

Challenges, developments and perspectives in intermittent river ecology

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- 1 Challenges, developments and perspectives in intermittent river ecology
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- 18 **Running head:** Intermittent river ecology
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26 Summary

27	1.	Although more than half the world's river networks comprise channels that
28		periodically cease to flow and dry (intermittent rivers [IRs]), river ecology was largely
29		developed from and for perennial systems. Ecological knowledge of IRs is rapidly
30		increasing, so there is a need to synthesize this knowledge and deepen ecological
31		understanding.
32	2.	In this Special Issue, we bring together 13 papers spanning observational case studies,
33		field and laboratory experiments and reviews to guide research and management in
34		this productive field of freshwater science. We summarize new developments in IR
35		ecology, identify research gaps and needs, and address how the study of IRs as highly
36		dynamic ecosystems informs ecological understanding more broadly.
37	3.	This series of articles reveals that contemporary IR ecology is a multifaceted and
38		maturing field of research at the interface between aquatic and terrestrial ecology. This
39		research contributes to freshwater and general ecology by testing concepts across a
40		range of topics, including disturbance ecology, metacommunity ecology and coupled
41		aquatic-terrestrial ecosystems.
42	4.	Drying affects flow continuity through time and flow connectivity across longitudinal,
43		lateral and vertical dimensions of space, which aligns well with the recent emphasis of
44		mainstream ecology on meta-system perspectives. Although most articles here focus
45		on the wet phase, there is growing interest in dry phases, and in the terrestrial
46		vegetation and invertebrate assemblages living in and along IR channels. We
47		encourage interdisciplinary studies on IRs to further blur the demarcation between
48		aquatic and terrestrial ecosystems and develop more integrated perspectives.
49	5.	As a result of climate change and human modification of landscapes and waterways,
50		flooding and drought are expected to become more extreme and widespread. Shifts in

streamflow regimes from perennial to intermittent may exacerbate the duration and
frequency of dry phases in IRs with serious implications for river ecosystems and the
quality and diversity of services they provide.

54

55 Introduction

56 River ecology initially developed from and for the study of perennial rivers and streams

57 (Datry *et al.*, 2014a). Yet more than half the channels comprising river networks globally

58 cease to flow or dry periodically (Acuña et al., 2014; Datry et al., 2014a), with the proportion

59 exceeding 80% in some regions (Levick *et al.*, 2005; Sabater & Tockner, 2010). Moreover, the

60 frequency and duration of flow cessation and drying are increasing in many rivers in response

61 to climate change and growing demands for fresh water (McKerchar & Schmidt, 2007;

62 Larned et al., 2011; Jäeger, Olden & Pelland, 2014). Several large and once-perennial rivers

63 no longer flow continuously to their deltas, including the Yellow (China), Colorado (U.S.A),

64 Amu and Syr Darya (Central Asia) and Nile (Egypt) (Gleick, 2003; Postel & Richter, 2012),

and the number of perennial systems becoming intermittent is expected to increase (Larned et

66 al., 2010; Döll & Schmied, 2012; Jaeger, Olden & Pelland, 2014). Rivers and streams without

67 continuous flow have been named by a variety of terms, such as non-perennial, temporary,

68 intermittent, ephemeral, seasonal and episodic (Uys & O'Keeffe, 1997; Williams, 2005;

69 Arthington et al., 2014). For sake of consistency, generality and simplicity, we refer to them

70 here as intermittent rivers (IRs): any river or stream that ceases to flow or dries at any point in

71 time and space (Acuña et al., 2014; Datry et al., 2014a), whether naturally or as a result of

72 human activities.

73

After years of relative obscurity, the ecological study of IRs is trending upward. The interest
 from river scientists and managers in IRs has grown rapidly in the past decade (Datry, Arscott

76	& Sabater, 2011, Arthington, Bernardo & Ilhéu 2014) and the number of IR studies published
77	in freshwater and ecology journals, including Freshwater Biology, has grown considerably
78	(Fig. 1). According to the results of a Web of Science search conducted by Leigh et al. (2016
79	a) in 2014, over 70% of IR literature published in these journals on the topics of fish and
80	invertebrate ecology, biogeochemistry, and hydrological and ecological-condition assessment
81	has appeared since the mid-2000s (Fig. 1). Ecological research on IRs now appears in high-
82	rank (e.g. Raymond et al., 2013; Jaeger et al., 2014; Acuña et al. 2014) and general ecology
83	journals (Boersma et al. 2014; Datry, Larned & Tockner, 2014a; Datry et al., 2014b; Cañedo-
84	Argüelles et al. 2015). Since 2008, almost every major freshwater conference has had a
85	special session on IRs in its program, producing special issues of journals on the topic of IRs
86	or related themes such as hydrological (dis)connectivity (e.g. Nadeau & Rains, 2007; Datry et
87	al. 2011; Arthington et al., 2014). Several international projects on IRs have been initiated,
88	including the MIRAGE European project (www.igb-berlin.de/mirage.html, 2009-2011) and
89	its continuation as a Life project (www.lifetrivers.eu, 2014-2016), the international
90	Intermittent River Biodiversity Analysis and Synthesis program (<u>www.irbas.fr.</u> 2013-2016),
91	the 1000 Intermittent River Project (<u>http://1000_intermittent_rivers_project.irstea.fr</u> , 2015-)
92	and a H2020 European COST Action (Science and Management of Intermittent Rivers and
93	ephemeral streams (<u>http://www.cost.eu/COST_Actions/ca/CA15113</u> , 2016-2020). This
94	growing interest in IRs from scientists and managers indicates the prevalence and specificities
95	of IRs, but also the scientific contribution that the study of IRs can make to general ecology
96	and the growing awareness of the social, cultural, economic and environmental values of
97	these systems.

99 From humble beginnings, the future of IR ecology is bright

100	In this Special Issue, we specifically seek to examine how IR ecology is maturing and how
101	some key questions in ecology can be addressed through the study of IRs as highly dynamic
102	ecosystems. We bring together 13 papers, including this introduction, spanning observational
103	case studies (Corti & Datry, 2016; Datry et al., 2016 a, b; Marshall et al., 2016; Rolls, Heino
104	& Chessman, 2016; Siebers et al., 2016; Stromberg & Merritt, 2016; Welter & Fisher, 2016;
105	Whitney et al., 2016), field (Vander Vorste, Malard & Datry, 2016) and laboratory
106	(Stubbington et al., 2016) experiments, and reviews (Leigh et al., 2016 a).
107	
108	Leigh et al. (2016 a) review the basic and applied ecological literature on IRs, document the
109	genealogy of ideas and highlight major advances across several research themes. While IR
110	studies were instrumental in the development of fluvial geomorphology theory (e.g., Schumm
111	& Hadley, 1953; Leopold & Miller, 1956), research on temporary standing waters (e.g.,
112	Wiggins, Mackay & Smith, 1980) and perennial systems (e.g., Harrel, Davis & Dorris, 1967)
113	was influencing early work on IRs. Persistent research themes that have informed ecological
114	theory and IR management include resistance and resilience to drying. Leigh et al. (2016 a)
115	show through their review that the recent boom in IR publications coincides with the
116	increasing need for science to inform policy concerning the legal status of IRs, and
117	recognition that the number and length of IRs are likely to increase with climate change and
118	growing human demands for fresh water. In aggregate, IR research has a strong future with
119	growing interest around the world and across multiple research themes.
120	
121	IRs are highly dynamic ecological systems: landscape mosaics, biogeochemistry and
122	biological organisation in IRs

- 123 Landscapes change over time through physical and biological drivers that are patchily
- 124 distributed, spatially and temporally (Bormann & Likens, 1979). Although terrestrial

125 landscapes and branching riverine networks differ in many respects, the terrestrially based 126 concept of shifting habitat mosaics applies well to rivers because of their dynamic nature, the 127 interrelationships between hydrology, fluvial geomorphology and the distribution of biota, 128 and water as a vector linking patches (Ward, Malard & Tockner, 2002; Stanford, Lorang & 129 Hauer, 2005; Altermatt, 2013). Datry et al. (2016 a) use hydrogeographic data collected by 130 citizen scientists to describe the spatio-temporal dynamics of IR habitats within five river 131 networks in west-central France, their hydrologic controls, and associated extinction risk and 132 colonisation for hypothetical fish and aquatic insects. Whereas general configuration and 133 compositional patterns of IR habitats were found among the networks studied, a high degree 134 of complexity was seen in landscape-metric variability over relatively short time frames and 135 among neighbouring catchments. Describing IRs with landscape metrics adds spatial 136 dimensions that are typically missed by flow metrics alone and may be particularly applicable 137 to internally coupled aquatic-terrestrial systems (i.e., dry, lentic and lotic conditions within 138 and across reaches). Datry *et al.* (2016 a) propose that a landscape approach not only 139 advances ecological understanding within IRs, but that this perspective will further advance 140 IRs as models for studying network dynamics in general.

141

142 The spatio-temporal variability in flow and mosaic composition of IR landscapes (dry, lentic 143 and lotic patches) poses particular challenges for determining the extent of material transport, 144 transformation and retention across networks (Malard, Tockner & Ward, 1999; Larned et al., 145 2010; Sponseller, Heffernan & Fisher, 2013). Transport of nutrients and organic matter from 146 terrestrial hillslopes into intermittent channels can generate hot spots of microbial activity, but 147 this may depend on the spatial extent of the hydrologic flow paths generated by rainfall pulses 148 and how long resources, including water, remain available. Welter & Fisher (2016) show how 149 storm characteristics influence the extent of hydrologic connectivity along a terrestrial-aquatic

150 flowpath in the Sycamore Creek catchment of Arizona, U.S.A., and thereby affect

151 denitrification and nitrogen retention. They show that the size, timing and intensity of storms

152 control hydrologic connectivity in small streams and that riparian terrace and hillslope soils

153 have the highest denitrification potential. Deep and moist channel sediments (i.e. the

154 hyporheic zone) have low denitrification potential but contribute to nitrogen loss for a longer

period of time overall. The study reveals important links between rainfall pulses, vertical and horizontal hydrologic connectivity and the magnitude of transport, storage and processing of

157 nutrient and organic material in arid IR landscapes.

158

159 IRs in many regions are renowned for their high levels of flow variability (e.g. Puckridge et 160 al., 1998; Poff et al., 2006, Bonada et al., 2007), which can profoundly affect biogeochemical 161 processes, creating 'hot spots' and 'hot moments' within IRs (Fisher et al., 2001; Datry et al., 162 2014a). As surface flows become discontinuous and pools contract, particulate and dissolved 163 organic matter (DOM) become concentrated and can subsequently change in composition 164 through the drying phase. Siebers et al. (2016) investigate the flow-related causes of such 165 changes in pools of dryland Australian IRs using a combination of δ^{18} O and δ^{2} H values of 166 surface and alluvial ground waters and DOM fluorescence excitation-emission spectroscopy. 167 Although the extent to which pools remain connected to alluvial ground water is strongly 168 linked to changes in fluorescence, the findings suggest that high variability in groundwater-169 surface water mixing contributes to substantial variability in compositional patterns among 170 pools. This study emphasizes that variation in biogeochemical processes and DOM 171 characteristics within these systems cannot be attributed to the presence or absence of surface 172 flow alone, highlighting the complexity of IR flow regimes and hydrological-(dis)connectivity 173 dynamics.

174

175 Landscape characteristics and hydrological-(dis)connectivity dynamics can also interact with 176 species traits to influence local extinction, connectivity and movement behaviour of biota 177 (Fausch et al., 2002; Datry et al., 2015). In many dryland IRs, for example, fish can be 178 confined to isolated waterholes for much of the year and are able to colonise the network only 179 during brief flow events. The ways fish use waterholes over time and space to either re-180 establish populations following dry periods (resilience) or maintain populations during dry 181 periods (resistance) are key determinants of population persistence (Jaeger *et al.*, 2014). 182 Marshall *et al.* (2016) tagged 215 individuals of three common species of large-bodied fish 183 and monitored their movements in and between waterholes of the Moonie River (Queensland, 184 Australia) for 3 years. Although some individuals of all species moved during flow events 185 without clear upstream or downstream preference, others remained within the same waterhole 186 throughout the study. Timing of flow resumption, independent of its magnitude or duration, 187 was a key trigger for fish movement, and many individuals returned to the same waterhole 188 after flow ceased. Maintaining such behaviour in a highly variable environment may improve 189 survival and management and conservation strategies should promote movement 190 opportunities for fish at large spatial scales to preserve population resilience. 191 192 Predicting responses of terrestrial vegetation in the riparian zones of IRs (and other river 193 types) to changes in hydrological (dis)connectivity and water availability is also vital for 194 effective river management and conservation. Terrestrial plant diversity, composition and 195 health not only are affected strongly by water availability but also can influence water 196 availability via their effects on evapotranspiration and the terrestrial water cycle (Stromberg et 197 al., 2007; Sterling, Ducharne & Polcher, 2013; Van Dijk et al., 2013). Stromberg & Merritt 198 (2016) use functional traits that capture the mechanisms underlying habitat preferences and 199 tolerances of dryland riparian plants to classify guilds associated with ephemeral, intermittent

and perennial reaches of Arizonan rivers. The use of functional traits presents a transferable
approach that goes beyond taxonomy and simple life-form or region-of-origin classification,
and can be used to predict changes in plant guild and trait composition in response to shifts in
flow intermittence and flood frequency. In particular, their findings emphasize the need to
conserve ephemeral and intermittent reaches, not just the rare perennial reaches, within river
networks of arid landscapes to maximize plant-guild diversity.

206

207 Revival of disturbance ecology: community resistance and resilience in IRs at the

208 aquatic-terrestrial interface

209 Many mechanisms contribute to the well known resilience of aquatic invertebrate

210 communities in IRs to periodic drying (Stanley et al., 1994; Fritz & Dodds, 2004; Vander

211 Vorste *et al.* 2015; Leigh *et al.*, 2016 a). Challenging the view that drift is the primary

212 mechanism, Vander Vorste et al. (2016) suggest that other processes such as colonization from

the hyporheic zone may be equally important. By manipulating both flow and drift, and

214 creating artificial dry spells in channels of a braided alluvial IR, they show that community

resilience is not significantly altered by the absence of drift. The results instead point to the

216 hyporheic zone as the main source of colonists. The importance of the hyporheic zone as a

- 217 vertical refuge from drying in IRs depends on multiple factors, including physical habitat
- 218 characteristics (e.g. sediment porosity) and biotic features (e.g. invertebrate behaviour and
- 219 morphology; Stubbington, 2012). Vander Vorste et al. (2016) emphasize the need to consider
- all potential sources of colonists as contributors to community resilience across the entire
- spectrum of IRs and invertebrate inhabitants.

222

223 Resistance as well as resilience is important in promoting biological persistence in IRs.

224 Stubbington et al. (2016) investigate one such resistance mechanism by examining and

225 rehydrating the 'seedbank' of IRs, which comprises the life stages of aquatic invertebrates 226 that remain viable in bed sediments during dry phases. Their results provide some support for 227 the hypothesis that longer dry phases decrease the abundance and richness of the seedbank but 228 also highlight the complexity of wet-dry cycling, seasonality and invertebrate life-cycle 229 relationships. Their study adds to the scant knowledge of IR seedbank ecology (e.g. Storey & 230 Quinn, 2013; Stubbington & Datry, 2013) and, we hope, will inspire researchers to devise 231 experiments to disentangle effects of environmental variables that influence biotic responses 232 to drying and survival in dry riverbeds.

233

234 Indeed, evidence indicates that dry riverbeds are not ecologically dead (e.g. Steward *et al.*, 235 2012; Stubbington et al., 2016; Vander Vorste et al., 2016) and there are likely complex 236 interactions between dry/flowing (or terrestrial/aquatic) riverbed communities and ecosystem 237 processes awaiting explanation. However, while there has historically been interest in 238 studying community patterns at terrestrial-aquatic interfaces (e.g., Reich, 1991; O'Callaghan 239 et al., 2013; Langhans & Tockner, 2014), characterization of community dynamics involving 240 complete terrestrial-aquatic shifts, as seen in IRs, has received considerably less attention (but 241 see Moon 1956; Plum, 2005; Steward et al. 2012). Corti & Datry (2016) describe terrestrial 242 and aquatic invertebrate communities from the active channel and the riparian zone of 243 intermittent and perennial reaches along a French river. Aquatic and terrestrial taxa at 244 intermittent reaches (during flowing and dry conditions, respectively) were largely subsets of 245 the taxa at perennial reaches and the adjacent riparian zone, respectively, indicating strong 246 longitudinal and lateral biological connections to upstream perennial reaches and bank 247 habitats, respectively. Among-reach variation in community composition contributed most to 248 overall terrestrial diversity, whereas within-reach variation was the greatest contributor to

249 overall aquatic diversity. These findings deny the assumption that the dry phase of IRs is a

250 lifeless period and emphasize that IRs are coupled terrestrial-aquatic ecosystems.

251

252 IRs as models to test and develop metacommunity concepts

253 Following current developments in community ecology and biodiversity research, 254 metacommunity perspectives are increasingly being applied in IR studies (e.g. Cañedo-255 Arguelles et al., 2015; Datry, Bonada & Heino, 2016). Patterns of nestedness and species 256 turnover can be due to two non-mutually exclusive processes. Common species of high 257 dispersal ability can colonise more patches in the network, leading to large population sizes 258 and ranges (Albanese, Angermeier & Peterson, 2009), than can species with limited dispersal 259 ability. Common species also tolerate harsh environmental conditions and persist throughout a wide range of habitats, resulting in low extinction rates and large population sizes and ranges 260 261 (Schlosser, 1990; Taylor & Warren, 2001). Although these processes were integrated in a 262 metacommunity model by Larned et al. (2010), stating that longitudinal patterns of nestedness 263 across connected stream habitats are caused by dispersal limitation, whereas temporal patterns 264 of nestedness within habitats during stream drying are caused by interspecies differences in 265 environmental tolerance, there is still little empirical evidence for this model (but see Datry et 266 al., 2014b). Whitney et al. (2016) quantify species abundance distributions and the 267 importance of nestedness and turnover to community dissimilarity in tallgrass prairie streams 268 of eastern Kansas, U.S.A. Few species were common but nestedness drove community 269 dissimilarity across sites, notably during the first phases of recolonisation. Although common 270 species were the first to colonise and the last to go extinct locally, the study raises the 271 question: is commonness predictive of rather than predicted by colonisation and extinction?

272

273 Datry et al. (2016 b) further examine the relative roles of dispersal and environmental filtering 274 in structuring biological communities in dynamic landscapes. When one thinks of IRs, 275 tributaries of the Amazon River typically don't come to mind. Datry et al. (2016 b) compare 276 the local and regional diversity of aquatic invertebrate and fish communities from intermittent 277 and perennial reaches of rivers in central Bolivia to see if natural drying has a similar effect 278 on the structure and composition of neotropical communities as has been observed in other 279 regions. Intermittent and perennial reaches had similar invertebrate assemblages but perennial 280 reaches had higher fish densities and richness than intermittent reaches. Using 281 metacommunity analyses, in combination with life history knowledge of the local fauna, the 282 authors were able to infer the relative importance of dispersal compared to local 283 environmental filtering in structuring aquatic invertebrate and fish communities across the 284 study sites. The study further suggests that drying may act as a relatively mild disturbance for 285 aquatic invertebrate communities of neotropical IRs.

286

287 While differences in the community composition among rivers with contrasting flow regimes 288 are a consistent pattern in many climate regions worldwide, it is not clear whether such 289 differences among flow regimes are consistent during and after periods of prolonged drought 290 (sensu Lake, 2003) and how much these differences are affected by organismal dispersal 291 modes and strengths (e.g. Leigh & Datry, 2016). Using biomonitoring data on aquatic 292 invertebrate and fish communities in subtropical rivers in Australia, Rolls et al. (2016) tested 293 whether beta diversity across naturally near-perennial, mildly intermittent and highly 294 intermittent streams were greater during a three-year drought characterized by low rainfall 295 than during a wetter post-drought phase. They also tested whether differences in beta diversity 296 between phases would decrease with increasing dispersal strength of organisms (Cañedo-297 Arguelles *et al.*, 2015; Datry *et al.*, 2015). Except for locally dispersing fish, beta-diversity

298 differed among flow regimes for all dispersal groups of invertebrates and fish. Within flow-299 regime groups, beta-diversity was higher during the drought than post-drought for most 300 invertebrate groups (but was not different for fish) and, for most dispersal groups, was driven 301 predominantly by replacement of taxa among samples rather than nestedness (except for 302 diadromous fish). Overall, the findings suggest that beta diversity was similar during both 303 phases, possibly due to consistent shifts in average community composition during recovery 304 from drought in all flow regime groups. These consistent shifts may reflect some community 305 resistance to the effects of drying-wetting cycles (Leigh et al., 2016 b) but also resilience and 306 the presence of perennial refuges (e.g., water holes) in many of the rivers studied.

307

308 Where to next? A roadmap for ecological research on intermittent rivers

309 Contemporary IR ecology is a multifaceted and maturing field of research at the interface 310 between aquatic and terrestrial ecology. The research findings in this Special Issue contribute 311 to freshwater and general ecology by testing and challenging concepts across a range of topics 312 in highly dynamic systems. This issue is also timely because information on IRs is being 313 sought by managers and policy makers faced with increasing flow intermittence in response to 314 climate change and growing demands for fresh water. There is ongoing debate regarding the 315 legal status of IRs in many countries despite the increasing recognition of their ubiquity, their 316 influence on perennial waters, and human-induced shifts increasing their prevalence and 317 prolonging droughts (Acuña et al., 2014; Datry et al., 2014b). From the insights provided by 318 the studies presented in this Special Issue, we draw several conclusions and proposals to 319 direct future research and management.

320

321 Drying imposes a series of strong local filters expressed as incremental

322 physicochemical and biological constraints selecting for biota and governing ecosystem

323	processes (Williams, 1996; Boulton, 2003). Much of what is known about IR ecology has
324	been informed by this perspective (Leigh et al., 2016 a). Comparative studies have used this
325	perspective to test predictions regarding biodiversity and ecosystem processes between
326	perennial and intermittent rivers as well as across continua of intermittence (Datry et al., 2016
327	b; Stromberg & Merritt, 2016; Rolls et al., 2016). Although this Special Issue defines "IRs" as
328	any channel that periodically ceases to flow and dries, we recognize that IRs really represent a
329	continuum with varying durations, frequencies, timing and expansion/contraction rates of
330	drying (Stanley, Fisher & Grimm, 1997; Hughes, 2005; Costigan et al., 2016). The study of
331	these continua can therefore provide a deeper understanding and predictive capability
332	regarding how drying influences biodiversity and ecosystem function. Experimental
333	approaches that manipulate different dimensions of intermittency (Vander Vorste et al., 2016)
334	and the use of models (Fisher et al., 1998; Datry et al., 2016 a) will also provide focused
335	mechanistic advances.
336	

337 Drying is increasingly recognized as affecting flow continuity through time and flow 338 connectivity through space (e.g. Larned et al., 2011; Bogan, Boersma & Lytle, 2013; Jaeger et 339 al., 2014). This aligns well with mainstream ecology's recent use of meta-system perspectives 340 (e.g., Loreau, Mouquet & Holt, 2003; Datry et al., 2016) and riverine ecologists' growing 341 interest in connectivity at the network scale (e.g., Carrara et al., 2012; Zimmermann et al., 342 2014). A common thread across most the papers in this Special Issue is the role of 343 hydrological (dis)connectivity in IR ecology, whether by affecting dispersal or material flows 344 (e.g. Whitney et al., 2016). Connectivity underlies processes across all scales, from local 345 within-habitat (Vander Vorste et al., 2016) to longitudinal segments (Marshall et al., 2016) to 346 entire networks (Datry et al., 2016 a; Rolls et al. 2016; Welter & Fisher, 2016). Longitudinal 347 connectivity along IRs may receive the most attention because surface water is the dominant

348 transport route for many riverine organisms and materials, however because of recurrent wet-349 dry cycles the vertical connectivity with ground water and atmosphere can be important 350 (Siebers et al., 2016; Vander Vorste et al. 2016; Welter & Fisher, 2016). In spite of the role 351 surface-groundwater interactions play in shaping drying patterns in IRs, disruption of vertical 352 connectivity has attracted little interest from ecologists (Leigh et al., 2016 a), reflecting the 353 lag in recognition of the functional significance of hyporheic zone in perennial systems 354 (Boulton et al., 2010). Although rivers are linear features, lateral connectivity is critical in 355 understanding resource subsidies (McClunev & Sabo, 2012) and community dynamics for 356 floodplain systems (Fernandes et al., 2014). Less studied has been lateral movement by 357 organisms that inhabit or use dry stream beds, and more research is needed to inform 358 understanding of their persistence in IRs (Corti & Datry, 2016). In addition to movement 359 across space, storage or the lack of movement can be critical in understanding the distribution 360 and abundance of materials and organisms in IRs. Numerous species inhabiting IRs have 361 resting stages and adaptations that enable them to withstand (resist) the drying conditions in 362 situ and resume growth and reproduction after rewetting (Stromberg & Merritt, 2016; 363 Stubbington et al., 2016). The importance of resistance or storage to the persistence or 364 dynamics of populations and communities in IRs relative to other ecosystems, such as 365 temporary pools, estuaries, or forest soils, is unknown. The papers presented here will 366 encourage modification of existing approaches and development of novel approaches for 367 investigating the (dis)continuity and (dis)connectivity in IRs and the degree to which these 368 factors, as well as environmental filtering, govern biodiversity and biogeochemical patterns 369 across ecosystems.

370

The study of IRs is situated at the interface of aquatic and terrestrial ecology. Most papers in this Special Issue emphasise the wet rather than the dry phase, however there is

373 growing interest in dry phases and the terrestrial vegetation and invertebrate assemblages 374 living in and along IR channels. Biogeochemistry and microbial studies are also areas where 375 the dry phase is being incorporated into a more complete understanding of IR ecosystems. 376 Interdisciplinary studies on IRs should be encouraged to further blur the demarcation between 377 aquatic and terrestrial ecology (Datry et al., 2014 a). IRs are also extreme examples of 378 systems where ecological patterns and processes are shaped by temporal environmental 379 fluctuation. Therefore, IRs can be considered useful models for understanding seasonal food 380 web shifts over time (McMeans et al., 2015) and where / how resource booms and busts 381 interact and alternate among members of flowing and dry channel communities (Balcombe et 382 al., 2015). The compression of interactions and flows of materials and energy to particular 383 points in time and space that may vary in their predictability is certainly an area where IR 384 research can contribute to aquatic and general ecology (Datry et al., 2016 a).

385

386 As a result of ongoing climate change and human modification of landscapes and 387 waterways, flooding and drought are expected to become more extreme and possibly more 388 frequent (Ledger & Milner, 2015). Although flooding can be detrimental to society and 389 riverine ecosystems, drying of formerly perennial rivers is expected to be more adverse and 390 more difficult to mitigate through management actions (Palmer et al., 2008). Shifts in 391 streamflow regimes from perennial to intermittent and extension of the duration and 392 frequency of dry phases in IRs could strongly affect river ecosystems and the services they 393 provide (Costigan et al. 2016; Datry et al., 2016). While this Special Issue testifies to recent 394 advances in IR ecology, scientific aspects are certainly yet to be uncovered, and foreseeable 395 challenges for their protection and restoration indicate there is still much room for growth and 396 innovation.

397

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- 410

411 References

412 Acuña V., Datry T., Marshall J., Barceló D., Dahm C. N., Ginebreda A., et al. (2014) Why

413 should we care about temporary waterways? *Science*, **343**, 1080-1081.

- 414 Albanese B., Angermeier P.L. & Peterson J.T. (2009) Does mobility explain variation in
- 415 colonisation and population recovery among stream fishes? *Freshwater Biology*, 54, 1444–
 416 1460.
- 417 Altermatt F. 2013. Diversity in riverine metacommunities: a network perspective. Aquatic
- 418 *Ecology*, **47**, 365 377.
- 419 Arthington A. H., Bernardo J. M. & Ilhéu M. (2014) Temporary rivers: linking ecohydrology,
- 420 ecological quality and reconciliation ecology. *River Research and Applications*, **30**, 1209-
- 421 1215.

- 422 Balcombe S.R., Turschwell M.P., Arthington A.H. & Fellows C.S. (2015) Is fish biomass in
- 423 dryland river waterholes fuelled by benthic primary production after major overland

424 flooding? Journal of Arid Environments, **116**, 71-76.

- 425 Boersma K. S., Bogan M. T., Henrichs B. A. & Lytle D. A. (2014) Top predator removals
- 426 have consistent effects on large species despite high environmental variability. *Oikos*, **123**,
- 427 807-816.
- 428 Bogan M. T., Boersma K. S. & Lytle D. A. (2013) Flow intermittency alters longitudinal
- 429 patterns of invertebrate diversity and assemblage composition in an arid-land stream

430 network. *Freshwater Biology*, **58**, 1016-1028.

- 431 Bormann, F.H. & Likens, G.E. (1979) Catastrophic disturbance and the steady state in
- 432 northern hardwood forests. *American Scientist*, **67**, 660-669.
- 433 Boulton A. J. (2003) Parallels and contrasts in the effects of drought on stream
- 434 macroinvertebrate assemblages. *Freshwater Biology*, **48**, 1173-1185.
- 435 Boulton, A. J., Datry, T., Kasahara, T., Mutz, M., & Stanford, J. A. (2010) Ecology and
- 436 management of the hyporheic zone: stream-groundwater interactions of running waters
- 437 and their floodplains. *Journal of the North American Benthological Society* **29**, 26-40.
- 438 Cañedo-Argüelles M., Boersma K. S., Bogan M. T., Olden J. D., Phillipsen I., Schriever T. A.
- 439 & Lytle D. A. (2015) Dispersal strength determines meta-community structure in a
- dendritic riverine network. *Journal of Biogeography*, **42**, 778-790.
- 441 Carrara F., Altermatt F., Rodriguez-Iturbe I. & Rinaldo A. (2012) Dendritic connectivity
- 442 controls biodiversity patterns in experimental metacommunities. *Proceedings of the*
- 443 *National Academy of Sciences*, **109**, 5761-5766.
- 444 Costigan K. H., Jaeger K. L., Goss C. W., Fritz K. M. & Goebel P. C. (2016) Understanding
- 445 controls on flow permanence in intermittent rivers to aid ecological research: integrating
- 446 meteorology, geology and land cover. *Ecohydrology*, doi/10.1002/eco.1712/

- 447 Corti R. & Datry T. (2016) Terrestrial and aquatic invertebrates in the riverbed of an
- intermittent river: parallels and contrasts in community organisation. *Freshwater Biology*,
 xx, xxx-xxx.
- 450 Datry T., Arscott D. B. & Sabater S. (2011) Recent perspectives on temporary river ecology.
 451 *Aquatic Sciences*, **73**, 453-457.
- 452 Datry T., Bonada N. & Heino J. (2016) Towards understanding the organisation of
- 453 metacommunities in highly dynamic ecological systems. *Oikos*, **125**, 149-159.
- 454 Datry T., Larned S. T., Fritz K. M., Bogan M. T., Wood P. J., Meyer E. I., et al. N. (2014b)
- 455 Broad-scale patterns of invertebrate richness and community composition in temporary
- 456 rivers: effects of flow intermittence. *Ecography*, **37**, 94-104.
- 457 Datry T., Larned S. T. & Tockner K. (2014a) Intermittent rivers: a challenge for freshwater
- 458 ecology. *BioScience*. doi: 10.1093/biosci/bit027
- 459 Datry T., Moya N.B., Zubieta J. & Oberdorff, T. (2016 b) Determinants of local and regional
- 460 communities in intermittent and perennial headwaters of the Bolivian Amazon.
- 461 *Freshwater Biology*, xx, xxx-xxx.
- 462 Datry T., Pella H., Leigh C., Bonada N. & Hugueny, B. (2016 a). A landscape approach to
- 463 advance intermittent river ecology. *Freshwater Biology*, xx, xxx-xxx.
- 464 Döll P. & Schmied H. M. (2012) How is the impact of climate change on river flow regimes
- 465 related to the impact on mean annual runoff? A global-scale analysis. *Environmental*
- 466 *Research Letters*, **7**, 014037.
- 467 Fausch K. D., Torgersen C. E., Baxter C. V. & Li H. W. (2002) Landscapes to riverscapes:
- 468 bridging the gap between research and conservation of stream fishes a continuous view of
- the river is needed to understand how processes interacting among scales set the context
- 470 for stream fishes and their habitat. *BioScience*, **52**, 483-498.

- 471 Fernandes I. M., Henriques-Silva R., Penha J., Zuanon J. & Peres-Neto P. R. (2014)
- 472 Spatiotemporal dynamics in a seasonal metacommunity structure is predictable: the case
- 473 of floodplain-fish communities. *Ecography*, **37**, 464–475
- 474 Fisher S. G., Grimm N. B., Martí E., Holmes R. M. & Jones J. B. (1998) Material spiraling in
 475 stream corridors: A telescoping ecosystem model. *Ecosystems*, 1, 19-34.
- 476 Fisher S. G., Welter J., Schade J. & Henry J. (2001) Landscape challenges to ecosystem
- 477 thinking: creative flood and drought in the American Southwest. *Scientia Marina* 65
- 478 **(Suppl. 2)**, 181-192.
- 479 Fritz K. M. & Dodds W. K. (2004) Resistance and resilience of macroinvertebrate
- 480 assemblages to drying and flood in a tallgrass prairie stream system. *Hydrobiologia*, **527**,
- 481 **99-112**.
- 482 Gleick P. H. (2003) Global freshwater resources: soft-path solutions for the 21st century.
- 483 *Science*, **302**, 1524-1528.
- 484 Harrel R. C., Davis, B. J. & Dorris T. C. (1967) Stream order and species diversity of fishes in
- 485 an intermittent Oklahoma stream. *American Midland Naturalist*, **78**, 428-436.
- 486 Jaeger K. L., Olden J. D. & Pelland N. A. (2014) Climate change poised to threaten
- 487 hydrologic connectivity and endemic fishes in dryland streams. *Proceedings of the*
- 488 *National Academy of Sciences*, **111**, 13894-13899.
- 489 Lake P. S. (2003). Ecological effects of perturbation by drought in flowing waters. Freshwater
- 490 *Biology*, **48**, 1161-1172.
- 491 Langhans S. D. & Tockner K. (2014) Edge effects are important in supporting beetle
- 492 biodiversity in a gravel-bed river floodplain. *PLOS One* **9**, e114415.
- 493 Larned S. T., Datry T., Arscott D. B. & Tockner K. (2010) Emerging concepts in temporary-
- 494 river ecology. *Freshwater Biology*, **5**, 717–738.

- 495 Larned S. T., Schmidt J., Datry T., Konrad C. P., Dumas J. K. & Diettrich J. C. (2011)
- 496 Longitudinal river ecohydrology: flow variation down the lengths of alluvial rivers.
- 497 *Ecohydrology*, **4**, 532-548.
- Ledger M. E. & Milner A. M. (2015) Extreme events in running waters. *Freshwater Biology*,
 60, 2455-2460. doi:10.1111/fwb.12673
- 500 Leigh C., Boulton A. J., Courtwright J. L., Fritz K., May C. L., Walker R. H., et al. (2016 a)
- 501 Ecological research and management of intermittent rivers: an historical review and future 502 directions. *Freshwater Biology*, xx, xxx-xxx.
- 503 Leigh C., Bonada N., Boulton A. J., Hugueny B., Larned S. T., Vander Vorste R., et al. (2016
- b) Invertebrate assemblage responses and the dual roles of resistance and resilience to
- 505 drying in intermittent rivers. *Aquatic Sciences*, **78**, 291-301.
- 506 Leigh C. & Datry T. (2016) Drying as a primary hydrological determinant of biodiversity in
- 507 river systems: a broad scale analysis. *Ecography*, DOI: 10.1111/ecog.02230
- 508 Levick L., Fonseca J., Goodrich D., Hernandez M., Semmens D., Stromberg J., et al. (2008).
- 509 The Ecological and Hydrological Significance of Ephemeral and Intermittent Streams in
- 510 *the Arid and Semi-arid American Southwest*. U.S. Environmental Protection Agency and
- 511 USDA/ARS Southwest Watershed Research Center, EPA/600/R-08/134, ARS/233046,
- 512 116 pp.
- Leopold L. B. & Miller J. P. (1956) Ephemeral streams hydraulic factors and their relation
 to the drainage net. *United States Geological Survey Professional Paper*, 282-A.
- 515 Loreau M., Mouquet N. & Holt R. D. (2003) Meta-ecosystems: a theoretical framework for a
- 516 spatial ecosystem ecology. *Ecology Letters*, **6**, 673-679.
- 517 Malard F., Tockner K, & Ward J. V. (1999) Shifting dominance of subcatchment water sources
- and flow paths in a glacial floodplain, Val Roseg, Switzerland. Arctic, Antarctic, and
- 519 *Alpine Research*, **31**, 135-150.

- 520 Marshall J. C., Menke N., Crook D. A., Lobegeiger J., Balcombe S, Huey J., et al. (2016) Go
- 521 with the flow: the movement behaviour of fish from isolated waterhole refugia during
- 522 connecting flow events in an intermittent dryland river. *Freshwater Biology*, xx, xxx-xxx.
- 523 McCluney K. E. & Sabo J. L. (2012) River drying lowers the diversity and alters the
- 524 composition of an assemblage of desert riparian arthropods. *Freshwater Biology*, **57**, 91-
- 525 103.
- McKerchar A. I. & Schmidt J. (2007) Decreases in low flows in the lower Selwyn River? *Journal of Hydrology (New Zealand)*, 46, 63-72.
- 528 McMeans B. C., McCann K. S., Humphries M., Rooney N. & Fisk A. T. (2015) Food web
- 529 structure in temporally-forced ecosystems. *Trends in Ecology & Evolution*, **30**, 662-672.
- 530 Moon H. (1956) Observations on a small portion of a drying chalk stream. Proceedings of the
- 531 Zoological Society of London, **126**, 327–334.
- 532 Nadeau T.-L. & Rains M. C. (2007) Hydrological connectivity between headwater streams
- and downstream waters: how science can inform policy. Journal of the American Water
- 534 *Resources Association*, **43**, 118–133.
- 535 O'Callaghan M. J., Hannah D. M., Boomer I., Williams M. & Sadler J. P. (2013) Responses to
- 536 river inundation pressures control prey selection of riparian beetles. *PLOS One*, **8**, e61866
- 537 Palmer M. A., Reidy Liermann C. A., Nilsson C., Flörke M., Alcamo J., Lake P. S., et al.
- 538 (2008) Climate change and the world's river basins: anticipating management options.
- 539 *Frontiers in Ecology and the Environment*, **6**, 81-89.
- 540 Perkins P. D. (1976) Psammophilous aquatic beetles in southern California: a study of
- 541 microhabitat preferences with notes on response to stream alteration (Coleoptera:
- 542 Hydraenidae and Hydrophilidae). *The Coleopterists Bulletin*, **30**, 309-324.
- 543 Plum N. (2005) Terrestrial invertebrates in flooded grassland: a literature review. Wetlands,
- **5**44 **25**, 721-737.

- 545 Poff N. L., Olden J. D., Pepin D. M. & Bledsoe B. P. (2006) Placing global stream flow
- variability in geographic and geomorphic contexts. *River Research and Applications*, 22,
 149-166.
- 548 Postel S. & Richter B. (2012) *Rivers for Life: Managing Water for People and Nature*. Island
 549 Press.
- 550 Puckridge, J. T., Sheldon F., Walker K. F. & Boulton A. J. (1998) Flow variability and the
- 651 ecology of large rivers. *Marine and Freshwater Research*, **49**, 55-72.
- 552 Raymond P. A., Hartmann J., Lauerwald R., Sobek S., McDonald C., Hoover M., et al.
- 553 (2013) Global carbon dioxide emissions from inland waters. *Nature*, **503**, 355-359.
- 554 Reich M. (1991) Grasshoppers (Orthoptera, Saltatoria) on alpine and dealpine riverbanks and
- their use as indicators for natural floodplain dynamics. *Regulated Rivers: Research and*
- 556 *Management*, **6**, 333-340.
- 557 Rolls R. J., Heino J. & Chessman B. (2016) Unravelling the joint effects of flow regime,
- 558 climatic variability and dispersal mode on beta diversity of riverine communities.
- 559 Freshwater Biology, xx, xxx-xxx.
- 560 Sabater S., & Tockner K. (2010) Effects of hydrologic alterations on the ecological quality of
- 561 river ecosystems. In: Water Scarcity in the Mediterranean: Perspectives under Global
- 562 *Change* (Eds S. Sabater & D. Barceló), pp. 15-39. Springer-Verlag Berlin Heidelberg.
- 563 Schlosser I. J. (1990) Environmental variation, life history attributes, and community structure
- in stream fishes: implications for environmental management and assessment.
- 565 *Environmental Management*, **14**, 621–628.
- 566 Schumm S. A. & Hadley R. F (1957) Arroyos and the semiarid cycle of erosion. American
- 567 *Journal of Science*, **255**, 161-174.

- 568 Siebers A. R., Pettit N. E., Skrzypek G., Fellman J. B., Dogramaci S. & Grierson P. F. (2016)
- 569 Alluvial ground water influences dissolved organic matter biogeochemistry of pools
- 570 within intermittent dryland streams. *Freshwater Biology*, xx, xxx-xxx.
- Sponseller R. A., Heffernan J. B. & Fisher S. G. (2013) On the multiple ecological roles of
 water in river networks. *Ecosphere*, 4, 1-14. http://dx.doi.org/10.1890/ES12-00225.1
- 573 Stanford J. A., Lorang M. S., & Hauer F. R. (2005) The shifting habitat mosaic of river
- 574 ecosystems. Verhandlungen der Internationalen Vereinigung für Theoretische und
- 575 Angewandte Limnologie, **29**, 123-136.
- 576 Stanley E. H., Fisher S. G., & Grimm, N. B. (1997) Ecosystem expansion and contraction in
- 577 streams. *BioScience*, **47**, 427-435.
- 578 Stanley E. H., Buschman D. L., Boulton A. J., Grimm N. B. & Fisher S.G. (1994)
- 579 Invertebrate resistance and resilience to intermittency in a desert stream. *American*
- 580 *Midland Naturalist*, **131**, 288-300.
- 581 Steward, A. L., von Schiller, D., Tockner, K., Marshall, J. C., & Bunn, S. E. (2012) When the
- river runs dry: human and ecological values of dry riverbeds. *Frontiers in Ecology and the*
- 583 *Environment*, **10**, 202-209.
- 584 Sterling S. M., Ducharne A. & Polcher J. (2013) The impact of global land-cover change on
- 585 the terrestrial water cycle. *Nature Climate Change*, **3**, 385-390.
- 586 Storey R. G. & Quinn J. M. (2013). Survival of aquatic invertebrates in dry bed sediments of
- 587 intermittent streams: temperature tolerances and implications for riparian management.
- 588 *Freshwater Science*, **32**, 250-266.
- 589 Stromberg J. C., Lite S. J., Marler R., Paradzick C., Shafroth P. B., Shorrock D., et al. (2007)
- 590 Altered stream-flow regimes and invasive plant species: the *Tamarix* case. *Global*
- 591 *Ecology and Biogeography*, **16**, 381-393.

- Stromberg J. C. & Merritt D. M. (2016) Riparian plant guilds of ephemeral, intermittent and
 perennial rivers. *Freshwater Biology*, xx, xxx-xxx.
- 594 Stubbington R. (2012) The hyporheic zone as an invertebrate refuge: a review of variability in

space, time, taxa and behaviour. *Marine and Freshwater Research*, **63**, 293-311.

- 596 Stubbington R. & Datry T. (2013) The macroinvertebrate seedbank promotes community
- 597 persistence in temporary rivers across climate zones. *Freshwater Biology*, **58**, 1202-1220.
- 598 Stubbington R., Gunn J., Little S., Worrall T. & Wood P. (2016) Macroinvertebrate seedbank
- 599 composition in relation to antecedent duration of drying and multiple wet-dry cycles in a
- 600 temporary stream. *Freshwater Biology*, xx, xxx-xxx.
- Taylor C. M. & Warren M. L. (2001) Dynamics in species composition of stream fish
- assemblages: environmental variability and nested subsets. *Ecology*, **82**, 2320–2330.
- 603 Uys M. C. & O'Keeffe J. H. (1997). Simple words and fuzzy zones: early directions for
- 604 temporary river research in South Africa. *Environmental Management*, **21**, 517-531.
- 605 Vander Vorste R., Corti R., Sagouis A. & Datry T. (2015) Invertebrate communities in
- gravel-bed, braided rivers are highly resilient to flow intermittence. *Freshwater Science*,
- **35**, 164-177.
- 608 Vander Vorste R., Malard F. & Datry T. (2016) Is drift the primary process promoting the
- resilience of river invertebrate communities? A manipulative field experiment in an
 intermittent alluvial river. *Freshwater Biology*, xx, xxx-xxx.
- 611 Van Dijk A. I. J. M., Beck H. E., Crosbie R. S., De Jeu R. A. M., Liu Y. Y., Podger G. M., et
- 612 *al.* (2013) The Millennium Drought in southeast Australia (2001–2009): Natural and
- 613 human causes and implications for water resources, ecosystems, economy, and society.
- 614 Water Resources Research, 49, 1040-1057.
- 615 Ward J. V., Malard F. & Tockner K. (2002) Landscape ecology: a framework for integrating
- 616 pattern and process in river corridors. *Landscape Ecology* **17** (Suppl. 1), 35-45.

- 617 Welter J. R & Fisher S. G. (2016) The influence of storm characteristics on hydrological
- 618 connectivity in intermittent channel networks: implications for nitrogen transport and
- 619 denitrification. *Freshwater Biology*, xx, xxx-xxx.
- 620 Whitney J. E., Gido K. B., Martin E. C. & Hase, K. J. (2016) The first to arrive and the last to
- 621 leave: colonisation and extinction dynamics of common and rare fishes in intermittent
- 622 prairie streams. *Freshwater Biology*, xx, xxx-xxx.
- 623 Wiggins G. B., Mackay R. J. & Smith I. M. (1980) Evolutionary and ecological strategies of
- 624 animals in annual temporary pools. *Archiv für Hydrobiologie Supplement*, **58**, 97–206.
- 625 Williams D. D. (2005) *The Biology of Temporary Waters*. Oxford University Press.
- 626 Williams, D. D. (1996) Environmental constraints in temporary fresh waters and their
- 627 consequences for the insect fauna. Journal of the North American Benthological Society,
- **15**, 634-650.
- 629 Zimmermann B., Zimmermann A., Turne, B. L., Francke T. & Esenbeer H. (2014)
- 630 Connectivity of overland flow by drainage network expansion in a rain forest catchment.
- 631 *Water Resources Research*, **50**, 1457-1473
- 632

633 Figure captions

- 634 Fig. 1: Number of IR-related publications on the combined topics of fish ecology, invertebrate
- 635 ecology, biogeochemistry, hydrological assessment and ecological-condition assessment
- 636 appearing in freshwater and ecology journals, and *Freshwater Biology* specifically, through
- 637 time. Freshwater journals include any journal with the character string *water or aquatic in the
- title, ecology journals include the string *ecol in their titles, and freshwater and ecology
- 639 journals include one or more of these three strings in their titles. Data taken from a Web of
- 640 Science literature search conducted by Leigh et al. (2016 a).
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