

Highlights:

- Conceptual review of hyporheic zones of rivers as natural anoxic-oxic bioreactors
- Interactions between organohalide respiration and biogeochemical cycling
- Aerobic vinyl chloride mineralisation during hyporheic mixing is conceptualised
- Field experience, challenges and characterisation technologies critically reviewed

1 **Natural attenuation of chlorinated ethenes in hyporheic zones: a review of**
2 **key biogeochemical processes and in-situ transformation potential**

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17 **Abstract**

18 Chlorinated ethenes (CEs) are legacy contaminants whose chemical footprint is expected to
19 persist in aquifers around the world for many decades to come. These organohalides have been reported
20 in river systems with concerning prevalence and are thought to be significant chemical stressors in urban
21 water ecosystems. The aquifer-river interface (known as the hyporheic zone) is a critical pathway for
22 CE discharge to surface water bodies in groundwater baseflow. This pore water system may represent a
23 natural bioreactor where anoxic and oxic biotransformation processes act in synergy to reduce or even
24 eliminate contaminant fluxes to surface water. Here, we critically review current process understanding
25 of anaerobic CE respiration in the competitive framework of hyporheic zone biogeochemical cycling
26 fuelled by in-situ fermentation of natural organic matter. We conceptualise anoxic-oxic interface
27 development for metabolic and co-metabolic mineralisation by a range of aerobic bacteria with a focus
28 on vinyl chloride degradation pathways. The superimposition of microbial metabolic processes
29 occurring in sediment biofilms and bulk solute transport delivering reactants produces a scale
30 dependence in contaminant transformation rates. Process interpretation is often confounded by the
31 natural geological heterogeneity typical of most riverbed environments. We discuss insights from recent
32 field experience of CE plumes discharging to surface water and present a range of practical monitoring
33 technologies which address this inherent complexity at different spatial scales. Future research must
34 address key dynamics which link supply of limiting reactants, residence times and microbial
35 ecophysiology to better understand the natural attenuation capacity of hyporheic systems.

36 **Keywords:** Chlorinated ethenes, hyporheic zone, natural attenuation, biogeochemistry,
37 biotransformation, heterogeneity.

38 Abbreviations:

39 1,1,1-TCA: 1,1,1-trichloroethane; DCA: 1,2-dichloroethane; Ac: acetate, CA: chlorinated ethane;
40 cDCE: *cis*-1,2-dichloroethene; CE: chlorinated ethene; Da: Damköhler number; DIC: dissolved
41 inorganic carbon; DNAPL: dense, non-aqueous phase liquid; DOC: dissolved organic carbon; DOM:
42 dissolved organic matter; DON: dissolved organic nitrogen; HCB: hexachlorobenzene; HCE: higher
43 chlorinated ethene; HFC: hyporheic flow cell; K: saturated hydraulic conductivity; LCE: lower
44 chlorinated ethene; OHR: organohalide respiration; OHRB: organohalide-respiring bacteria; OM:
45 organic matter; PCE: tetrachloroethene; PCR: polymerase chain reaction; POM: particulate organic
46 matter; SCFA: short chain fatty acid; SOM: sedimentary organic matter; TCE: trichloroethene; TCM:
47 trichloromethane; TEA: terminal electron acceptor; TEAP: terminal electron accepting process; VC:
48 vinyl chloride.

49 1 Introduction

50 Legacy soil and groundwater contamination by chlorinated ethenes (CEs) remains a global
51 environmental concern (Burston et al., 1993; Jackson, 1998; Rowe et al., 2005; Rivett et al., 2006, 2012;
52 2014). The higher CEs (HCEs), tetrachloroethene (PCE) and trichloroethene (TCE) are parent
53 compounds employed historically as non-polar solvents for industrial cleaning and degreasing
54 applications which enter aquifers as dense, non-aqueous phase liquids (DNAPLs) (Pankow and Cherry,
55 1996; Rivett et al., 2014) and release dissolved-phase plumes (e.g. Benker et al., 1996; Rivett and
56 Feenstra, 2005; Koch and Nowak, 2015). Diffusive retention of HCEs in low-permeability and fractured
57 formations is increasingly recognised as an important driver of plume longevity from the resulting back-
58 diffusive flux (Parker et al., 1994; Seyedabbasi et al., 2012; Damgaard et al., 2013; Yang et al., 2014).
59 Under predominantly anoxic conditions, HCEs are reductively transformed to the lower CEs (LCEs)
60 *cis*-1,2 dichloroethene (cDCE) and vinyl chloride (VC) (Squillace and Moran, 2007; Chambon et al.,
61 2010; Huang et al., 2014; Lu et al., 2015). CEs have been detected in surface water with alarming
62 frequency around the world (e.g. Christof et al., 2002; McGuire et al., 2004; Ellis and Rivett, 2007;
63 Yamamoto et al., 2014 Wittlingerová et al., 2016). Groundwater baseflow input of CEs has been
64 implicated as a contributory factor in the global impoverishment of surface water ecological quality

65 known as urban stream syndrome (Meyer et al., 2005; Roy and Bickerton, 2011; Roy et al., 2017). The
66 aquifer-river interface, inclusive of the hyporheic zone (Fig. 1) forms a key diffuse pathway for
67 discharge of groundwater plumes hydrologically connected with local surface water bodies (Table 1)
68 (Conant et al., 2004; Chapman et al., 2007; McKnight et al., 2012; Weatherill et al., 2014; Puigserver et
69 al., 2014; Simsir et al., 2017; Sonne et al., 2017).

70 The hyporheic zone of streams and rivers has been defined by various disciplines from alternative
71 perspectives (e.g. Boulton et al., 1998; Buss et al., 2009; Krause et al., 2011; Cardenas, 2015). In this
72 review, we conceptualise the hyporheic zone as a dynamic volume of the saturated zone where the pore
73 water domain is physically influenced by exchanges across the sediment–water interface (Cardenas,
74 2009). This spatially and temporally variable volume may encompass shallow benthic sediments and
75 adjacent parts of the alluvial aquifer (Boano et al., 2014). The transition zone is frequently heterogeneous
76 with a wide range of saturated hydraulic conductivities (K) and associated scaling of solute residence
77 times (Haggerty et al., 2002; Sawyer and Cardenas, 2009; Gomez-Velez et al., 2014). Fluvial sediments
78 are sites of organic matter (OM) retention with high interstitial surface area linked to enhanced microbial
79 metabolic activity and characterised by sharp physical and biogeochemical gradients (Krause et al.,
80 2013, 2014; Boano et al., 2014; Atashgahi et al., 2015; Narango et al., 2015; Briggs et al., 2016). A
81 generic scenario is depicted in Fig. 1 where OM present in the sediment sequence induces anoxia along
82 discharging groundwater flow paths through the interface. The CE plume depicted may be composed
83 solely of parent HCEs (e.g. McKnight et al., 2010) or a mixture of LCEs (e.g. Sonne et al., 2017; Rønde
84 et al., 2017). Shallow oxic zones are created where local hyporheic exchange flows with the river
85 penetrate the underlying anoxic zone (Hester et al., 2013; 2017, Trauth et al., 2014). Heterotrophic
86 biogeochemical cycling of carbon, nitrogen, sulfur and metals will take place along anoxic sediment
87 flow paths (Baker et al., 2000; Mermillod-Blondin et al., 2005; Lautz and Fanelli, 2008; Rahimi et al.,
88 2015). Fluvial sediments can form niche environments for anaerobic biotransformation of CEs by
89 reductive dechlorination and much interest has arisen in their intrinsic capacity to assimilate
90 groundwater CE loads in baseflow (Table 1) (Conant et al., 2004; Abe et al., 2009; Hamonts et al., 2009;
91 Weatherill et al., 2014; Atashgahi et al., 2015; Freitas et al., 2015; Simsir et al., 2017). Discharging

92 groundwater flow paths through fluvial sediments may enter into mixing zones around infiltrating
93 surface water during hyporheic exchange flows (Boano et al., 2014; Gomez-Velez et al., 2014; Hester
94 et al., 2017). Dynamic pore water oxygen (O_2) gradients are induced in mixing zones as a result (Trauth
95 et al., 2014; Vieweg et al., 2015; Cardenas et al., 2016). These mixing zones provide a separate niche
96 environment for aerobic mineralisation of LCE daughter products originating in the anoxic zone or in
97 partially dechlorinated plumes (e.g. Rønde et al., 2017). Hence, hyporheic zones of groundwater-fed
98 rivers may constitute a unique natural ‘biobarrier’ for complementary anaerobic and aerobic biological
99 treatment of diffuse baseflow CE pollution (Hamonts et al., 2007; Abe et al., 2009; Atashgahi et al.,
100 2017a; Simsir et al., 2017).

101 The aim of this review is to provide a critical evaluation of the current understanding of
102 synergistic anoxic-oxic biotransformation processes and their environmental supporting conditions in
103 typical riverbed sediment sequences. We consider the fate of parent HCEs in anaerobic heterotrophic
104 food webs driven by OM fermentation and observed metabolic interactions with key biogeochemical
105 cycles. We conceptualise the development of anoxic-oxic interfaces in surface water mixing zones
106 which support aerobic bacteria capable of assimilating LCEs via metabolic and cometabolic pathways
107 with a particular focus on VC. We discuss challenges in integrating microbially-mediated transformation
108 processes occurring in sediment biofilms with larger scale hydrological fluxes observable at field scale.
109 We critically review available in-situ monitoring technologies and support our discussions with insights
110 from recent field experience. This process-based understanding is intended to underpin the risk-based
111 management of legacy CE contamination. The coupled biogeochemical and transport processes
112 understanding is highly relevant for river restoration and prospective ecological engineering applications
113 aimed at reducing the impact of organohalide stressors in urban streams (e.g. Lawrence et al., 2013;
114 Rasmussen et al., 2016; Roy et al., 2017).

115 **2 Biotransformation of chlorinated ethenes in riverbed sediments**

116 Biotransformation of CEs in hyporheic zones will occur when requisite bacteria, a circumneutral
117 pH, nutrients and electron donors/acceptors are present and where solute residence time equals or
118 exceeds the transformation timescale (half-life) of the contaminant (Meckenstock et al., 2015). In anoxic

119 zones, CEs may undergo anaerobic reductive dechlorination (hydrogenolysis) by heterotrophic bacteria
120 where chlorine atoms are sequentially replaced by hydrogen (Fig. 2) in organohalide respiration (OHR)
121 (Vogel et al., 1987; Hug et al., 2013; Leys et al., 2013). LCEs produced by OHR may reach oxic
122 interfaces where metabolic and co-metabolic aerobic pathways lead to complete natural attenuation.
123 This part of our review profiles key advances in the understanding of these synergistic biotransformation
124 process for which hyporheic zones present a unique capacity.

125 **2.1 Organohalide respiration**

126 Under anoxic conditions, CEs are terminal electron acceptors (TEAs) for energy conservation
127 (Fig. 2) during oxidation of an electron donor (usually hydrogen) in the presence of a carbon source
128 (usually acetate) and a nitrogen source (ammonium) (Mohn and Tiedje., 1992; Smidt and de Vos, 2004).
129 OHR is an energy-conserving process where the chemically stable carbon-halide bond of organohalides
130 is unlocked by replacing the halogen atom with hydrogen and liberating it as a halide (Fig. 2) (Maymo-
131 Gatell et al., 1997; McCarty, 1997; Aulenta et al., 2002). The reductive cleavage of carbon-halide bonds
132 is catalyzed by membrane-bound, cobalamin-containing enzymes known as reductive dehalogenases
133 (Bommer et al., 2014). OHR is mediated by organohalide-respiring bacteria (OHRB) belonging to
134 distinct bacterial groups i.e., *Chloroflexi*, *Proteobacteria* and *Firmicutes* (Atashgahi et al., 2016).
135 Members of the genus *Dehalococcoides mccartyi*, *Dehalogenimonas* and *Dehalobacter* spp. are
136 restricted to OHR for their metabolism, although fermentative growth is also shown in the latter (Justica-
137 Leon et al., 2014; Lee et al., 2012; Yang et al., 2017). In contrast, *Geobacter*, *Desulfuromonas*,
138 *Anaeromyxobacter*, *Desulfomonile*, *Desulfoluna*, *Desulfovibrio*, *Sulfurospirillum* and
139 *Desulfitobacterium* genera are facultative OHRB with versatile metabolisms including but not restricted
140 to OHR (Maphosa et al., 2010; Atashgahi et al., 2016).

141 **2.2 Potential heterotrophic food webs in anoxic zones**

142 **2.2.1 Carbon flow in the hyporheic zone**

143 Dissolved organic matter (DOM) is a key source of electron donors and nutrients for OHR and
144 other heterotrophic terminal electron accepting processes (TEAPs) in hyporheic zones (Pusch and
145 Schwoerbel, 1994; Fischer et al., 2005; Zarnetske et al., 2011; Stegen et al., 2016). Anaerobic DOM
146 mineralisation is coupled to the reduction of TEAs as follows (Donn and Barron, 2013):

147 DOM + TEA → DIC + metabolite + NH₄

148 Where TEA (in descending potential metabolic energy yield) is O₂ > NO₃ > PCE > Fe(III) > TCE >
149 cDCE > SO₄ > VC > CO₂, DIC is dissolved inorganic carbon, metabolite is N₂, CO₂, Mn²⁺, LCEs, Fe²⁺,
150 ethene, ethane, S²⁻, CH₄ and acetate) and ammonium (NH₄). Biogeochemical cycling and OHR will take
151 place when DOM concentrations exceed the stoichiometric requirements (non-limiting) for each TEAP
152 where the reaction timescales are less than the exposure time along reactive sediment flow paths (Haest
153 et al., 2011; Zarnetske et al., 2012; Abbot et al., 2016). Hyporheic sediments are often enriched in
154 complex organic matter (Fig. 1) derived from allochthonous terrestrial ecosystem (such as soil particles
155 and leaf litter) and anthropogenic sources of dissolved-phase and suspended particulate material (e.g.
156 urban runoff and wastewater discharges) (Stelzer et al., 2014; Atashgahi et al., 2015). The nature of the
157 organic material that accumulates in riverbed sediments may also influence the effective residence time
158 of the plumes through hydrophobic partitioning to sedimentary organic matter (SOM) (Conant et al.,
159 2004; Allen-King et al., 2002; Wang et al., 2013).

160 Anaerobic decomposition of particulate organic matter (POM) is initiated by extracellular
161 hydrolytic enzymes produced by primary fermenters which release soluble DOM components capable
162 of traversing bacterial cell walls (Fig. 2) (Mani et al., 2016). Hydrolysis is frequently the rate-limiting
163 step in sedimentation zones with high POM loading rates, large particle size ranges or where highly
164 polymerised complex POM (e.g. lignin) are present that are not readily degradable (Gavala et al., 2003;
165 Nogaro et al., 2007; Atashgahi et al., 2014). DOM hydrolysis is a source of labile dissolved organic
166 carbon (DOC) and dissolved organic nitrogen (DON) species including organic acids and the soluble
167 oligomers and monomers of proteins and carbohydrates which influence the onset and rate of anaerobic
168 respiration processes (Jakobsen and Postma, 1999; Zarnetske et al., 2011; Mineau et al., 2013; Helton
169 et al., 2015). Groundwater typically contains <0.15 mM of DOC (Lapworth et al., 2009; Chapelle et al.,
170 2012). Of this, the labile fraction represents only 0.5–5% of carbon present which is often below
171 thresholds required for induction of catabolic genes (Egli, 2010). In contrast, hyporheic zone pore water
172 DOC concentrations often exceed 1 mM (e.g. Lewandowski and Nützmann, 2010) of which 10–30% is
173 labile and bioavailable (Baker et al., 1999; Vidon and Hill, 2004; Romani et al., 2006).

174 2.2.2 Molecular hydrogen

175 Free dissolved hydrogen (H₂) is produced from internal proton reduction during primary and
176 secondary DOC fermentation (Fig. 2) (Nath and Das 2004; Hallenbeck et al., 2009). H₂ serves as the
177 strict electron donor for known obligate OHRB (such as *D. mccartyi*) as well as many facultative OHRB
178 (Jugder et al., 2016). A generalised reaction for H₂ production during fermentation of DOC is given as
179 follows:



181 Where SCFA is a short-chain fatty acid which is discussed in the next section. In anoxic sediment zones,
182 DOC fermentation is a metabolic process utilised by common fermentative bacteria such as
183 *Desulfovibrio* (Walker et al., 2009), *Syntrophomonas* (Sieber et al., 2010), and *Clostridium* (Wu et al.,
184 2012), among others. H₂ is also produced by homoacetogens (Diekert and Wohlfarth, 1994), certain
185 acetoclastic methanogens (Heimann et al., 2007) and *Geobacter* spp. (Löffler and Sanford, 2005). Low
186 extracellular H₂ concentrations are required for DOC fermentation to be energetically favourable. This
187 is often maintained by unique syntrophic interspecies H₂ transfer relationships (Fig. 3) amongst
188 anaerobes which express membrane-bound hydrogenases (Walker et al., 2009; Morris et al., 2013;
189 Sieber et al., 2014; Jugder et al., 2015). As a result, the reaction timescales for H₂ uptake are typically
190 on the order of minutes (Heimann et al., 2009). Under electron-donor limiting conditions, ambient H₂
191 concentrations in sediments are often at steady-state that is determined by the hydrogenotrophs with the
192 lowest threshold concentration (Lovely and Goodwin, 1988; Fennel et al., 1997; Löffler et al., 1999;
193 Hoelen and Reinhard, 2004). H₂ threshold ranges have been reported for specific TEAPs (Fig. 4) ranging
194 from <0.1 nM for denitrification to >350 nM for homoacetogenesis. From Fig. 4 it can be observed that
195 HCE reduction (0.6–0.9 nM) occurs at an overlapping H₂ threshold to Fe(III) reduction (0.1–0.8 nM)
196 whereas LCE reduction overlaps with sulfate reduction and methanogenesis (Paul et al., 2016). Overlap
197 of TEAP H₂ thresholds are commonly observed in experimental studies (e.g. Jakobsen and Postma,
198 1999; Aulenta et al., 2008; Paul et al., 2016) with lower levels than predicted from thermodynamic
199 considerations (Heimann et al., 2009). This phenomenon has been attributed to kinetic effects arising
200 from the relative efficiencies in H₂ producers and consumers in microbial consortia (Richardson, 2016).

201 2.2.3 Short chain fatty acids

202 Short-chain fatty acid (SCFAs) are labile DOC species such as butyrate, propionate and lactate
203 produced as intermediates during primary DOC fermentation (Fig. 2) and are accompanied by an
204 increase in pore water H^+ ion concentrations. SCFAs readily undergo secondary fermentation to produce
205 electron donors (Giovannini et al., 2016). SCFA turnover is often the rate-limiting step in the anaerobic
206 decomposition of complex DOM (Mani et al., 2016) and release of reducing equivalents for OHRB and
207 other heterotrophs (Atashgahi et al., 2014). The most frequently reported SCFA in hyporheic zones is
208 acetate which is both a terminal metabolite and intermediate of DOC fermentation (Fig. 2) (Baker et al.,
209 1999; Baker and Vervier, 2004; Hlaváčová et al., 2005). It is an easily assimilated carbon source and
210 possible direct electron donor for OHRB which may be independent of extracellular H_2 (Richardson,
211 2016). Under high pore water H_2 concentrations (>350 nM) acetate is also produced from other labile
212 DOC compounds (i.e. methanol) or H_2/CO_2 by homoacetogens such as *Acetobacterium*, *Sporomusa*,
213 *Spirochaeta* (Ziv-El et al., 2012).

214 Acetate has been widely shown to sustain respiration of HCEs as far as cDCE by facultative
215 OHRB groups such as *Geobacter* and *Desulfuromonas* (Sharma and McCarty, 1996; Krumholz et al.,
216 1996; Löffler et al., 2000; Wagner et al., 2012). For example, in contaminated aquifer media, Lee et al.
217 (2007) found that whilst H_2 could sustain dechlorination of PCE as far as ethene, with uptake of acetate,
218 dechlorination stalled at cDCE and proceeded at less than half the rate of H_2 alone. However, the role
219 of acetate in OHR appears to be more complex. He et al. (2002) found that several organohalide-
220 containing sediment microcosms amended with acetate alone were capable of sustaining PCE
221 dechlorination to ethene. Their findings suggest a potential syntrophic partnership between cDCE and
222 VC dechlorinators and acetate-oxidising bacteria (such as *Clostridium ultunense*) because of the low H_2
223 concentrations maintained by OHRB. Wei and Finneran (2013) also showed that acetate could sustain
224 TCE, cDCE and VC dechlorination to ethene. Little difference in dechlorination rates were observed
225 between amendments with 10 times more acetate. Similarly, higher production of acetate from solid
226 organic polymeric materials used to stimulate cDCE in riverbed sediment microcosms did not lead to
227 higher dechlorination rates as the excess organic loading was channelled to methanogenesis (Atashgahi
228 et al., 2014). These results highlight the complex role acetate plays in anaerobic food webs with

229 sometimes contradictory effects observed in relation to OHR rates.

230 **2.3 Interactions between OHR and alternative anaerobic TEAPs**

231 Interspecies H₂ transfers between fermenters and anaerobic TEAPs (Fig. 3) including denitrification,
232 metal reduction, sulfate reduction and methanogenesis (Fig. 4) may influence the rate and extent of OHR
233 in anoxic zones (Aulenta et al., 2007). Toxic metabolites generated may also inhibit steps of the
234 dechlorination process (Berggren et al., 2013). In this section, we review the current understanding of
235 these competitive interactions between OHR steps and biogeochemical cycling in the hyporheic zone.

236 **2.3.1 Nitrate reduction**

237 Groundwater nitrate has become elevated over the last century as a result of the 6.4-fold increase
238 in global inorganic fertiliser production between 1961 and 1999 (Peng et al., 2002). Heterotrophic
239 denitrification producing nitrous oxide (N₂O) or dinitrogen (N₂) and dissimilatory nitrate reduction to
240 ammonium (NH₄) occur in suboxic conditions (<125 μM O₂) with ambient H₂ levels (<0.1 nM) which
241 are below the level at which OHR becomes favourable for HCEs (~0.6 nM) (Fig. 4). Nitrate respiration
242 has only a slightly lower energy yield than that of oxygen with reaction timescales in the order of hours
243 to days (Rivett et al., 2008a; Jahangir et al., 2012) and has been widely documented in hyporheic
244 sediments (e.g. Ullah et al., 2014; Heppell et al., 2014). The interactions between OHR and nitrate
245 respiration is apparently complex. Yu et al. (2014) have shown that nitrate stimulated OHR of
246 pentachlorophenol at concentrations below 1 mM (likely by providing OHRB with a nitrogen source for
247 growth) but became inhibitory at concentrations greater than 1 mM. This was supported by Chen et al.
248 (2002) where 3 mM of nitrate inhibited hexachlorobenzene respiration under lactate-fermentation
249 conditions in river sediments. Nelson et al. (2002) found PCE reduction inhibition from only 0.6 mM of
250 nitrate and sulfate in a mixed culture where H₂ was continuously fed at non-limiting aqueous
251 concentrations (0.4–0.8 mM). In that study, the metabolite N₂O was found to be an inhibitor of OHR at
252 13 μM. These results are consistent with capture of available electrons by denitrifiers and reduction of
253 ambient H₂ levels below the minimum threshold necessary for OHRB as a key inhibition mechanism
254 which is also in line with thermodynamic predictions and observed H₂ thresholds (Fig. 4).

255 2.3.2 Iron(III) reduction

256 Iron oxides and oxyhydroxides such as haematite (Fe_2O_3) and goethite ($\text{FeO}(\text{OH})_3$) are abundant
257 primary and secondary minerals in streambed sediments which contain Fe(III) as a solid or colloidal
258 phase (Liu et al., 2014). Similarly, manganese oxide minerals such as pyrolusite (MnO_2) contain the
259 TEA Mn(IV) which behaves comparably to Fe(III) in redox processes (Ng et al., 2016). Some facultative
260 OHRB including *Desulfuromonas michiganensis* and *Geobacter lovleyi* (Sung et al., 2006) can also use
261 Fe(III) as a TEA. Dissimilatory Fe(III) reduction to Fe(II) has been observed at a similar H_2 threshold
262 range (0.1–0.8 nM) to HCE reduction (Fig. 4) and direct competition for electron equivalents is therefore
263 possible over reaction timescales measured months. Inhibition of TCE reduction as a result of Fe(III)
264 was demonstrated in microcosm studies prepared from dolomite aquifer materials (Yager et al., 1997).
265 Dupont et al. (2003) observed general inhibition of TCE reduction due to a large pool of bioavailable
266 Fe(III) minerals in microcosms prepared from contaminated aquifer sediment. In contrast, Wei and
267 Finneran (2011) demonstrated simultaneous reduction of TCE to ethene and Fe(III) to Fe (II) in
268 microcosms dominated by *D. mccartyi* and *Geobacter* with an excess of electron donor. Fe(III) was
269 observed to facilitate OHR by maintaining the ambient H_2 concentration at a level favourable for the
270 OHRB. Using the ‘KB-1’ organohalide-respiring enrichment culture (Duhamel et al., 2002), Paul et al.
271 (2013) showed that pH and iron mineral structure play an important role in inhibition of OHR in assays
272 of 14 different synthetic Fe(III) minerals using an excess of formate as an electron donor. High-surface
273 area, poorly crystalline minerals (such as ferrihydrite) have a greater Fe(III) fraction which is
274 bioavailable for electron transfer than highly crystalline minerals such as haematite and goethite. These
275 studies suggest the inhibition of OHR by Fe(III) in hyporheic sediments is complex and will be strongly
276 linked to pH, specific bioavailable iron mineral composition, surface area and sediment particle size
277 rather than total sediment Fe(III) content (Paul et al., 2013; 2016).

278 2.3.3 Sulfate reduction

279 Sulfate is a major oxyanion in groundwater derived from chemical denudation of sedimentary
280 rocks, oxidation of sulfide minerals and anthropogenic point sources. Dissimilatory sulfate reduction to
281 sulfide occurs at similar observed H_2 thresholds (1–15 nM) to HCE and LCE respiration (Fig. 4). The
282 presence of sulfate has been widely implicated where incomplete dechlorination leads to the

283 accumulation of LCEs (e.g. Bagley and Gossett, 1990; Pavlostathis and Zhuang, 1993; Lorah et al.,
284 2007) although laboratory studies of interactions between sulfate reduction and OHR are inconclusive.
285 For example, sulfate reduction was reported to affect rates of VC respiration and to a lesser extent cDCE
286 respiration (Boopathy and Peters, 2001; Aulenta et al., 2008; Pantazidou et al., 2012). Hoelen and
287 Reinhard (2004) observed respiration of TCE at sulfate concentrations of 1–2.6 mM in sediment
288 microcosm containing CEs and alkyl benzenes. TCE reduction was found to inhibit sulfate reduction
289 under an ambient H₂ concentration of 0.7 nM. Sulfate reduction at higher H₂ levels (2.5 nM) was
290 observed to compete with reduction of cDCE and VC which was sustained at a H₂ concentration of 1.6
291 nM. With constant H₂ concentrations (2–4 nM) sustained by lactate fermentation, Berggren et al. (2013)
292 found that the addition of sulfate affected the cDCE to VC respiration step the greatest (67%) in
293 comparison to VC to ethene (25%) and TCE to cDCE (8%). This work showed that sulfate addition
294 could produce a shift in the community structure of *D. mccartyi* leading to a reduction in OHR rates. In
295 contrast, an absence of any inhibitory effect under non-limiting electron donor conditions were shown
296 (e.g. Hoelen et al., 2006; Aulenta et al., 2007) and even sulfate addition was reported to strongly enhance
297 TCE respiration (Harkness et al., 2012). This may be a result of syntrophic partnership between sulfate
298 reducers and OHRB through provision of essential nutrients such as corrinoids (Men et al., 2012, Sutton
299 et al., 2015; Lu et al., 2017 Atashgahi et al., 2017a). A recent study on isolates, constructed consortia
300 and enrichments containing *D. mccartyi* showed that rather than sulfate, sulfide inhibited the growth of
301 *D. mccartyi* and its syntrophic partner, *Syntrophomonas wolfei* (Mao et al., 2017). Interestingly, in a co-
302 culture of *D. mccartyi* and sulfate-reducing *Desulfovibrio vulgaris* Hildenborough, a high sulfate
303 concentration (5 mM) was not inhibitory to OHR under electron donor (lactate) limitation likely due to
304 strong electron scavenging capacity of *D. mccartyi*. Moreover, at low sulfate levels (2 mM), sulfate
305 reduction was not inhibitory even under excess of electron donor (Mao et al., 2017).

306 2.3.4 Methanogenesis

307 Methanogenesis represents the final phase of anaerobic DOM metabolism (Figs 2 and 4) with
308 methane production timescales in riverbed sediments reported in the order of 200 days (Sela-Adler et
309 al., 2017). Its importance as a metabolic process in hyporheic zones is becoming increasingly recognised
310 (e.g. Sanders et al., 2007; Shelley et al., 2014; Brablková et al., 2015). Methanogens are dependent on

311 the same key substrates and electron donors as OHRB including acetate (via acetoclastic
312 methanogenesis) and H₂ (through hydrogenotrophic reduction of CO₂). The effects of acetoclastic
313 methanogenesis on OHRB is less well understood than hydrogenotrophic methanogenesis. Heimann et
314 al. (2006) suggested that acetoclastic methanogenesis may enhance VC dechlorination by providing
315 additional H₂ during acetate cleavage. The observed H₂ thresholds for hydrogenotrophic methanogenesis
316 (5–100 nM) are considerably greater than OHR (0.1–2.5 nM) (Paul et al., 2016) and OHRB have been
317 shown to out-compete hydrogenotrophic methanogens for reducing equivalents at low H₂ concentrations
318 (Ballapragada et al., 1997; Yang and McCarty, 1998; Azizian et al., 2010). CEs on the other hand have
319 been shown to have direct inhibitory effect on methanogenesis through enzyme inactivation although
320 the effect is less pronounced than for other organohalides such as chlorinated methanes (Chan and
321 Radom, 2011). Yu and Smith (2000) found that 137 μM of TCE could inhibit methanogenesis but no
322 effect was observed from PCE at 87 μM. In enrichment cultures containing fermenters, *D. mccartyi*
323 strain 195, homoacetogens and methanogens, Men et al. (2013) found that 22 μM of TCE caused
324 inhibition of hydrogenotrophic methanogenesis with an increase of electron flow to OHR by an average
325 of 7% under non-limiting H₂ concentrations (7–12 nM). Their results suggested that TCE could inhibit
326 homoacetogenesis at H₂ levels well above the optimal range for OHRB. That study showed that non-
327 methanogenic cultures generated significantly more ethene faster than methanogenic cultures and
328 suggested that fermenters (*Clostridium* spp.) may also supply essential nutrients (corrinoids) in addition
329 to H₂.

330 2.3.5 Co-mingled organohalides

331 An important consideration in the anaerobic transformation potential of hyporheic zones for CEs
332 is the presence of other organohalides in co-mingled groundwater plumes. The chlorinated ethane 1,1,1-
333 trichloroethane (1,1,1-TCA) introduced as an alternative to TCE and the chlorinated methane,
334 chloroform, (TCM) are frequent co-contaminants (Scheutz et al., 2011; Simsir et al., 2017). TCM is also
335 produced from natural processes in soils at low concentrations (<10 μM) (Laternas et al., 2002). Both
336 1,1,1-TCA and TCM have been implicated in the accumulation of LCEs and VC in particular (Duhamel
337 et al 2002; Chung and Rittmann, 2008; Chan et al., 2011). TCM has also shown to inhibit dechlorination
338 of cDCE by *D. mccartyi*. (Maymo-Gatell et al., 2001). Conversely, Grostern et al. (2009) showed that

339 VC and cDCE to lesser extent can cause inhibition in the respiration of chlorinated ethanes (CAs) (1,1,1-
340 trichloroethane and 1,1-dichloroethane) Mayer-Blackwell et al. (2016) found that prolonged exposure
341 to 1,2-dichloroethane (DCA) in a VC-respiring culture caused selective changes in *D. mccartyi*
342 community structure and reduced VC-respiration capacity. Strong inhibition of DCA respiration was
343 observed by cDCE. At field scale, Simsir et al. (2017) observed no inhibitory effect on the CE
344 dechlorination capacity of hyporheic sediments due to the spatial segregation of CE and CA
345 dechlorinators (demarcated by expression of the *cfrA* gene associated with CA respiration) along
346 discharging groundwater flow paths. CEs themselves may also competitively inhibit steps of the OHR
347 sequence (Chambon et al., 2013). In a kinetic study using two mixed anaerobic cultures, Yu et al. (2005)
348 found that HCEs generally inhibited the respiration of LCEs whilst the LCEs weakly inhibited the
349 dechlorination of the HCEs. These studies highlight the complex inter-relationship between different
350 organohalide respiration steps and their metabolites.

351 **2.4 Development and implications of anoxic-oxic interfaces**

352 The higher halogen substitution of HCEs produces electrophilicity with resistance to electrophilic
353 attack by oxygenase enzymes of aerobic bacteria. Thus, oxidation of the carbon backbone of the ethene
354 molecule is energetically unfavourable (Vogel et al., 1987). Schmidt et al. (2014) recently reported
355 aerobic TCE oxidation as a sole carbon source which may be evidence of a novel metabolic process not
356 witnessed before. Nonetheless, susceptibility to the reductive pathway decreases proportionately with
357 the ratio of chlorine to carbon substituents. As such, LCEs may be amenable to transformation via
358 aerobic mineralisation pathways during hyporheic exchange flows (Atashgahi et al., 2013). From a
359 natural attenuation perspective, mineralisation reactions are attractive in that no stable toxic
360 intermediates generated (Tiehm and Schmidt, 2011).

361 **2.4.1 Hyporheic oxygen gradients and mixing zones**

362 A unique feature of hyporheic zones is the establishment of oxic domains located where river water
363 infiltrates bringing dissolved O₂ into riverbed pore water at saturation concentrations (Knapp et al.,
364 2015). A common scenario illustrating development of a shallow oxic zone within a predominantly
365 anoxic groundwater regime is depicted in Fig. 5 based on the conceptual framework of Figs 1 and 2. In

366 this situation, river water advection is induced by bed topography in a process known as hyporheic
367 exchange flow where water currents infiltrate on the upstream side of a topographic feature
368 (downwelling zone) and return back to surface water a short distance downstream along with
369 groundwater discharge (upwelling zone) (Harvey and Bencala, 1993; Gomez-Velez et al., 2014; Trauth
370 et al., 2015). These bi-directional hyporheic flow cells (HFCs) will develop at various scales around bed
371 and channel topographic features such as gravel bars, riffles, weirs, debris dams and meanders (Krause
372 et al., 2014; Boano et al., 2014; Fox et al., 2014).

373 A thin mixing zone (Fig. 5); governed by dispersive processes is thought to develop along the
374 boundary between HFCs and the surrounding groundwater-dominated pore water regime which is
375 characterised by unidirectional vertical and lateral flow paths (Zarnetske et al., 2012; Briggs et al., 2013;
376 Binley et al., 2013; Boano et al., 2014). Numerical simulations of conservative tracer transport through
377 hydrologically gaining hyporheic zones (Hester et al., 2013) have suggested that the thickness of this
378 mixing zone is greatest with homogeneous high K sediment (e.g. well-sorted sand or gravel) and reduced
379 groundwater discharge (Fig. 5). Conversely, in zones of elevated groundwater discharge, low stream
380 velocity and smooth bed topography the mixing zone may be reduced or absent entirely (Trauth et al.,
381 2015). During high river flows, HFCs and associated mixing zones may transiently penetrate into deeper
382 sediment layers which is accompanied by a temporary surface water invasion of the groundwater regime
383 at depth (e.g. Cuthbert et al., 2010; Hamonts et al., 2012; Byrne et al., 2013; Freitas et al., 2015). These
384 transient flow events may be particularly acute for urban rivers where peak flows are accentuated by
385 impervious catchment areas and channel canalisation (Meyer et al., 2005).

386 Advection of river water solutes including O_2 and DOM into bed sediments induces zones of
387 aerobic respiration (Fig. 5) where the reaction timescale is often rapid with zero-order rates reported
388 from 9 and 75 $\mu\text{M}/\text{h}$ in a gravel bar (Vieweg et al., 2016). Where mixing occurs, an anoxic-oxic interface
389 will develop which shifts dynamically in response to the balance of surface water downwelling and
390 groundwater discharge pressures. Approximately 0.25–0.33 mM of DOC is required to deoxygenate
391 downwelling river water and other reductants from the anoxic zone (Fig. 4) such as Fe(II) and reduced
392 sulfur species may also consume O_2 . Hence, conditions for a spatially fluctuating ‘reactive fringe’

393 (Trauth et al., 2014) where complementary aerobic co-metabolic or growth-coupled assimilation of
394 cDCE and VC may be favoured. As a result, hyporheic zones can offer a spatially variable secondary
395 aerobic 'treatment' zone facilitating complete mineralisation of LCE metabolites (Atashgahi et al., 2013;
396 2017a).

397 **2.4.2 cDCE mineralisation**

398 Although cDCE is amenable to aerobic biotransformation, cDCE oxidation is not widely reported.
399 Bradley and Chapelle (1998a) were among the first to examine the potential for aerobic cDCE
400 mineralisation in microbial communities indigenous to riverbeds. Their study demonstrated biological
401 aerobic mineralisation of [¹⁴C]-DCE (4:1 *cis-to-trans* isomers) to CO₂ in microcosms with recovery of
402 [¹⁴C]-CO₂ ranging from 17–100% after just eight days. However, this study used natural sediment media
403 potentially containing a range of other substrates where co-metabolic processes could not be ruled out
404 (Bradley and Chapelle, 2000). Other work has shown aerobic cDCE mineralisation to proceed much
405 more slowly in aquifer media e.g. Klier et al. (1998) where just 3–10% mineralisation was achieved
406 after 180 days incubation. Moreover, Abe et al. (2009) found no evidence of cDCE removal in aerobic
407 riverbed microcosms incubated over 1.5 years. Such patchy metabolic cDCE aerobic degradation is
408 illustrated by the fact that only a single isolate, *Polaromonas* sp. JS666 has unequivocally shown to
409 grow by cDCE assimilation as the sole carbon and energy source (Coleman et al., 2002). With the advent
410 of compound-specific stable isotope analysis (CSIA), kinetic isotope fractionation of carbon ($\delta^{13}\text{C}$ -
411 cDCE) has provided an additional line of evidence that cDCE mineralisation may be a growth-linked
412 assimilatory process. Reported $\delta^{13}\text{C}$ enrichment factors for aerobic uptake of cDCE range from -7.1‰
413 to -22.4‰ for both mixed and pure cultures (Tiehm et al., 2008; Abe et al., 2009; Schmidt et al., 2010).
414 Biomass yields from cDCE uptake are reported to range from 6.1 to 12.5 g/M cDCE (Coleman et al.,
415 2002; Schmidt et al., 2010).

416 **2.4.3 VC mineralisation**

417 VC mineralisation is the most thermodynamically attractive CE oxidation process due to the
418 presence of only one chlorine substituent. Reaction timescales tend to be relatively rapid as a result. In
419 a radiolabelled microcosm study prepared from riverbed sediments (Bradley and Chapelle, 1998a), it

420 was shown that greater than 90% of VC was consumed after 12 days over a concentration range of 0.2
421 – 57 μM with uptake rates most adequately described by Michaelis-Menten kinetics. Bacteria capable
422 of aerobic VC oxidation as a sole carbon and energy source belong to a range of genera associated with
423 aerobic ethene assimilation (Mattes et al., 2010; Atashgahi et al., 2017a). Aerobic degradation of VC
424 and ethene are initiated by an alkene monooxygenase (EtnABCD) and an epoxyalkane-coenzyme
425 enzyme (EtnE) (Mattes et al., 2010). Various isolates have been identified including *Mycobacterium*
426 (Hartmans et al., 1985; Hartmans and DeBont, 1992; Fullerton et al., 2014; Le and Coleman, 2011),
427 *Pseudomonas* (Verge et al., 2000; 2001; Atashgahi et al., 2017a), *Ochrobactrum* (Danko et al., 2004,
428 Atashgahi et al., 2017a), *Nocardioides* (Coleman et al., 2002) and a *Ralstonia* (Elango et al., 2006).
429 Besides these classical VC-assimilators, stable isotope probing coupled to high-throughput 16S rRNA
430 gene sequencing and quantitative polymerase chain reaction (qPCR) extended the potential range of VC
431 assimilators to *Sediminibacterium*, *Aquabacterium*, *Variovorax*, *Brevundimonas*, *Tissierella*, and
432 *Rhodoferax*. (Paes et al., 2015; Wilson et al., 2016). Gossett (2010) found that biological VC respiration
433 could be sustained by *Mycobacterium* at extremely low (microaerophilic) extracellular O_2
434 concentrations (e.g. 0.3–0.6 μM). This finding raises the important question as to whether anoxic
435 conditions classically defined by O_2 concentrations less than 3 μM (Chapelle and McMahon, 2006) are
436 in fact still capable of supporting aerobic metabolism. Microaerophilic VC oxidation may also play a
437 role in the apparent ‘stalling’ of the dechlorination sequence at cDCE. A lack of VC detection may in
438 fact reflect scenarios where VC mineralisation rates equal or exceed the production rate under hypoxic
439 conditions (Bradley and Chapelle, 2011). In a recent microcosm study by Fullerton et al. (2014) subject
440 to inadvertent O_2 contamination, VC disappearance was observed without ethene formation. A strictly
441 aerobic *Mycobacterium* was identified in high numbers in the groundwater microcosm which was
442 inadvertently exposed to O_2 . Atashgahi et al. (2013, 2017a) used 16S rRNA of *D. mccartyi* and genes
443 encoding for reductive dehalogenase enzymes (*vcrA* and *bvcA*) and the genes *etnC* and *etnE* involved in
444 aerobic mineralisation of VC and ethene to track the fate of VC in microcosms prepared from riverbed
445 sediments obtained from anoxic-oxic interface. Using a combination of chemical analysis and qPCR,
446 the authors demonstrated co-occurrence and co-activity of aerobic VC degraders and anaerobic *D.*
447 *mccartyi* in hyporheic sediments of the eutrophic Zenne River in urban Belgium (Atashgahi et al.,

448 2017a).

449 **2.4.4 Aerobic co-metabolism**

450 CEs can be transformed under aerobic conditions by co-metabolic reactions initiated by common
451 enzymes produced by a wide range of aerobes (Arp, 1995). No known benefit is gained through
452 acquisition of metabolic energy or biomass production (Horvath 1972; Wacket, 1988; Semprini et al.,
453 1994). Co-metabolism is a widespread process by microbes containing non-specific monooxygenase or
454 dioxygenase enzymes which catalyze the initial step in oxidation of a growth-supporting substrate in the
455 presence of O₂ (Suttinun et al., 2013). A lack of enzyme specificity leads to competition between CEs
456 and growth-supporting substrates for active sites of oxygenases leading to incorporation of O₂ into the
457 chloroethene molecule via reactive epoxide formation (Mattes et al., 2010). TCE, cDCE and VC are all
458 amenable to co-metabolic aerobic oxidation by the action of oxygenases. PCE was originally thought to
459 be recalcitrant to aerobic oxidation under environmental conditions, however it is shown to be
460 degradable by the enzyme toluene-*o*-xylene monooxygenase (Ryoo et al., 2000; Shim et al., 2001).

461 In hyporheic sediments, methane and ammonium are often the most abundant potential substrates
462 for co-metabolism produced from the in-situ anaerobic decomposition of DOM (Figs 2 and 5) (Donn
463 and Barron, 2013; Atashgahi et al., 2013; Brablcová et al., 2015; Simsir et al., 2017). Extensive
464 hyporheic methanotrophy has been documented in association with fine-grained sediments under
465 vegetation stands in streams (Sanders et al., 2007) and in well-oxygenated coarse-grained sediments
466 (Trimmer et al., 2010). The enzyme methane monooxygenase is produced by the methanotroph
467 *Methylosinus trichosporium* and has been shown to oxidise TCE, cDCE, VC and ethene (Oldenhuis et
468 al., 1991; Forrester et al., 2005; Findlay et al., 2016). Conrad et al. (2010) observed evidence of co-
469 metabolic TCE oxidation at field-scale in a contaminated aquifer with an increase in *M. trichosporium*
470 abundance accompanied by a decline in TCE and dissolved methane concentrations. A co-metabolic
471 process was verified in a supporting microcosm study prepared from aquifer media and isotopically-
472 labelled TCE (¹³C-TCE). Simsir et al. (2017) presented evidence of possible hyporheic zone co-
473 metabolism with declines in cDCE and VC associated with strong vertical methane gradients (Fig. 6)
474 although no O₂ data are presented for comparison. Their study also documented the presence of

475 methanotrophs (*Methylococcaceae* spp.) coincident with the change in methane concentrations in the
476 sediment sequence. Nitrifying bacteria which oxidise ammonium produced from DOM metabolism (Fig.
477 2) such as *Nitrosomonas europaea* can also inadvertently oxidise TCE and LCEs (Arciero et al., 1989;
478 Vannelli et al., 1990; Rasche et al., 1991; Kocamemi and Cecen, 2005). Nitrification is well documented
479 in association with hyporheic flow cells (Jones, 1995; Briggs et al., 2013). Kocamemi and Cecen (2005)
480 found that ammonium and TCE compete for the same active sites of the enzyme ammonia
481 monooxygenase. In a mixed culture study containing *Nitrosomonas europaea*, Kocamemi and Cecen
482 (2010) found that the transformation yield for TCE was strongly dependent on initial ammonium and
483 TCE concentrations, with a minimum initial level of 3 μM of ammonium required to transform the
484 maximum initial TCE concentration of 7 μM . We are not aware of any studies which have considered
485 hyporheic nitrification as a natural attenuation process for CEs under field conditions.

486 **2.4.5 Interaction between aerobic and anaerobic VC transformation pathways**

487 Ethene-assimilating bacteria (ethenotrophs) can oxidise ethene as the sole carbon and energy
488 source and simultaneously transform VC co-metabolically whereas VC-assimilating bacteria can use
489 VC as the sole carbon and energy source (Mattes et al., 2010). Therefore, when anoxic groundwater
490 containing VC and ethene migrates through the mixing zones around HFCs, there is a potential for
491 assimilation of both substrates by ethenotrophs and VC-assimilators and also co-metabolic VC
492 degradation by ethenotrophs growing on ethene (Atashgahi et al., 2017a). Such synergetic interactions
493 can be further enhanced by methanotrophs that can oxidize both ethene and VC. In line with this, a
494 recent study using groundwater microcosms showed that when methane, ethene and VC were added to
495 microcosms, the rate of VC removal was faster than with either methane or ethene alone, consistent with
496 the idea that methanotrophs stimulate ethenotrophic removal of VC (Findlay et al., 2016). Moreover,
497 recent studies have shown that reductive dechlorination can impact aerobic VC degradation pathways
498 in hyporheic zones. Microcosms prepared from fine-grained, SOM-rich sediments with high reductive
499 dechlorination capacity did not have the potential for metabolic aerobic VC oxidation (Atashgahi et al.,
500 2013, 2017a). Under atmospheric O_2 conditions, *D. mccartyi* was protected from O_2 toxicity by the
501 sediment structure and grew by reductive dechlorination of VC. No ethene accumulation was noted
502 indicating the activity of ethenotrophs (Atashgahi et al., 2013, 2017a) which can, in turn, co-

503 metabolically oxidise VC (Findlay et al., 2016). In contrast, metabolic aerobic VC activity was observed
504 from microcosms prepared from sediments with low SOM, coarse grain size and low reductive
505 dechlorination potential. The study showed that local sediment geochemistry and reductive
506 dechlorination highly impact metabolic versus co-metabolic aerobic VC degradation pathway. Where
507 low reductive potential was noted, ethenotrophs exposed to continuous VC flow in oxic sediment layers
508 might adapt to metabolic aerobic VC degradation (Atashgahi et al., 2017a). These findings are consistent
509 with upon extended exposure to VC, ethenotrophic strains of *Mycobacterium* (Jin and Mattes, 2008) and
510 *Pseudomonas* (Verge et al., 2001) transit from co-metabolic to growth-linked VC mineralisation.

511 **3 Field-scale conceptualisation challenges and potential solutions**

512 **3.1 Spatial variability of potential transformation zones**

513 **3.1.1 OHRB occurrence in riverbed sediments**

514 qPCR approaches targeting 16S rRNA genes and reductive dehalogenation catabolic genes (*rdhA*)
515 (Lu et al., 2015) have been used to track the presence and activity of OHRB in hyporheic zones
516 (Vandermeeren et al., 2014). These biomarkers have revealed a close association with OHRB, SOM
517 content and grain-size distribution in organohalide-impacted hyporheic sediments (Atashgahi et al.,
518 2013; Kranzioch et al., 2013; Hamonts et al., 2014; Atashgahi et al., 2015). In a survey of OHRB in
519 relation to hexachlorobenzene (HCB) fate in 15 riverbed sediment locations across four European
520 catchments, Taş et al., (2011) detected *D. mccartyi* 16S rRNA genes at 80% of sites though their counts
521 did not surpass 0.1% of total bacterial 16S rRNA gene copies. HCB half-lives ranged from a few days
522 to up to one month. A relatively weak correlation was observed between decay rate and number of *D.*
523 *mccartyi* suggesting additional OHR activity from other OHRB. Atashgahi et al. (2013) used 16S rRNA
524 of *D. mccartyi* and genes encoding for VC reductive dehalogenase enzymes (*vcrA* and *bvcA*) to track
525 dominant VC biotransformation processes in microcosms prepared from hyporheic sediments. *D.*
526 *mccartyi* had a dominant role on VC removal in microcosms prepared from coarse-grained bed
527 sediments with an abundance of SOM. A subsequent in situ study at the same site showed a reduced VC
528 respiration potential due to construction of a wastewater treatment plant upstream of the study area that
529 resulted in a reduction of organic matter loading to the riverbed sediments (Atashgahi et al., 2015).

530 Similarly, PCR detection of *D. mccartyi* was reported to be closely linked with SOM content and
531 sediment particle size (Abe et al., 2009). This work showed that *D. mccartyi* presence in the sediment
532 profile was spatially correlated with SOM where it exceeded 2% (g/g) with stronger signals occurring
533 at greater depth. A recent study by Simsir et al. (2017) investigated CEs discharging to a creek in
534 Tennessee from organohalide-contaminated fractured bedrock (Table 1). Over a 300 m river section,
535 16S rRNA genes of *D. mccartyi* and *Dehalobacter* were found to be more abundant in deeper sediment
536 layers (50 cm) coincident with higher concentrations of cDCE and VC. Their study revealed the presence
537 of multiple strains of *D. mccartyi* occurring together and expressing functional genes involved in
538 respiration of CE (e.g. *tceA*, *bvcA* and *vcrA*) and chlorinated ethane/methane (*dcpA*).

539 3.1.2 Plume-scale discharge and transformation controls

540 Groundwater plume discharge and transformation patterns are controlled by the spatial
541 distribution and continuity of sediment domains with distinct hydraulic properties along river corridors
542 (Fleckenstein et al., 2006; Wondzell, 2011). The available field investigations (Table 1) have shown that
543 low-K sediments (e.g. clay and silt) are the most important local controls on contaminant discharge in
544 heterogeneous sediment sequences. For example, at the Pine River PCE plume site in Ontario (Conant
545 et al., 2004; Abe et al., 2009) and for a TCE plume discharging to the River Tern in the UK (Weatherill
546 et al., 2014), the lateral and longitudinal continuity of semi-confining silty deposits were shown to
547 control the location and magnitude of plume discharge (Table 1, Fig. 6). Where these low-K sediments
548 also contain SOM, conditions for enhanced biotransformation may also occur where residence times are
549 greatly extended due to hydrophobic partitioning to the organic carbon present. This is demonstrated by
550 Conant et al. (2004) and Abe et al. (2008) for the Pine River site with extensive in-situ transformation
551 of PCE or TCE to cDCE (Fig. 6) reported. The importance of low-K layers in contaminant
552 transformation has been illustrated in a modelling study by Gomez-Velez et al. (2014). Their work has
553 shown that the interfaces of sediments with large K contrasts can sequester upwelling groundwater by
554 creating stagnation zones which facilitate mixing of waters with different residence times. Hyporheic
555 exchange patterns arising from riverbed topography have also been shown to influence patterns of in-
556 situ transformation of discharging groundwater plumes. With the benefit of chloride as a conservative
557 surface water tracer, Freitas et al. (2015) observed TCE transformation associated with coarse grained

558 sediment along a riffle sequence of the River Tame (Birmingham, UK) where river water infiltration
559 carrying DOC was inferred from chloride profiles. In one location (Fig. 6) up to 80% TCE conversion
560 to ethene was reported. In-situ dechlorination was also shown to be largely absent along the investigation
561 corridor where riverbed topography was subdued.

562 The presence of preferential pathways with short residence times (such as springs) afford little
563 opportunity for in-situ natural attenuation (e.g. Fryar et al., 2000; LaSage et al., 2008; Rønde et al.,
564 2017). This mode of contaminant discharge arises where discontinuities occur in low-permeability
565 alluvial architecture that act as high-flux ‘geological windows’ (Conant et al., 2004; Weatherill et al.,
566 2014). For example, through temperature mapping, Conant (2004) found that nearly a quarter of the
567 baseflow accretion along a reach of the Pine River in Angus occurred from just 7% of the riverbed area.
568 The available case studies (Table 1) highlight the patchy nature of natural attenuation zones where the
569 mass fluxes to surface water are dominated by small areas through which most groundwater baseflow
570 discharge is concentrated.

571 3.1.3 Pore-scale mass transport and processing

572 Microbial metabolic activity and diversity in riverbeds has been reported to be concentrated in
573 the uppermost 50–60 cm of hyporheic sediment sequences (Franken et al., 2001; Freixia et al., 2016).
574 Heterotrophic microbes (including OHRB) are predominantly found attached to sediment surfaces in
575 epilithic biofilms (Davey and O’Toole, 2000). Hyporheic biofilms comprise diverse assemblages of
576 fungi, algae and microbial consortia enveloped by a matrix of extracellular polymeric substances (EPS)
577 through which diffusive transport of solutes predominates (Storey et al., 1999; Battin et al., 2016).
578 Biofilm structures have been shown to retain and process complex DOM in sediments (Fischer et al.,
579 2005; Bengtsson et al., 2014) and may be important sites of SCFA fermentation (Rulík and Hereka,
580 1998; Rulík et al., 2000). Biofilm accumulation in interstitial spaces is known to decrease the bulk K
581 and effective porosity of sediments in downwelling zones, thereby reducing hydrodynamic exchange
582 with the overlying water column (Battin et al., 1997; 2003; 2008; Mermillod-Blondin et al., 2005) and
583 inducing redox gradients (Boano et al., 2014). Biofilm development may be facilitated in downwelling
584 streambed zones during the clogging of coarse sediment pore spaces (colmation) by non-settlable POM

585 originating from surface water sources (Navel et al., 2012).

586 The convolution of flow paths through heterogeneous biofilm-enveloped sediments produces a
587 scale dependence in chemical reaction rates where microbial processes are superimposed on larger-scale
588 advective solute fluxes that deliver reactants (Sobczak and Findlay, 2002; Findlay et al., 2003; Nogaro
589 et al., 2013). Mendoza-Lera et al. (2017) propose a conceptual framework unifying hyporheic flow and
590 biogeochemical transformation capacity in riverbeds which is applicable for both anaerobic and aerobic
591 biotransformation of CEs. They propose that biogeochemical processing is controlled by the mass
592 transfer of key reactants (TEAs, fermentable DOM and nutrients) and the sediment surface area
593 available for biofilm colonisation. Mass transfer is characterised in three stages, the slowest of which
594 will be the rate-limiting step for biotransformation: (1) transfer from bulk groundwater or surface water
595 to the hyporheic environment (2) transfer from sediment pore water into microbial biofilms and (3)
596 membrane transfer from the extracellular biofilm matrix into individual cells. The sediment particle size
597 controls the surface area for colonisation which is inversely proportional to K . High K sediments (such
598 as gravel bars) exhibit high mass transfer rates but low processing capacities. Conversely, low K
599 sediments (e.g. silts and clays) have very high areas for colonisation due to the huge surface area to
600 sediment volume ratio present but mass transport is slow and diffusion-limited. Their model assumes
601 that solute uptake rates follow Michaelis-Menten kinetics where microbial activity increases
602 proportionally with the mass transfer rate until a saturation point is reached (Ribot et al., 2013). This
603 framework supports field observations where enhanced dechlorination activity and OHRB are reported
604 in sediment domains where diffusive transport is locally dominant (Conant et al., 2004; Abe et al., 2009;
605 Damgaard et al., 2013; Atashgahi et al., 2015; 2017a). Biofilm colonisation is also influenced by changes
606 in pore water temperature, pH and light exposure as well as sediment surface roughness (Gette-Bouvarot
607 et al., 2015; Voisin et al., 2016).

608 Delivery of CEs into reactive diffusion-dominated pore water systems will be dependent on the
609 presence, connectivity and residence time of macropore flow paths which facilitates advective transfer
610 from bulk upwelling groundwater (e.g. Bohlke et al., 2007; Chambon et al., 2010; Menichino and Hester,
611 2015). Heterogeneity in grain size distribution, particle packing and sphericity can induce locally anoxic

612 or hypoxic microzones where mass transfer rates from bulk water to biofilms vary at the scale of
613 sediment pores (Briggs et al., 2013; 2014). Anoxic microzones have been widely implicated where
614 mixed TEAPs and fermentation reactions appear to occur simultaneously (e.g. Storey et al., 1999; Baker
615 et al., 2000; Mermillod-Blondin et al., 2005) and may explain observed co-activity of aerobes and
616 anaerobes (e.g. Atashgahi et al., 2017a). Using two-dimensional dual-domain pore network models of
617 advective-diffusive transport, Briggs et al. (2015) showed that pore-scale heterogeneity (variation in
618 pore throat size) leads to zones of increased and decreased solute mobility in hyporheic in hyporheic
619 sediments. In the less mobile domain (which is responsible for observed tracer breakthrough tailing),
620 anoxia develops where O_2 consumption exceeds the re-supply rate. These microzones may be
621 cumulatively important in determining the overall transformation capacity of hyporheic sediments. For
622 example, LCEs produced anaerobically in the low-mobility domain are mineralised in the more mobile
623 aerobic domain. This is particularly important in the case of VC which can require only trace oxygen
624 levels to sustain aerobic uptake (Gossett, 2010; Fullerton et al., 2014).

625 **3.2 Multi-scale in-situ characterisation technologies**

626 **3.2.1 Evaluating hydrological connectivity**

627 The natural flow regime heterogeneity of most plume discharge zones makes the design and
628 implementation of effective in-situ monitoring programs non-trivial (Kalbus et al., 2006; Rivett et al.,
629 2008b; Roy and Bickerton, 2010; Burk and Cook, 2015). Heterogeneous sediment sequences present a
630 common challenge for the development of conceptual models of plume fate (Ellis and Rivett, 2007).
631 Direct sediment core sampling can provide intact samples in cohesive sequences but is impractical where
632 granular deposits occur. Freeze-coring methods have been applied to characterise fluvial sediment
633 sequences with mixed grain size distributions (Freitas et al., 2015). However, this extractive technique
634 necessitates disruption of sediment microstructure and associated transport pathways (Descloux et al.,
635 2010). Non-invasive hydrogeophysical 'imaging' technologies have emerged which offer promising
636 insights into the in-situ spatial variability of mass transport pathways at <1 m to >10 m scales. For
637 example, Mermillod-Blondin et al. (2014) used ground-penetrating radar (GPR) to map biologically
638 active domains within a gravel bar sequence of the River Rhone in France. Electrical resistivity imaging
639 technology has been used to image flow-controlling sedimentary structures and electrically conductive

640 solute plume migration at reach-scale (Nyquist et al., 2008; González-Pinzón et al., 2015). Recently,
641 Briggs et al. (2014) applied geoelectrical hysteresis techniques to characterise pore-scale rate-limiting
642 mass transport through dual domain porous media (mobile and less mobile porosity fractions).

643 Temperature mapping using contact probes or continuous optical fibres and has been used to
644 evaluate patchy sub-metre scale connectivity between groundwater and surface water in plan-view when
645 a seasonal thermal gradient is present (Conant, 2004; Tristram et al., 2014; Rosenberry et al., 2016).
646 Groundwater baseflow fluxes through riverbeds have been modelled using one dimensional advective-
647 diffusive heat flow as a surrogate for Darcian flow in grids of riverbed temperature profiles (Schmidt et
648 al., 2006; 2007) where upwelling flow predominates. Both upwelling and downwelling fluxes have more
649 recently been obtained from vertical profiles of riverbed temperature time series (e.g. Anibas et al.,
650 2011; 2016; Munz et al., 2016). Heat pulse injections have been used to understand three-dimensional
651 water fluxes through shallow sandy hyporheic sediments at centimetre scales (Angermann et al., 2012).
652 Recently, point-velocity probing (PVP) has been used to map hydrological connectivity with a
653 comparable resolution to temperature mapping (Rønne et al., 2017). Overall, hydrological connectivity
654 will be highly site-specific and the most useful results will be obtained where multiple techniques are
655 employed with overlapping nested scales (e.g. Milosevic et al., 2012).

656 **3.2.2 Resolving in-situ biogeochemical gradients**

657 Delineation of in-situ vertical chemical gradients has been accomplished using depth-discrete
658 profiling in riverbeds (Fig. 6) over vertical length scales of 0.1 to 3 m (Conant et al., 2004; Rivett et al.,
659 2008b; Krause et al., 2013; Heppel et al., 2014). Multi-port monitoring wells have achieved finer vertical
660 sampling resolution (<0.1 m) in contaminated aquifers (e.g. Jobelius et al., 2011). However, technical
661 challenges and cost implications are often likely to prohibit monitoring installations in riverbeds which
662 require a drilling rig. Instead, hand-portable lightweight installations utilising drive-point and direct-
663 push methods usually offer the most cost-effective solutions (Rivett et al., 2008b; Roy and Bickerton,
664 2010). CE fate in hyporheic systems has been investigated using the 'waterloo profiler' (Pitkin, et al.,
665 1999; Conant et al., 2004) and other pore water samplers (e.g. Hamonts et al., 2012), multi-level sampler
666 bundles (Conant et al., 2004), nested piezometers (Lorah and Olsen, 1999) and drive-point mini-

667 piezometers (Conant et al., 2004; Ellis and Rivett, 2007; Roche et al., 2008). Rivett et al. (2008b) and
668 Typical drive-point methods utilise a temporary casing to protect screens and sampling inlets during
669 installation. Collapse of adjacent sediments is relied upon to provide sufficient sealing action to prevent
670 any preferential flow along this ‘short-circuit’ pathway. This limitation is overcome with direct-push
671 methods such as that described by Roy and Bickerton (2010). They developed a miniaturised Waterloo
672 profiler optimised for rapid acquisition of riverbed sediment pore water using a portable hammer drill.
673 From Fig. 6 and Table 1, it can be seen that the selection of appropriate vertical monitoring scales is
674 site-specific and must be informed by an initial conceptual understanding of larger scale water fluxes.
675 Recent field experience suggests vertical resolutions of 0.1 – 0.2 m may be sufficient to capture bulk
676 gradients in pore water chemistry (Fig. 6).

677 A potential monitoring gap exists at μm – mm scales below the sediment-water interface where
678 preservation of natural stratification in pore water chemistry is impossible with extractive sampling
679 techniques. Transformation boundary zones at thin fine-grained sediment strata and critical mixing
680 zones may be overlooked by sampling at coarser resolutions (Hester et al., 2017). In-situ passive
681 sampling technology utilising diffusive pore water equilibration offers a possible means to address this
682 scale gap. Dialysis ‘peepers’ can provide high-resolution solute profiles using a chambered sampler
683 design (Lewandowski et al., 2002; Tan et al., 2005) and have been used to investigate CE plume fate at
684 centimetre-scale (Lorah and Olsen, 1999; LaSage et al., 2008). Peepers require up to several weeks for
685 concentration equilibration to be achieved between ambient pore water and dialysis cell water during
686 which the sampler is vulnerable to disturbance and transient effects (Tan et al., 2005). Simsir et al.
687 (2017) used direct push diffusion samplers (Fig. 6) comprised of glass vials with polyethylene or non-
688 woven porous fabric deployed into shallow creek sediment in Tennessee which were equilibrated for
689 two weeks. Passive sampling technology using hydrogel media requires much shorter equilibration
690 times (24–72 hours) and offers a promising way forward to address the mm monitoring scale gap.
691 Diffusive equilibration or gradient sampling in thin films (DET/DGT) has provided mm-scale depth
692 profiles of inorganic nitrogen species, phosphorus and metals in stream sediments (Davison et al., 1994;
693 Palmer-Felgate et al., 2010; Ullah et al., 2012). This technology has recently been applied to studies of

694 enhanced attenuation of groundwater nitrate in riverbeds influenced by in-stream vegetation growth
695 (Ullah et al., 2014). As Briggs et al. (2015) duly point out, extractive sampling procedures tend to always
696 bias the mobile porosity domain of reactive sediment zones which is overcome by diffusive thin film
697 approaches. This technology offers the best opportunity to capture vertical scales approaching the size
698 of pore networks where key reactive transport features such as anoxic microzones may be resolved. As
699 such, this technology can provide a nested observational capability when combined with larger scale
700 solute profiles for a more integrated understanding of plume behaviour.

701 Given that microbial biomass occurs predominantly as biofilms, microbial sampling in the
702 hyporheic zone is sensitive to particle size distribution given that the surface area available for
703 colonisation decreases with increasing particle size (Mendoza-Lera et al., 2017). Direct sampling of fine
704 sediment fractions for molecular analyses of microbial composition have been taken from surficial
705 sediment layers using piston corers (Atashgahi et al., 2012; Hamonts et al., 2014; Vandermeeren et al.,
706 2014) as well as high resolution sub-sampling of deeper cores (Abe et al., 2009; Atashgahi et al., 2014).
707 These methods are well suited to homogenous silt and clay sediments but are subject to the same
708 limitations for coring in granular deposits discussed above. Microbial 'trapping' using an emplaced
709 porous medium for biofilm colonisation is an established approach in aquifer media (Voisin et al., 2016).
710 At the Tennessee creek, Simsir et al. (2017) were able to measure in-situ changes in microbial
711 community structure at high spatial resolution using Bio-Sep bead traps (www.microbe.com) arranged
712 along a 55 cm vertical profile. By combining microbial and pore water sampling (Fig. 6), their work
713 permitted a comprehensive picture of microbial community structure and biogeochemical gradients to
714 be formed. This approach can offer unique insights on linkages between chemical and microbial factors
715 which govern in-situ biotransformation capacity and is suitable for deployment at a range of scales.

716 **4 Conclusions and outlook**

717 **4.1 Conclusions**

718 Our review has established the degree to which hyporheic zones can serve as natural bioreactors
719 capable of reducing mass fluxes of CEs to surface water from groundwater sources. Although OHRB
720 appear to be relatively widespread in riverbeds, field experience to date has shown that in-situ anoxic

721 biotransformation tends to be patchy (Table 1) where fine-grained or SOM-rich sediments occur (Fig
722 1). Observed transformation extents range from partial TCE to cDCE dechlorination to near-complete
723 stoichiometric conversion of TCE to ethene (Fig. 6). In DOM-dominated foodwebs, OHR requires
724 residence times along discharging groundwater flow paths to exceed the reaction timescales for higher-
725 energy TEAs such as O₂ and nitrate. Here, extracellular H₂ will be closely regulated by interspecies
726 transfers between fermenters and heterotrophs (Fig. 3) such as OHRB and methanogens. Competitive
727 interactions between dechlorinators and mineral reducing TEAPs (particularly sulfate) may inhibit OHR
728 steps, particularly cDCE to VC and VC to ethene (Fig. 4). When H₂ is limiting, OHR may compete with
729 these TEAPs for reducing power. Toxic metabolite formation (e.g. sulfide) may cause inhibition when
730 H₂ is unlimited. Conversely, syntrophic partnerships may develop between OHRB and other
731 hydrogenotrophs which may enhance in-situ biotransformation.

732 Thin mixing zones developing between discharging groundwater and downwelling surface water
733 carrying O₂ may create a dynamic anoxic-oxic interface potentially supporting ethenotrophs,
734 methanotrophs and ammonia oxidisers (Fig. 5). Here, a secondary niche environment for VC
735 mineralisation and to some extent, cDCE via direct metabolism and co-metabolism (from the action of
736 mono-oxygenase enzymes) can complete a coupled anoxic-oxic natural 'treatment' process. The oxic
737 potential of CE attenuation in hyporheic zones has only in the last few years become appreciated but is
738 not widely demonstrated at field scale. The efficacy of this step in the process will be dependent on the
739 presence of suitable aerobes and the thickness of mixing zones relative to discharging groundwater flow
740 paths transporting LCEs and co-substrates such as methane and ammonium produced from DOM
741 metabolism.

742 For both anoxic and oxic zone processes, mass transformation will be controlled by the sediment
743 surface area available for biofilm colonisation, delivery rates of critical reactants (e.g. H₂ or O₂), their
744 reaction timescales and exposure time in reactive zones. Diffusion-limited mass transfer from bulk pore
745 water to biofilm domains is likely to be the most important rate-limiting step governing in-situ
746 biotransformation efficacies. At pore scales, heterogeneity in pore throat size can induce dual mobility
747 porosity domains. Solute exposure timeframes are increased in the less mobile domain where anoxic

748 microsite formation may play an important role in the cumulative impact of processes.

749 Effective in-situ monitoring of hyporheic processes in plume discharge zones necessitates a
750 detailed conceptual understanding of the spatial and temporal variability of water flows and residence
751 times. This is best accomplished using non-invasive hydrogeophysical technologies and the application
752 of heat as a surrogate for advection where overlapping spatial scales may be observed simultaneously.
753 Bulk chemical gradients may be delineated using multi-level profiles with vertical resolutions of 0.1–
754 0.2 m often proving sufficient. Nested monitoring is warranted at critical boundary zones where the
755 local properties of pore networks may exert important cumulative controls on observed
756 biotransformation capacities. Simultaneous microbial and pore water sampling can demonstrate that
757 niche conditions for biotransformation are occurring in-situ and is advocated further.

758 **4.2 Future research opportunities**

759 Additional field experience is required to better address spatial and temporal biotransformation
760 variability and allow predictive models to be developed. Directions to improve the current conceptual
761 understanding are outlined as follows:

- 762 • Integrated field studies addressing coupled flow and reactive transport along pathways at
763 multiple scales are needed (Oldham et al., 2013; Pinay et al., 2015; Abbot et al., 2016). New in-
764 situ tools have emerged which can quantify residence times at high spatial resolution (e.g. Rønde
765 et al., 2017). Dual reactive-conservative ‘smart’ tracer studies such as acetate-bromide (Rinehart
766 et al., 2015), acetate-resazurin-resorufin (Briggs et al., 2013) which can quantify both residence
767 time and reaction timescales for TEA-DOC couples offer an integrated way forward.
- 768 • Capturing in-situ spatial and temporal dynamics of O_2/H_2 gradients along discharging
769 groundwater flow paths. These key end-members alone can provide the critical diagnostic
770 information on redox status and indications of key tipping points and priming events which
771 govern in-situ anaerobic-aerobic biotransformation efficacies.
- 772 • Finally, these recent advances should be coupled with microbiological studies to better
773 understand the ecophysiology of anaerobic and aerobic CE-transforming microbes and their

774 interaction with biotic and abiotic factors. Overall, interdisciplinary efforts are necessary to
775 enhance understanding of hydrological, chemical, physical and microbial interactions and
776 critical gradients that influence the fate of CE in hyporheic zones.

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1586 **Fig. 1.** Generic conceptual model of a groundwater CE plume discharging through heterogeneous
1587 hyporheic zone sediments with a range of solute residence times. The plume may be composed of HCEs,
1588 LCEs or a mixture of both. Dashed line indicates the approximate boundary between an upper oxic zone
1589 and lower suboxic/anoxic zone induced by buried sedimentary organic matter. OM: organic matter; K:
1590 sediment hydraulic conductivity. Brown areas in inset circles represents buried sedimentary organic
1591 matter (adapted from Gomez-Velez et al., 2014).

1592 **Fig. 2.** Schematic representation of hyporheic zone electron flow derived from complex organic matter
1593 metabolism (adapted from Heimann et al., 2009) to organohalide-respiring bacterial communities in
1594 anoxic riverbed sediments. Processes depicted include: orange arrow: hydrolysis; green arrows:
1595 fermentation; brown arrows: carbon assimilation; dashed red arrow: ammonification; dashed green
1596 arrow: methanogenesis; blue arrows/lines: electron transport and red arrows: organohalide respiration.
1597 Ac: acetate; DIC: dissolved inorganic carbon; ETH: ethene. Representative Gibb's free energy (ΔG°)
1598 values (kJ/M) indicated for sequential reduction of PCE to ethene taken from Dolfig and Beurskens
1599 (1995) with representative Eh values from Wiedemeier et al. (1998) under standard conditions reported
1600 therein.

1601 **Fig. 3.** Influence of extracellular hydrogen concentrations on growth rates of fermenters and
1602 hydrogenotrophs (redrawn from Ter Meer et al., 1999).

1603 **Fig. 4.** Idealised ecological succession of terminal electron acceptors and their reduced metabolites in
1604 hyporheic zones based on observed molecular hydrogen (H_2) threshold concentrations. Ac: acetate;
1605 ETH: ethene. Gibb's free energy (ΔG°) values (kJ/M) from Heimann et al. (2009) with representative
1606 Eh values from Wiedemeier et al. (1998) under standard conditions reported therein. Observed minimum
1607 hydrogen threshold ranges: (1) Cord-Ruwish et al. (1988); (2) Haring et al. (1991); (3) Sung et al. (2006);
1608 (4) Lu et al. (2001); (5) Mazur et al. (2001); (6) Lovely et al. (1989); (7) Caccavo et al. (1992); (8)
1609 Luijten et al. (2004); (9) Lovely et al. (1985).

1610 **Fig. 5.** (a) Conceptual illustration of potential metabolic (black arrows) and co-metabolic mineralisation
1611 of cDCE and VC (red arrows) at the dispersive mixing zone between upwelling anoxic groundwater
1612 containing DOM by-products (ammonium and methane) and oxic hyporheic flow cells (blue arrows).

1613 DIC: dissolved inorganic carbon; ETH: ethene. Note that reactive intermediates from co-metabolism are
1614 omitted for clarity. (b) Numerical simulations of oxygen concentration and uptake rate under gaining
1615 flow conditions in a dune bedform (adapted from Trauth et al., 2014). (c) Simulation of bedform mixing
1616 zone thickness with varying sediment hydraulic conductivity (K) and vertical groundwater flux (q_z)
1617 using a conservative groundwater tracer dilution approach (adapted from Hester et al., 2013).

1618 **Fig. 6.** Vertical pore water chemical gradients from selected case studies illustrating a range of in-situ
1619 transformation capacities (refer to Table 1 for profile context and supporting information). Black
1620 diamonds: TCE; open squares: cDCE; open triangles: VC; green circles: ethane; blue circles: ethane;
1621 orange squares: methane. Redrawn from: (a) Weatherill et al. (2014); (b) Abe et al. (2009); (c) Freitas
1622 et al. (2015) and (d) Simsir et al. (2017).

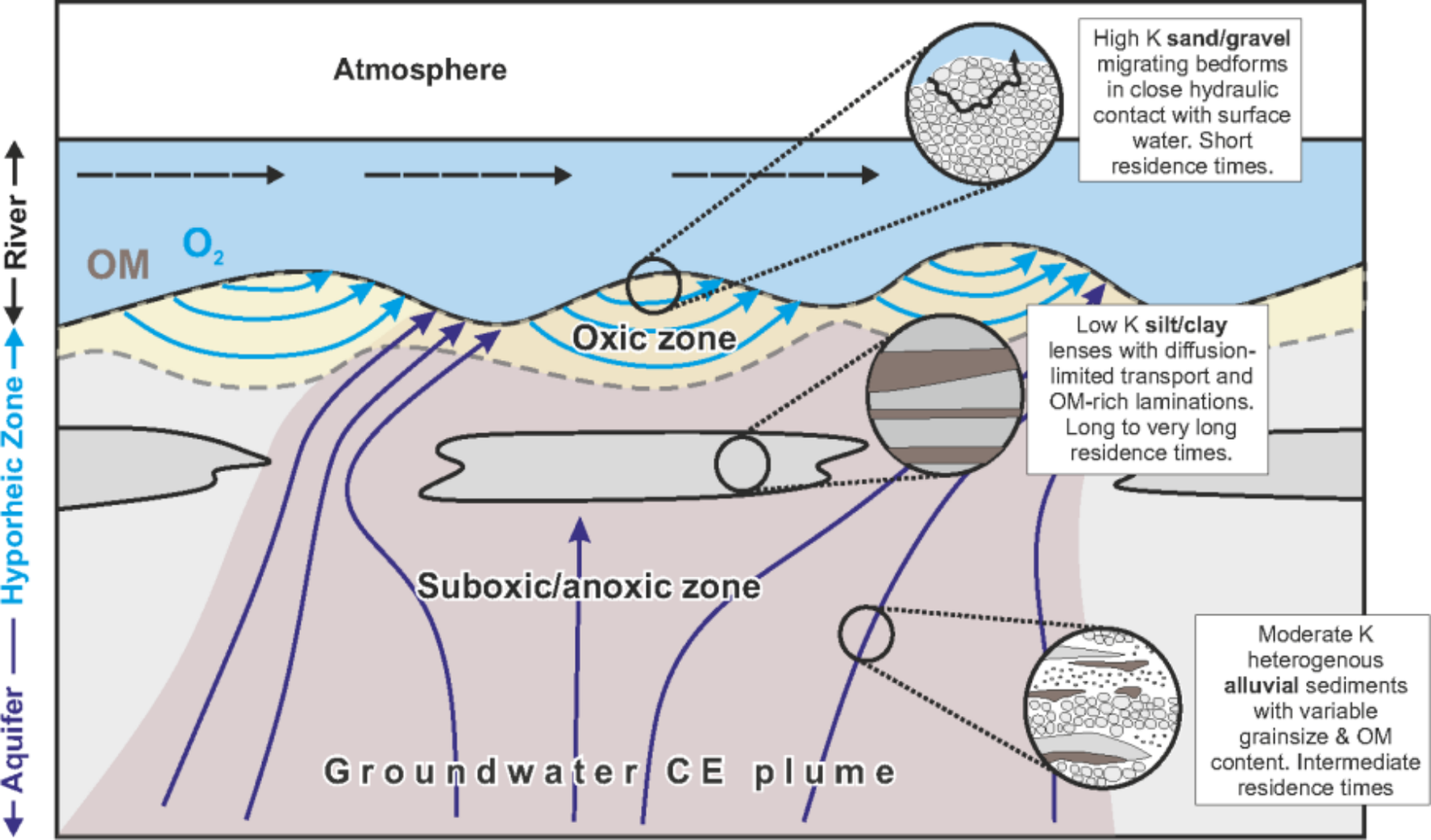
Table 1: Field experience of chlorinated ethene plume discharging to surface water receptors illustrating a range of in-situ transformation potentials from selected case studies (continued overleaf).

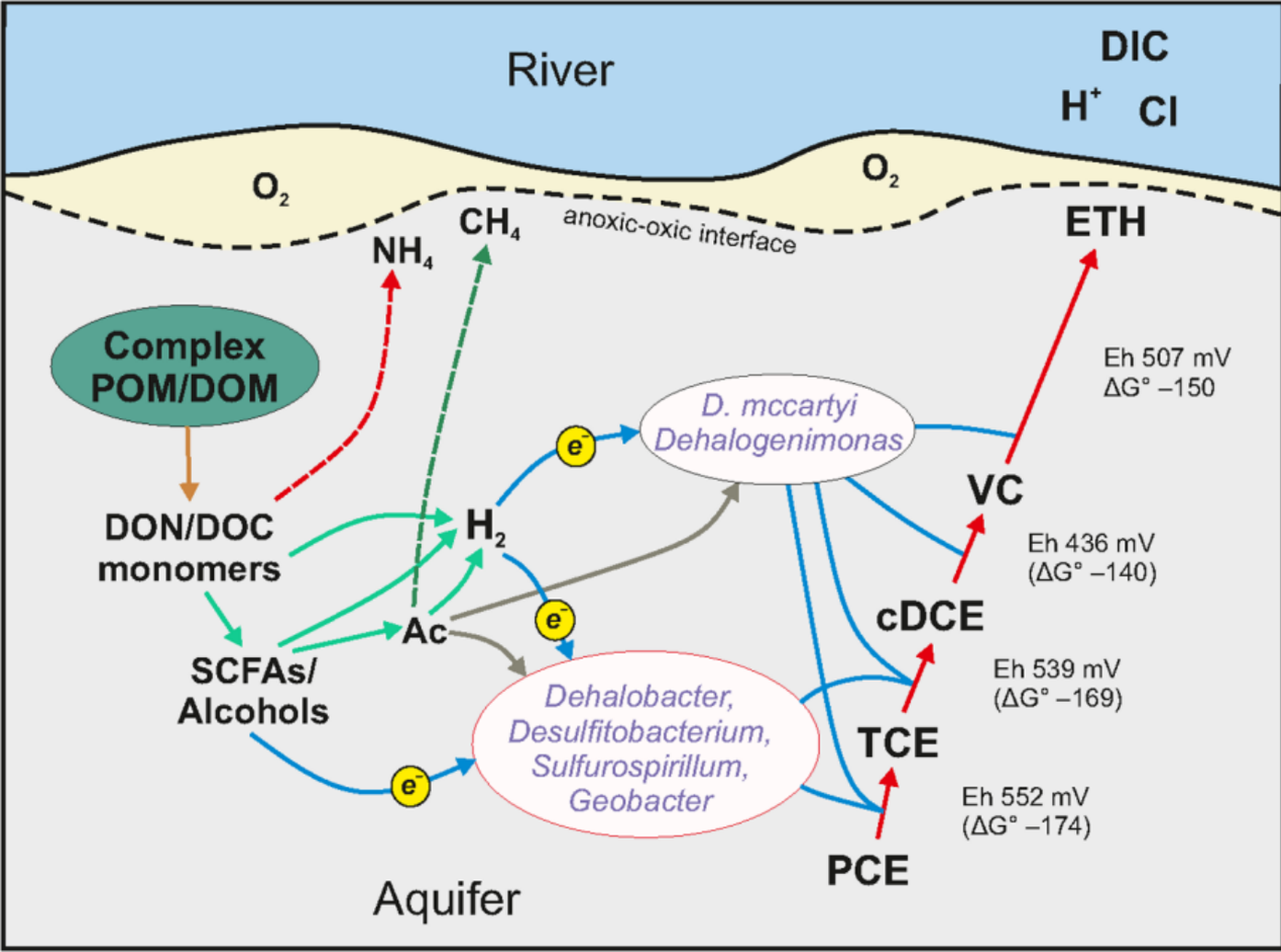
Receptor	CE source	Aquifer	Plume	Mass flux to surface water (kg/y)	Residence time (d/m)	In-situ biotransformation potential	References
Grindsted Stream , Jutland, Denmark.	Pharmaceutical factory in operation from 1914 to 1999.	Quaternary and Tertiary sands, underlain by thick regional clay layer, natural gradients.	cDCE (47 μM) and VC (72 μM). Co-mingled hydrocarbons.	45–123 (cDCE), 42–123 (VC)	0.4–10	Low , although not specifically addressed. CE-impacted stream, also impacted by metals and other organic contaminants.	Rønne et al., 2017; Sonne et al., 2017.
Third Creek , Knoxville, Tennessee, USA.	Spills associated with historic metal processing facility operated for over a century.	Fractured/karstic dolomite. Overlain by silty-clay saprolite, natural gradients.	PCE and TCE (6–60 μM). Co-mingled 1,1,1-TCA, TCM, hydrocarbons.	16.1 (TCE + cDCE)	58	High . No CEs detectable in surface water. Stoichiometric cDCE transformation in SO_4^- -reducing conditions and potential CH_4 cometabolism. Hydrocarbon co-contaminants source of SCFAs. Well distributed <i>D. mccartyi</i> population.	Simsir et al., 2017.
River Tame , Birmingham, UK.	Multiple historic urban/industrial sources, City of Birmingham.	Unconfined Permo-Triassic sandstones. Complex urban environment. Gradients historically affected by abstractions.	TCE + cDCE (0.25–9.5 μM). Measured in riverbed piezometers.	20–200 (TCE) over 7.4 km city reach	12–100 (sorption modified)	Transiently high , low-moderate overall. Stoichiometric TCE transformation to ethene in riffle sequence under Mn(IV) reduction. HEF and buried SOM a source of DOC driving redox. Inhibition by high SO_4 (5 mM) suspected.	Freitas et al., 2015; Roche et al., 2009; Ellis and Rivett, 2007.

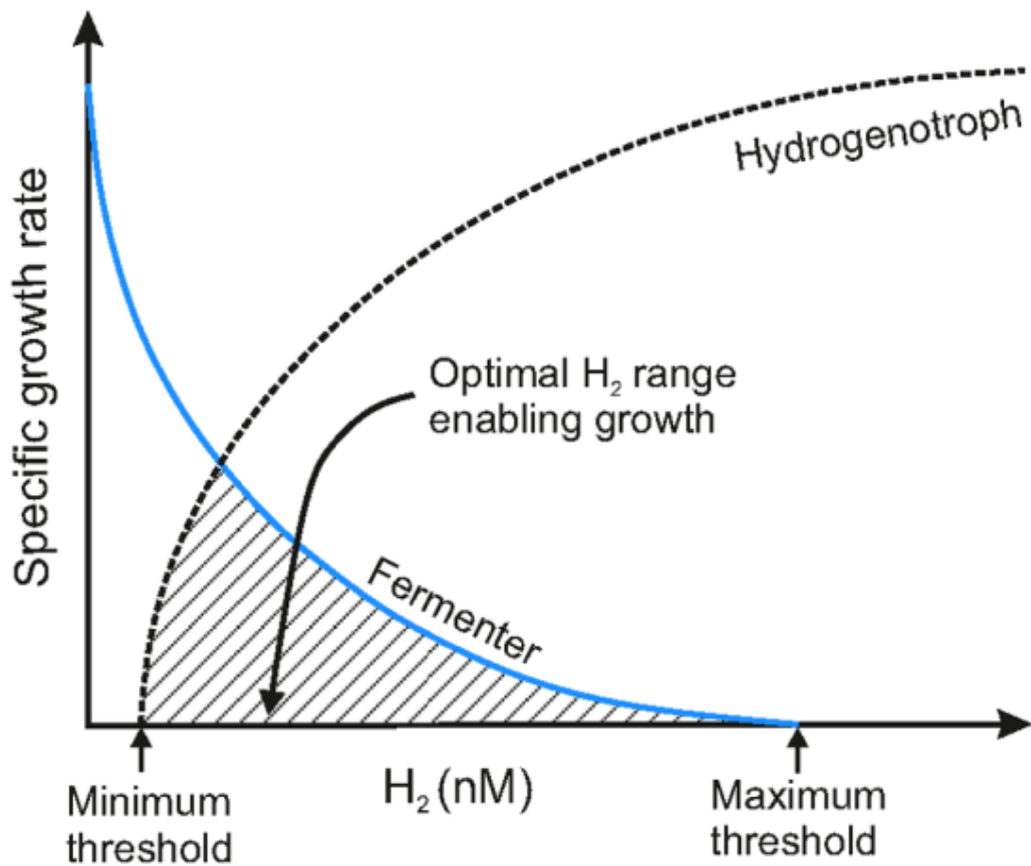
Zenne River , Vilvoorde, Belgium	Multiple industrial sources, Vilvoorde area.	Unconfined superficial gravel aquifer in urban setting. Gradients affected by retaining walls and canal water seepage.	VC (35 μM), cDCE (12 μM). 1.4 km wide plume.	Not addressed.	12–29	High. Transformation of cDCE and VC in up to 75% of locations. SOM-enriched sediments due to eutrophic conditions. <i>D. mccartyi</i> populations associated with SOM. VC/Ethene oxidisers present in SOM-poor zones.	Hamonts et al., 2009; 2012; Atashgahi et al., 2013; 2017a.
River Tern , Shropshire, UK.	Unknown. Suspected to be associated with a military airfield 1.5 km away.	Unconfined Permian sandstone. Natural gradients, occasional abstractions, agricultural setting.	TCE (0.1–1.4 μM).	0.15–0.55 (TCE)	0.5–1081 (sorption modified)	Low to moderate (patchy). Mass flux dominated by preferential pathways. Local TCE to cDCE hotspots associated peat lenses and surficial sediments. Elevated NO_3 in aerobic aquifer.	Weatherill et al., 2014; Weatherill, 2015.
Skensved Stream , Sjaelland, Denmark.	Leaking TCE storage tank, auto lacquer shop in operation since 1974. Leak discovered in 1993.	Interbedded soil, sand and gravel overlying bryozoan limestone. Pump-and-treat plume control in operation.	Max TCE (0.9 μM)	33 (TCE modelled from DNAPL source)	Not addressed	Low , aerobic to denitrifying conditions in hyporheic zone. Stream CE impact controlled by pump-and-treat system (as of 2009).	McKnight et al., 2010.
Pine River , Ontario, Canada.	Dry cleaning facility 195 m from river. PCE DNAPL contamination originated in the 1970s.	Semi-confined interbedded sand and clays in urban setting. Natural gradients.	Max PCE (0.25 mM)	3–53 (PCE) during 1995–1999.	858–71175 (sorption modified in silt/clay). Locally 3–23.	High. 52% of plume transformed to cDCE in riverbed. Stoichiometric ethene production in Fe(III)/SO_4 reducing SOM zones supporting <i>D. mccartyi</i> populations. Widespread aerobic VC mineralisation potential.	Conant et al., 2004; Abe et al., 2009.
Little Bayou Creek , McCracken County, Kentucky, USA.	Uranium enrichment facility, TCE used from 1953 to 1996 for cleaning. Residual DNAPL volume of 795 m^3 .	Regional gravel aquifer locally confined by silt and clay. Channelised river flow regime.	TCE near solubility limit near source (8.4 mM). 0.12–13 μM in riverbed. Co-mingled ^{99}Tc .	3–57 (TCE)	Not addressed	Low , TCE impacted surface water. No LCEs detectable in aerobic riverbed conditions with high mass flux through springs. Strong seasonal variability in stream water TCE concentrations.	LaSage et al., 2008; Fryar et al., 2000.

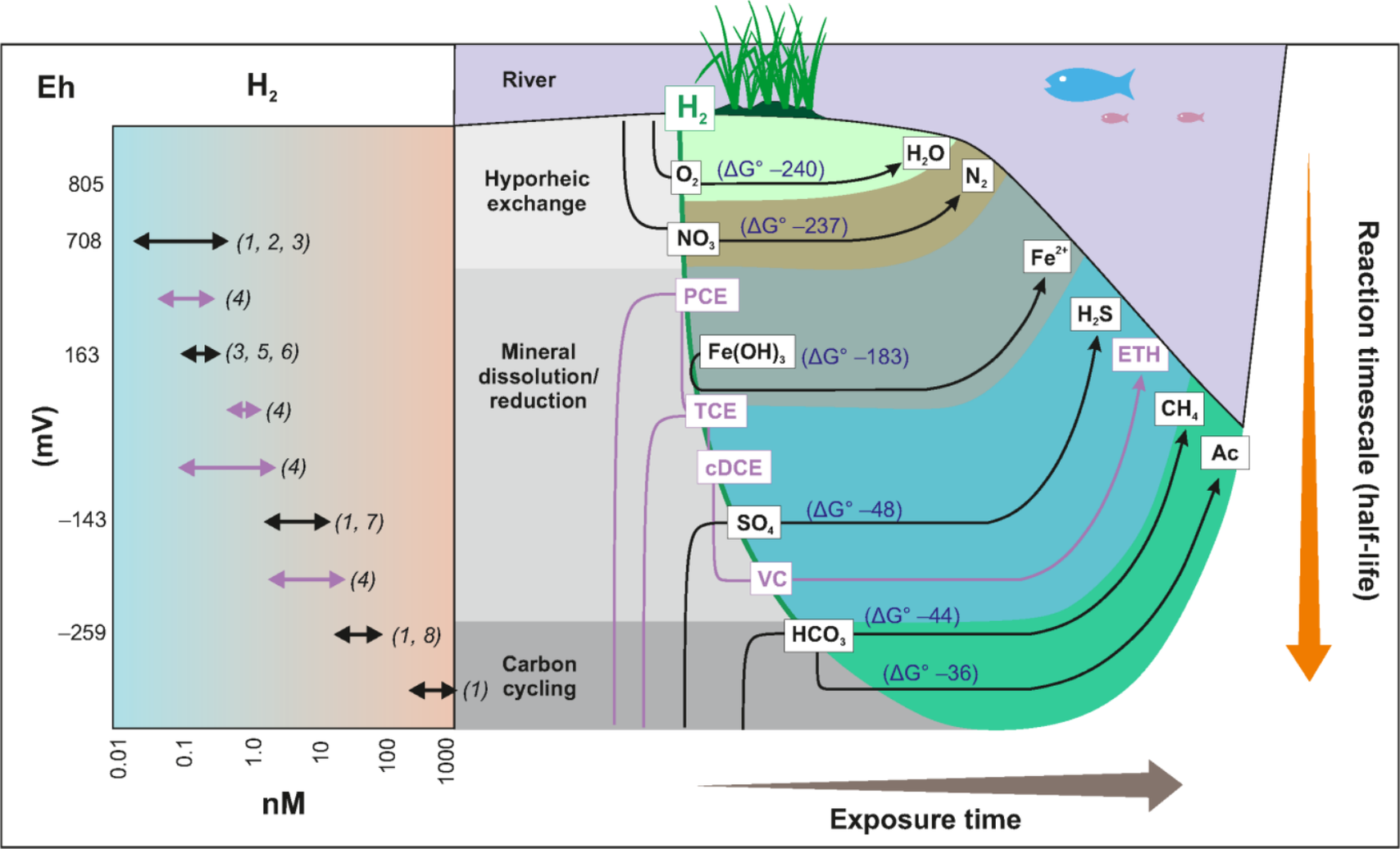
<p>Unnamed Tributary Streams, Industrial Facility, Conneticut, USA.</p>	<p>Manufacturing facility, 1970s underground degreaser tanks. TCE DNAPL source isolated in 1994. Plumes sustained by back-diffusion.</p>	<p>Unconfined surficial aquifer underlain by extensive aquitard. Steep topography, visible groundwater seepage zones.</p>	<p>TCE (18–24 μM), cDCE (5 μM) at Transect 2.</p>	<p>10 (TCE + cDCE) from Transect 2.</p>	<p>Not addressed</p>	<p>Unknown. Hyporheic zone processes in the streambeds not specifically investigated.</p>	<p>Chapman et al., 2007.</p>
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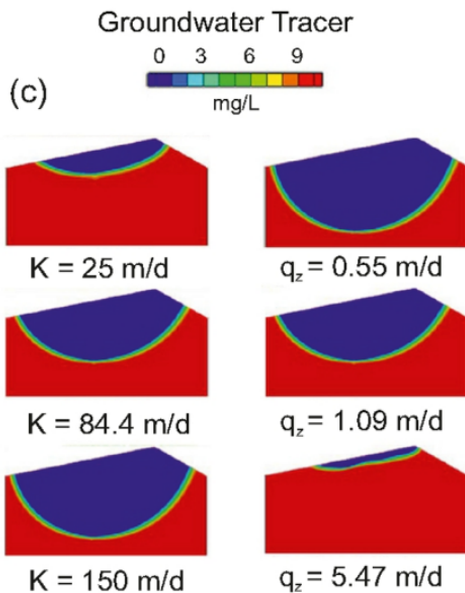
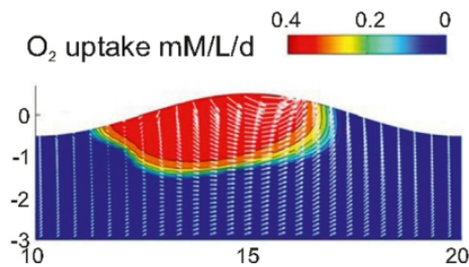
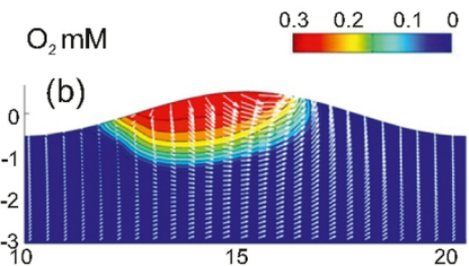
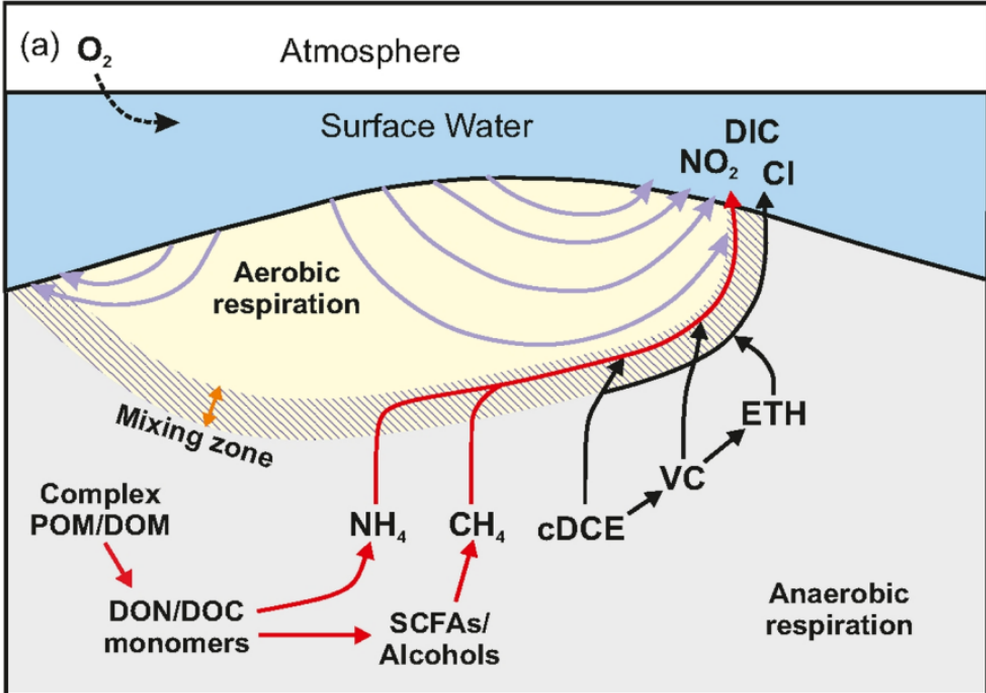
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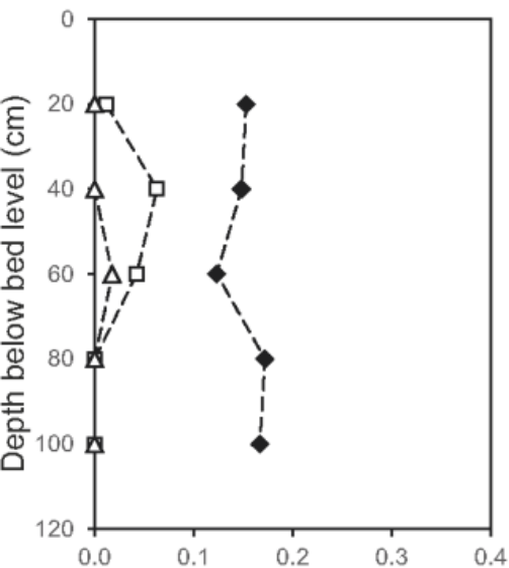




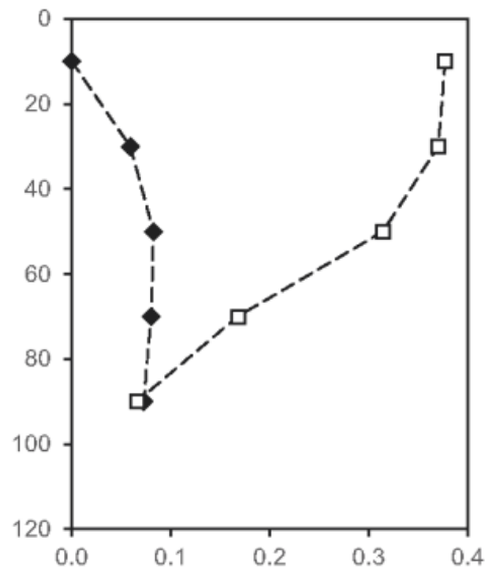




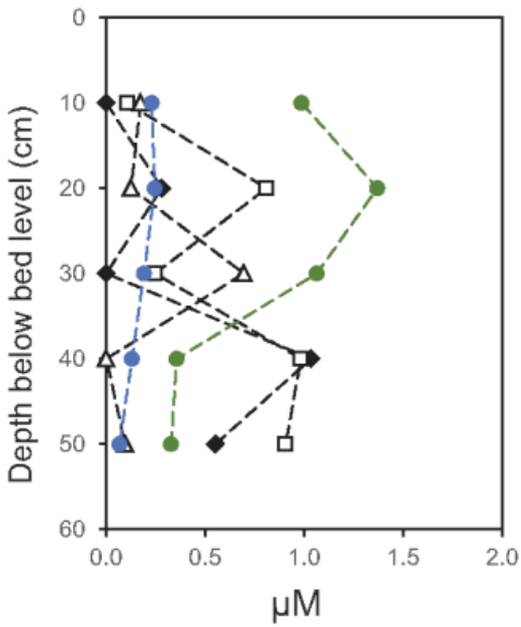
(a) River Tern, Shropshire, UK



(b) Pine River, Ontario, Canada



(c) River Tame, Birmingham, UK.



(d) Third Creek, Tennessee, US.

