1 Towards understanding the organisation of metacommunities in highly

2 dynamic ecological systems

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- 32
- 33 Abstract

34 Community ecology recognises today that local biological communities are not only affected by local biotic interactions and abiotic environmental conditions, but also by 35 36 regional processes (e.g. dispersal). While much is known about how metacommunities 37 are organised in space in terrestrial, marine and freshwater ecological systems, their 38 temporal variations remain poorly studied. Here, we address the question of the 39 dynamics of metacommunities in highly variable systems, using intermittent rivers 40 (IRs), those rivers which temporarily stop flowing or dry up, as a model system. We 41 first review how habitat heterogeneity in space and time influences metacommunity 42 organisation. Second, we compare the metacommunities in IRs to those in perennial 43 rivers (PRs) and develop the idea that IRs could undergo highly dynamic shifts due to 44 the temporal variability in local and regional community processes. Third, we develop the idea that in IRs, metacommunities of the wet and dry phases of IRs are closely 45 46 intertwined, thereby increasing even more their respective temporal dynamics. Last, we 47 provide a roadmap to stimulate further conceptual and empirical developments of 48 metacommunity research and identify possible applications for improving the 49 management of IRs and other highly dynamic ecological systems.

51 **1. Introduction**

52 Community ecology has progressed rapidly in recent years owing to the recognition 53 that local communities are not spatially closed and temporally stable (Leibold et al. 54 2004; Ricklefs 2008). Current views thus emphasise that local communities are not only 55 affected by local abiotic environmental conditions and biotic interactions, but also by 56 processes external to local ecological systems and operating at a regional scale, such as 57 speciation, extinction, immigration and emigration (Hubbell 2001). Although the 58 foundation of this idea dates back several decades (MacArthur & Wilson 1967), a shift 59 from purely local views of community organisation to those that also acknowledge the 60 importance of regional processes has been increasingly evident in the past years 61 (Hubbell 2001; Leibold et al. 2004). Sets of local communities linked by dispersal, or 62 metacommunities (Table 1), have been studied intensively in various ecological 63 systems (Logue et al. 2011), including terrestrial (e.g. Meynard et al. 2013), marine (e.g. 64 Moritz et al. 2013), and freshwater (e.g. Heino 2013) systems, and the number of 65 studies on the topic continues to increase rapidly (Heino et al. 2015). The 66 metacommunity framework has thus become a conceptually sound and empirically 67 well-explored framework to study the spatio-temporal organisation of communities.

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As an outcome of developments in spatial and dispersal ecology, different metacommunity models have been proposed to explain how both environmental filtering (i.e. local biotic interactions and abiotic environmental conditions, **Table 1**), and dispersal processes (i.e. the movement of individuals from one site to another within a region, **Table 1**), interact to shape local community structure (Vellend 2010, Logue et al. 2011; Winegardner et al. 2012). For example, "mass effects" models predict that high rates of dispersal can obscure the effect of environmental filtering, 76 while the "patch dynamic" models assume that the best dispersers arrive first and 77 occupy the patches (sensu Logue et al. 2011) as long as no more competitive species 78 have arrived (Logue et al. 2011). Although the potential differences between these 79 various models have often been emphasised (Leibold et al. 2004; Logue et al. 2011; 80 Winegardner et al. 2012), very few observational studies have succeeded in offering 81 unambiguous explanations for their relative importance in nature (Cottenie 2005; 82 Beisner et al. 2006; Bonada et al. 2012). One explanation stems from the fact that it is 83 highly difficult to measure dispersal directly and that separating the effects of 84 environmental filtering vs. dispersal processes is complex since most environmental 85 factors are spatially autocorrelated (Legendre et al. 2005; Tuomisto & Ruokolainen 86 2006; Jacobson & Peres-Neto 2010, Gilbert & Bennett 2010; Bonada et al. 2012). 87 Another explanation is that a number of metacommunity studies considered spatial 88 patterns of local communities as static within a given landscape (Leibold et al. 2004; 89 Presley et al. 2010), while communities can be very dynamic with abrupt and constant 90 change in richness and composition over very short scales (Chesson & Huntly 1989; 91 Azeria & Kolasa 2008; Erös et al. 2012; Aiken & Navarette 2014; Fernandes et al. 92 2014).

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Logue et al. (2011) suggested that broadening the types of focal habitats and organisms in metacommunity studies would provide a better understanding of the variability of metacommunity organisation (**Table 1**). They specifically pointed out that metacommunity studies should go beyond those focusing on insular habitats with discrete boundaries and those using large organisms as models. Different types of aquatic systems (e.g. marine, coastal, temporary ponds, estuaries, and running waters) provide excellent opportunities for such additional studies because they harbour highly

101 disparate organismal groups, vary widely in their degree of connectivity, and exhibit 102 wide spatial and temporal variability in local habitat conditions (Heino et al. 2015). 103 However, previous focus on purely aquatic communities has provided limited 104 perspectives on highly dynamic ecological systems, such as intermittent rivers (IRs), the 105 rivers which cease to flow or dry up in time and space. IRs provide especially suitable 106 arenas for examining metacommunity organisation in highly dynamic ecological 107 systems because they are mosaics of aquatic and terrestrial habitats shifting constantly 108 in time and space (Larned et al. 2010; Datry et al. 2014a). This dynamism, combined 109 with the fact that nearly 50% of lengths of rivers across the globe are characterized by 110 intermittent flow (Acuña et al. 2014; Datry et al. 2014a), calls for addressing 111 metacommunity organisation in IRs and other highly dynamic systems.

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113 Here, we address the question of how communities are organized in time and space 114 in highly dynamic systems. We use IRs as model ecological systems because of their 115 high dynamism and apply the metacommunity concept because of the possibility to 116 distinguish between local and regional processes in river systems (e.g. Brown & Swan 117 2010; Logue et al. 2011; Heino 2013). We first review the effects of habitat variability 118 on metacommunities in various dynamic systems. Second, we compare 119 metacommunities in IRs to those in perennial rivers (PRs) and develop the idea that IRs 120 could undergo constant and severe shifts due to the variability in environmental filtering 121 and dispersal processes. Third, we develop the idea that in IRs, metacommunities of the 122 wet and dry phases of IRs are closely intertwined, which enhances their respective 123 temporal dynamics. Last, we provide a roadmap to stimulate further conceptual and 124 empirical developments of metacommunity research and identify possible applications 125 for improving the management of IRs and other highly dynamic ecological systems.

127 2. The importance of spatial and temporal dynamics for metacommunity128 organisation

129 In the metacommunity framework, landscapes are considered as networks of 130 interconnected focal habitats in a matrix of unsuitable habitats, i.e. they are spatially 131 heterogeneous (Leibold et al. 2004; Holyoak et al. 2005). However, many conceptual 132 and empirical developments have considered metacommunities as relatively stable 133 entities (e.g. Leibold et al. 2004; Presley et al. 2010; Carrara et al. 2012; Altermatt 134 2013), although local communities and their environments can be temporally variable 135 (Azeria & Kolasa 2008; Erös et al. 2012; Aiken & Navarette 2014; Fernandes et al. 136 2014). Factors that control metacommunity organisation, including habitat availability, 137 local environmental conditions, and spatial connectivity, vary in space and time (Aiken 138 & Navarette 2014). The most extreme aquatic systems are arguably those that alternate 139 between aquatic and terrestrial conditions on a short time scale, for which high 140 variability may be the predominant rule. This category includes rocky marine 141 shorelines, tidal zones, small freshwater rock pools, temporary wetlands, vernal pools, 142 floodplains, and IRs. In such systems, environmental conditions vary both spatially and 143 temporally on a short-term basis. For example, the surface areas and connectivity of 144 floodplain aquatic habitats vary considerably during periods of weeks to months 145 between alternating wet and dry phases (Fernandes et al. 2014). Tide cycles generate 146 short pulses of aquatic habitat expansion and contraction on a daily basis (Kirwan & 147 Murray 2007). In such systems, and more generally in other systems exhibiting high 148 environmental variability, a static view of metacommunity organisation is likely to be 149 inaccurate.

151 In dynamic systems (i.e., systems experiencing constant and severe changes) the 152 various mechanisms shaping local communities, as well as their respective importance, 153 are constantly varying. For example, the contribution of dispersal will increase heavily 154 after a disturbance to allow for recolonization of a given patch (defined as a discrete 155 area with favourable environmental conditions, typically a flowing section in a river for 156 a fish or aquatic invertebrates, but see Cavanaugh et al. 2014). Yet, after most colonists 157 have reached the patch, environmental filtering will become more important in 158 explaining the local community organisation in the absence of mass effects. In systems 159 experiencing variable environmental conditions, such shifts between environmental 160 filtering vs. dispersal processes may be common and generate dynamic patterns in 161 community structure including short-term instabilities (Drake 1990; Chesson & Huntly 162 1989; Aiken & Navarrete 2014).

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164 Understanding and predicting the dynamics of metacommunities will be essential to 165 managing, conserving, and restoring biodiversity in all ecological systems, including in 166 freshwater ecological systems which have been drastically altered by global changes 167 (Dudgeon et al. 2006). Because extreme climatic events and disturbances to ecological 168 systems are occurring more frequently (Easterling et al. 2000; Parmesan, 2006), current 169 static views of community organisation are becoming less and less useful and, given 170 these limitations, we may fail to predict biodiversity loss accurately in disturbed 171 systems. Moreover, most current management, conservation and restoration applications 172 do not fully recognize the fact that maintaining the spatial and temporal dynamics of 173 entire ecological systems is essential (Heino 2013; Tonkin et al. 2014). This recognition 174 is certainly very important in the context of highly dynamic systems, where 175 communities are restructured again and again within a short period of time.

177 3. Intermittent rivers as model systems to explore the dynamics of 178 metacommunities

179 IRs occur under all types of climates on all continents, including Antarctica, and 180 make up the majority of river networks in terms of length in many regions (Acuña et al. 181 2014; Datry et al. 2014a). Globally, IRs represent 69% of the low-order streams south of 60°N latitude and from 30 to 40% of the larger river basins (Raymond et al. 2013). In 182 183 the coming decades, the number and length of IRs will increase in many regions that 184 experience drying trends due to climate and land-cover change, and increasing water 185 abstraction for irrigation and other economic uses (Palmer et al. 2008; Larned et al. 186 2010).

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188 Most conceptual and empirical developments in freshwater ecology, including the 189 application of metacommunity models (e.g. Brown & Swan 2010; Altermatt et al. 2011; 190 Heino et al. 2015), have emerged from and been produced for perennial river systems 191 (Datry et al. 2014a). Consequently, metacommunity ecology is still in its infancy in IRs, 192 and this contrasts with temporary lentic systems which have been widely used to test 193 predictions from metapopulation and metacommunity ecology (e.g. Kolasa & Romanuk 194 2005; Vanschoenwinkel et al. 2007) or have provided new insights into these fields (e.g. 195 Smol & Douglas 2007; Vanschoenwinkel et al. 2010). One important difference with 196 temporary lentic systems is that rivers occur as dendritic networks, which consist of 197 linearly-arranged, hierarchical and branching habitat elements (Fagan 2002; Grant et al. 198 2007; Altermatt 2013). Headwaters are more isolated than mainstem reaches and this 199 possibly results in spatial differences in the relative importance of community assembly 200 processes (Brown & Swan 2010). Moreover, unidirectional flow and directionally201 biased dispersal can exacerbate the effects of disturbance (drying) on species202 coexistence (Altermatt et al. 2011).

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204 IRs are dynamic shifting habitat mosaics of flowing, non-flowing and dry patches 205 (Figure 1), the extent and connectivity of which constantly vary across drainage basins 206 in response to river discharge and groundwater levels (Stanley et al. 1997; Jaeger et al. 207 2014; Datry et al. 2015, Figure 2). This spatially complex and temporally dynamic 208 habitat template is inhabited by both aquatic (e.g. fish, invertebrates, microbes, fungi, 209 algae) and terrestrial (e.g. birds, mammals, arthropods, microbes, fungi) organisms, the 210 communities of which alternate, coexist, interact, and experience extreme disturbances 211 in terms of drying and rewetting phases (Stanley et al. 1997; Datry et al. 2014a). 212 Aquatic and terrestrial communities can show intense biotic interactions. Predation by 213 terrestrial organisms on aquatic organisms trapped in drying pools can be an important 214 mortality factor in addition to interactions among fully aquatic organisms (Larned et al. 215 2010). Some large, lentic specialist predators, such as giant water bugs or diving 216 beetles, can also colonize the pools to prey on the stranded organisms, thereby 217 enhancing aquatic interactions (Gasith & Resh 1999; Boulton 2003). Altogether, local 218 environmental conditions, biotic interactions and accessibility to patches vary 219 continuously in IRs, challenging the current static views of metapopulation and 220 metacommunity approaches. Here, we make the case that IRs provide suitable arenas to 221 explore the temporal dynamics of metacommunities and notably the idea that 222 communities can experience highly dynamic shifts in structure and composition due to 223 the variability in environmental filtering and dispersal processes.

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4. Uniqueness of intermittent rivers and implications for metacommunitydynamics

IRs have some unique features that challenge current views of metacommunity organisation in river systems, which are mostly derived from research in PRs. While **Table 2** synthesises these features and their possible implications for metacommunity organisation, the different hydrological phases through which IR communities have to persist are presented below.

232 Flow cessation: shifts from lotic to lentic conditions

233 The most striking difference between PRs and IRs is that the flow ceases 234 periodically in IRs, and surface water can disappear completely from IR channels 235 (Figure 1, Table 2). Flow cessation gradually converts flowing river channels into 236 chains of disconnected pools of standing waters, which face increased water 237 temperatures and solute concentrations, and decreased pH and dissolved oxygen levels 238 (Boulton 2003). As these pools decrease in size, the densities of organisms can increase 239 dramatically, leading to strong intra- and interspecific interactions, such as competition 240 or predation. Some large pools can persist throughout dry phases and represent essential 241 refugia for aquatic organisms (Sheldon et al. 2010) and can be important "stepping 242 stones" facilitating the continued existence of populations and communities across an 243 IR network by connecting refuges and allowing recolonization of re-wetted sites by 244 actively flying or passively dispersing aquatic organisms (Bogan & Boersma 2012: 245 Cañedo-Argüelles et al. 2015). These shifts from lotic to lentic conditions occurring in 246 many IRs challenge the view that riverine systems, in general, are dominated by 247 physical constraints related to unidirectional flow (e.g. water velocity, shear stress) 248 (Fagan 2002; Altermatt et al. 2011). During the lentic phase, biotic interactions may be the dominant processes structuring aquatic communities (Gasith & Resh 1999; Boulton2003).

251 Streambed drying

252 In many IR systems, riffles dry first, after which remnant pools progressively dry 253 up, thus converting river channel into terrestrial habitat and leading to the disappearance 254 of aquatic communities (Table 2). Physiological tolerance to loss of water has been 255 well documented in organisms of temporary ponds and pools (Wiggins et al. 1980) and 256 a variety of species have also evolved physiological resistance strategies to cope with 257 desiccation in IRs (Table 2). This "invertebrate seedbank" contributes to community 258 recovery upon flow resumption, although its efficiency in such contribution is variable 259 and not completely understood (Warner & Chesson 1985; Snyder 2006; Stubbington & 260 Datry 2013). Described as the storage effect (Table 1), this mechanism contributes 261 strongly to species coexistence when resources are limiting and recruitment fluctuates 262 (Warner & Chesson 1985; Snyder 2006), thus promoting the resistance (defined as the 263 capacity to persist unchanged through a disturbance) of IR communities to drying. 264 During these dry phases, dry riverbeds are also being colonised by rich and abundant 265 terrestrial biotas, including microbes, plants, arthropods, birds, and mammals (Steward 266 et al. 2012). In spite of a recent increase in research on dry riverbed communities, they 267 still represent *terra incognita* from an ecological perspective (Steward et al. 2012).

268 Rewetting

Rewetting of previously dry patches is often sudden and unpredictable and can take the form of impressive flood bores (a video can be found here: www.irstea.fr/en/datry). Myriads of terrestrial invertebrates have been reported on these advancing fronts, with densities typically reaching > 5000 individuals/m³ (Corti & Datry 2012; Rosado et al. 2014). Many of these organisms have the ability to float or resist submersion for short

periods of time and are thus deposited alive further downstream. Such events could bemass dispersal events for terrestrial organisms.

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7 5. A dynamic view of metacommunities

278 The above features (Table 2) and resulting effects on environmental filtering and 279 dispersal processes can be used to build a new conceptual model of metacommunity 280 organisation in highly dynamic systems. We propose that (1) metacommunities are very 281 dynamic owing to temporal variations of habitat availability, environmental 282 heterogeneity, and connectivity between patches; (2) these dynamics vary spatially 283 within systems, depending on where environmental variability is the highest, where 284 potential sources of colonists are located, and how much their dispersal is limited by 285 distances between patches; and (3) metacommunities from aquatic and terrestrial phases 286 interact and may have intertwined dynamics.

287

288 5.1. Metacommunity organisation can be very variable

289 Alternating wet and dry cycles create contrasting terrestrial and aquatic habitat 290 phases in IRs (Figure 3a). As presented above, each of these phases is associated with 291 the predominance of community processes operating locally or regionally. For example, 292 shifts from lotic to lentic conditions are followed by an immediate increase in the 293 importance of environmental filtering processes, including adaptions to lentic 294 conditions, enhanced biotic interactions within contracting pools and very strong 295 predation pressure by terrestrial organisms (Table 2). Later, the relative importance of 296 dispersal processes to explain community structure and composition increases with the 297 arrival of large specialist predators such as dragonflies (Odonata), diving beetles 298 (Coleoptera) and some true bugs (Heteroptera), which are generally strong fliers and

299 colonize suitable habitats rapidly (Bogan & Boersma 2012; Bonada et al. 2012). 300 Conversely, upon rewetting, dispersal processes may first predominate to allow 301 colonisation of rewetted habitats by dispersal from refuges (Datry et al. 2014b). Soon 302 after, the communities may be influenced by storage effects from the emergence of 303 desiccation-resistant forms from the rewetted sediments (Stubbington & Datry 2013). 304 After the initial colonisation events, environmental filtering should become gradually 305 more important (Figure 3b). High dynamism may also characterize terrestrial 306 communities in IRs, although community-structuring processes involved during each 307 phase shift have been inadequately explored.

308

309 Over time, the structure of metacommunities should show imprints of the high temporal variability in the relative roles of community assembly processes. The most 310 311 obvious example is probably the respective portions, at a given location, of lotic, lentic, 312 and terrestrial species (Figure 3c). Community structure may thus vary sharply during 313 the different phases, with notable dominance by lotic species during flowing phases, 314 dominance by lentic species during non-flowing phases, and dominance by terrestrial 315 species during dry phases (Figure. 3c). Although this is speculative, there are some 316 datasets supporting these ideas in IRs (e.g. Bonada et al. 2007; Anna et al. 2008; Corti 317 & Datry 2015). For example, Corti & Datry (2015) described how aquatic and terrestrial 318 successions alternate following hydrological phases in one French IR. Other metrics of 319 community structure, such as taxonomic richness, species abundance, or the proportion 320 of predatory species should also change abruptly and include "a step-change" following 321 phase shift.

322

323 The strong temporal variability in the relative roles of community assembly 324 processes should also generate predictable spatial patterns of metacommunities. During 325 phases dominated by dispersal processes from patches to other patches, communities 326 should be predominantly nested, particularly for weak to moderate dispersers, while 327 species turnover may dominate in phases dominated by environmental filtering 328 operating locally to determine species coexistence (Figure 3d). However, these patterns 329 are likely to alternate on short time scales, jeopardizing attempts to infer on 330 communities processes from snap-shot spatial views of metacommunities. Lines of 331 evidence from various systems support these predictions. For example, frequent 332 hurricanes temporarily reduce the degree of nestedness of gastropod assemblages in 333 tropical wet forests (Bloch et al. 2007). Fish in seasonal floodplains show differences in 334 the metacommunity structure between the initial and late phases of the flooding period 335 in response to a shift in the importance of connectivity versus local environmental 336 conditions in structuring local communities (Fernandes et al. 2014). In PRs, different 337 metacommunity models apply along a gradient of disturbance level (Campbell et al. 338 2015). In temporary ponds, shifts from terrestrial to aquatic phases modulate 339 community processes, and the importance of environmental filtering decreases with 340 inundation time (Vanschoenwinkel et al. 2010).

341

342 **5.2.** The temporal dynamics of metacommunities in dendritic structures

In dendritic structures experiencing drying events, the spatial scale of metacommunities can be defined through the competitive process and/or the spatial scale of the disturbance, if wet and dry phases occur more rapidly than competitive exclusion (Massol et al. 2011). In this context, the dynamics of metacommunities likely varies spatially within ecosystems with hierarchical structure, notably in case of

348 directionally-biased dispersal (Brown & Swan 2010; Altermatt et al. 2011). For 349 example, in the case of IR networks, the temporal variability of communities may differ 350 according to where drying events (i.e. disturbance) prevail. There are five types of 351 spatial drying configurations in river systems (Figure 4), each being potentially 352 associated to a different spatial structuration of the temporal community dynamics. 353 Contrary to the paradigm that headwater communities are being driven purely by 354 environmental filtering and those of downstream, lowland reaches by mass effects due 355 to convergence of all branches and downstream water flow (Brown & Swan 2010), 356 more complex patterns may emerge for each drying configuration. For example, rivers 357 drying completely or partly in their headwaters should have headwater communities 358 driven essentially by dispersal (and perhaps by storage effects), as the source of 359 colonists may be located downstream or in the saturated or dry underlying sediments 360 (see section 3). For the different drying configurations, the degree of connectivity and 361 resulting dispersal rates between patches is probably a key factor, as very high dispersal 362 rates may decouple communities from purely local environmental control (Mouquet & 363 Loreau 2003; Ng et al. 2009; for riverine systems, see also Heino & Peckarsky 2014). 364 At the river network scale, the communities may show contrasting spatial patterns, with 365 community nestedness being more predominant in mid-reach or downstream drying 366 systems (e.g. Datry et al. 2014b), and environmental filtering dominating in headwaters 367 or complete drying systems (e.g. Grant et al. 2007; Clarke et al. 2008; Brown & Swan 368 2010).

369

370 **5.3.** Terrestrial and aquatic metacommunities have intertwined dynamics

Biotic interactions between species forming aquatic and terrestrial communities inIRs are localised in space (i.e. drying and rewetting sections) and time (i.e. drying and

373 rewetting phases) (see section 4). These discrete and punctuated interactions may 374 enhance the temporal variability in the structure of both aquatic and terrestrial 375 metacommunities. For example, aquatic species in drying reaches may be heavily 376 preyed upon by terrestrial predators (e.g. beetles, spiders), further reducing the number 377 of species able to survive flow cessation events (McHugh et al. 2014). Conversely, the 378 myriad of terrestrial invertebrates colonising dry riverbeds are entrained by advancing 379 rewetting fronts to downstream river sections, where they may provide subsidies to 380 aquatic food webs and influence aquatic community dynamics (Corti & Datry 2012; 381 Rosado et al. 2014). From a terrestrial perspective, these mass dispersal events could 382 also homogenize terrestrial metacommunities, which could thus be dominated by 383 dispersal processes in IRs rather than by local processes. Yet, aquatic and terrestrial 384 metacommunities show contrasting spatial organisations in IRs, with aquatic 385 metacommunities being longitudinally organised along river networks and terrestrial 386 metacommunities being laterally organised by riparian and upland processes away from 387 river channels (Corti & Datry 2015). Understanding the complex interaction between 388 the respective temporal dynamics and spatial organisation of aquatic and terrestrial 389 metacommunities offers a unique opportunity for integrating aquatic and terrestrial 390 ecology (Datry et al. 2014a).

391

392 6. A roadmap for future research on metacommunities in intermittent rivers and 393 other highly dynamic ecological systems

As a first step, the questions associated with our conceptual models shown above should be tested, particularly because datasets may be now or will be soon available following the growing interest in IRs (e.g. Larned et al. 2010; Datry et al. 2014a, Acuña et al. 2014). Below, we list specific research questions and indicate their relevance to furthering our understanding of metacommunities in IRs. Most of these researchquestions could be also adapted to other highly dynamic ecological systems.

400

401 How temporally variable are metacommunities of dynamic systems?

402 Fine-scale descriptions of the temporal dynamics of metacommunities during 403 typical flowing/non-flowing/dry/flowing phase cycles (Figure 3a) are necessary to 404 identify if high dynamism of local communities and metacommunities is a general rule. 405 This would be useful for both aquatic and terrestrial communities and address current 406 limitations in the static view of metacommunities (Erös et al. 2012, 2014). Such 407 temporal descriptions of communities across multiple systems would help to determine 408 if thresholds in phase duration, frequency or timing leading to alternative states of 409 community organisation exist (Bogan & Lytle 2011). Describing and understanding 410 such thresholds is crucial to predict biodiversity change in the context of climate change 411 (Parmesan, 2006; Palmer et al. 2008; Jaeger et al. 2014). Empirical developments 412 should in turn foster conceptual development of metacommunity models suitable for 413 highly dynamic systems.

414

415 *How the dendritic nature of river systems influences the dynamics of metacommunities?* 416 Previous attempts to explore the spatial dynamics of communities in ecosystems, 417 including rivers (e.g., Brown & Swan 2010) have been limited due to the low power of 418 distance matrix-based approaches (Legendre et al. 2005). The development of spatial 419 modelling in the context of constrained ordination (Cottenie 2005; Legendre et al. 2005; 420 Muneepeerakul et al. 2007; Heino et al. 2015) and in particular those accounting for 421 directionally-biased dispersal (Blanchet et al. 2008) provide better opportunities to 422 understand the spatial organisation of communities in dynamic ecosystems. Yet,

423 constrained ordination and associated spatial models assume a sort of equilibrium in
424 communities, and further developments, such as cost distance-based methods used to
425 model the dispersal of large terrestrial species (Larkin et al. 2004; LaRue & Nieslen
426 2008; Erős & Campbell Grant 2015), represent a more relevant alternative approach.

427

428 How significant is the storage effect in promoting community dynamics in IRs?

429 The contribution of the invertebrate seedbank to community recovery has been 430 shown to be significant, although highly variable across IR systems (Stubbington & 431 Datry 2013). Yet, its role in driving metacommunity dynamics in IRs is still unknown 432 and notably, whether or not it can compensate for dispersal is an open question (Snyder 433 2006). Manipulative approaches, such as common-garden experiments manipulating the 434 invertebrate seedbank and/or the dispersal of organisms in water and the air could be 435 helpful to address this question. Alternatively, cross-system comparisons of 436 metacommunity dynamics in systems with contrasting drying patterns (i.e. with and 437 without perennial refuges) may help disentangle the respective role of storage effects 438 and dispersal in promoting community dynamics. From a modelling perspective, 439 mechanistic approaches, for example using multi-occupancy models (e.g. Lamy et al. 440 2013) applied to more than one species, would allow accounting for storage effects in 441 estimating colonization and persistence rates, thus assessing the extent of cryptic 442 dormant stages in metacommunities.

443

444 Do interactions between aquatic and terrestrial metacommunities alter their respective445 dynamics?

446 Understanding how biotic interactions between aquatic and terrestrial447 metacommunities influence their respective temporal dynamics requires further

quantification in IRs. Yet, studies describing synoptically aquatic and terrestrial metacommunities in IRs or other aquatic-terrestrial systems (e.g. wetlands, tidal marshes) remain scarce (but see Corti & Datry 2015). Considering simultaneously aquatic and terrestrial metacommunities in IRs would provide a way forward to account for the linkages between these two components, as well as to bring terrestrial and aquatic ecologists together in metacommunity research.

454

455 *Is there mass dispersal of terrestrial organisms during rewetting events?*

456 While recent studies have reported how dramatic rewetting events in IRs can be and 457 how much large quantities of terrestrial organisms are being entrained downstream of a 458 river network, it is still unknown how strong roles these events play in terrestrial or 459 aquatic community dynamics (Corti & Datry 2012; Rosado et al. 2014). Typically, such 460 events could produce a mass effect for downstream riparian communities, thereby 461 obscuring local environmental filtering processes. To address this question, field 462 experiments could monitor the fate of the entrained organisms on downstream 463 communities using stable isotopes or molecular approaches. Also, the amount and type 464 of terrestrial inputs to these rewetting fronts could be manipulated to examine their 465 effects on aquatic and terrestrial communities.

466

467 How can the metacommunity perspective help managers to conserve the biodiversity of468 IRs?

The metacommunity perspective is a mechanistic route to relating biodiversity patterns to landscape features and exploring biodiversity conservation plans (Economo 2011). Identifying the relative roles of environmental filtering and dispersal processes in metacommunities can help managers to better conserve these dynamic ecological

473 systems. If local communities are mainly governed by environmental filtering 474 processes, management initiatives to maintain local habitats should be prioritized. If 475 dispersal processes are predominant instead, management strategies should maintain 476 landscape connectivity and natural disturbance regimes to promote "source" patches of 477 biodiversity (Bengtsson 2010). In the specific case of IRs, this would require preserving 478 the natural mosaic of shifting habitat types (Datry et al. 2015) and the local conditions 479 of perennial headwater patches, which are considered as important sources of 480 biodiversity (Cañedo-Argüelles et al. 2015). Additionally, the metacommunity 481 framework applied to IRs will help to pinpoint keystones species or habitat-patches in 482 the mosaic, on which managers should focus conservation and restoration efforts 483 (Mouquet et al. 2013). Empirical studies analysing the metacommunity processes in IRs 484 are thus urged to implement conservation actions specifically tailored to these dynamic 485 systems.

486

487 **6.** Conclusions

488 While community ecology has progressed rapidly in recent years owing to the 489 recognition that local communities are not spatially closed but form metacommunities 490 (Logue et al. 2011; Winegardner et al. 2012), empirical studies have often reported 491 ambiguous findings about the relative importance of underlying processes (Cottenie 492 2005; Beisner et al. 2006; Logue et al. 2011). This could be partly due to the common 493 consideration in metacommunity research that both biological communities and their 494 habitats as relatively stable, whereas many ecological systems are actually highly 495 dynamic in nature. Because IRs are common ecological systems across the globe and 496 are dynamic shifting mosaics of lotic, lentic, and terrestrial habitats, they are ideal 497 arenas for addressing the spatio-temporal variability of metacommunities in highly

498 dynamic settings. As developed here for IRs, the respective importance of 499 environmental filtering and dispersal processes may shift abruptly or gradually over 500 time in highly dynamic ecological systems, leading local community structure to vary 501 constantly. This temporal variability is certainly enhanced by discrete and punctuated 502 biotic interactions between aquatic and terrestrial communities in IRs and other coupled 503 aquatic-terrestrial systems. In the case of dendritic systems with directionally-biased dispersal (Fagan 2002; Grant et al. 2007; Altermatt et al. 2011), the location and spatial 504 505 extent of disturbances, such as drying, may interact with the temporal variations in 506 community assembly processes to produce complex spatio-temporal variability in local 507 community structure. We contend that addressing these questions in IRs may 1) 508 substantially advance the metacommunity theory (Erös et al. 2012, 2014); 2) offer a 509 unique opportunity for bridging the gap between aquatic and terrestrial community 510 ecology (Datry et al. 2014a; Soininen et al. 2015); and 3) improve the management of 511 ecological systems (Heino 2013; Tonkin et al. 2014). In the context of increasing 512 extreme climatic events and ecosystem disturbances (Easterling et al. 2000; Parmesan 513 2006), understanding how metacommunities are organised in highly dynamic systems is 514 becoming a key research topic. Also, translating such research into efficient 515 management guidelines is urgently needed.

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742 Table 1. Definitions of terms used throughout the manuscript.

| Term | Definition |
|--------------------------------|---|
| Community (or local community) | The individuals of all species that potentially interact within a single patch or local area of habitat (Leibold et al. 2004). |
| Dispersal | A regional process influencing metacommunity dynamics involving the movement of individuals from one site (i.e., emigration) to another (i.e., immigration) within a region (see also Leibold et al. 2004). |
| Storage effect | A local process involving the recruitment of many individuals from a single generation, allowing species coexistence in systems prone to disturbance (see also Warner & Chesson 1985; Snyder 2006). |
| Flow intermittence | Proportion of a given period, generally a year, during which a site in a river network is either under lentic or terrestrial phases (Datry et al. 2014). |
| Local community structure | A general term used to describe community characteristics resulting from the processes involved in its formation (e.g. species richness, dominance, etc.). |
| Metacommunity | A set of local communities that are linked by dispersal of multiple potentially interacting species (Leibold et al. 2004). |
| Metacommunity dynamics | The dynamics that arise within metacommunities; these consist of spatial dynamics, temporal dynamics, and community dynamics (multispecies interactions or the emergent properties arising from them within communities), and the interaction of these three dynamics (see also Leibold et al. 2004). |
| Metacommunity organisation | A term that refers to the processes that explain metacommunities, i.e., environmental filtering or dispersal (Heino et al. 2015). |
| Environmental filtering | A local, niche-based process influencing metacommunity dynamics and encompassing (i) the effects of local abiotic factors on species survival and (ii) local species interactions (see also species sorting in Leibold et al. 2004). |
| Nestedness | The species of communities with smaller numbers of species are subsets of species-richer communities, reflecting a non-random process of species loss or gain as a consequence of any factor that promotes the orderly disaggregation (or aggregation) of community (Baselga 2010). |
| Turnover | The replacement of some species by others between communities is a consequence of environmental filtering or spatial and historical constraints. Different from nestedness, the gain of one species is necessarily accompanied by a loss of a second species (Baselga 2010). |

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