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Molecular approaches reveal weak sibship aggregation and a high dispersal propensity in a non-native fish parasite

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Abstract

Inferring parameters related to the aggregation pattern of parasites and to their dispersal propensity is important for predicting their ecological consequences and evolutionary potential. Nonetheless, it is notoriously difficult to infer these parameters from wildlife parasites given the difficulty in tracking these organisms. Molecular-based inferences constitute a promising approach that has yet rarely been applied in the wild. Here, we combined several population genetic analyses including sibship reconstruction to document the genetic structure, patterns of sibship aggregation and the dispersal dynamics of a non-native parasite of fish, the freshwater copepod ectoparasite *Tracheliastes polycolpus*. We collected parasites according to a hierarchical sampling design, with the sampling of all parasites from all host individuals captured in eight sites spread along an upstream-downstream river gradient. Individual multilocus genotypes were obtained from 14 microsatellite markers, and used to assign parasites to full-sib families and to investigate the genetic structure of *T. polycolpus* among both hosts and sampling sites. The distribution of full-sibs obtained among the sampling sites was used to estimate individual dispersal distances within families. Our results showed that *T. polycolpus* sibs tend to be aggregated within sites but not within host individuals. We detected important upstream-to-downstream dispersal events of *T. polycolpus* between sites (modal distance: 25.4 km; 95% CI [22.9, 27.7]), becoming scarcer as the geographic distance from their family core location increases. Such a dispersal pattern likely contributes to the strong isolation-by-distance observed at the river scale. We also detected some downstream-to-upstream dispersal events (modal distance: 2.6 km; 95% CI [2.2 – 23.3]) that likely result from movements of infected hosts. Within each site, the dispersal of free-living infective larvae among hosts likely contributes to increasing genetic diversity on hosts, possibly fostering the evolutionary potential of *T. polycolpus*.

Keywords

Parasite dispersal; Parentage analysis; Full-sibs; genetic structure; transmission; *Tracheliastes polycolpus*
1. **Introduction**

Dispersal is a major process influencing ecological and evolutionary dynamics, including the dynamics and persistence of populations, as well as local adaptation and speciation (Clobert et al., 2012; Dieckmann et al., 1999). In parasites, dispersal determines the evolution of life-history traits such as their transmission dynamics and their virulence (Barrett et al., 2008; Clayton & Tompkins, 1994; Criscione et al., 2005; Gandon & Michalakis, 2002; Huyse et al., 2005). Parasite dispersal is a complex process that can result from the combination of their own movements (when free-living stages exist) and that of their intermediate and/or definitive hosts (e.g., McCoy, 2008; Witsenburg et al., 2015). Over large geographical scales, parasite dispersal is generally considered as being mostly driven by the movements of their hosts/vectors (Blasco-Costa et al., 2012; Feis et al., 2015; Prugnolle et al., 2005; but see Mazé-Guilmo, Blanchet, McCoy, et al., 2016). Yet, dispersal of parasites among hosts also contributes to the overall observed dispersal pattern as soon as a free-living stage occurs, specifically at small spatial scales (e.g., Sire et al., 2001). The individual dispersal among hosts depends on both the intrinsic characteristics of free-living stages, including mobility and survival time, and the environment in which free-living stages are released (Barrett et al., 2008; Boxaspen, 2006; Samsing et al., 2015; Viney & Cable, 2011).

Estimating dispersal of parasites is fundamental to better document and predict their spread, as well as to identify potential source and sink populations of infection (Barrett et al., 2008; Blasco-Costa et al., 2012). From a practical perspective, the above information is useful to design management plans to limit parasite propagation and mitigate their impacts, notably in the case of emergent parasites. The most straightforward -yet challenging- approach to investigate dispersal consists in directly tracking individual movements. Although commonly used for large organisms (Broquet & Petit, 2009; Cayuela et al., 2018; Wikelski et al., 2007), these direct methods are generally unsuited for parasites, notably because of their small size and the difficulty to make them traceable (but see Rieux et al., 2014; Zohdy et al., 2012). Accurate spatio-temporal occurrence data can also be used to indirectly infer dispersal patterns of parasites. This approach is commonly used in epidemiology to retrace and predict the spatiotemporal dynamics of well monitored parasites and/or pathogens (Ostfeld et al., 2005; Pullan et al., 2012).
The advent of molecular approaches has greatly contributed to our understanding of parasite dispersal (Blasco-Costa et al., 2012; Giraud, 2004; Mazé-Guilmo, Blanchet, Rey, et al., 2016; Karen D. McCoy et al., 2003; Prugnolle et al., 2005). Molecular tools have mainly been used to infer parasite dispersal indirectly through the use of population genetic structure approaches and/or through phylogenetic analyses (Archie et al., 2009; Lymbery & Thompson, 2012). The examination of parasite population genetic structure at different hierarchical levels of organisation, i.e., within hosts, among hosts from the same site and among sites, is particularly valuable to assess the respective contribution of parasite transmission and host/vector movements to global parasite dispersal (Agola et al., 2009; Bruyndonckx et al., 2009; Dharmarajan et al., 2010; Mazé-Guilmo, Blanchet, Rey, et al., 2016; McCoy, 2009; Sire et al., 2001). However, these methods often rely on the presence of strong genetic signatures (Faubet & Gaggiotti, 2008; Holderegger & Gugerli, 2012) and may fail to provide accurate estimates of the geographical distances covered by parasites. Alternatively, molecular sibship reconstruction can be used to assign each parasite to at least one of their parents, their families or their populations of origin based on their multilocus genotypes (Manel et al., 2005). The membership of each parasite to a group, either a population or a family, constitutes individual traceable marks that can be used to explore the distribution of geographical dispersal distances covered by parasites in a way similar to the analyses of “dispersal kernels” (Cayuela et al., 2018; Clobert et al., 2012; Pinsky et al., 2017). Surprisingly, this approach has rarely been used for estimating dispersal parameters of parasite populations (Dubé et al., 2020; Lu et al., 2010).

Here, we empirically tested the value of combining sibship reconstruction to other population genetic tools to assess parasite dispersal and to tease apart the respective contribution of both free-living stages and host-driven dispersal in structuring parasite populations in natural landscapes. We focused on populations of the freshwater ectoparasite copepod *Tracheliastes polycolpus* and its principal local host, the rostrum dace *Leuciscus burdigalensis* (a cyprinid fish), in the Viaur River in southwestern France. We analysed the distribution of full-sib families and the genetic structure of *T. polycolpus* at different scales, by hierarchically sampling all parasites from all hosts captured within eight sites along the upstream-downstream gradient of the Viaur River. Based on the ecological information available for *T. polycolpus* and its host (see section Biological model), we built several non-mutually exclusive
predictions. After hatching, the free-living larvae of *T. polycolpus* released into the water column almost instantaneously develop into an infectious stage (Copepodid instar, see Figure 1) allowing a rapid infection of hosts (within a few days; Mazé-Guilmo et al., 2016). Moreover, daces are relatively gregarious and often behave in shoals. We thus expected that parasites from the same clutch would mostly infect their natal hosts and/or new hosts from their natal population and would thus mostly aggregate within sites. Alternatively, *T. polycolpus* free-living larvae could passively disperse with the waterflow (i.e., upstream-to-downstream biased dispersal) over “large” distances until encountering a new host. We thus expected that parasites from the same clutch would drift, infecting hosts from downstream non-natal populations. Finally, because daces are relatively sedentary and their dispersal particularly constrained by several artificial weirs and dams in the Viaur River (Blanchet et al., 2010; Clough, 1997; Clough & Beaumont, 1998), we expected that host-driven upstream-directed dispersal movements of *T. polycolpus* would only occur over short distances.
2. **Material and methods**

2.1. Biological model

*Tracheliastes polycolpus* is a freshwater ectoparasite copepod that was recently introduced in Western Europe (Rey et al., 2015) and that threatens local populations of daces (*Leuciscus sp.*) and, to a lower extent, several other cyprinid fish species (e.g., chubs, gudgeons, minnows; Loot et al., 2004; Lootvoet et al., 2013). The principal host of *T. polycolpus* *Leuciscusburdigalensis*(the rostrum dace), with a high prevalence (10% to 90%) when compared to the average prevalence on alternative hosts (1% to 10%; Lootvoet et al., 2013). *Tracheliastes polycolpus* is monoxenous, i.e., it requires a single host to fulfill its life cycle. The post-embryonic development involves three main stages: nauplius, copepodid, and chalimus (Piasecki, 1989). Nauplius is the free-living pre-infective stage. It contains an already formed copepodid inside, whose release can be very quick after hatching (almost immediately or after a few seconds or minutes). A short pre-infective phase is generally considered as an adaptation in parasitic copepods to reach the infective stage as soon as possible, hence maximizing time for infective larvae to encounter and attach on a susceptible host (Piasecki, 1989). The free-living infective copepodid (Figure 1) displays modest ability to swim and, not adapted to feeding, can live freely for about five days under laboratory conditions (Mazé-Guilmo, Blanchet, Rey, et al., 2016; Piasecki, 1989). Once attached to a host, it transforms into chalimus within five hours. Both sexual dimorphism and mating occur at this stage. Males are dwarf and able to crawl over the host body in search of a female. Females are much larger and attached to the fins of host, feeding on the mucus and epithelial cells and hence causing lesions and gradually leading to the total destruction of hosts’ fins (Loot et al., 2004). The species is monogamous, the female vaginal pore being sealed after fertilization (Piasecki, 1989; sell also Appendix S1). While males usually die very soon after mating (Kabata, 1986), females can live up to 89 days (Piasecki, 1989) and produce two egg sacs each containing up to 165 eggs (Loot et al., 2011).

2.2. *Sampling design and collection of genetic data*
We focused our study on the Viaur River, a 169 km-long river located in the Adour-Garonne drainage basin in southwestern France (Figure 2). Eight sites scattered over 80.5 km of the whole river upstream-downstream gradient were sampled during the summer 2006 (Figure 2; Table 1). Parasites were exclusively sampled on *L. burdigalensis*. At each site, daces were sampled using electric-fishing along a 50-200 m-long transect using a DEKA 7000, generating 200–500 V with an intensity range of 1 to 3 A. A total of 126 daces was captured and each was anaesthetized using clove oil (30-50 mg/L). The attached parasites to each fin, if any, were counted before being collected using forceps and stored in ethanol for subsequent genetic analyses. All host individuals were then returned alive to their original sampling site.

Individual DNA extractions were performed on parasite trunks to avoid any contamination with genetic material from eggs, following a standard salt protocol (Aljanabi & Martinez, 1997). Individual multilocus genotypes were obtained at 16 microsatellite markers (Appendix S2) for each of the 1207 parasites. The 16 microsatellite loci were co-amplified by PCR in two multiplex batches using the QIAGEN® Multiplex PCR Kit (Qiagen). The two PCR were carried out in a 10 μl final volume containing 5-20 ng of genomic DNA, 5 μl of 2× QIAGEN Multiplex PCR Master Mix, and locus-specific optimised combination of primers (Appendix S2). Both multiplex PCRs were performed in a Mastercycler PCR machine (Eppendorf®) under the following conditions: 15 min at 95 °C followed by 30 cycles of 30s at 94 °C, 90s at 56 °C and 60s at 72 °C and finally followed by a 45 min final elongation step at 60°C. The resulting PCR products were separated by electrophoresis on an ABI3730 at the GenoToul (Toulouse France). Allele scoring was performed using GENMAPPER version 4.0.

2.3. Preliminary genetic analyses

We first checked for anomalies owed to the genotyping procedure (e.g. large allele dropouts; null alleles) using Microcheckerv2.2 (Van Oosterhout et al., 2004). We then tested for linkage disequilibrium between loci and departure from Hardy-Weinberg equilibrium within each sampling site and for each locus using GENEPOP (Raymond & Rousset, 1995), with sequential Bonferroni correction to account for multiple related tests (Rice, 1989). Two markers (TRA12 and TRA66) displayed either strong deficit in heterozygosity, most likely because of the presence of null alleles, or linkage disequilibrium with
several other markers. These two loci were therefore discarded from the database in subsequent analyses. Forty individuals were genotyped twice and showed a 100% match in allele scoring at the 14 retained microsatellite markers.

2.4. Genetic diversity and structure

Genetic diversity within each of the eight sampling sites was estimated over all loci by computing the unbiased expected heterozygosity ($H_e$) using GENETIX (Belkhir et al., 2004), the standardized allelic richness ($A_r$; minimum sample size of 66; Table 1) using FSTAT (Goudet, 2001) and the $F_{IS}$ index using GENEPOP. Genetic differentiation was assessed by computing the Meirmans’ $F_{ST}$ (Meirmans, 2006) overall sites and pairwise $F_{ST}$ between sites using the rmod R-package (Winter, 2012). The effective population size $N_e$ of *T. polycolpus* within the Viaur River (all individuals combined) was estimated using NeEstimator v.2.1 (Do et al., 2014) based on a linkage disequilibrium method and setting the lowest allele frequency to 5%, considering monogamous mating and using 95% confidence intervals based on Jackknife resampling. We expected $N_e$ to be small, since metazoan parasites generally have smaller effective population sizes than free-living species (Criscione & Blouin, 2005).

We then explored how the genetic diversity of *T. polycolpus* was genetically and spatially structured among sampling sites along the Viaur River using three independent approaches. First, we tested whether the global spatial pattern of genetic differentiation between sites along the Viaur River followed a pattern of isolation-by-distance. To do so, we performed a mantel test between matrices of pairwise measures of genetic differentiation and geographical riparian distances (i.e., geographical distances along the water course; Blanchet et al., 2010) between sites using the R-package vegan (Oksanen et al., 2020). The Mantel correlation $r$ was computed and the associated $P$-value was calculated using 10,000 random permutations. Additionally, we performed a non-directional Mantel correlogram (Borcard & Legendre, 2012; Smouse & Peakall, 1999) using the R-package ecodist (Goslee & Urban, 2007) with one-sided Mantel tests with 1,000 permutations and geographical riparian distance classes defined every 10 km (up to 80 km). Secondly, we performed a discriminant analysis of principal components (dAPC) using the R-package adegenet (Jombart et al., 2010) for a visual assessment of between-sites differentiation. Finally,
we performed an analysis of molecular variance (AMOVA) using ARLEQUIN V.3.5 (Excoffier & Lischer, 2010) to measure the amount of overall genetic variance of T. Polycolpus explained by each of the three hierarchical structure levels considered within the Viaur River: (i) within hosts, (ii) among hosts within sites and (iii) among sites.

2.5. Reconstruction of full-sib families

Full-sibs families of T. polycolpus were reconstructed using the full-likelihood approach implemented in COLONY2.0 (Jones & Wang, 2010) based on the 1207 individual multilocus genotypes. Briefly, COLONY 2.0 implements full-pedigree likelihood methods, i.e., with likelihood considered over the entire pedigree, to infer sibship among individuals. We assumed that both sexes are monogamous and we allowed for possible inbreeding. All individuals were considered as offspring in COLONY 2.0 and we defined no a priori candidate parental genotypes (neither males nor females). Allele frequencies were directly determined from the genetic dataset using COLONY version 2.0. Only the full-sib families with associated inclusion probability higher than 95% were retained for further analyses.

2.6. Distribution of full-sib families

We first assessed whether full-sib individuals were rather clumped within the same site or randomly distributed across sites. To do so, we built two binary matrices that respectively included (i) the membership status of each pair of individuals to the same family (i.e., 1: parasites are full-sibs, 0: parasites are not full-sibs; hereafter called the sibship matrix) and (ii) the membership status of each pair of individuals to the same site (i.e., 1: parasites share the same site, 0: parasites come from different sites; hereafter called the site matrix). Based on our observed dataset, we computed the proportion of full-sib pairs sharing the same site (i.e., pairs of individuals displaying values of 1 in the two matrices) given the total number of full-sib pairs over the river (i.e., pairs of individuals that displayed value of 1 in the sibship matrix). This observed proportion was compared to a series of expected proportions under the null hypothesis of a random distribution of full-sib pairs among sites, using 10,000 random
permutations of the site matrix to compute the probability of correctly rejecting the null hypothesis (Legendre & Legendre, 1998).

Similarly, we assessed whether full-sib individuals were rather clumped on the same host or randomly distributed across hosts. Because hosts were not distributed homogeneously among the eight sampling sites, we considered each site independently. For each site, we first built two binary matrices that respectively included (i) the membership status of each pair of individuals to the same family (sibship matrix) and (ii) the membership status of each pair of individuals to the same host (i.e., 1: parasites are on the same host, 0: parasites are on different hosts; hereafter called the host matrix). We then computed the proportion of full-sib pairs sharing the same host (i.e., pairs of individuals displaying values of 1 in the two matrices) given the total number of full-sib pairs within the considered site and compared this observed proportion to a series of expected proportions under the null hypothesis of a random distribution of full-sib pairs among hosts, using 10,000 random permutations of the host matrix.

2.7. Estimation of T. polycolpus dispersal

To investigate the dispersal of T. polycolpus along the Viaur River, we focused on a subset of full-sib families including at least 5 full-sibs (N = 94 families). For each of these 94 families, we first determined a “family core location” as the mode of the kernel distribution of the geographical distance of each family member to the river source using the R-package stats (R core team; 2014). Next, we computed for each of the 94 families (i) a “downstream maximal dispersal distance” estimated as the difference between the estimated “family core location” and the distance of the most downstream family member to the river source, and (ii) an “upstream maximal dispersal distance” estimated as the absolute value of the difference between the estimated “family core location” and the distance of the most upstream family member to the river source. We then calculated the modes of the distributions of both the downstream and the upstream maximal dispersal distances across the 94 families. These modes provide a proxy of the most common maximal downstream and upstream distances covered by T. polycolpus from the family core location. We computed 95% confidence intervals about these upstream and downstream distance modes using 10,000 bootstrap replicates. Finally, we tested whether the upstream and
downstream maximal dispersal distances from the family core location were significantly different using a non-parametric Wilcoxon test implemented in the R-package stats (R core team; 2014).
3. Results

3.1. Genetic diversity and structure

From 126 captured daces, 114 were infected by *T. polycolpus* (parasite prevalence over all sites of 90.5%) with a parasite load of 13.4 ± 13.2 (mean ± SD). A total of 1207 parasites were sampled from infected hosts. Over all sampling sites, $H_e$ was 0.52 ±0.01 (mean ± SD), Arranged from 3.43 to 4.05 and $F_{IS}$ ranged from -0.02 to 0.01 (Table 1). The effective population size Ne of *T. polycolpus* at the river scale was 537.6 (95% CI [334.2, 885.1]). The mean genetic differentiation estimated overall sites and overall loci was $\phi_{ST} = 0.08$ and pairwise $\phi_{ST}$-values between sites ranged from 0 to 0.21, suggesting weak to moderate genetic structure in the Viaur River. We found however a strong and significant correlation between pairwise $\phi_{ST}$ and pairwise riparian distances between sites ($r = 0.90$; P-value < 0.001) as expected under an isolation-by-distance pattern (Figure 3A). Additionally, the non-directional Mantel correlogram indicated that parasites from sites distant by less than 20 km tend to be more genetically similar than expected by chance (Figure 3B). These results are in accordance with the high overlap observed between sites within the retained dAPC parameter space (two first components, together explaining 92.3% of variance) and the slight upstream-to-downstream gradient along the first component (Figure 3C).

According to the AMOVA analysis, most of the genetic variation in *T. polycolpus* in the Viaur River was actually observed within individual hosts (i.e., 97.9%; $\Phi_{ST} = 0.021$; $p$-value < 0.01; Table 2). The “among site” level explained a weak (yet significant) amount of total genetic variation (2.17%; $\Phi_{CT} = 0.022$; $p$-value < 0.01; Table 2), whereas no partition of the total genetic variation was attributed to the “among hosts within sites” level ($\Phi_{SC} = 0$; $p$-value = 0.52; Table 2).

3.2. Reconstruction and distribution of full-sibs families

Overall, 1075 out of the 1207 genotyped parasites were assigned to 160 full-sib families with a probability higher than 95%. On average, reconstructed full-sib families were composed of 6.8 individuals (ranging from 1 to 35; Appendix S3).

We found that 21.0% of the 5450 full-sib pairs reconstructed over the Viaur River belonged to the same site (Figure 4A). This proportion, although moderate, was significantly higher than the expected
theoretical proportion (14.3%) under the null hypothesis (i.e., pairs of full-sibs are distributed randomly across the eight sampling sites; \( \chi^2 = 200.75, \) d.f. = 1, P-value < 0.001). Moreover, the proportion of full-sib pairs belonging to the same site and infecting the same host differed slightly (but significantly) between sites, and ranged from 5.7 to 23.6% (Figure 4B). Yet, none of these local proportions significantly differed from the expected theoretical proportions under the null hypothesis (i.e., pairs of full-sibs are distributed randomly over the sampled hosts within each site; Appendix S4).

3.3. Estimation of T. polycolpus dispersal

The family core location estimated for each reconstructed full-sib families with more than five full-sibs ranged from 48.6 to 127.2 km from the river source (mean = 68.3 km; Appendix S5). The downstream maximal dispersal distance from the family core location ranged from 0 to 77.9 (mode = 25.4 km, 95% CI[22.9, 27.7]; Figure 5). The upstream maximal dispersal distance from the family core location was significantly lower than the downstream distance (P-value < 0.01) and ranged from 0 to 78.6 km (mode = 2.6 km, 95% CI[2.2, 23.3]; Figure 5).
4. Discussion

By combining sibship reconstruction with more classical population genetic tools, we were able to estimate various dispersal parameters in the parasite *T. polycopus*. Hereafter we will discuss the dispersal dynamics of *T. polycopus* among hosts and among sites at a river scale. We will also discuss the relative contribution of the passive copepodid dispersal and of the host-driven chalimus dispersal in shaping the genetic structure of populations as well as the possible evolutionary outcomes of such dispersal dynamics for both parasite and host populations.

We found that most of the full-sib family members of *T. polycopus* do not infect the same host—and hence not their natal hosts—as a clump but are rather scattered over several host individuals. Consequently, at each generation, the dispersal of free-living copepodid among hosts probably contributes to the genetic mixing of unrelated adult breeders within hosts. Accordingly, the AMOVA revealed that most of the genetic variability of *T. polycopus* along the river occurs within hosts (Table 2). The research of a sexual partner by males occurring once on the host, this dispersal strategy may contribute to limit the probability of mating between related individuals (random mating) and to minimize the possible detrimental effects resulting from inbreeding depression. Theoretical models predict that multi-infection of hosts by parasites from distinct strains can increase parasite virulence (Buckling & Brockhurst, 2008; López-Villavicencio et al., 2011). Local freshwater fish species and specifically daces from the Viaur River may thus suffer from virulent *T. polycopus* variants. This is in line with a previous study showing that the pathogenic effects induced by *T. Polycopus* in the Viaur River is severe (Loot et al., 2004), and, combined with high prevalence, that they might have been responsible for the serious demographic decline of daces locally observed over the last decade (Mathieu-Bégné et al., 2019).

At the site level, a substantial (and significant) fraction of the overall reconstructed full-sib pairs was found to be ‘aggregated’ within sites (i.e., 21.0%). This pattern of within site ‘aggregation’ strongly suggests that some recently hatched *T. polycopus* infective larvae are able to persist on their natal site by infecting susceptible hosts in the close neighbourhood of their natal hosts. Two non-exclusive ecological factors may account for such a pattern. First, the very short lifetime of the non-infective nauplius stage of *T. polycopus* (Piasecki, 1989) is likely to facilitate their attachment to host individuals
neighbouring their natal hosts as soon as they are released into the water column. Second, daces are gregarious and commonly form shoals (Keith et al., 2011). Local congregations and frequent social interactions between dace hosts may improve host-to-host transmission of parasites within sites (Johnson et al., 2011), all the more so as *T. polycolpus* has recently been shown to preferentially occur at very specific microhabitats that maximize encounter rate and create hotspots of infection (Mathieu-Bégné et al., 2020). Parasite transmission between neighbouring hosts inhabiting the same location is expected to homogenize the genetic variation among hosts at the site level (e.g., Bruyndonckx et al., 2009). Accordingly, the AMOVA revealed that the “among hosts within sites” level did not contribute significantly to the overall genetic variation of *T. polycolpus* in the Viaur River.

At the river level, and despite the significant within-site ‘aggregation’ pattern of full-sibs, the overall genetic structure was weak and reconstructed families were generally disseminated over several sites, indicating successful dispersal events. The overall genetic structure was characterized by a strong isolation-by-distance pattern (Figure 3A) that suggests, according to Hutchison and Templeton (1999), that populations of *T. polycolpus* were at migration-drift equilibrium. This isolation-by-distance pattern also conforms to the results obtained from the AMOVA, which indicates that a significant fraction of the overall genetic variability of *T. polycolpus* along the river occurs among sites. With dispersal among hosts facilitating random mating and dispersal among sites resulting in gene flow, the hierarchical dispersal strategy of *T. polycolpus* probably contributes to maintaining high genetic diversity (high expected heterozygosity *He* and low *F*<sub>IS</sub> values; Table 1) despite limited effective population sizes (Criscione & Blouin, 2005) and may explain the reported invasion success of *T. polycolpus* (Mathieu-Bégné et al., 2019; Mathieu-Bégné et al., 2020; Rey et al., 2015).

Determining the respective contribution of free-living copepodid dispersal and host-driven chalimus dispersal is challenging. Yet, several lines of evidence may help disentangling these two modes of dispersal. As for most riverine free-living organisms with low dispersal ability, copepodids are expected to drift passively downstream their hatching sites due to the unidirectional water flow (Paz-Vinas & Blanchet, 2015). We accordingly detected an upstream-to-downstream dispersal bias from the estimated core location of *T. polycolpus* families, with the majority of downstream dispersal events...
occurring over the first 25.4 km. It is noteworthy that this direct estimate of downstream dispersal distance is highly congruent with the Mantel correlogram (Figure 3b), with demes becoming genetically differentiated as soon as they are distant from more than ~20 km. Host-driven dispersal of fixed adult parasites is also likely to contribute to the overall dispersal of *T. polycolpus* along the river. However, daces are relatively sedentary, spending extended periods in a single site before moving towards surrounding sites within a mean radius of two kilometres and up to ten kilometres over the year (Clough, 1997; Clough & Beaumont, 1998). Moreover, dispersal of daces in the Viaur River is highly limited given the important number of obstacles (weirs and dams) that scatter the river (~1 obstacle every 2-3 kilometres in average; Blanchet et al., 2010). This suggests that host-driven dispersal of *T. polycolpus* either downstream or upstream may regularly occur, but may be limited over short geographical distances. Thus, we argue that long upstream-to-downstream dispersal events of *T. polycolpus* (twice the distance covered by their hosts annually; Blanchet et al., 2010; Clough, 1997) likely result from the drift of free-living infectious larvae with waterflow. At smaller geographical scale, dispersal of *T. polycolpus* may be driven by the combination of both free-living and host-driven movements. Interestingly we also detected some downstream-to-upstream dispersal events that mostly occur over short geographical distances (i.e., 2.6 km; Figure 5). The swimming ability of copepodid is clearly insufficient to overcome the water flow of the Viaur River (Piasecki, 1989). Thus, the downstream-to-upstream dispersal of *T. polycolpus* detected testifies the frequent although spatially constrained host-driven movements from downstream to upstream sites once the infective larvae are fixed to their host.

Overall, these conclusions about *T. polycolpus* dispersal strategy are based on the use of an original methodological framework that was made possible by the specific life history traits of both the considered parasite and its host: *T. polycolpus* is a strictly aquatic and monoxenous parasite (i.e., a single host is required to fulfill its life cycle) that is mostly found on *L. burdigalensis* in the studied system. The latter showing both small population sizes and spatially limited movements in the studied system (Mathieu-Bégné et al., 2019), we probably sampled a representative proportion of both hosts and parasites at each sampling site (total sample size twice as high as estimated total effective population size). Furthermore, the monogamous mating system of the parasite strongly facilitated the reconstruction
of family groups. We acknowledge that this approach might be more difficult to implement in other host-parasite systems, such as in terrestrial habitats or with species showing more complex life history traits.

5. **Conclusion**

   Documenting the hierarchical genetic structure and quantifying the dispersal of parasites is crucial to better understand their evolutionary potential and dynamics. By combining various population genetic tools including sibship reconstruction, we found that *T. polycopus* sibs tend to be aggregated within sites but not within hosts. This pattern may contribute to maintain high genetic variation on each host through random mating, with possible positive evolutionary outcomes in terms of individual fitness and/or parasitic virulence. We also deciphered the relative importance of free-living dispersal of *T. polycopus* and host-driven dispersal of fixed adults along the river. Our results suggest that *T. polycopus* displays a substantial ability to disperse throughout its lifetime, through passive downstream dispersal at the copepodid stage and through host-driven upstream dispersal at the chalimus stage. This hierarchical dispersal strategy may contribute to maintaining high genetic diversity despite limited effective population sizes and is probably one of the various traits that may explain the invasion success of *T. polycopus* since its recent introduction within the Viaur River and most likely overall French watersheds (Mathieu-Bégné et al., 2020; Rey et al., 2015).
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Authors’ Contributions

JGP: Formal analysis (Equal), Investigation (Equal), Methodology (Equal), Writing—original draft (Equal), Writing—review & editing (Equal). KSP: Formal analysis (Equal), Investigation (Equal), Methodology (Equal), Writing—original draft (Equal), Writing—review & editing (Equal). SB: Conceptualization (Supporting), Methodology (Supporting), Project administration (Supporting), Supervision (Equal), Writing—review & editing (Equal). GL: Conceptualization (Equal), Funding acquisition (Lead), Methodology (Supporting), Project administration (Supporting), Supervision (Equal), Writing—review & editing (Equal). OR: Conceptualization (Equal), Funding acquisition (Supporting), Investigation (Equal), Methodology (Equal), Project administration (Lead), Supervision (Equal), Writing—original draft (Equal), Writing—review & editing (Equal).

Ethics Statement

Fieldwork was conducted with adequate administrative permits for electrofishing (Permit #2005-34-4 delivered by the “Direction Départemental de l’Aveyron”) and fish were treated in accordance with the French Law (Use of live animals for scientific purposes; Articles R214-87 to R214-137 of the rural code).

Data Accessibility Statement
Microsatellite data are available on Figshare: 10.6084/m9.figshare.14038901.

Conflict of Interest

The authors declare no conflict of interest.

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Table 1: Sampling sites of *T. polycolpus* over the River Viaur and genetic diversity estimated across loci at each sampling site or averaged across sites (ALL). Ar: Mean standardized allelic richness; He: expected heterozygosity.

| Sampling site | Locality | Distance from the source (km) | N_{Hosts} | N_{Parasites} | Ar  | He  | F_{IS} |
|---------------|----------|-----------------------------|-----------|---------------|-----|-----|--------|
| V01           | Bannes   | 48.61                       | 14        | 231           | 3.93| 0.53| 0.012  |
| V02           | Capelle  | 52.14                       | 18        | 257           | 3.95| 0.53| -0.014 |
| V03           | Fuel     | 67.23                       | 12        | 108           | 4.05| 0.52| -0.029 |
| V04           | Serres   | 69.44                       | 15        | 100           | 3.87| 0.51| -0.012 |
| V05           | Serres   | 69.44                       | 15        | 100           | 3.87| 0.51| -0.012 |
| V06           | Albinet  | 75.15                       | 18        | 200           | 3.76| 0.52| -0.016 |
| V07           | Navech   | 93.77                       | 17        | 136           | 3.73| 0.50| -0.025 |
| V08           | Just     | 99.97                       | 12        | 109           | 3.73| 0.52| -0.007 |
| ALL           |          | 14.25                       | 150.88    | 3.81          | 0.52| -0.01|

Table 2: Results of the Analysis of Molecular Variance (AMOVA). D.f.: degrees of freedom.

| Source of variation | D.f. | Sum of squares | Variance component | % of variation |
|---------------------|------|----------------|--------------------|----------------|
| Among sites         | 7    | 191.93         | 0.08               | 2.17           |
| Among hosts within sites | 106  | 382.14         | -0.002             | -0.07          |
| Among individuals within hosts | 2300 | 8417.65        | 3.66               | 97.90          |
**Figure legends**

**Figure 1.** Pictures of *Tracheliastes polycolpus* at different stages. A: parasitic adult females at chalimus stage (indicated by white arrows) attached to a host (*Leuciscus burdigalensis*). B: mature parasitic adult female carrying two eggs sacs. C: eggs of *T. polycolpus* enclosed within a maternal egg sac. D: recently hatched free-living copepodid larva ready to infect a new host.

**Figure 2.** Localisation of the eight sampling sites along the River Viaur in France. Tributaries are in light grey.
Figure 3. A: Scatterplot and best fit linear trend of the Mantel test relating pairwise estimates of genetic differentiation $\phi_{ST}$ and pairwise riparian geographical distances between sites. B: Scatterplot of the non-directional Mantel correlogram, representing Mantel correlation values (r) obtained between pairwise estimates of genetic differentiation $\phi_{ST}$ and pairwise riparian geographical distances between sites, with riparian distances classes defined every ten kilometres. Grey points stand for significant (or very close to significance) p-values. Error bars bound the 95% confidence interval about r values as determined by bootstrap resampling. C: Scatterplot of individuals along the two first components of the dAPC and barplot of eigenvalues; each color (points and ellipse) of the scatter plot represent a sampling site.
**Figure 4.** Percentage of the reconstructed full-sib pairs sharing the same host (black boxes), the same site (grey boxes) and different sites (white boxes) along the whole river (A) and within each sampling site (B). The lower case letters in B indicate sites that do not differ statistically in the percentage of full-sib pairs sharing the same host.

![Figure 4 Diagram]

**Figure 5.** Distribution of the upstream and downstream maximal distances covered by individuals from the core location of their family. Only families with more than five full-sibs were considered (n = 94). The modes of the upstream and downstream maximal distances distributions are indicated by dotted lines. The 95% confidence intervals around the upstream and downstream distance modes are highlighted in shaded grey.

![Figure 5 Diagram]
APPENDICES

**Appendix S1.** As a part of preliminary work for a previous study (Loot et al., 2011), we genotyped 20 eggs (5 eggs per egg sac per female for 2 females) at several polymorphic microsatellite loci and found a maximum of 2 alleles per locus across eggs from the same female, clearly indicating that all eggs originate from the same unique father (unpublished data).
Appendix S2. Main characteristics of the 16 microsatellite markers used in *Tracheliastespolycolpus* and composition of the two PCR multiplexes. Discarded loci are in italics. He: expected heterozygosity; Ho: observed heterozygosity; Na: number of alleles.

| PRIMER NAME | MULTIPLEX (and DYE) | PCR PRODUCT SIZE (min-max) | PRIMER LEFT SEQUENCE | PRIMER RIGHT SEQUENCE | MOTIF | He | Ho | Na |
|-------------|----------------------|-----------------------------|----------------------|-----------------------|-------|----|----|----|
| TRA5        | B (FAM)              | 151-159                     | CAGTGGGCAACAAAGAAAAT | AGTTTGGGGAAAACCTGCT   | ATTG  | 0.130 | 0.131 | 4  |
| TRA6        | B (HEX)              | 193-197                     | AAAGGAAAGGCATTTGCC   | GGAATGTTGCAAAGGGAT    | CATT  | 0.485 | 0.504 | 2  |
| TRA12       | A (FAM)              | 164-184                     | CTCAGTTACACACCAGTTTC | GAAAGGCTCACAGCAAAAAGTC | TATT  | 0.434 | 0.439 | 6  |
| TRA20       | B (FAM)              | 195-197                     | CAGGATAGGTGATGATG    | CAGTTTCCTTGAGTTGCA    | AG    | 0.362 | 0.367 | 2  |
| TRA33       | B (ATTOS50)          | 188-196                     | CAGTCGGACAGCGCAATA   | ATTTTGAGCCGACTCCGGA   | AG    | 0.669 | 0.678 | 5  |
| TRA42       | A (FAM)              | 264-280                     | CACAAATGTGCAGGAAATG  | GGCTCTGIAAGTGATACG    | GA    | 0.640 | 0.663 | 5  |
| TRA44       | A (HEX)              | 275-279                     | CTTGAAATCGAATGAAACCA | GGAAGATGTAATGGAATAATC | GA    | 0.638 | 0.639 | 3  |
| TRA49       | B (HEX)              | 288-294                     | AGGAACTTCGGAGCTATGA  | CACACGACACAGACACACTCA | GA    | 0.511 | 0.505 | 5  |
| TRA53       | B (ATTOS50)          | 314-322                     | CCCACAAAATAGCTTGGT   | CATTTTTAGAAGCTTGGCT   | TC    | 0.572 | 0.566 | 5  |
| TRA58       | A (FAM)              | 316-330                     | TTACGCGGGAAGGACACTG  | GAGGTTTTACGCGACCTGC   | GA    | 0.513 | 0.507 | 5  |
| TRA59       | A (HEX)              | 330-336                     | GCACAGCTAATAGTTTGT   | CTCTGAGCTTTACGGCATT   | TC    | 0.479 | 0.497 | 4  |
| TRA62       | B (FAM)              | 344-366                     | CGGCTAGAGGGAGCGAAGA  | TCCATTTTCTGATGGCATCA  | TC    | 0.691 | 0.686 | 6  |
| TRA66       | B (HEX)              | 351-361                     | TCCCTGATGGCCACACACA  | TCTCAATTAATATTTTCTTCT | AG    | 0.393 | 0.210 | 4  |
| TRA76       | A (ATTOS50)          | 115-139                     | CAGCAACTTATAAGATATTACAC | ACTGCGCTTAAACACAAAG | TA    | 0.681 | 0.698 | 12 |
| TRA81       | A (FAM)              | 111-115                     | TGCTACCTTCCTGATGCCC  | AGGCCCTCTCTGCTATTT    | AG    | 0.417 | 0.423 | 3  |
| TRA90       | A (ATTOS50)          | 189-199                     | AGATGTCAAACCTGCGGATG | TTCCATAACCCCAACGGGAC  | TC    | 0.473 | 0.486 | 5  |

Appendix S3. Distribution of the number of full-sib members obtained in the 160 reconstructed full-sib families of *T. polycolpus*.
Appendix S4. Observed proportion of full-sibs sharing the same host within each site (Obs) compared to the series of expected proportions of full-sibs infecting the same host within each site under the null hypothesis (i.e., pairs of full-sibs are distributed randomly among hosts within each site) obtained after 10,000 permutations of the host matrix.
Appendix S5. Distribution of the family core location estimated for the reconstructed full-sib families of *T. Polycolpus* including more than five full-sibs.