An uncertainty and sensitivity analysis applied to the prioritisation of pharmaceuticals as surface water contaminants from wastewater treatment plant direct emissions

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ABSTRACT

In this study, the concentration probability distributions of 82 pharmaceutical compounds detected in the effluents of 179 European wastewater treatment plants were computed and inserted into a multimedia fate model. The comparative ecotoxicological impact of the direct emission of these compounds from wastewater treatment plants on freshwater ecosystems, based on a potentially affected fraction (PAF) of species approach, was assessed to rank compounds based on priority. As many pharmaceuticals are acids or bases, the multimedia fate model accounts for regressions to estimate pH-dependent fate parameters. An uncertainty analysis was performed by means of Monte Carlo analysis, which included the uncertainty of fate and ecotoxicity model input variables, as well as the spatial variability of landscape characteristics on the European continental scale. Several pharmaceutical compounds were identified as being of greatest concern, including 7 analgesics/anti-inflammatories, 3 β-blockers, 3 psychiatric drugs, and 1 each of 6 other therapeutic classes. The fate and impact modelling relied extensively on estimated data, given that most of these compounds have little or no experimental fate or ecotoxicity data available, as well as a limited reported occurrence in effluents. The contribution of estimated model input variables to the variance of freshwater ecotoxicity impact, as well as the lack of experimental abiotic degradation data for most compounds, helped in establishing priorities for further testing. Generally, the effluent concentration and the ecotoxicity effect factor were the model input variables with the most significant effect on the uncertainty of output results.

Keywords: Pharmaceuticals, Multimedia fate model, Freshwater ecotoxicity, Ionisable organics, Uncertainty analysis, Wastewater treatment plants

1. Introduction

The presence of pharmaceuticals in the environment and their potential to induce adverse biological effects have been known for many years (Aherne and Briggs, 1989; Tabak and Bunch, 1970). The most common environmental contamination pathways are the emission of pharmaceutical compounds from wastewater treatment plants (WWTPs) and the application of livestock manure as a top soil dressing (without previous wastewater treatment). Livestock manure contains veterinary drugs that are likely to contaminate the soil and groundwater, which, after rainfall incidents, can reach surface waters from contaminated soil by run-off. The main sources of emission for these environmental contamination pathways are the urinal and faecal excretion products of medically treated human and animals. Other less important sources of contamination include industrial wastewater and drugs disposed of with domestic waste in landfill sites, which...
could lead to groundwater contamination by leaching (Ternes, 1998). The pathways of contamination after excretion and passage through municipal sewage systems include the infiltration of sewage from leakages in drains, the application of biosolids from WWTPs on agricultural areas and landscapes, and, due to incomplete removal, the disposal of WWTP effluents and raw sewage into surface waters and as reclaimed water into agricultural fields and landscapes by irrigation. Regarding these emission pathways from WWTPs, we distinguish between direct and indirect emissions to the freshwater compartment. The application of biosolids and effluents into agricultural soils and landscapes can also lead to the migration of contaminants to surface waters via run-off (Borgman and Chefetz, 2013; Sabourin et al., 2009); therefore, such emissions are defined here as indirect emissions to freshwater.

Although much research has been conducted on the topic of direct emissions of pharmaceuticals from WWTPs, past studies examining the prioritisation of pharmaceuticals (e.g., Besse and Garric, 2008; Christen et al., 2010; Sanderson et al., 2004) do not account for spatial variations of the environmental landscape, or include a comprehensive uncertainty and sensitivity analysis of the results when most of fate and impact data are estimated, nor do most of them account for the ionising properties of pharmaceuticals when most of these compounds are acids or bases.

The aim of this study is to prioritise pharmaceutical compounds from WWTP direct emissions in their impact to freshwater ecosystems, identifying gaps of knowledge and relevant fate and impact issues in order to establish topics for further research. To provide a holistic view of the pharmaceuticals of greatest concern, we collected data concerning pharmaceutical occurrence in 179 WWTPs in Europe. A multimedia model representing the European continental scale was applied to prioritise pharmaceuticals according to their probabilistic impact on freshwater ecosystems, computed by means of Monte Carlo analysis, from WWTP direct emissions. Generally, experimental fate input variables, such as partitioning coefficients or degradation rates, and ecotoxicity data are scarce for most pharmaceuticals; therefore, estimation methods must be applied in an assessment. Research topics on monitoring in WWTP effluents, degradation in the environment or in ecotoxicology effects were prioritised for the compounds of most concern by indentifying important gaps of knowledge, as well as by computing the contribution of estimated model input variables’ uncertainty and variability to the impact variance. Currently, a similar assessment is being performed for indirect emissions to the freshwater compartment.

The multimedia model USEtox (Rosenbaum et al., 2008) was chosen as the basis for this comparative impact assessment because it results from a consensus building effort, under the auspices of UNEP and SETAC, amongst modellers; hence, its underlying principles reflect common and agreed recommendations from these experts. In comparative impact assessment methodologies, the conversion of emissions to ecotoxicological impacts comprises a fate and an effect analysis step (van Zelm et al., 2007). The fate factor describes the marginal increase in environmental concentration per unit of emission. The ecotoxicity effect factor (EEF) addresses the marginal increase in effect (toxic pressure on ecosystems) per unit of chemical concentration. An assessment factor (AF) based on the predicted no effect concentration (PNEC) approach is recommended in generic risk assessment according to the TGD (EC, 2003); however, a potentially affected fraction (PAF) of species approach based on the average toxicity was considered in the present study as a basis for the EEF calculation, as adopted in the USEtox model. Both approaches have advantages and drawbacks (Larsen and Hauschild, 2007a,b); however, a PAF-based approach has two main advantages that better serve the purposes of this study: 1) a PNEC approach targets the protection of the most sensitive species; therefore, the risk of bias is high when scarce ecotoxicity data are available, which is the case for pharmaceuticals; and 2) the assessment of the mean impact (AMI) on ecosystem method, a PAF-based approach, allows the quantification of uncertainty, giving an indication of the
reliability of the results. The AMI method is based on the hazardous concentration (HC) at which the effect concentration (with an endpoint of, for example, mortality) affecting 50% of tested individuals (EC50) is exceeded for 50% of the included species; this is also called HC50EC50 (Payet, 2004, 2005; Payet and Jolliet, 2005). Two statistical estimators can be used to estimate the toxicity of a substance to biological species and the associated confidence interval: a non-parametric estimator using the median as the HC50EC50 combined with bootstrap statistics to estimate its uncertainty (Payet and Jolliet, 2005) or a parametric estimator based on the assumption of a lognormal distribution of data using the geometric mean as HC50EC50 and Student’s t-statistics for its confidence interval (Payet, 2004, 2005).

2. Methodology

2.1. Emission data

A survey of the occurrence of pharmaceuticals in the effluents of European WWTPs was performed to compute concentration probability distributions. The survey is based on a recent review conducted by Verlicchi et al. (2012) on the global occurrence of pharmaceuticals in urban wastewater. For this Europe-focused study, 54 peer-reviewed publications were identified from the cited review covering 179 WWTPs located in Austria, Denmark, Finland, France, Germany, Greece, Italy, Spain, Sweden, Switzerland, and the UK, with capacities ranging from 6000 to 2500000 population equivalents. Effluent concentration data included 82 drugs pertaining to 15 different classes: 19 analgesics/anti-inflammatory (including 1 metabolite), 15 antibiotics, 12 β-blockers, 7 psychiatric drugs, 7 lipid regulators (including 2 metabolites), 4 hormones, 4 β-agonists, 3 receptor antagonists, 3 anti-neoplastics, 2 antihypertensives, 2 diuretics, 1 proton-pump inhibitor, 1 antiseptic, 1 contrast agent, and 1 antifungal (Supplementary data, Table S3). The quality of effluent concentration data reported in the literature has been confirmed according to the EC Technical Guidance Document (TGD) on Risk Assessment (EC, 2003). Therefore, the references included in the survey feature a description of the analytical methodology and the quality assurance programme used for sampling, analysis and elaboration. Considering only WWTPs with data available on population served (number of the inhabitants in the catchment), the effluent concentration in each WWTP was weighted by the population served and the geometric mean and the geometric standard deviation of the effluent concentration, in mg/l, in European WWTPs were computed. Aggregated data on a compound concentration in several WWTPs effluents were weighted using the aggregated data on population served. The probability distribution, assuming a lognormal distribution, of the effluent concentration of each compound was used as input into a multimedia fate and transport model, assuming steady-state concentrations, to assess the comparative impact to freshwater ecosystems.

2.2. Fate modelling

Fate factors describing the marginal increase in environmental concentration of pharmaceuticals per unit of emission were computed by a model based on the multimedia model USEtox (Rosenbaum et al., 2008) and described in detail in Morais et al. (2013a,b). The difference between models refers to the inclusion of regressions to estimate pH-dependent fate parameters if no suitable experimental values are available, such as the solid-water partitioning coefficient normalised by the organic carbon, KOC (Franco and Trapp, 2008) and the bioconcentration factor in fish, BCFFish (Fu et al., 2009). Over 60% of pharmaceuticals are acids or bases that are fully or partially dissociated at environmental pH (Avdeef, 2003); hence, conventional non-polar regressions cannot be applied without considering the ionisation of pharmaceuticals (Escher et al., 2011; Tarazona et al., 2010). For the environmental compartments evaluated, the landscape characteristics of the USEtox European continental scale were applied. The fate model accounts...
for inter-media transport processes, intramedia partitioning and degradation in the environment (Fig. 1) and is further described in Supplementary data, sections S1 and S2.

Abiotic degradation mechanisms in the freshwater compartment are important elimination processes for pharmaceuticals (Andreozzi et al., 2003; Doll and Frimmel, 2003). However, the USEtox model does not address estimation procedures for these mechanisms; therefore, to estimate direct and indirect photodegradation rates, a number of models and assumptions were applied and are described in detail in Morais et al. (2013a,b). A short description is provided in Supplementary data, section S2.

2.3. Ecotoxicity effect factor

The EEF indicator, i.e., $0.5/\text{HC50}_{\text{EC50}}$, in $\text{PAF m}^3 \text{kg}^{-1}$, focuses on the trophic structure by including the EC50 values of at least 3 trophic levels: primary producers (algae), primary consumers (crustaceans), and secondary consumers (fish) (Supplementary data, Table S5). The low environmental concentrations but constant introduction to the environment indicate that pharmaceuticals are more likely to have chronic rather than acute toxic effects on aquatic biota (Carlsson et al., 2006; Fent et al., 2006; Quinn et al., 2008); hence, chronic EC50 values are preferred as well as, due to the comparative context of the assessment, standard tests/test conditions and standard test organisms. However, the ecotoxicological data on pharmaceuticals remain scarce, and there are not enough chronic experimental ecotoxicity data available to perform an assessment (Escher et al., 2011). An acute–chronic ratio of 2 was applied to extrapolate chronic HC50 EC50 values from acute HC50 EC50 values, as recommended by Larsen and Hauschild (2007b), and was applied in the USEtox model (Huijbregts et al., 2010). However, the best estimate AFs for this extrapolation have not yet been developed, and research is needed in this area (Larsen and Hauschild, 2007b), particularly in the context of micropollutants. Even acute ecotoxicity data are only available for a very limited set of pharmaceuti- cals (Escher et al., 2011); therefore, EC50 values are completed by extrapolation from the lowest observed effect concentration (LOEC) or no observed effect concentration (NOEC) values, according to the best-estimate AFs from Payet (2004). To determine missing experimen- tal data, quantitative–structure activity relationship (QSAR) data were included using the software programme ECOSAR v1.00 (Nabholz and Mayo-Bean, 2009). For estimated data, a conservative approach was followed by considering the chemical class with the highest potency (i.e., the lowest concentration predicted to cause the toxic effect).
except in the case of the neutral organics class if a compound is completely dissociated at environmentally relevant pH values.

The baseline toxicity, or narcosis, is the addressed toxic mode of action (TMoA) in most generic risk assessment or impact assessment methodologies. Previous studies have shown that most pharmaceuticals produce their environmental adverse effect via narcosis (Sanderson and Thomsen, 2007). However, some pharmaceuticals, which are designed to be bioactive (with the exception of contrast agents), also exhibit a therapeutic effect in non-target aquatic life, such as the estrogenic effects caused by hormones in fish (Santos et al., 2010), or they act via a specific TMoA, such as the inhibition of photosynthesis caused by β-blockers in algae (Escher et al., 2006). As a change in sex ratio apparently relates directly to the reproduction of a fish population, this endpoint is considered more relevant than vitellogenin in an impact assessment context (Larsen et al., 2010). Hence, the endpoints used for the average toxicity calculation include the inhibition of growth and photosynthesis for algae, mortality or immobility (Daphnia) for invertebrates, and mortality or change in sex ratio for fish.

2.4. Uncertainty and sensitivity analysis

The propagation of the uncertainty and variability of model input variables in the output results was quantified by Monte Carlo analysis. The model output is the ecotoxicity impact on freshwater, in PAF•m$^3$•d. The parameters included in the analysis are described in the Supplementary data, Table S6. The analysis includes the following factors:

1. The variability of effluent concentrations, direct photolysis rates ($k_{\text{photo, fw}}$) and continental-scale environmental parameters (freshwater pH, rainfall, freshwater concentration of suspended matter, dissolved organic carbon, and $^1$OH). For each pharmaceutical, the maximum and minimum of the uniformly distributed photodegradation rate parameter were obtained considering results for the winter and summer seasons, the latitudes of 40 and 60, and experimental quantum yields (see Supplementary data, section S2)

2. The uncertainty of the EEF. Uncertainty distributions of HC50EC50 values were estimated according to the parametric estimator, as recommended by Payet (2004). Moreover, the parametric estimator is based on the geometric mean, which is the most robust average estimator for HC50EC50 (Larsen and Hauschild, 2007b). However, the uncertainty of extrapolating average chronic
toxicity, i.e., chronic HC50EC50, from average acute toxicity was not addressed in the present study, nor was the uncertainty of extrapolating and estimating individual endpoints.

(3) The uncertainties associated with the regression equations adopted in the model to estimate partition coefficients (KOC and KOW), bioconcentration factors in fish (BCFfish) and bio-transformation rates (kbio, fw). The procedure to compute the uncertainty descriptors of regressions equations is described in detail in Morais et al. (2013a, b). In short, the training and validation sets used to derive the regression methods applied in the present study (Franco and Trapp, 2008; Fu et al., 2009; USEPA, 2008a, 2009) were used to derive mean residual errors and their uniformal distributions and were fit into the regressions.

(4) The uncertainty associated with experimental parameter values (partition coefficients, biotransformation half-lives, and kOH). The geometric mean and the geometric standard deviation of experimental values were set as uncertainty descriptors, assuming a lognormal distribution.

The identification of relevant parameters to the impact variance, performed by a sensitivity analysis, enables setting research priorities. The contribution to the variance provides an approximation of the percentage of the variance or uncertainty of an output result caused by the variability or uncertainty of a given model parameter. The contribution was calculated by squaring the correlation coefficients between model input variables and impact results, for a given number of trials, and normalising them to 100%.

3. Results and discussion

Fig. 2 shows the comparative ecotoxicological impact of pharmaceuticals emitted directly from WWTP effluents to the freshwater compartment. In order to rank compounds for further discussion the contribution of each pharmaceutical's uncertainty to the variance of total ecotoxicity was calculated. A general condition for impact assessment methodology is that the impact indicator be additive (Larsen and Hauschild, 2007b); however antagonistic–synergistic interactions in mixtures of pharmaceuticals are not accounted for in such approaches. The total ecotoxicity impact is $6.51 \times 10^{-2}$ PAF m$^3$ d per m$^3$ of effluent (95% confidence interval = $2.84 \times 10^{-2}$–$6.61 \times 10^{-1}$). The contribution of each pharmaceutical's uncertainty to the variance of total ecotoxicity can be computed; these results are shown in Fig. 3. It is assumed that the pharmaceuticals of most concern are those totalling a 90% contribution to the total impact variance. For the pharmaceuticals of most concern, the combination of model input variables to the variance of the results of Fig. 2 is shown in Fig. 4. Generally, for the substances of most concern, the HC50 parameter is the most relevant one for the statistical spread of impact results shown in Fig. 2. For most substances, the parametric quantification of HC50 uncertainty is based on only 3 data values, which typically produces wide confidence limits (Larsen and Hauschild, 2007a), making the statistical differentiation between substances ambiguous. Compounds of most concern are further discussed by dividing the most relevant therapeutical classes into sections.

3.1. Antineoplastics

The antineoplastic tamoxifen displays the highest median ecotoxicity impact (Fig. 2). The uncertainty of the HC50 parameter contributes 93.8% of the variance of the tamoxifen impact results (Fig. 4). Only 2 experimental acute EC50 values, covering 1 trophic level, were obtained in the present study (Supplementary data, Table S5). The ecotoxicological datum on algae was estimated by ECOSAR. The quantification of this QSAR method's uncertainty is not considered in the present study, as stated in Section 2.4; therefore, its influence on impact results is unclear. In addition, the EC50 value for crustaceans was extrapolated from the NOEC. The inherent uncertainty of extrapolating ecotoxicological endpoints is also not considered in the present study. Overall, a more comprehensive ecotoxicological study is needed. Moreover, the
Fig. 2. Probability distribution median and 95% confidence interval of ecotoxicity impact, in PAF m$^3$ d, of pharmaceuticals on freshwater per m$^3$ of WWTP effluent.
calculated impact of tamoxifen is based on very limited data on measurements in WWTP effluents (e.g., Roberts and Thomas, 2006). According to the outcome of the present study, tamoxifen should be subject to monitoring in WWTP effluents for more conclusive results. The neutral form of tamoxifen, with an estimated log Kow of 6.30 (USEPA, 2008b), is highly hydrophobic. Moreover, tamoxifen is predominantly found in the basic form at pH 7 (pKa = 8.52); therefore, electrostatic interactions may play a significant role in its partitioning into negatively charged sorption sites of particles and, consequently, in its removal from WWTPs. The environmental occurrence of tamoxifen is, however, common (e.g., Hilton and Thomas, 2003; Roberts and Thomas, 2006; López-Serna et al., 2012). Another issue of concern, and a subject for further study, is the depletion of tamoxifen, which may be underestimated in the aquatic environment because no data on indirect photolysis are available in the literature, such as bimolecular rate constants for the reaction between the compounds and chemical transients. This compound, which has double bonds and aromatic rings, may react with chemical transients generated by natural water constituents under sunlight, especially with...
Fig. 4. Contribution of model input variables to impact variance of pharmaceuticals of most concern.
the extremely reactive hydroxyl radical that can abstract hydrogen from saturated organics, add to double bonds or add to aromatic rings. In contrast, the chronic ecotoxicity of tamoxifen derivatives produced by direct photolysis revealed no significant differences in comparison to the parental compound (DellaGreca et al., 2007); therefore, the overall impact of tamoxifen may be underestimated, given that photoproducts were not included in the present study.

3.2. Analgesics/anti-inflammatories

Mefenamic acid is a widely used non-steroidal anti-inflammatory compound and is commonly found in WWTP effluents (e.g., Barron et al., 2009; Radjenovic et al., 2009; Rosal et al., 2010; Tauxe-Wuersh et al., 2005). Literature data on mefenamic acid concentrations in WWTP effluents vary by 3 orders of magnitude, from 0.005 (Kasprzyk-Hordern et al., 2009) to 3.0 μg/l (Tauxe-Wuersh et al., 2005). The variability of the effluent concentration contributes 97.7% of the variance of the impact of mefenamic acid (Fig. 4). However, the calculated effect of this anti-inflammatory may be overestimated, given that Werner et al. (2005) suggested that photosensitisation by ex- cited triplet-state DOM may contribute to the environmental degrada- tion of mefenamic acid. The influence of this degradation mechanism on the calculated effect remains unknown in the present study.

The calculated impact of aminopyrine is based on estimated ecotoxicological data. Even excluding the uncertainty of ecotoxicity data estimation, the HC50 parameter has a contribution of 97.4% to the vari- ance of the impact of aminopyrine. Moreover, this compound is not commonly detected in WWTP effluents (e.g., Ternes, 1998; Andreozzi et al., 2003). Poor sorption to particles in WWTPs may be expected, given that the predominant neutral form of this basic compound at pH 7 (pKa = 5.0) has an estimated log KOW of 0.6 (USEPA, 2008b). There- fore, depending on the role of its biotransformation in WWTPs, a very low influent concentration or non-existent discharge may have been observed in WWTPs; nevertheless, no data on influent concentrations were reported in the literature. In fact, the human clinical use of amino- pyrine is widely banned due to the risk of agranulocytosis and due to its potential to produce carcinogenic nitrosamines (U.N., 2003); hence, its presence in WWTP discharges may be caused by low levels of appli- cation in veterinary medicine or by industrial release (Ternes, 1998).

No abiotic degradation data are available; however, aminopyrine is ex- pected to be susceptible to indirect photolysis. In addition, it contains chromophores that absorb at wavelengths N 290 nm and may therefore also be susceptible to direct photolysis; hence, the residence time of aminopyrine in the aquatic environment may be overestimated.

The concentration reported in the literature on the occurrence of the opiate codeine in WWTP effluents varies by 3 orders of magnitude (Gómez et al., 2007; Wick et al., 2009), from 0.022 to 15.59 μg/l. This variability of the effluent concentration contributes 66.6% of the variance of impact results. The HC50 parameter contributes 32.8%; furthermore, estimated data were applied and, as stated above, the quantification of uncertainty in endpoints estimation was not consid- ered in the present study; therefore, the influence of using estimated data on impact variance is unclear. Codeine is expected to be susceptible to indirect photolysis and contains chromophores that absorb at wavelengths N 290 nm; therefore, it may also be susceptible to direct photolysis.

The concentration of tramadol in WWTP effluents reported in the literature varies by 3 orders of magnitude, from 0.02 to 97.62 μg/l (Kasprzyk-Hordern et al., 2009; Wick et al., 2009). The effluent concen- tration of tramadol contributes 75.3% to the impact variance. The HC50 parameter represents 24.1% of the tramadol impact result variance; fur- thermore, for experimental EC50 values of 2 trophic levels, crustaceans and fish, the species were not specified in the literature. In terms of environmental occurrence, tramadol was detected in 2 rivers in South Wales, UK at a maximum concentration of 5970 ng/l (Kasprzyk- Hordern et al., 2009).
3.3. β-Blockers

The HC50 parameter contributes between 70 and 86% to the impact results for variance of the β-blockers betaxolol, oxprenolol, and propanolol. Moreover, oxprenolol ecotoxicity data have been estimated for all trophic levels, and in the case of betaxolol, only 1 acute EC50 value is experimental. Propanolol is commonly measured in WWTP effluents (e.g., Alder et al., 2010; Maurer et al., 2007; Wick et al., 2009); however, limited data are available on the occurrence of oxprenolol and betaxolol in WWTP effluents (Ternes, 1998; Andreozzi et al., 2003). According to the molecular structures of β-blockers, indirect photolysis may play a role in their persistence in the aquatic environment; however, except for propanolol, no experimental data on photosensitisation were found in the literature; therefore, its residence time in the aquatic environment may be overestimated. In fact, in terms of environmental occurrence, no literature data on oxprenolol were found; however, betaxolol was not detected in 29 rivers in Germany, even when it was present in WWTP effluents (Ternes, 1998), nor was it detected in the Ebro River in Spain (López-Serna et al., 2012).

3.4. Psychiatric drugs

In addition to the high contribution of the HC50 parameter to the variance impact of the tricyclic antidepressant amitriptyline (95.9%), the experimental ecotoxicological data were limited to crustaceans, with chronic EC50 values for 4 species. In the case of other trophic levels, ECOSAR values were applied. Both the high sorption potential of the neutral form, with an estimated log KOW of 4.95 (USEPA, 2008b), and the predominance of the basic form at pH 7 (pKa = 9.4) indicate significant removal in WWTPs. Nevertheless, the literature data (both on measurements of amitriptyline in WWTP effluents and on the fate of amitriptyline in WWTPs) are too limited for conclusive results. In addition, according to its molecular structure, amitriptyline may be susceptible to indirect photolysis; hence, its residence time in the aquatic environment may be overestimated in the present study.

The impact of diazepam is comparatively significant for the higher concentrations in WWTP effluents that have been reported in the literature (Supplementary data, Table S3). This parameter has a contribution of 92.5% to the impact variance. The concentration ranges 3 orders of magnitude, from 0.04 to 19.3 μg/l (Suárez et al., 2005; Ternes, 1998); however, measurements of this compound in WWTP effluents are very scarce in the literature.

The HC50 parameter contributes 43.3% to impact variance of the serotonin reuptake inhibitor fluoxetine. Three acute experimental HC50s covering 3 trophic levels were applied. An acute LC50 value was applied for fish; however, for fluoxetine, other TMOAs such as endocrine disruption may be relevant, given that Mennigen et al. (2008) suggested that fluoxetine may have the potential to affect sex hormones and modulate genes involved in the reproductive function of fish. The effluent concentration has a contribution of 39.4% to the impact variance. Although direct photolysis could potentially limit the persistence of fluoxetine in surface waters, Lam et al. (2004) suggested that its degradation by indirect photolysis would be the limiting degradation mechanism.

3.5. Other therapeutical classes

The statistical spread of the antifungal clotrimazole's impact is also mainly due to the HC50 parameter, with a 95% contribution to the variance. According to its molecular structure, clotrimazole is expected to be susceptible to indirect photolysis; therefore, its residence time in the aquatic environment may be overestimated. The neutral form of clotrimazole, which predominates at pH 7 (pKa = 5.22), is highly hydrophobic, with an estimated log KOW of 6.26 (USEPA, 2008b); therefore, significant partitioning to particles in WWTPs may be observed. There are limited data on the occurrence of this topical product in WWTP effluents at detectable concentrations (OSPAR, 2005); however,
clostrimazole is a widely used over-the-counter antifungal agent. Moreover, in terms of environmental occurrence, clostrimazole was the most frequently detected of 14 pharmaceuticals analysed in UK estuaries, with median concentration of 7 ng/l (Hilton and Thomas, 2003); in addition, it was detected with a median concentration of 21 ng/l in the River Tyne, UK (Roberts and Thomas, 2006); nevertheless, it was not detected in the Elbe and Saale Rivers in Germany at any of the measured points (OSPAR, 2005).

The HC50 parameter contributes 59% to the impact variance of the antihypertensive receptor diltiazem. This parameter is of even greater concern, given that only 1 experimental EC50 value was found in the literature (Supplementary data, Table S5).

No abiotic degradation data are available. However, diltiazem is expected to be susceptible to indirect photolysis since it has double bonds and aromatic rings, and because it has chromophores that absorb at wavelengths N 290 nm, it has the potential to be degraded by direct photolysis; therefore, the depletion of diltiazem in the aquatic environment may be underestimated.

The HC50 parameter contributes 95% to the impact variance of the proton-pump inhibitor omeprazole. Moreover, only 1 experimental EC50 value was found in the literature (Supplementary data, Table S5). Very limited data on measurements of omeprazole in WWTP effluents are available in the literature (Rosal et al., 2010); nevertheless, it is one of the most widely prescribed pharmaceuticals. Omeprazole is expected to undergo hydrolysis in the environment due to the presence of functional groups that hydrolyse under environmental conditions, and it may also be susceptible to direct and indirect photolysis (DellaGreca et al., 2006); however, no experimental data were found in the literature.

In addition to the high contribution of HC50 to the variance of the macrolide antibiotic azithromycin’s impact result (52.2%) all EC50 values were estimated. The effluent concentration contributes 46.7% to the impact variance. The photodegradation of azithromycin was shown to be enhanced in the presence of nitrates and humic acids (Tong et al., 2011), which indicates the role of indirect photolysis in the persistence of this compound in the environment. However, indirect photodegradation was not included in the present study for this compound; therefore, the impact of azithromycin is most likely overestimated.

Concentrations of the hormone 17β-estradiol in WWTP effluents reported in the literature vary by 2 orders of magnitude, from 0.0007 to 0.0180 µg/l (Baronti et al., 2000; Clara et al., 2004). The variability of effluent concentration represents 63.6% of the impact variance. The HC50 parameter contributes 29.2% to the variance. Experimental acute EC50 values for 6 species were applied; however, the EC50 value for algae was estimated.

The diuretic bendroflumethiazide, the anti-inflammatories 5- aminosalicylic acid and ketorolac, and the lipid regulator clofibrate are discussed in the Supplementary data (Section 5).

3.6. Additional considerations

Table 1 summarises future research topics for the pharmaceuticals of greatest concern. These topics can be related to 3 issues: a) the fate of pharmaceuticals in WWTPs, b) substance-specific modelling parameters, and c) lack of spatial and time resolution models. The first topic includes compounds with very limited data on measurements or detection in WWTP effluents, such as tamoxifen or amitriptyline. These sub- stances should be subject to further monitoring in WWTPs, depending on geographical usage patterns, for more conclusive results. This category should also include compounds whose impact result would be most sensitive to variations of the emission concentration. Ideally, a comparatively well characterised drug from an impact perspective would account for low variance of output results due to environmental fate and transport modelling parameters, either estimated or experimental, and due to EEF characterisation. The uncertainty of its impact result, from a modelling perspective, would be related mainly to the variability of the concentration in WWTP effluents, depending on geographical and seasonal usage patterns, treatment technologies, and operation conditions. The focus of research for these compounds should be detailed ecological risk assessments possibly leading to research and development on the operation and design of WWTPs to improve the reduction of

Table 1
Research topics for pharmaceuticals of most concern. Three arrows denote a research topic of higher concern, two arrows denote a research topic of moderate concern, and one arrow denotes a research topic lower concern.

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<th>Parameter</th>
<th>Abiotic degradation</th>
<th>Derivative</th>
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<td>5-Aminosalicylic</td>
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<td>Codeine</td>
<td>↓↓</td>
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</tr>
<tr>
<td>Diazepam</td>
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<tr>
<td>Diltilazem</td>
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<tr>
<td>Fluoxetine</td>
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<td>Ketornlac</td>
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<tr>
<td>Mefenamic acid</td>
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<tr>
<td>Omeprazole</td>
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<tr>
<td>Oxprenolol</td>
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<tr>
<td>Prunanolol</td>
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<tr>
<td>Tamoxifen</td>
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<tr>
<td>Tramadol</td>
<td>↓↓</td>
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</table>

a: more than 10 peer-reviewed publications; ↓: between 5 and 10 peer-reviewed publications; ↓↓: between 2 and 5 peer-reviewed publications; ↓↓↓: only one peer-reviewed publication.

b: more than 3 acute EC50s covering 3 trophic levels; ↓↓↓: 3 acute EC50s covering 3 trophic levels; ↓↓↓: at least 1 estimated or extrapolated EC50.

c: Number of possible abiotic degradation mechanisms not included in the assessment (hydrolysis, direct and indirect photolysis); (↓↓) denotes a specific degradation pathway with some evidence of occurrence in the literature but with no data available.

d: Number of possible degradation mechanisms generating derivatives (hydrolysis, photolysis and biodegradation); (↓↓) denotes a specific degradation pathway with evidence of derivatives toxicity in the literature.
the compounds' effluent concentrations. However, the compounds most sensitive to the emission concentration, such as diazepam or mefenamic acid, have other research priorities either because of limited data on their occurrence or incomplete modelling parameters.

The second issue includes drugs whose impact results are mostly sensitive to the uncertainty of substance-specific modelling parameters, such as degradation rates or partitioning coefficients, or to EEF characterisation. It also includes drugs whose impact result may be affected by modelling incompleteness, either from the lack of abiotic degradation data (such as for omeprazole or azithromycin) or from the exclusion of degradation products (such as for tamoxifen). These compounds should be subjected to further experimental research according to the most sensitive parameters because of a lack of precise knowledge regarding those parameters. The third issue, the lack of spatial and time resolution models, addresses the variability of landscape parameters, such as freshwater pH or [OH], and the seasonal variation of direct photolysis rates. However, for the compounds of greatest concern, only the spatial variability is somewhat significant, and only in the case of [OH]. The large scale applied in the present study displays a great variety of landscape characteristics; nevertheless, the uncertainty regarding the HC50 parameter and the variability of the effluent concentration predominate in terms of the contribution of variance to the output results.

3.7. Model limitations

It should be noted that other sources of uncertainty not included in the Monte Carlo analysis may be important. Some have already been discussed above, such as the uncertainty of ecotoxicological data estimation, the extrapolation of endpoints, the lack of abiotic degradation data for several compounds, and the exclusion of abiotic and biotic derivatives of parent compounds. This last source of uncertainty may be relevant in the case of tamoxifen, as already mentioned; however, substances that do not appear in the ranking of compounds of most concern may have their comparative impact substantially increased by the inclusion of their derivative impact. For example, some researchers have suggested that the phototransformation products of triclosan, diclofenac or hydrochlorothiazide have a higher toxicity potential than their parent compounds (Han et al., 2000; Schmitt-Jansen et al., 2007). Nevertheless, the inclusion of phototransformation product impact is possible, if the chemical structures are identified, by applying the method proposed by van Zelm et al. (2010).

In addition, the uncertainty of the influence of pH on direct and indirect photolysis rates, the uncertainty of the application of a linear dose–response curve for the calculation of EEFs, and the lack of spatial variation of background impacts in the AMI method remain unclear. For example, for uncertainty of the influence of pH on the abiotic degradation, the literature data on the direct phototransformation of triclosan (pK = 8.1) applied in the present study are based on its anionic form (Tixier et al., 2002), which is the dominant photochemical degradation pathway. Therefore, by disregarding the influence of pH on the direct photolysis rate, the residence time of triclosan in the freshwater environment may be underestimated for lower pH values.

A first screening approach to deal with the uncertainty of speciation of an organic compound could be based on a uniformal distribution using the lowest and highest degradation rates amongst all the species involved in the speciation as the minimum and maximum. Therefore, quantum yields and experimental molar absorption coefficients in function of the UV/VIS wavelength range of all the species involved must be experimentally obtained and applied to models that compute direct photolysis rates and half-lives of pollutants in the aquatic environment. A similar approach can be applied for indirect photolysis by obtaining experimental rate constants between chemical transients and all the chemical species involved in the speciation.
4. Conclusions

Despite the high uncertainties of the pharmaceutical impact results, which range up to 12 orders of magnitude, and the model’s limitations and parameter incompleteness, the outcome of the present study allows priorities to be set for further experimental testing. Several pharmaceutical compounds were identified as being of greatest concern, including 7 analgesics/anti-inflammatories, 3 β-blockers, 3 psychiatric drugs, and 1 each of 6 other therapeutic classes.

Notably, some pharmaceuticals identified as of greatest concern, such as tamoxifen, clotrimazole and oxprenolol, have rarely been investigated previously with regard to their ecotoxicity, their occurrence in WWTPs, or their degradation in the environment. Theoretically, the relevant pharmaceuticals may be susceptible to abiotic degradation. However, in general, no experimental data are available; hence, the persistence of these pharmaceuticals in the freshwater compartment is estimated to be comparatively higher than that of well-researched pharmaceuticals that were not included on the ranking of priority compounds, such as triclosan, diclofenac or ibuprofen.

Ecotoxicity data remain to be the most critical issue affecting impact or risk assessments of pharmaceuticals. The present assessment is based on only 3 data values for most of priority pharmaceuticals (only approximately 4% of these compounds have more than 3 EC50 values) that produced wide confidence limits. Moreover, approximately 58% of the pharmaceuticals of priority have at least 1 estimated or extrapolated EC50.

In short, this study identified several pharmaceuticals both for further WWTP monitoring and for testing their ecotoxicity and their persistence in the environment.

Conflict of interest

All the authors of manuscript don’t have any actual or potential conflict of interest including any financial, personal or other relationships with other people or organisations within three years of beginning the submitted work that could inappropriately influence, or be perceived to influence, their work.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.scitotenv.2014.04.082.

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