



The duration of channel drying affects survival of *Gammarus pulex* (Amphipoda: Gammaridae) within subsurface sediments: an experimental flume study

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Abstract Field studies have demonstrated that benthic fauna use hyporheic sediments during drying events in temporary rivers, but the factors influencing the survival of fauna in subsurface sediments remain poorly quantified. Laboratory mesocosm experiments were conducted to determine how the length of drying events (1, 7, 14 and 21 days) influenced the survivorship of *Gammarus pulex* (L.) (Amphipoda: Gammaridae). The water level was reduced to 5 cm below the substrate surface during drying experiments and held at 5 cm above the sediment surface during control experiments. The results demonstrate that *G. pulex* survivorship was reduced with increasing length of the experiment, particularly in the drying treatment compared to the control treatment. We show that *G. pulex* can persist in subsurface sediments for up to 21 days during surface drying. In view of interacting climatic

drivers and water resource pressures that are increasing the spatial and temporal occurrence of streambed drying, our results have important implications for stream management. Our results also highlight the utility of mesocosm-based studies for elucidating the abiotic controls of macroinvertebrate survival during stream drying events.

Keywords Intermittent rivers · Ephemeral streams · Streambed drying · Mesocosm experiment · Drought · Survivorship

Introduction

Temporary streams experience periods of reduced hydrological connectivity as surface water levels decline, often resulting in sections of the streambed becoming dry (Boulton, 2003; Larned et al., 2010). The ecological effects of surface water level reduction and drying on aquatic organisms potentially increase over time as abiotic conditions become increasingly unfavourable (Lake, 2003; Dewson et al., 2007; Lake, 2011). Flow cessation and surface drying events typically result in a reduction in the abundance of rheophilic taxa (Chessman, 2009; Verdonschot et al., 2015), an increased dominance of generalist taxa able to persist in isolated pools (Robson et al., 2011) and the persistence of a limited number of desiccation-tolerant taxa in drying sediments (Stubbington et al.,

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2009; Datry, 2012). An increased duration of flow intermittence and channel drying can reduce benthic and hyporheic faunal density and richness as taxa reach their physiological limits (Datry et al., 2007; Arscott et al., 2010; Lancaster & Ledger, 2015), although recovery following flow resumption may occur quickly depending on refuge availability (Chester & Robson, 2011). Studies investigating temporary streams have indicated that the duration of drying represents a primary control on the structure and diversity of communities (e.g. Feminella, 1996; Datry et al., 2014; Storey, 2016). Adaptations of temporary stream fauna that enhance resistance and resilience to surface drying include behavioural responses, for example movement into saturated subsurface sediments (Vander Vorste et al., 2016a; Vadher et al., 2017), life history adaptations, for example aestivation and diapause (Strachan et al., 2015) and physiological adaptations including desiccation tolerance of egg, juvenile or adult life stages (Stubbington et al., 2016).

Subsurface sediments represent a potential habitat and refuge for benthic macroinvertebrates during adverse environmental conditions (Williams & Hynes, 1974; Stubbington, 2012). Although some studies have demonstrated that lotic benthic macroinvertebrates move into saturated subsurface sediments in response to severe low flows and surface water loss (e.g. Fenoglio et al., 2006; Wood et al., 2010; Vander Vorste et al., 2016a), others have reported limited or no evidence of subsurface refuge use (e.g. del Rosario & Resh, 2000; Young et al., 2011). These equivocal results illustrate that abiotic conditions in subsurface sediments are heterogeneous and that their use by benthic fauna depends on factors including substrate characteristics (Stubbington, 2012; Descloux et al., 2013; Vadher et al., 2017) and water quality (Lawrence et al., 2013). Such variability makes generalisation difficult, since predicting lotic community responses to drying is currently constrained by a lack of species-specific information. In light of predicted future increases in the frequency and duration of stream drying associated with climate change and other anthropogenic pressures (Ledger & Milner, 2015; Pyne & Poff, 2017), improved knowledge surrounding the influence of such changes on benthic invertebrate communities is becoming increasingly important. To control for heterogeneity within temporary streams, ex situ laboratory flume and mesocosm investigations have been undertaken to

examine the effects of surface drying on aquatic invertebrate fauna (e.g. Ledger et al., 2011; Poznańska et al., 2013; Vadher et al., 2015; Vander Vorste et al., 2016b).

Gammarus pulex (L.) (Amphipoda: Gammaridae) is an ecologically important amphipod which functions as a processor of coarse particulate organic matter (Navel et al., 2010), a predator, and as prey of both invertebrates and fish across northwest Europe (MacNeil et al., 1997). It has been extensively used in laboratory mesocosm investigations (e.g. McGrath et al., 2007; Mathers et al., 2014; Stubbington et al., 2017) and occurs widely in benthic and subsurface habitats of perennial and temporary streams (Stubbington et al., 2009; Wood et al., 2010; Verdonshot et al., 2015). *Gammarus pulex* has been observed to actively migrate into subsurface sediments in response to surface drying in both field (e.g. Vander Vorste et al., 2016a) and laboratory (e.g. Vadher et al., 2015; Vander Vorste et al., 2016b; Vadher et al., 2017) investigations.

We examined the effect of increased duration of surface water loss and sediment drying on the survivorship of *G. pulex* within mesocosms in a recirculating flume. The mesocosms allowed flow velocity, water level, water quality (conductivity, dissolved oxygen, pH and temperature) and sediment characteristics to be controlled. Our aim was to determine the extent to which *G. pulex* can survive in sediments during drying events and the effect of dry period duration on survivorship.

Materials and Methods

Development of the flume mesocosm

The experiment was conducted using a laboratory flume (219 cm long × 56 cm wide × 56 cm deep; adapted from Vadher et al., 2015). The flume was constructed using a 300 L capacity tank linked to a water cooler and pump to create a closed recirculating system held at a temperature of 12 ± 0.5 °C (Fig. 1). Five 20 cm³ mesocosms constructed from 3 mm aperture steel mesh were lined with 150 µm mesh nylon netting and filled to a depth of 10 cm with pre-washed coarse fluvial gravel (10–20 mm diameter). The mesocosms were placed adjacent to each other on a fine gravel base (2–4 mm diameter; Fig. 1). Water

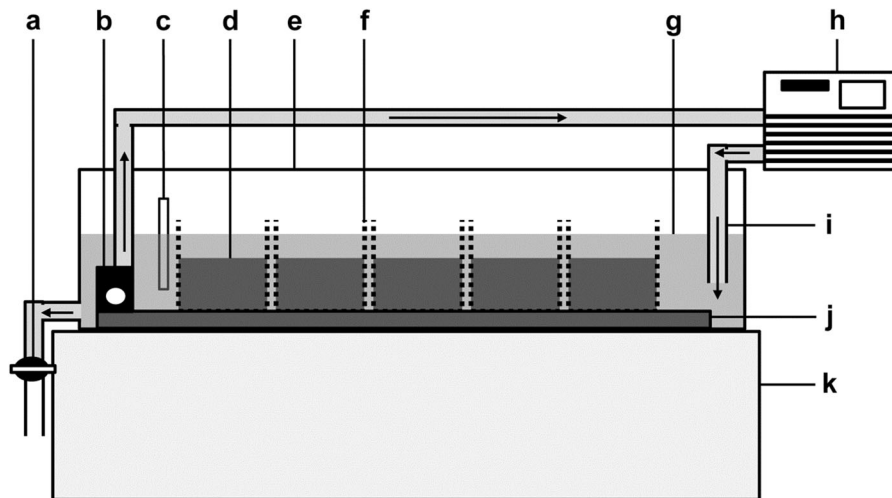


Fig. 1 Schematic cross section through the experimental flume. (a) ball valve to control water level; (b) water pump; (c) ruler; (d) coarse gravel (10–20 mm) matrix; (e) experimental tank; (f) mesocosm (20 cm × 20 cm × 20 cm); (g) water level

(5 cm above coarse gravel matrix); (h) water cooler; (i) pipe; (j) gravel base; (k) platform. Arrows represent direction of flow. Not to scale (adapted from Vadher et al., 2015)

level was controlled using a ball valve and monitored (0.25 mm accuracy) using a ruler fixed to the inside wall of the flume (Fig. 1).

Experimental procedure

Twenty-four hours prior to each experiment, each mesocosm and its sediments were washed and the experimental tank filled with dechlorinated tap water to 5 cm above the substrate surface (Fig. 1) before being circulated until a temperature of 12 °C was recorded. *Gammarus pulex* was collected using a standard kick net (900 µm mesh, 230 mm × 255 mm frame, 275 mm bag depth) from a riffle on Black Brook (52°45'46.7"N, 1°19'19.1"W) in Loughborough (Leicestershire, UK). Specimens used in experimental trials were carefully removed from the net using tweezers and those > 5 mm in length were placed into a 5-l container of stream water for immediate transport to the laboratory. Mixed pre-conditioned native leaf litter from the stream was thoroughly washed in the stream channel to remove attached stream fauna and returned to the laboratory to serve as a food source during experiments. Between four and six leaves (depending on size) were shredded and distributed through the gravel subsurface of each mesocosm. Ten *G. pulex* were then transferred into each of the five mesocosms (population density in

each mesocosm = 50 *G. pulex* m⁻²) and left to acclimatise for approximately 30 min prior to water level reduction. This acclimation period has been demonstrated to be sufficient to allow individuals to complete initial exploratory movements and seek refuge between sediment clasts (Vadher et al., 2015).

To examine the effect of drying duration (1, 7, 14 and 21 days) on *G. pulex* survivorship, three replicate experiments were conducted for each duration in both drying (water level reduction) and control conditions (water level maintained at 5 cm above the coarse gravel surface), providing a total of 24 experiments (5 mesocosms per experiment; $n = 120$ mesocosms). A water pump recirculated water through the tank at a constant rate of 41.7 l min⁻¹ for the duration of all experiments (drying and control conditions). After the initial faunal acclimation period, water level was reduced over a 2 h period from 5 cm above the gravel surface to 5 cm below the surface (dewatering rate of 5 h cm⁻¹) by opening the ball valve and reducing water level by 1.25 cm every 15 min. The rate of water level reduction was directly comparable to previous mesocosm (Vadher et al., 2015) and field research (Vadher et al., 2018) and provided a 5 cm refuge retaining free water in the subsurface sediments of the mesocosm. At the end of the water level reduction period, the flume was maintained at the

same water level (5 cm saturated gravel), temperature and flow rate for the duration of each experiment.

Assessing *G. pulex* survivorship

Upon completion of each experiment, mesocosms were individually removed and the number of surviving *G. pulex* was determined. The contents of each mesocosm were carefully excavated into a steel tray (50 cm diameter) and rinsed so that all *G. pulex* could be removed and transferred into a beaker of dechlorinated tap water. The survivorship of *G. pulex* was determined by the number of live (moving) *G. pulex* present. Whole, non-mobile individuals were considered to have died during the experiment and absent individuals or body parts were assumed to reflect predation (McGrath et al., 2007) or decomposition following stranding above the waterline. Body parts (e.g. legs, antennae) were not recorded unless the head was observed.

Abiotic parameters

Prior to the start of each experiment, pH, dissolved oxygen (mg l^{-1}), temperature ($^{\circ}\text{C}$) and conductivity ($\mu\text{S cm}^{-1}$) were measured in the free water around each mesocosm using laboratory metres (Hanna Instruments, Bedfordshire, UK). At the end of each experiment, abiotic parameters were re-measured to determine the stability of abiotic conditions throughout the experiment.

Statistical analysis

To determine if each mesocosm could be regarded as independent, a preliminary Repeated Measures (RM) ANOVA was used to examine the effect of both experiment condition (control/drying) and duration (1, 7, 14, 21 days) on the percentage survival (survivorship) of *G. pulex*. Mesocosm (1–5) was defined as a within-subject effect and experiment condition, duration and their interaction, were both defined as between-subject effects. Mauchly's tests were used to verify the RM ANOVA assumption of sphericity and the results of Greenhouse–Geisser tests were consulted when this assumption was violated. Mesocosm (RM ANOVA, Greenhouse–Geisser, $F_{2.6, 41.53} = 2$, $P = 0.136$) and its interaction with condition ($F_{2.6, 41.53} = 2.348$, $P = 0.095$), duration

($F_{7.8, 41.53} = 2.348$, $P = 0.065$) and their interaction ($F_{7.8, 41.53} = 2.348$, $P = 0.086$) did not affect *G. pulex* survivorship and mesocosms could therefore be considered independent. General Linear Models (GLM) were used to examine the effect of experiment condition, duration and the change in abiotic parameters (end–start) on *G. pulex* survivorship. Experiment condition and duration were defined as fixed factors with survivorship defined as the dependent variable. Dissolved oxygen, pH, water temperature and conductivity were defined as covariates. The effect of experiment condition and duration on these abiotic covariates was examined using GLM models by defining the mean abiotic parameters as dependent variables and experiment condition and duration as fixed factors. All analyses were conducted in IBM SPSS Statistics (version 23, IBM Corporation, New York).

Results

Overall, abiotic parameters during the experiments were similar (Table 1). Water pH and dissolved oxygen did not vary significantly between control and drying experiments (Table 2). Water temperature was marginally higher and conductivity was lower in drying experiments (Tables 1, 2). Water temperature, conductivity and pH varied among experiment durations (Table 2).

Gammarus pulex survivorship decreased with increasing duration for both experiment conditions, but was reduced to a greater extent during the drying experiments (interaction term; GLM, $F_{3, 112} = 5.562$, $P = 0.001$; Fig. 2). Within control experiments, *G. pulex* survivorship was similar after 1 and 7 days (Fishers LSD, $P = 1$). Mean survivorship ($\pm 2.08\%$ SE) following control experiments was 100% after both 1 and 7 days and was reduced to 93% and 85% after 14 and 21 days. Mean survivorship following drying experiments was 100% after 1 day, but reduced to 89, 79 and 70% after 7, 14 and 21 days, respectively. When individual abiotic parameters were considered, change in pH (GLM, $F_{1, 108} = 0.701$, $P = 0.404$), dissolved oxygen ($F_{1, 108} = 0.114$, $P = 0.736$), water temperature ($F_{1, 108} = 0.167$, $P = 0.684$) or conductivity ($F_{1, 108} = 1.929$, $P = 0.168$) did not influence *G. pulex* survivorship.

Table 1 Mean (± 1 SE) values of pH, dissolved oxygen, water temperature and conductivity after experiments in the control and drying condition

Parameter	Duration (days)			
	1	7	14	21
Control condition				
pH	8.4 \pm 0.0	8.4 \pm 0.0	8.2 \pm 0.0	8.2 \pm 0.0
Dissolved oxygen (mg l ⁻¹)	7.4 \pm 0.0	7.3 \pm 0.0	7.2 \pm 0.0	7.3 \pm 0.0
Water temperature (°C)	12.1 \pm 0.0	12.1 \pm 0.0	12.4 \pm 0.0	12.3 \pm 0.0
Conductivity (μ S cm ⁻¹)	513 \pm 1.2	509 \pm 2.1	572 \pm 6.9	572 \pm 2.4
Drying condition				
Ph	8.3 \pm 0.0	8.3 \pm 0.0	8.1 \pm 0.0	8.3 \pm 0.0
Dissolved oxygen (mg l ⁻¹)	7.1 \pm 0.1	7.2 \pm 0.1	7.2 \pm 0.1	6.9 \pm 0.1
Water temperature (°C)	12.2 \pm 0.0	13.1 \pm 0.1	13.0 \pm 0.1	12.7 \pm 0.1
Conductivity (μ S cm ⁻¹)	512 \pm 3.5	520 \pm 6.4	485 \pm 4.5	507 \pm 1.9

Table 2 General linear model (GLM) analysis of variation in abiotic parameter between control and drying experiments (experiment condition) and duration (1, 7, 14 and 21 days)

Abiotic parameter	Experiment condition			Duration		
	<i>df</i>	<i>F</i>	<i>P</i>	<i>df</i>	<i>F</i>	<i>P</i>
pH	1	2.714	0.102	3	6.896	< 0.001
Dissolved oxygen	1	1.517	0.221	3	1.580	0.198
Temperature	1	44.458	< 0.001	3	9.894	< 0.001
Conductivity	1	38.873	< 0.001	3	10.598	< 0.001

Significant terms are emboldened

Discussion

Survivorship was lower for drying experiments compared to experiments in which surface water persisted. Our findings support those of previous laboratory studies, which have reported a reduction in gammarid survival in response to drying (Poznańska et al., 2013; Vander Vorste et al., 2016b). The limitation of these studies has often been the short experimental durations, from hours (Vadher et al., 2015) to approximately a week (e.g. Poznanska et al., 2013; Vander Vorste et al., 2016b), leaving the effect of longer drying durations on gammarid survival uncharacterized. Vadher et al. (2015) observed limited vertical movements and stranding above the water line in 2 h dewatering experiments as fine sediment reduced sediment porosity in mesocosms of identical design to those used in this experiment. By extending the experimental duration, the current study extends our understanding of *G. pulex* response to surface water loss. The increased experimental duration may have

enabled any stranded individuals to move vertically within the moist subsurface sediments back into the saturated sediments. *Gammarus pulex* survivorship remained relatively high following 7 days of surface water loss compared to Vander Vorste et al. (2016b), who reported survival of up to 71% after 7 days in mesocosms experiencing water level reduction to 30 cm below the substrate surface. Survivorship in our experiments may have been higher due to the relatively short vertical distance required to move for *G. pulex* individuals to remain submerged.

Medium to coarse gravel particle sizes used in our study were unlikely to have impeded vertical movement of *G. pulex* into the subsurface (Vadher et al., 2017) and the excess leaf litter provided a plentiful food source. Subsurface leaf litter in coarse gravels has been reported as an important food source for detritivores such as gammarids (Navel et al., 2010). At the end of the experiments we observed an excess of leaf litter in the sediments which combined with the high survivorship in both the control and drying conditions,

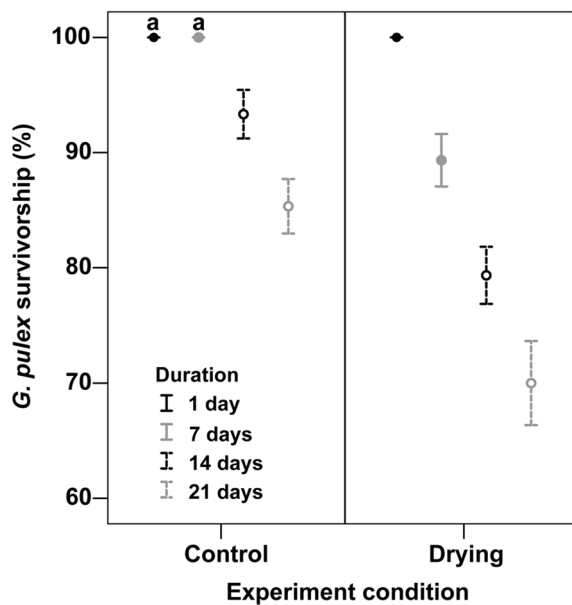


Fig. 2 Mean (\pm 1 SE) *Gammarus pulex* survivorship after each experiment duration. The left pane shows the control condition, and the right pane shows the drying condition. Letter ‘a’ indicates values which are not statistically different in the control experiment (Fisher’s LSD, $P > 0.05$). Y axis starts at 60%

which suggests that intraspecific predation was limited if it occurred (McGrath et al., 2007; Stubbington, 2012). Given that abiotic conditions and food resource availability were comparable (with the exception of water level) our results clearly illustrate the effect of channel drying on the survivorship of *G. pulex*.

Our experiments provide clear evidence that increasing drying duration reduces survivorship. This finding supports our observations made as part of a mesocosm-based field experiment investigating the effect of identical drying durations (7, 14 and 21 days) on the survivorship of *G. pulex* (Vadher et al., 2018). Under field conditions, *G. pulex* survivorship was approximately 65, 60 and 51% following 7, 14 and 21 days of dry periods, respectively (Vadher et al., 2018). In the current investigation, the effect of drying duration on survivorship was reduced compared to this field study (the lowest mean survivorship recorded was 70% after 21 days of drying in the current experiment), almost certainly reflecting the relative stability of other abiotic variables within the laboratory flume compared to field conditions. This study provides supporting evidence that the hyporheic zone functions as a refuge for benthic invertebrates if

environmental parameters (e.g. water quality and food resources) remain favourable (Hose et al., 2005; Fenoglio et al., 2006; Vander Vorste et al., 2016a). However, the relationship between survivorship and drying duration appears to be linear (Fig. 2). This indicates that subsurface sediments may be limited in their long-term refuge potential for taxa typically associated with perennial waterbodies, such as *G. pulex*, although some taxa which display adaptations to such conditions, such as some adult Coleoptera, have been recorded to persist in an active state for much longer durations in the field (e.g. Fenoglio et al., 2006). Further study is required to explore the effect of the duration of drying on macroinvertebrate taxa associated with temporary waters so that species-specific responses to drying can be quantified. This should include species from streams that dry irregularly or partially (e.g. into isolated pools—Hill & Milner 2018) and should also consider patterns of drying and proximity of perennial waterbodies and refuges (Wood et al., 2005; White et al., 2018).

The results of this study support field investigations highlighting the effect of streambed drying on instream faunal densities (e.g. Clarke et al., 2010; Datry et al., 2014; Bogan et al., 2015; Verdonshot et al., 2015). Specifically, our results show that surface drying reduces survivorship within subsurface sediments, and that this reduction increases over time. However, field studies often highlight the effect of other, usually interacting, abiotic parameters associated with channel drying (e.g. low dissolved oxygen, high conductivity and elevated water temperatures; Bond et al., 2008; Verdonshot et al., 2015). This laboratory-based mesocosm study demonstrates that ex situ experiments can facilitate understanding of the influence of abiotic factors by specifically manipulating them or holding them constant (Vadher et al., 2015, 2017; Vander Vorste et al., 2016b, 2017).

The results of this experimental study have wider implications for the study of temporary streams since they demonstrate that some common benthic fauna, such as *G. pulex*, can persist in subsurface sediments and the hyporheic zone following channel drying for at least 21 days and potentially longer. The experimental study provides further empirical evidence to support for the Hyporheic Refuge Hypothesis (Williams & Hynes, 1974) under low flow and channel drying conditions (Hose et al., 2005; Fenoglio et al., 2006; Wood et al., 2010) where the presence of free water

and abiotic conditions remain suitable for taxa to persist. However, the results highlight that knowledge of the habitats used, faunal traits that confer persistence, and the mechanisms employed by generalist and temporary water specialist taxa during channel drying events remain poorly quantified (Stubington et al., 2016). Many temporary stream specialist taxa do not typically co-exist at sites with large numbers of ubiquitous taxa, such as *G. pulex* (White et al., 2018). The present study results suggest that this lack of coexistence probably reflects the local exclusion of ubiquitous taxa, such as *G. pulex*, when drying duration increases, although the ability of taxa to persist longer periods of drying requires further research.

The predicted effects of climate change and increasing demands on water resources are likely to increase the frequency and duration of stream drying events (Ledger & Milner, 2015; Pyne & Poff, 2017). The effect of increasing drying duration on *G. pulex* survivorship could have consequences that reverberate throughout food webs if processing of coarse organic matter is reduced (Ledger et al., 2011). Given the linear relationship between *G. pulex* survivorship and drying duration, longer experiments are required to advance understanding of the effects of drying duration on survivorship and potential consequences for ecosystem functioning. This study provides further evidence of the need for effective refuge restoration and management in streams and the maintenance of streambed sediments to enhance vertical connectivity (Boulton et al., 2010). Quantifying the effects of drying and refuge use is therefore important, and future research should consider combined field and mesocosm-based approaches to facilitate greater understanding.

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