</th>
<th>SEC discharges</th>
<th>SAM discharges</th>
<th>SAV discharges</th>
<th>SEM discharges</th>
</tr>
</thead>
<tbody>
<tr>
<td>TOC (mgC/L)</td>
<td>32</td>
<td>7.8</td>
<td>-</td>
<td>7.3</td>
<td>7.2</td>
<td>-</td>
<td>5.5</td>
</tr>
<tr>
<td>DOC (mgC/L)</td>
<td>27</td>
<td>7.7</td>
<td>6.6</td>
<td>6.8</td>
<td>6.6</td>
<td>11</td>
<td>5.2</td>
</tr>
<tr>
<td>UV-254 (cm(^{-1}))</td>
<td>0.859</td>
<td>0.203</td>
<td>0.120</td>
<td>0.150</td>
<td>0.173</td>
<td>0.237</td>
<td>0.140</td>
</tr>
<tr>
<td>Fluo ( \alpha ) (UA)</td>
<td>1 879</td>
<td>455</td>
<td>812</td>
<td>1 522</td>
<td>822</td>
<td>2 097</td>
<td>701</td>
</tr>
<tr>
<td>Fluo ( \alpha' ) (UA)</td>
<td>2 019</td>
<td>442</td>
<td>741</td>
<td>1 262</td>
<td>712</td>
<td>2 041</td>
<td>730</td>
</tr>
<tr>
<td>Fluo ( \beta ) (UA)</td>
<td>1 926</td>
<td>440</td>
<td>742</td>
<td>1 371</td>
<td>761</td>
<td>2 018</td>
<td>657</td>
</tr>
<tr>
<td>Fluo ( \delta ) (UA)</td>
<td>4 949</td>
<td>1 080</td>
<td>665</td>
<td>1 348</td>
<td>678</td>
<td>1 685</td>
<td>622</td>
</tr>
<tr>
<td>Fluo ( \gamma ) (UA)</td>
<td>4 194</td>
<td>907</td>
<td>354</td>
<td>1 077</td>
<td>448</td>
<td>590</td>
<td>231</td>
</tr>
<tr>
<td>HIX</td>
<td>1.0</td>
<td>1.2</td>
<td>3.4</td>
<td>3.2</td>
<td>3.6</td>
<td>5.0</td>
<td>6.4</td>
</tr>
<tr>
<td>BIX</td>
<td>0.85</td>
<td>0.90</td>
<td>0.97</td>
<td>0.96</td>
<td>0.99</td>
<td>0.95</td>
<td>0.96</td>
</tr>
</tbody>
</table>

SEC = Seine Centre WWTP; SAM = Seine Amont WWTP; SAV = Seine Aval; SEM = Seine Morée.
BF = biofiltration; CAS = conventional activated sludge; MBR = membrane bioreactor: C = carbon removal; N = nitrogen removal (nitrification + denitrification).
DOC = dissolved organic carbon, UV-254 = UV absorbance at 254 nm, Fluo = fluorophore, HIX = humification index, BIX = biological activity index.

### 3.4. Influence of the residual methanol concentration in the WWTP discharges

A residual methanol concentration is present in both SEC and SAV discharges because of the post-denitrification stage.

Indeed, denitrifying microorganisms needs a carbon source and when the denitrification is performed after the carbon removal and the nitrification, as in both SEC and SAV, the carbon content is too low (Rocher et al. 2012). Methanol is
then added during this stage as carbon source, but its consumption by the microorganisms is not complete. Operationally speaking, it is interesting to study the impact of this residual concentration on the micropollutant adsorption by activated carbon. Thus, the performances obtained with the SEC discharges have been compared to performances obtained with the same water spiked with methanol (6 and 12 mgC/L). The adsorbability of methanol has also been studied and preliminary tests demonstrated that no adsorption of methanol occurs when contacting 12 mgC/L of methanol in ultra-pure water with 10 mgPAC/L.

3.5. Influence of the ferric chloride

The CarboPlus® process operating in PAC configuration requires the injection of FeCl$_3$ to stabilize the PAC bed and prevent any leakage. This substance is known to destabilize particles by neutralization of the surface charges, which could impact the adsorption of micropollutants. To evaluate the impact of FeCl$_3$, 1 L of SEC discharges have been contacted with PAC alone (10 mg/L), FeCl$_3$ alone (2.5 gFeCl$_3$/m$^3$) and both PAC and FeCl$_3$ under strong mixing during 45 min.

RESULTS AND DISCUSSION

1. Influence of PAC characteristics on its efficiency

Structural and morphological properties of the 4 tested PACs are given in the Table 2. The removals of the micropollutants with a PAC dose of 10 mg/L during 45 min are also provided.

<table>
<thead>
<tr>
<th>Table 2 - Characterization and performances of the 4 PACs studied</th>
</tr>
</thead>
<tbody>
<tr>
<td>Producer</td>
</tr>
<tr>
<td>-----------</td>
</tr>
<tr>
<td>Raw material</td>
</tr>
<tr>
<td>Bulk density (g/cm$^3$)</td>
</tr>
<tr>
<td>Specific BET surface (m$^2$/g)</td>
</tr>
<tr>
<td>Pore size distribution (micro, meso, macroporous)</td>
</tr>
<tr>
<td>Pore volume (micro + meso) (mL/g)</td>
</tr>
<tr>
<td>Particle size distribution (µm)</td>
</tr>
</tbody>
</table>
Among the 4 PACs, the PB 170® and WP 235® have a similar mesoporous structure with close pore size distribution and specific BET surface (900-1000 m²/g), while W 35® and especially PC 1000® have different pore size distributions and lower specific BET surfaces (supporting material - Figure S3; Table 2). W 35® is strongly mesoporous with 45% of the BET surface resulting from mesopores, while PC 1000® is microporous with 59% of the surface resulting from micropores. 3 of the tested PACs have similar bulk densities (0.30-0.40 g/cm³) while PC 1000® is heavier (0.54 g/cm³). Regarding the particle size distribution, PB 170® and WP 235® are comparable while the two other PACs have a larger particle size distribution and a higher median diameter. This is particularly notable for the PC 1000® which is characterized by median and d90 values twice higher than those of the PB 170® and WP 235®.

Even if only 1 punctual experiment was performed, a clear trend can be observed with most of the compounds. For individual compounds, the PB 170® allows the highest removals for 15 of the 18 micropollutants and the PC 1000® is always the worst efficient. The WP 235® and W 35® seem to have similar performances, slightly weaker than PB 170®. Overall, the sum of the 16 pharmaceuticals is removed by 61%, 53%, 51% and 32% respectively by the PB 170®, WP 235®, W 35® and PC 1000®.

The micropollutant removal is well correlated with the specific BET surface (Figure 1): the higher the specific BET surface, the higher the micropollutant removal. In addition, the specific BET surface is also correlated to the bulk density of the PAC, a low bulk density corresponding to a high BET surface. This relationship has been verified with 7 other PACs differing by their nature, as the bulk density is very easy to measure and could therefore be a proxy of the BET surface. Considering the 11 PAC, this link is still observed (Figure 1). Thus, the bulk density could be used as an
indicator to select activated carbons and estimate their BET surface and efficiency for micropollutants removal from wastewater.

The link between the efficiency of activated carbons and their specific BET surface (600-1500 m²/g) has already been observed in the literature (Çeçen & Aktas 2012), but the distribution of this surface between the different categories of pores (micro < 2 nm, meso 2-50 nm and macropores > 50 nm) has also to be considered. This is particularly important in presence of organic matter because it competes with micropollutants for adsorption, directly or by pore blocking (Newcombe et al. 2002; Yu et al. 2008). Thus, a mesoporous structure with a large pore size distribution should limit the competition with the DOM (Ebie et al. 2001).
Figure 1 - Correlations between the specific BET surface and the micropollutant removals obtained with a PAC, and link with its bulk density

2. Adsorption of micropollutants on PAC

2.1. Influence of fresh PAC dose and adsorption kinetic

The Figure 2 displays the influence of the contact time and the PAC dose on the removal of micropollutants from Seine Centre discharges by PAC.

According to the results (Figure 2), a higher dose of PAC logically results in higher micropollutant removals, i.e. carbamazepine (32-54-80%; removals at 5-10-20 mgPAC/L), diclofenac (22-36-60%), propranolol (56-81-95%) or sulfamethoxazole (51-61-80%). However, the relationship is not linear as doubling the PAC mass doesn’t result in a doubling of the removals. Actually, the relationship is logarithmic resulting in the reaching of a plateau when increasing...
the dose, as displayed in supporting material - Figure S4. Thus, the PAC dose in wastewater has a great influence on the removal, as already observed in the literature (Snyder et al. 2007; Boehler et al. 2012; Margot et al. 2013) or at large scale with the CarboPlus® pilot (Mailler et al. 2015). Considering the individual results of the micropollutants, this positive influence of the dose is statistically significant (test of Shapiro-Wilk, Fischer and Student - p-value < 0.05) between 5 and 10 mgPAC/L (p-value = 0.016) and between 10 and 20 mgPAC/L (p-value = 0.0005).

Furthermore, the removals notably increase with the contact time (Figure 2), i.e. ciprofloxacin (29-40-62-66-71%; removals at 5-10-30-45-60 min contact times) or diclofenac (19-23-32-36-39%). However, the PAC is very fine (< 50 µm), resulting in a relatively fast adsorption kinetic. As a consequence, although the equilibrium is not reached after 60 min, differences of removal between 30, 45 and 60 min contact time for a same PAC dose are limited (< 10%) indicating the proximity of the equilibrium. Thus, the short contact times (30-60 min) employed in PAC tertiary treatments are suitable for an efficient adsorption of micropollutants. For several compounds (8/15), such as atrazine (26-46%; removals after 5 and 60 min), norfloxacin (38-71%), ofloxacin (39-64%) or sulfamethoxazole (40-55%), the removal achieved after 5 min represents more than half of the removal after 60 min, confirming the very fast kinetic. In contrary, erythromycin (7-39%) or roxithromycin (3-38%) seem to have slower kinetics, probably because of their high molecular weight. These compounds will then be more sensitive to changes of contact time in adsorption processes.

2.2. Influence of the matrix

2.2.1. Dissolved organic carbon

The Figure 3 displays the removals of micropollutants observed with different types of wastewater, sorted from the highest to the lowest DOC value.
Figure 3 - Pharmaceuticals removals by PAC adsorption from different types of wastewater

Removals observed with the SEC settled water are overall lower than those from the three other wastewaters. In particular, they are significantly lower (p-value = 0.001) than in the SEC discharges (water from the same WWTP), considering all the compounds displayed on Figure 3. Comparable removals are overall achieved with the 3 other waters, which have undergone an intense biological carbon treatment. For instance, the carbamazepine is removed at 18% in SEC settled water, against 38 to 42% in the other matrixes. Similarly, the trimethoprim removal increases from 5% in settled water to 39-50% in the other waters.

These results indicate that DOC value has a negative impact on micropollutant adsorption, as the removals observed in the SEC settled water, featuring the highest DOC concentration (27 mgC/L), are the lowest compare to the other waters. The negative influence of DOC on micropollutant adsorption from wastewater was also observed in the literature (de Ridder et al. 2011; Margot et al. 2013; Altmann et al. 2014). In contrary, DOC value seems to have a limited impact on propranolol behavior, a compound which is easily adsorbed by activated carbon as positively charged (Mailler et al. 2015).

According to what is previously stated, micropollutant removals should be lower in SAV discharges compared to those
in the SEC carbon biofiltration effluent and discharges, regarding the differences of DOC (11 vs. 6.6–6.8 mgC/L). This is observed for bezafibrate, lorazepam and metronidazole. However, removals are similar for the other compounds, in spite of slightly higher influent concentrations in the SAV discharges. Besides, considering the individual removals of compounds from Figure 3, the removals from SEC and SAV discharges are significantly similar (p-value = 0.175). The organic matter of SAV discharges should then be less competitive than organic matter from other discharges, what counterbalances the higher DOC.

This result indicates that the sole DOC level isn’t sufficient to explain the variations of removals from water to water, the organic matter composition and nature have also to be considered. Considering that, the DOM nature and adsorption was assessed before and after contact with PAC for these 4 waters, as well as 3 additional wastewater matrixes.

2.2.2. Composition and adsorption of the organic matter

Figure 4 displays the composition (indexes from 3D fluorescence analysis) of the organic matter from 7 wastewater matrixes, as well as the organic matter removal by PAC adsorption (10 mgPAC/L - 45 min).
First of all, the 4 WWTP discharges (SEC, SAM, SEM and SAV) and the SEC carbon biofiltration effluent have a similar distribution of DOM with a large fraction of humic-like organic matter (peaks $\alpha$, $\alpha'$ and $\beta$; indexes I1 and I2), and a smaller fraction of organic matter resulting from bacterial activity ($\gamma$ and $\delta$; I3 and I4), as shown on Figure 4 and in Table
1. In contrary, the DOM distribution is different in the SEC settled water with very high I3 and I4 indexes, characterizing protein molecules, compared to I1 and I2 indexes, characterizing the humic-like substances. This indicates that carbon biological treatments significantly remove DOC and fluorophores δ and γ (see SEC settled water and carbon biofiltration effluent in Table 1), resulting in a significant modification of the DOM distribution (Figure 4). Together with this difference of DOM composition, the SEC settled water has higher DOC (27 mgC/L) and UV-254 (0.859 cm⁻¹) values, highlighting a higher quantity of DOM.

In addition, the HIX increases along the different steps of wastewater treatment, as shown by the low HIX in the SEC settled water (1.0) and the higher HIX after the biological treatments (> 3). This suggests that small protein molecules (δ and γ) are transformed into humic-like substances (α, α’ and β) during biological treatments, explaining the HIX increase. The HIX from SAM and SEC discharges, as well as SEC carbon biofiltration effluent, are similar (3.2-3.6), while those from SAV and SEM discharges are higher (5.0-6.4), resulting from higher hydraulic retention times.

Regarding the removal of DOM by activated carbon, DOC and UV-254 removals are notably higher in the SEC settled water compared to the other ones. This is consistent as DOC and UV-254 are much higher in this water. This suggests that in this water, less sites will be available for micropollutants as more DOM is sorbed, what is consistent with micropollutants results (Figure 3). For the other wastewaters, the DOM removal is rather comparable (10-25% for DOC and 5-10% for UV-254).

In addition, fluorophores removals are rather similar in all wastewaters except in the SEC settled water. In the WWTP discharges, all the fluorophores are removed between 10 and 30%. In contrary, in SEC settled water, fluorophores δ and γ are predominant (I3 and I4) and are better removed (35-50%) than the other fluorophores. They are also better removed than in the other water. In consequence, these fluorophores will have the highest impact on the competition with micropollutants for adsorption. This is consistent with the study of (de Ridder et al. 2011) which identified the small size DOM as the most competing ones for adsorption. Moreover, the fluorophores removal pattern is the same in SEC diluted settled water, confirming that the distribution of DOM is as important in the adsorption process as the DOC value.

In this context, and considering that the carbon biological treatment significantly reduces the DOC and the quantity of fluorophores δ and γ (the most competing DOM), the lower removals of micropollutants by PAC in the SEC settled water compared to the other waters are then consistent. Indeed, the competition is stronger in this water resulting from i) a higher quantity of DOM (DOC and UV-254) and ii) a different DOM distribution, with more small protein-like substances and less humic-like substances. In addition, I3 and I4 are lower in the SAV discharges than in SEC carbon biofiltration
effluent and discharges, together with a higher HIX. This means that this water has less very competitive DOM what
counterbalances the higher DOC, resulting in similar micropollutants removals.

It can be concluded that, besides a low DOC, low I3 and I4 indexes, together with a high HIX, are positive for the
minimization of DOM competition during the micropollutants adsorption.

2.2.3. Presence and concentration of methanol

The micropollutant and organic matter removals obtained in presence of a residual concentration of methanol are
presented in Figure 5.

![Figure 5 - Impact of a residual concentration of methanol in the WWTP discharges on the adsorption of
pharmaceuticals by PAC](image)

The comparison of removals obtained in the carbon biofiltration effluent and in the SEC discharges allows assessing the
impact of the methanol presence in the wastewater on the PAC adsorption (Figure 5). Its presence doesn’t decrease the
performances as the removals observed in the SEC discharges are similar or higher than those in the carbon biofiltration
The better removals in the SEC discharges could be explained i) by the methanol itself, and ii) by the carbon and nitrogen treatment achieved in the second and third biofiltration stages. but further experiments are needed to bring an answer. Moreover, no clear trend is observed between the increase of the residual concentration of methanol and the micropollutant removals, except for sulfamethoxazole and naproxen for which the removal decrease and increase respectively. However, the results indicate that the increase of the methanol concentration doesn’t lead to a significant degradation of the PAC performances for most compounds.

Thus, results have shown that the residual concentration of methanol doesn’t affect negatively the removal of micropollutants by activated carbon (Figure 5). In contrary, its presence in biological treatment effluents could favor a biological activity in the tertiary treatments by activated carbon.

2.2. Influence of the injection of FeCl₃

The ferric chloride influence on the fate of micropollutants in presence of PAC is illustrated on figure 7.
Three types of behaviors are observed for micropollutants. For most compounds, such as carbamazepine, beta blockers or diclofenac, and for DOM (DOC and UV-254), a slightly higher removal (+ 5-15%) is obtained by coupling PAC and ferric chloride, probably by coagulation of the colloidal fraction. These differences are not significant taking uncertainties into account, but 10 compounds in 13, as well as DOC and UV-254, have a greater removal in presence of FeCl₃. In contrary, no clear influence is observed for lorazepam and bezafibrate. Finally, sulfamethoxazole seems to be the only one with a lower removal (- 30%) in presence of ferric chloride.

(Margot et al. 2013) also observed a positive effect of the coagulant on micropollutant removals. However, the mechanism at the origin of this improvement is not identified. Supplementary tests are required to determine if coagulation, complexation or direct decrease of the competition with organic matter are involved.

CONCLUSIONS

The Parisian public sanitation service (SIAAP) and the Water Environment and Urban Systems laboratory (LEESU) study the CarboPlus® process at large scale, in collaboration with the SAUR teams. In parallel to the study of the large scale pilot, complementary laboratory scale experiments were carried out to improve the understanding of the micropollutants adsorption on activated carbon, in particular PAC.

These experiments allowed displaying the links between structural properties of activated carbons and their efficiency to remove micropollutants. In particular, it has been shown that the PAC performances are closely related to the specific BET surface, which can easily be estimated through the bulk density measure.

The strong influence of the PAC dose has also been confirmed, as well as the suitability of the adsorption kinetic (less than one hour) for WWTP application (operational requirements). In an operational point of view, the injection of FeCl₃ doesn’t disrupt the adsorption process. A slight improvement is even observed for most compounds (+ 10-15%), except for sulfamethoxazole, probably through coagulation of the colloidal fraction of the DOM. In contrary, the residual concentration of methanol present in the WWTP discharges due to the post denitrification process seems to have no impact on the fate of most pharmaceuticals.
Then, this study highlighted the differences of micropollutants adsorption in the various wastewater matrixes. The DOC concentration is important considering the competition of adsorption but it isn’t always sufficient to explain the performances reductions from water to water. It has been shown that the distribution of the different molecular components of the DOM should be considered in addition to the quantity of DOM. Indeed, small protein-like fluorophores have been identified as the most problematic for adsorption competition in wastewater, while a high HIX, highlighting the humification of the DOM, is favorable to micropollutants adsorption. The reduction of protein-like substances in wastewater is then a way to maximize the adsorption of micropollutants by activated carbon.

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