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Modelling Illicit Drug Fate in Sewers for Wastewater-Based Epidemiology

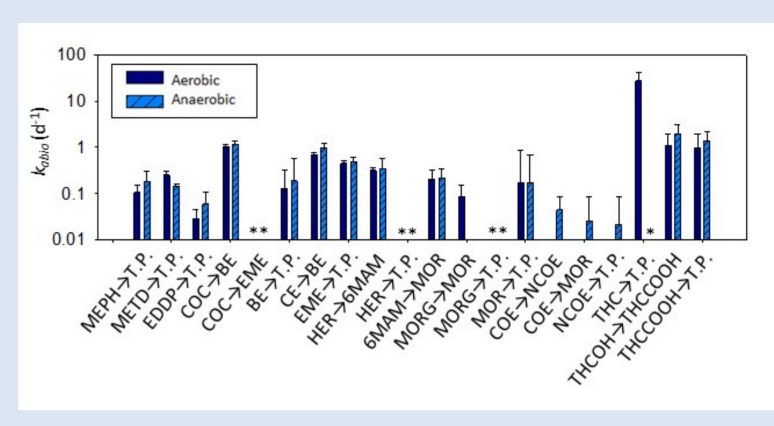
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The sewer as a bioreactor

Sewer pipelines, although primarily designed for sewage transport, is in fact a bioreactor where the primary substrates (e.g. natural organic fractions) and trace organic substrates (e.g. excreted drug biomarkers) can undergo significant transformation. Hence, the occurrence of illicit drug biomarkers can be potentially influenced by physicochemical and biological processes (fate processes) in the sewer

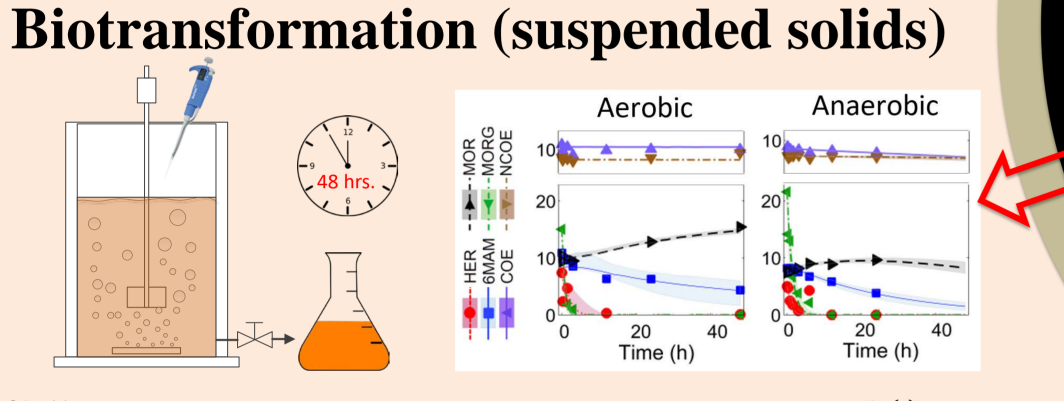
Abiotic transformation Aerobic Anaerobic



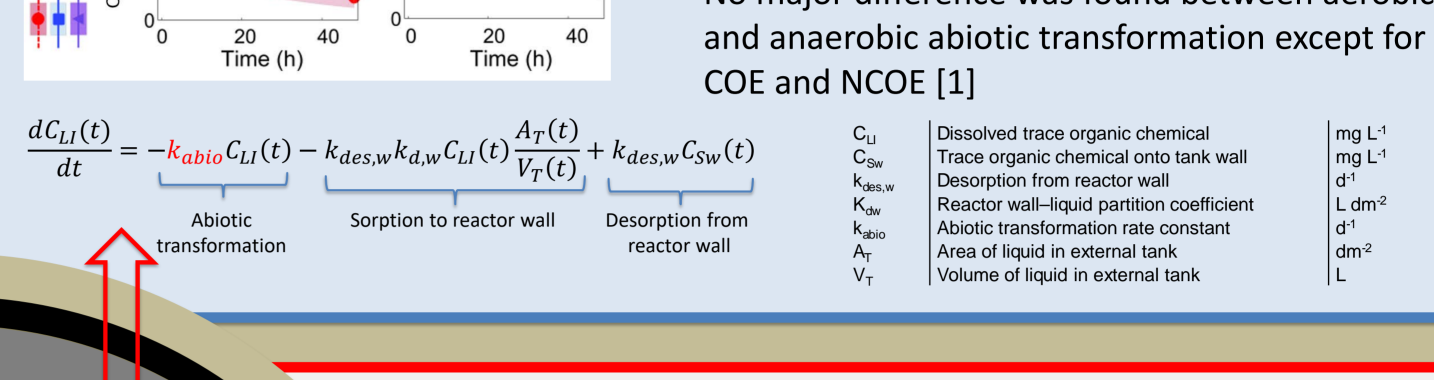
- Experiments were performed with mineral water
- First order equation was used for predictions
- Abiotic transformation is significant (>1 d⁻¹) for many of the compounds specially for COC, CE, THC, THCOH and THCOOH
- No major difference was found between aerobic

Aims

- Providing new evidences on drug transformation and sorption in the sewer
- Predicting the fate processes using mathematical models and statistical analysis
- Primarily focusing on batch experiments representing the sewer as well as preliminary analysis in catchments



 $\frac{dC_{LI}(t)}{dt} = -\frac{k_{bio}}{C_{LI}(t)X} - k_{abio}C_{LI}(t) - k_{des}k_dC_{LI}(t)X_{SS} + k_{des}C_{LI}(t) - k_{des,w}k_{d,w}C_{LI}(t)\frac{A_T(t)}{V_{T}(t)} + k_{des,w}C_{Sw}(t)$



Sorption (biofilm)	
NaN ₃ ¹	
Image: state	
$K_{df} = \frac{C_{SL,eq} - C_{loss}}{(C_{LI,eq} + C_{loss})X_{SS}}$ $C_{loss} = C_{LI,t=0} - C_{LI,t=end}$	

Compound	<i>K_{df}</i> (L g⁻¹) aerobic	<i>K_{df}</i> (L g⁻¹) anaerobic
MEPH	0.2	0
METD	0.32	0.55
EDDP	0	0.15
BE	0.9	0.62
EME	0	1.59
ТНСОН	2.81	1.68
ТНССООН	0	1.06

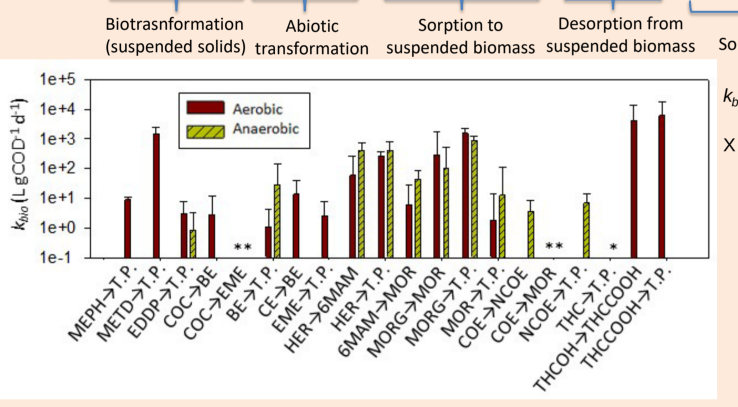
mg L⁻¹

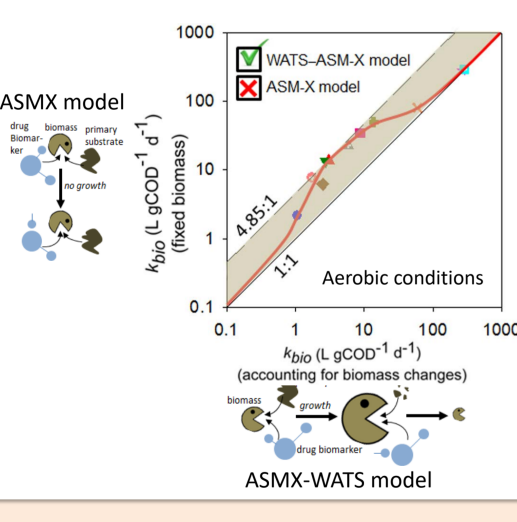
mg L⁻¹

L dm⁻²

dm⁻²

- Sorption tests were performed with suspended biofilm [4]
- Sorption to biofilm was observed for few compounds



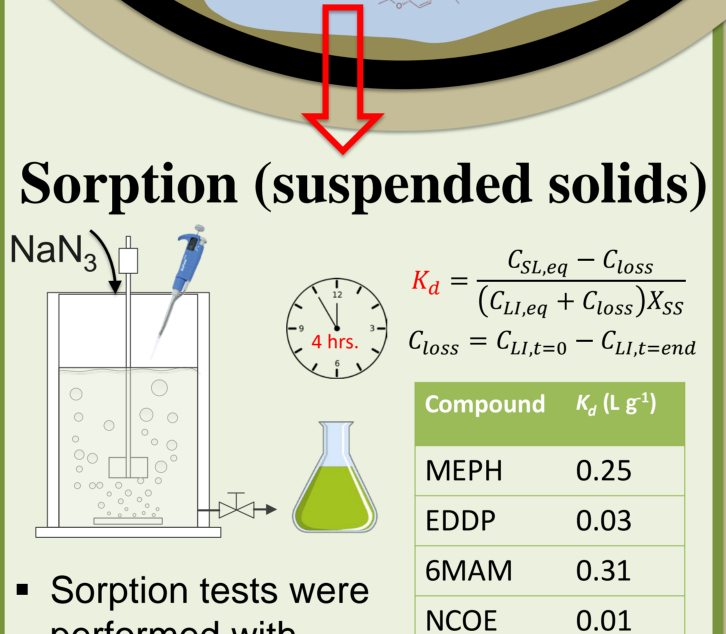


Desorption from Sorption to reactor wa reactor wal L gCOD⁻¹ d⁻¹ rate constant Total particulate dm⁻² solids/biomass Suspended solids/biomass play and important role in biotransformation of most of the selected compounds [1]

Redox condition has significant impact on biotransformation rates for nearly all selected drugs Accounting for biomass growth to estimate biotransformation rate under aerobic conditions is crucial Biomass growth do not play a major role in estimation of

anaerobic biotransformation rate

Fate of the drug biomarkers was assessed using two modelling approaches I- Describing the biokinetics only for secondary metabolic substances (drug biomarkers) assuming that biomass is constant (based on ASM-X [2]) II- Modelling the primary metabolic processes (related to COD fractions) (based on WATS [3]) combined with secondary metabolic processes (WATS—ASM-X model [1])



- performed with THCOH washed-off diluted THCCOOH 0.8 primary sludge [1]
- Sorption to reactor wall and hydrolysis (loss) were accounted for in calculation of k_d using blank experiments
- Sorption to suspended solids was found to be relevant for a few compounds

Abiotic processes

Catchment A Catchment B Catchment (

Biotransf. (raw wastewater)

Biotransf. (sewer biofilms)

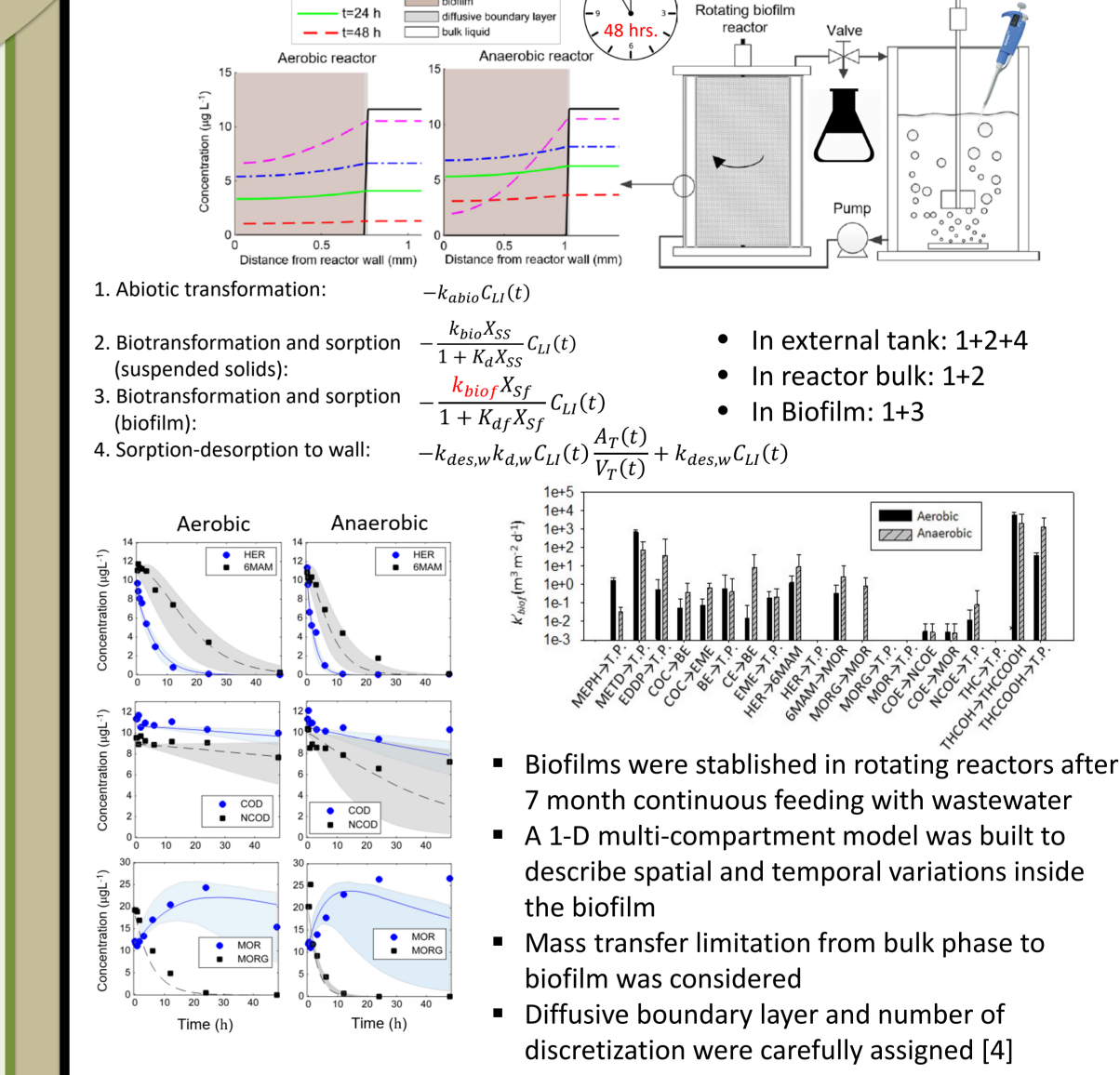
THC was found to completely sorped to reactor wall

80 🛞

to tal

20

Average



Biotransformation (biofilm)



Drug biomarkers loss in three hypothetical

A: 10000 PE, HRT=1.4 h

B: 50000 PE, HRT=2.8 h

C: 200000 PE, HRT=5.5 h

catchments (simulation study) [5]

Final remarks

0.75

- Through different experimental assessments, it was found that drug biomarkers can potentially undergo significant sorption and transformation in the sewers
- Sorption and transformation rates are the key indicators of *drug stability* in the sewer which can be estimated using mathematical models and statistical analysis
- In the sewer, different drug biomarkers can transform to each other, hence pathway identification is crucial when modelling transformations [6] Ignoring in-sewer fate processes for drug biomarkers can be a significant source of bias for wastewater-based epidemiology
- [1] Ramin, P., Brock, A.L., Polesel, F., Causanilles, A., Emke, E., de Voogt, P., Plósz, B.G., 2016. Transformation and sorption of illicit drug biomarkers in sewer systems : understanding the role of suspended solids in raw wastewater. Environ. Sci. Technol. 50, 13397–13408.
- [2] Plósz, B. G.; Reid, M. J.; Borup, M.; Langford, K. H.; Thomas, K. V. Biotransformation kinetics and sorption of cocaine and its metabolites and the factors influencing their estimation in wastewater. Water Res. 2013, 47, 2129-2140.

80

60

40

20

a: MOR

b: MORG

c: 6MAM

- [3] Hvitved-Jacobsen, T.; Vollertsen, J.; Nielsen, A. H. Sewer Processes: Microbial and Chemical Process Engineering of Sewer Networks, Second Edition; CRC Press, 2013 [4] Ramin, P., Brock, A.L., Causanilles, A., Valverde Pérez, B., Emke, E., de Voogt, P., Polesel, F., Plosz, B.G., 2017. Transformation and sorption of illicit drug biomarkers in sewer biofilms. Environ. Sci. Technol. 51, 10572–10584. [5] Ramin, P., Vezzaro, L., Mikkelsen, P.S., Plósz, B.G. Back-calculating illicit drug abuse rates in urban areas – Quantifying uncertainties associated with in-sewer transformation and sampling. Submitted [6] Ramin, P., Valverde Pérez, B, Polesel, F., Locatelli, L., Plósz, B.G., 2017b. A systematic model identification method for chemical and biochemical transformation pathways combined with reaction kinetics – the case of heroin biomarkers. Sci. Rep. 7, 9390.
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