First scientific support plan for a sustainable development policy (SPSD I)

Programme “Sustainable management of the North Sea”

Development of methods for the analysis of hydrocarbons and organic micro-pollutants in the marine environment

Summary of the research

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Introduction

Since few years, significant advances have been realised in the field of dioxins analysis. The combined use of new extraction [1] and clean-up [2,3] techniques as well as latest developments in the area of mass spectrometry [4] made the development of integrated strategies possible. The reference methods still require labor intensive sample preparation as well as high-cost high resolution mass spectrometers (HRMS) only available in some well specialized laboratories limiting the number of samples within an acceptable cost for monitoring [5]. In order to be compatible with higher sample throughput, operating procedures (protocols) have to be simplified or alternative methods should be validated. In that field, the leading ideas are 1) fast, 2) screening and 3) low cost.

As potential tools for screening method, biological assays (based on antibodies or cells response) focused attention during last few years with the emergence of a battery of bio and immuno-assays presenting advantages and drawbacks [6]. The pros of these assays are not only the low cost (generally 5 times lower than classical HRMS) but also the possibility of parallel processing of samples. However, since the assays can also be activated by other chemicals present in the mixture in often higher concentration than the analytes of interest, sample preparation steps are still required to reduce the risks of false positives and they become the bottleneck of the procedure. In addition they involve delicate solvent exchanges, due to the need of performing the assays in aqueous-type media [7]. Cross-reactivity of the various congeners (based on 2,3,7,8-TCDD) can significantly alter the TEF values. Knowing that final TEQ estimation for many matrices mainly rest on the relative contribution of few congeners, these disparities regarding to the TEF can introduce uncertainties on the TEQ estimation.

On the side of the development of these biological methods, advances in physico-chemical analysis tools have also reach an interesting level [8]. It was therefore worthwhile evaluating their potentialities in term of screening for dioxins on a selected congeners basis [9].

Discussion

From reviewing recent available literature concerning congeners distribution in food and human matrices, some tendencies can be outlined. It appears that for the hundreds of computed samples, 1,2,3,7,8-PeCDD and 2,3,4,7,8-PeCDF are the major contributors to the WHO-TEQ (Table 1). These relative contributions are quite constant over matrices types.

Table 1 : Relative contributions (%) of selected penta congeners.

<table>
<thead>
<tr>
<th>Food</th>
<th>Mean 1,2,3,7,8-PeCDD</th>
<th>Mean 2,3,4,7,8-PeCDF</th>
<th>Sum SD Range</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Food</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Beef</td>
<td>26</td>
<td>45</td>
<td>71 3 [ 68-74 ]</td>
<td>10-12</td>
</tr>
<tr>
<td>Veal</td>
<td>39</td>
<td>35</td>
<td>74 n.a. n.a.</td>
<td>10</td>
</tr>
<tr>
<td>Pork</td>
<td>39</td>
<td>36</td>
<td>75 11 [ 68-83 ]</td>
<td>10-12</td>
</tr>
<tr>
<td>Lamb</td>
<td>47</td>
<td>34</td>
<td>77 9 [ 71-84 ]</td>
<td>10,12</td>
</tr>
<tr>
<td>Horse</td>
<td>29</td>
<td>36</td>
<td>65 9 [ 56-73 ]</td>
<td>10-12</td>
</tr>
<tr>
<td>Chicken</td>
<td>21</td>
<td>46</td>
<td>67 2 [ 65-69 ]</td>
<td>10-12</td>
</tr>
<tr>
<td>eggs</td>
<td>22</td>
<td>43</td>
<td>65 n.a. n.a.</td>
<td>11</td>
</tr>
<tr>
<td>cheese</td>
<td>23</td>
<td>50</td>
<td>73 n.a. n.a.</td>
<td>11</td>
</tr>
<tr>
<td>Creme</td>
<td>62</td>
<td>16</td>
<td>78 n.a. n.a.</td>
<td>12</td>
</tr>
<tr>
<td>Butter</td>
<td>19</td>
<td>59</td>
<td>78 1 [ 77-78 ]</td>
<td>11,12</td>
</tr>
<tr>
<td>Milk</td>
<td>25</td>
<td>43</td>
<td>68 10 [ 51-78 ]</td>
<td>11-18</td>
</tr>
<tr>
<td>Prawn</td>
<td>13</td>
<td>42</td>
<td>55 n.a. n.a.</td>
<td>12</td>
</tr>
<tr>
<td>Trout</td>
<td>16</td>
<td>41</td>
<td>57 n.a. n.a.</td>
<td>12</td>
</tr>
<tr>
<td>Salmon</td>
<td>21</td>
<td>41</td>
<td>62 n.a. n.a.</td>
<td>19</td>
</tr>
<tr>
<td>Mackerel</td>
<td>13</td>
<td>30</td>
<td>43 1 [ 42-44 ]</td>
<td>11,15</td>
</tr>
<tr>
<td>Herring</td>
<td>28</td>
<td>46</td>
<td>74 n.a. n.a.</td>
<td>15</td>
</tr>
<tr>
<td>Plaice</td>
<td>24</td>
<td>49</td>
<td>73 n.a. n.a.</td>
<td>15</td>
</tr>
<tr>
<td>Rice</td>
<td>26</td>
<td>11</td>
<td>37 n.a. n.a.</td>
<td>20</td>
</tr>
<tr>
<td>Badley</td>
<td>28</td>
<td>19</td>
<td>47 n.a. n.a.</td>
<td>20</td>
</tr>
<tr>
<td>Bean</td>
<td>32</td>
<td>8</td>
<td>40 n.a. n.a.</td>
<td>20</td>
</tr>
<tr>
<td>Spinach</td>
<td>36</td>
<td>22</td>
<td>58 6 [ 54-62 ]</td>
<td>21</td>
</tr>
<tr>
<td>Human</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood</td>
<td>29</td>
<td>26</td>
<td>55 10 [ 37-70 ]</td>
<td>11,22-37</td>
</tr>
<tr>
<td>Breast milk</td>
<td>32</td>
<td>31</td>
<td>63 11 [ 45-78 ]</td>
<td>13,23,24,31,32,38</td>
</tr>
</tbody>
</table>
The standard deviations are relatively low for all types of matrices, indicating that these contributions are representative of the general situation in good proportions. This observation has different consequences, 1) it is of prime necessity for a screening tool to be able to consider these two congeners as accurately as possible, 2) these two congeners can be use as "screening congeners" for first sorting out of samples before relevant HRMS analysis. Since biological tools sometimes have relative response factors (especially for these two congeners) which does not match with the WHO-TEF values [39] and that these assays only give a global response, another tool would be appropriate. Knowing that even for assays, relevant clean-up methodology is required, it is quite conceivable to use GC to separate the congeners of interest and MS sensitive detector for screening. Recent improvement in the quadrupole ion storage tandem mass spectrometers (QISTMS) sensitivity present them as valuable detector [40,41,42]. They are easy to operate, their cost is acceptable, they have low picogram detection limits in isotopic dilution mode and they permit recovery rates calculations without any standards compatibility problems. Using adequate parameters and after time compression of the GC run, analyses can be carried out very rapidly. Fig. 1 illustrates an example of fast run (cycle time of less than 10 min) obtained with a benchtop ion trap mass spectrometer coupled to a classical GC system (A:1,2,3,7,8-PeCDF, B:2,3,4,7,8-PeCDF, C:1,2,3,7,8-PeCDD).

In order to avoid false-negative production, a safe screening condition is to use the mean value for the sum of the two PeCDD/Fs as representative value for each type of matrix with a confidence of 20 %. Calculations from the obtained quantities would then allow the evaluation of the total WHO-TEQ and eventually the complementary analysis on HRMS. A crucial point here is that the samples which need further injection in HRMS are already available and does not require any additional preparation before injection, greatly improving the overall speed and cost of the process. Considering the global control of the screening approach, as in the case of assays [43], certain amount (10%) of declared negative samples can systematically be confirmed by HRMS. This approach is currently under investigations to evaluate its robustness and it's already clear that such a strategy would not yields to the production of greater amounts of false negatives than other screening processes.

In addition to a fast GC-MS analysis time, automated clean-up allows the preparation of large number of samples in parallel in a short amount of time. The global process considering milk samples for example can be as fast as few hours for batches of 10 samples, also allowing isolation of the PCBs and persistent pesticides fractions.

We plan to adapt the strategy to marine matrices. The clean-up step is of course the key step on which efforts have to be made since most of the screening cost results of that. A promising alternative being for us the use of disposable solid phase extraction (SPE) pre-packed cartridges that can easily be combined to produce clean extracts [44]. The
optimization as well as the transposition of this to automated systems using the new 96-well SPE technology for high sample throughput preparation would then also really be adapted to micro-plates bio-assays screening capabilities for the analysis of many samples in parallel.

Conclusions
Screening capability is one of the most wanted criteria for large number samples analysis in order to reduce the time spent to process samples containing negligible analyte levels. However, since the analysis of trace levels of dioxins require complex clean-up procedure, high sample throughput biological tools are currently not exploited at their optimum level. The production of such a number of samples has to include simple and/or automated processes which then produce extracts presenting levels of cleanness compatible with the GC. The approach suggested here rests on the screening out of negative samples, before expensive GC-HRMS analysis, using quantification of selected representative congeners isolated by automated clean-up and analyzed by FGC-QISTMS. This method is versatile, the “screening congeners” are still representative in different contamination types (TCDD can be added if necessary) and the correlation between their concentration and the TEQ is easier than in the case of marker PCBs analysis for dioxins levels evaluation.

This strategy can be seen as a cost effective “dioxin-dedicated” physico-chemical screening method complementary to a powerful biological tool capable of estimating the total toxicity of complex mixtures of large numbers of different halogenated aromatic hydrocarbons contained in samples.

Acknowledgements
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