

The power of numbers: dynamics of hatching and dormant egg production in two populations of the water flea Daphnia magna

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Abstract Many organisms that live in inland standing waters produce dormant life stages that can accumulate in propagule banks to survive temporarily unfavourable periods. These egg banks have important effects on the ecology of populations and communities in terms of phenology, population densities, the probability of extinction, species diversity and habitat connectivity in time and space. They also have important consequences for the evolutionary versatility of populations. Although diapause and dormant

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egg banks in freshwater zooplankton have been studied for several decades, little is known about the quantitative contribution of egg production and hatching to yearly egg bank budgets and their seasonality in natural ponds and lakes. Here we quantified inter- and intra-annual variation in hatching and dormant egg production in the water flea Daphnia magna in two natural shallow ponds in Flanders (Belgium) using high-intensity sampling and *in situ* measurements. Hatching started in spring and occurred in several bouts (April–July), accumulating to a yearly average total of 3.5×10^3 hatchlings m⁻². Dormant egg production occurred in one-to-three bouts mainly during late spring and summer (May–August), resulting in a total yearly production ranging from 1.2×10^4 up to 17.3×10^4 ephippia m⁻². In both years, there was an average surplus of 3.14 \times 10⁴ and 15.24×10^4 ephippia produced m⁻² for ponds OM2 and OM3, respectively, contributing to the accumulation of the persistent egg bank. We discuss the ecological and evolutionary consequences of both the high number of ephippia that are produced and the high number of hatchlings at the start of each growing season.

Keywords Dormant egg bank dynamics · Propagule bank · Dormancy · Zooplankton · Seasonality · Daphnia

Introduction

To survive periods of unsuitable conditions, many organisms have developed resistant life stages such as seeds, dormant eggs or spores (Harper [1977;](#page-12-0) Wiggins et al. [1980](#page-13-0)). Dormancy—a metabolic resting phase allows species to increase their generation time: from decades to even centuries (Hairston [1996\)](#page-12-0). The propagule bank can be an effective refuge against unfavourable conditions and thus facilitates population persistence (Hairston [1996\)](#page-12-0), but also contributes to the genetic resilience of populations (Latta et al. [2010\)](#page-12-0) and to the persistence of species and genes at the metapopulation level through facilitating dispersal (Gyllström and Hansson 2004). The presence of a dormant reservoir of genotypes and species in the sediment has important consequences for population and community dynamics (Brendonck and De Meester [2003;](#page-12-0) Gyllström and Hansson [2004;](#page-12-0) Orsini et al. [2016\)](#page-13-0). It influences species and genotype diversity through the storage effect (Chesson [1994](#page-12-0); Chesson and Warner [1981\)](#page-12-0) and impacts ecological dynamics not only in terms of reducing extinction rates of local populations but also by initiating a new population and augmenting population sizes at the start of the growing season (Bozelli et al. [2008\)](#page-12-0). A mixed egg bank with delayed hatching on the one hand potentially increases rates of evolution through an increased evolutionary potential but can, on the other hand, slow down evolution in response to directional selection (Hairston and De Stasio Jr [1988](#page-12-0)). Banks of resistant and dispersal-prone propagules can also increase connectivity at the metapopulation and metacommunity level by increasing dispersal rates (Figuerola and Green [2002;](#page-12-0) Vanschoenwinkel et al. [2007;](#page-13-0) Waterkeyn et al. [2010\)](#page-13-0).

In zooplankton populations inhabiting lakes and ponds, the dormant stages collect in the, often layered, sediment and form a dormant propagule bank (Brendonck and De Meester [2003](#page-12-0)). The size of propagule banks in lakes and ponds at a certain point in time is thought to be determined by a number of processes that together also determine the budget of the propagule bank. These include the amount of dormant eggs produced, the proportion of these that hatch in the subsequent and later seasons and their mortality due to predation, infection or senescence (Gyllström and Hansson [2004\)](#page-12-0). Most studies suggest the amount of dormant propagules in ponds and lakes can be very

high, from thousands to millions per square metre (Brendonck and Riddoch [2000;](#page-12-0) Cousyn and De Meester [1998;](#page-12-0) Hairston [1996\)](#page-12-0). Generally it is expected that the amount of dormant eggs in the sediment reflects the population size during previous' reproduction periods (De Stasio [1989;](#page-12-0) Verschuren and Marnell [1997](#page-13-0)).

How do egg banks build up in the sediments of lakes and ponds? The number of dormant eggs that are transported among habitats, such as separate lakes, tends to be low compared to the number of resident individuals in an established population (Badosa et al. [2017;](#page-12-0) Green et al. [2005](#page-12-0); Hulsmans et al. [2007;](#page-12-0) Louette et al. [2007;](#page-13-0) Vanschoenwinkel et al. [2008](#page-13-0)). Local propagule production is thus by far the dominant process that determines whether seasonal losses due to hatching and mortality are compensated (Gyllström and Hansson [2004](#page-12-0)). Little is known, however, about the phenology of hatching and dormant egg production of Daphnia under field conditions, likely because it is a laborious endeavour to collect such data. Relevant data should also be collected in the field, as it has been shown that there can be large differences between laboratory and field measurements of hatch-ing rates (Cáceres and Schwalbach [2001\)](#page-12-0). Thus farfield estimates of egg production are only available for a limited number of species (Montero-Pau et al. [2017](#page-13-0)). A well-known example is the freshwater copepod Diaptomus sanguineus, where a net accumulation of dormant eggs in the sediment was found over a 3-year period in Bullhead Pond, a permanent shallow body of water (De Stasio [1989,](#page-12-0) [1990](#page-12-0)). In Oneida Lake, a large shallow lake in NY, Cáceres ([1998](#page-12-0)) quantified both hatching and dormant egg production of Daphnia galeata mendotea and Daphnia pulicaria over a 3-year period (1992–1995). She documented strong seasonality in the hatching and dormant egg production processes, showing that dormancy played a key role for the long-term persistence of the daphnid population in Oneida Lake. Similarly, Gyllström (2004) (2004) measured hatching and dormant egg production of Ceriodaphnia, Daphnia and Bosmina for 16 months in Lake Krankesjön, a shallow lake in southern Sweden. These studies used inverted funnel traps to quantify hatching, and zooplankton nets $(75 \mu m)$ and sediment core samples (Cáceres [1998;](#page-12-0) De Stasio [1989](#page-12-0)) to quantify dormant egg production.

We studied the dynamics of hatching and egg production of Daphnia magna, a large-bodied cladoceran that holds a central position in aquatic food webs as the main grazer of unicellular algae and the preferred prey of fish and macro-invertebrate predators (Miner et al. [2012](#page-13-0)). D. magna has a cyclical parthenogenetic life cycle (De Meester et al. [2004](#page-12-0); Hebert [1987](#page-12-0)). In general, this entails that at the start of the growing season, which is early spring in the study region, female D. magna individuals hatch from dormant eggs. As these eggs have been produced sexually, this results in a genetically diverse population. These individuals give rise to clonal lineages that reproduce asexually as long as conditions are favourable. Sexual reproduction is induced when conditions start to deteriorate: dormant eggs are produced that are encapsulated by a protective membrane called an ephippium (Mitchell et al. [1998](#page-13-0)).

In this study, we aimed to reconstruct the pattern of hatching, population densities and dormant egg production of *D. magna*. To this end, we engaged in a detailed sampling of two shallow ponds over a period of 3 years. We tested three hypotheses: (1) that hatching is characterized by a pronounced hatching peak in early spring, perhaps followed by several smaller hatching peaks later in the season (as observed by De Stasio [1989;](#page-12-0) De Stasio and Hairston [1992\)](#page-12-0); (2) that dormant egg production is high just after the population peak in summer as resources are depleted (Alekseev and Lampert 2001 ; Gyllström 2004) as well as in autumn in response to cues indicating the onset of the unfavourable winter conditions (Gyllström and Hansson [2004](#page-12-0)); and (3) that the egg budget shows a positive balance that would lead to a gradual accumulation of dormant eggs in the mixed egg bank.

Material and methods

Study sites

Both study systems (OM2 and OM3) are small (OM2: 2.53 ha; OM3: 1.84 ha) shallow (average depth 0.71 m) ponds located in Abdij van 't Park (Heverlee, Belgium; 50°51'48"N, 4°43'16"E). They are part of a group of four neighbouring eutrophic ponds that were created in the eighteenth century for fish production, which continued until 1993. Overall, the ponds have a simple morphometry, with littoral zones that are partly covered by reed and quite rapidly reach a depth of 0.5 to 1 m. At the time of the sampling, a fish community was still present, consisting mainly of carp (Cyprinus carpio), perch (Perca fluviatilis), Prussian carp (Carassius gibelio) and tench (Tinca tinca) (Decaestecker et al. [2005](#page-12-0)). The D. magna populations and dormant egg banks in these ponds have been used in earlier studies, including a resurrection ecology study (Decaestecker et al. [2007\)](#page-12-0) and population genetic analyses (Vanoverbeke et al. [2007;](#page-13-0) Vanoverbeke and De Meester [2009\)](#page-13-0). From this background information, we know that the ponds have a welldeveloped dormant egg bank and that there are yearly hatching bouts replenishing genetic diversity at the start of the growing season (De Meester et al. [2006](#page-12-0)). These studies have also shown that individuals may survive during mild winters and that populations can remain active in the water column for prolonged periods of time, from several months to years (Vanoverbeke and De Meester [2009\)](#page-13-0).

Quantifying hatching

We quantified hatching throughout the year in both 2002 and 2003. Starting in February, the ponds were monitored daily for hatchlings using a Schindler– Patalas zooplankton trap of 12 L. Each day, four 12-L samples were taken at eight sampling points at random locations in the pond (total: 384L). Starting late February, eight sediment cores of each 21.24 cm² (total surface area: 169.92 cm^2 , total volume: 339.8 cm³) were taken every week at different, randomly chosen locations per pond but avoiding sites that had been sampled before to avoid disturbance. Sediment sampling was intensified to once every 4 days after first hatching was observed. The total volume of mud sampled for each pond was 6,117.1 cm³ in 2002 and 10,195.2 cm³ in 2003. To quantify hatching as it would occur in the pond, the top 2 cm of each core sample was mixed and exposed in the laboratory to optimal hatching conditions of 20 $^{\circ}$ C and a light regime of 14:10 L:D. Given that resting egg development time at 20 $^{\circ}$ C is 2.69 days (Bottrell et al. [1976\)](#page-12-0), all animals that hatched within 48 h after isolation were considered to have also been ready to hatch in the field. Hatching for each time period was calculated based on the sum of the hatchlings for all eight sediment cores. Average hatching per 24 h was calculated by summing the number of hatchling over the 2 days and then dividing by two. In this way, every 4 days we obtained an estimate of the number of animals that would have hatched during a period of 24 h in the field from a surface area of 169.92 cm^2 . By multiplying the resulting number by 58.89, we calculated the amount of hatchlings per $m²$ of surface sediment per 24 h (Fig. [1](#page-4-0)). Calculations for total hatchlings (per pond and per day) and for hatchlings per litre (per pond and per day) were calculated in the following way:

Total hatchlings =
$$
\left(\frac{hatchlings}{m^2}\right) \times \text{pond area}(m^2)
$$

$$
Hatchings per litre = \frac{Total hatchlings}{pond volume (litre)}
$$

To explore the relationship between pond temperature and hatching, we obtained average daily temperature over the study period from the Uccle weather station of the Royal Meteorological Institute of Belgium (KMI).

Quantifying ephippial production

During three consecutive years (2001, 2002 and 2003), we sampled the active zooplankton community of both ponds weekly from February to October using a 12-L Schindler–Patalas trap at each of eight different locations, i.e. 96 L per week in each pond. These samples were filtered over $250 \mu m$ and visually screened for the presence of ephippia-bearing Daphnia and subsequently preserved in EtOH for population density counts of Daphnia magna. Whenever ephippial females were discovered, a 384-L sample was taken from each pond $(4 \times 12-L)$ Schindler– Patalas samples at each of eight sampling points) each week. The animals caught per pond were kept in a container, and after 24 h the number of deposited ephippia was counted. As ephippia take 2–3 days to develop, all ephippia deposited during the first 24 h after isolation were considered to be the result of natural induction in the pond. This method allowed us to quantify ephippia production at weekly intervals during the growth season, with the number of ephippia produced during a 24-h period every week. At three separate moments during the sampling seasons of 2001 and 2002, 100 newly produced ephippia were decapsulated to check for empty ephippia or damaged dormant eggs. On average, the percentage of damaged resting eggs or empty ephippia was 15% for OM2 and 23.5% for OM3. This percentage was used to correct the collected data: the number of ephippia produced was multiplied by the percentage of viable ephippia. The numbers of ephippia production presented here always refer to the corrected numbers. Ephippia production was next quantified as ephippia per $m²$ (the total amount of ephippia were calculated for the total volume and divided by the area of each pond). Both monitoring of hatching and ephippia production were stopped when in late summer or autumn the populations disappeared from the water column.

Dormant egg budget calculation

Assuming that all eggs that hatched were deposited in the preceding season, we can calculate the percentage of eggs produced in a given season (2001 and 2002) that hatched in the subsequent growing seasons (2002 and 2003, respectively). In this way, we also calculated an approximated dormant egg budget for the two studied growth seasons. In the overall balance, we neglected unmeasured factors such as egg mortality in the egg bank. Our egg budget calculations are based on detailed data on both the number of hatchlings and ephippia production, but need to be interpreted by taking into account that we did not quantify egg mortality in the sediments. For our calculations, we assumed a 1:1 relation between ephippia and dormant eggs. This is a conservative assumption since we know that D. magna usually deposits two dormant eggs in one ephippium. Using the measured production and hatching data, interpolations were done in order to have data points for each calendar date.

Results

Hatching

In 2002, a first quite pronounced hatching peak of 118 hatchlings m^{-2} per 24 h (reflecting a total of two hatchlings per day when summed over eight sediment cores) was observed in the first half of April for OM2. During the second half of April, a peak of 88 hatchlings m^{-2} (i.e. reflecting a total of 1.5 hatchlings per day over eight sediment cores) was observed for OM3. The second and third hatching peaks occurred simultaneously in May and July (Fig. [1](#page-4-0)a). The peak hatching rate observed was 206 hatchlings m^{-2} per

Fig. 1 Number of hatchlings m⁻² in OM2 (filled circles, full line) and OM3 (open circles, full line) plotted together with ambient temperatures (grey line) for a 2002 and b 2003

24 h (i.e. reflecting a total of 3.5 hatchlings per day over eight sediment cores) in OM2 and 177 hatchlings m^{-2} per 24 h (i.e. based on average hatching per 24 h of three hatchlings over eight sediment cores) in OM3. The total amount of hatchlings during April, May and July were 1266, 1118 and 1595 hatchlings m^{-2} , respectively, in OM2 and 442, 1680 and 880 hatchlings m^{-2} , respectively, in OM3.

In 2003, hatching started earlier, with a small hatching burst at the end of March and a pronounced peak in both ponds in April, with a maximum number of hatchlings of 265 m^{-2} per day (i.e. 4.5 hatchlings on a total of eight sediment cores) in OM2 and 177 m^{-2} per day (i.e. 3 hatchlings on a total of eight sediment cores) for OM3 (Fig. 1b). For OM3, this hatching bout lasted almost the entire month of April. In the beginning of May, both ponds showed simultaneous hatching peaks, but these were smaller compared with the early spring peak. The total amount of hatchlings during March, April, May and June were

74, 2186, 524 and 132 hatchlings m^{-2} , respectively, in OM2 and 191, 2657, 213 and 795 hatchlings m^{-2} , respectively, in OM3.

Based on the raw data, we counted an average of 11.75 hatchlings per year and per pond hatching from the sediment samples, i.e. a total of 47 hatchlings over all sediment samples in the study. Per unit of volume water in the ponds, we estimated that in 2002 a total of 1.2 animals hatched per litre water in OM2 and a total of 0.6 animals hatched per litre water in OM3 (assuming an average depth of 0.55 m for OM2 and 0.87 m for OM3). In 2003, a total of 1.3 and 1.0 animals hatched per litre water of OM2 and OM3, respectively.

In Figure S1, we plotted the hatching peaks per year (hatching m^{-2}) for each pond and the corresponding ambient air temperatures (see Supplementary material). There was no relationship between spring temperatures and hatching rates. It is important to note that we found active populations in both ponds and all studied years at the start of every sampling season (March). This means we missed overwintering populations and possibly early hatching events.

Ephippia production

At the end of May 2001, both study ponds had a peak production of ephippia (3835 ephippia m^{-2} for OM2, 1544 ephippia m^{-2} for OM3; see Fig. 2a). OM2 also had relatively smaller ephippia production peaks in mid-July (855 ephippia m^{-2}) and mid to end of August (30 ephippia m^{-2}), while OM3 had a small production peak at the end of June (309 ephippia m^{-2}). The raw data for 2001 reveal that we counted a total of 5388 and 987 ephippia in our samples from OM2 and OM3, respectively. In 2002, both ponds had a peak ephippia production in mid-June. The D. magna population of OM3 showed a peak production of over 11 000 ephippia m^{-2} per day, while for OM2 this was less than 1000 ephippia m^{-2} (Fig. 2b). The latter pond, however, showed a second peak of ephippia production in mid-July, reaching 1975 ephippia m^{-2} per day. Based on the raw data for 2002, a total number of 3371 and 5752 ephippia were counted for OM2 and OM3, respectively. In 2003, in both OM3 and OM2, an ephippia production peak occurred during mid-June, with a production of up to 20 000 ephippia m^{-2} per Fig. 2 Log₁₀ of ephippia production m^{-2} plotted per year (a 2001, b 2002, and c 2003) and per pond (OM2 and OM3). Ephippia production (black circle, full line) plotted against amount of ephippia per female (grey triangle, full grey line) and total abundance of D. magna per litre (thick grey line). Note that due to a high percentage of ephippia per females in OM3 in 2003, the range of the y-axis (Ephippia per D . magna female) is different in this plot (c: OM3 2003)

day in OM3 (Fig. 2c). There was no peak in ephippium production associated with the end of the growth season. Based on the raw data for 2003, a total number of 2221 and 10118 ephippia were counted for OM2 and OM3, respectively. In 2001 and 2002, the active population gradually disappeared during the month of August, but without a peak of dormant egg production. Rather, ephippium production peaked earlier in the season. The total amount of ephippia produced per pond and per year ranged from 6.3×10^8 to 10.6×10^8 for OM2 and from 2.1×10^8 to 32.0×10^8 for OM3 (see Table [1\)](#page-8-0).

Hatching fractions and egg budget

Assuming that all eggs that hatched were deposited in the preceding season, we can calculate the percentage of eggs produced in a given season (2001 and 2002) that hatched in the subsequent growing season (2002 and 2003, respectively). Reconstructed hatching percentages in 2002 and 2003 were comparable for both ponds and averaged 11.7% (Table [1\)](#page-8-0).

In the spring-summer growth season of 2002, the total reconstructed dormant egg budget provides an assessment of the degree to which the dormant egg bank of the local populations increased in size (assuming on average only one dormant egg per ephippium). For 2001–2002, OM2 showed a positive balance of 4.1 \times 10⁴ dormant eggs m⁻² while this was 13.5×10^4 for OM3; for 2002–2003, these numbers were 2.2×10^4 and 17.0×10^4 , respectively. Given the surface area of the ponds, this translates into a large number of dormant eggs that were added to the ponds: 10.3×10^8 and 24.9×10^8 for OM2 and OM3, respectively, in 2001–2002 and 5.6×10^8 and 31.2×10^8 for OM2 and OM3, respectively, in 2002–2003.

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 $\mathbf c$

Fig. 2 continued

Table 1 Hatching and ephippia production in ponds OM2 and OM3 during 2001, 2002 and 2003: Total ephippia (Eph) produced m^{-2} , total hatching m^{-2} , yearly average *Daphnia*

density (individuals/litre) and the hatching percentage (percentage hatchlings compared to the number of ephippia produced in the previous year)

The data were calculated based on our observed counts and interpolations based on these counts (to estimate numbers on days on which no observations were done). For the percentage hatching, we make two assumptions: we assume that each ephippium contains only one egg (average of 50% filling rate, the average filling rate reported by Cousyn and De Meester ([1998\)](#page-12-0) across multiple eutrophic ponds including OM2), and we assume that only eggs that were produced in the previous growing season hatched. When relaxed (higher filling rate or some hatchlings are derived from older eggs), both these assumptions would result in lower hatching percentages (i.e. lower than 3.8–25% reported here)

Discussion

We quantified ephippia production, hatching and their phenology in two natural D. magna populations during two (three for ephippia production) consecutive years. We discovered large numbers of hatchlings and a very high number of ephippia produced in these populations from moderately sized habitats. Both egg production and hatching showed marked seasonality. The surplus of dormant eggs being produced compared to the number of hatchlings resulted in the production of a persistent dormant egg bank.

Hatching rate

We used a reliable method to quantify hatching of induced eggs in the laboratory as it would occur in the field for two ponds and two consecutive years and collected detailed data for the whole period starting late winter until late summer, when the active populations disappeared. While earlier studies used hatching traps to monitor seasonal patterns in hatching, this method did not work properly in the shallow ponds we studied, as the hatchlings did not immediately swim upwards but rather built up populations in the traps (KM, pers. obs.). As the method we used reliably quantifies hatchlings whose development was induced in the field, we could assess the amount of animals that would hatch on a given day per unit surface area. Our approach might, however, have resulted in an overestimation of daily hatching rates especially in early spring. This is because we incubated the eggs at 20° C. Whereas part of the development of induced eggs will have taken place in the field at ambient temperatures, final development until hatching took place at an accelerated rate in the laboratory. Assuming that on average half of the development of the eggs took place in the field and half in the laboratory and taking into account the typical development time of Daphnia resting eggs (see Bottrell et al. [1976;](#page-12-0) Vanoverbeke and De Meester [2009\)](#page-13-0), this overestimation is probably marginal for ambient temperatures above 15° C. For ambient temperatures around 10 $^{\circ}$ C or lower (April 2002 and 2003), hatching rates may have been increased considerably. We note, however, that in all cases all eggs that we observe to hatch were initiated in the field and would have hatched anyway. As such, our assessment of hatching peaks in terms of the amount of eggs that hatched in total based on one set of sediment cores is correct. Yet, our estimates of daily hatching rates might be overestimated for early spring samples.

Each sampling day we quantified hatching for a total surface area of 169.92 cm^{-2} , and summing the surface area of all cores from a given year, we sampled on average 0.31 m^2 in 2002 and 0.51 m^2 in 2003. Extrapolating our results to the total surface area of each pond resulted in a very high total number of hatchlings in the ponds, ranging from 5.5×10^7 up to 10.8×10^7 depending on the pond and year (see Table [1\)](#page-8-0). While the error on these estimates is obviously high given that we need to extrapolate over a much larger surface area than sampled, our results at least indicate that, in total, the number of hatchlings that initiate a population is very high. Even if we consider only the individuals that hatch during the first month of the growing season, the numbers range from 1.9×10^6 to 3.2×10^7 (total hatchlings per pond). This very high number has important consequences. First, given that all dormant eggs are the product of sexual recombination, all hatchlings are genetically unique. This implies that clonal variation at the start of the growing season, even if only the first month of hatching is taken into consideration, is very high. This clonal variation provides the raw material for clonal selection to act upon (Pfrender and Lynch [2000](#page-13-0); Vanoverbeke and De Meester [2010\)](#page-13-0). Even if clonal erosion is severe (Pfrender and Lynch [2000](#page-13-0); Vanoverbeke and De Meester [2010\)](#page-13-0), it is likely that there will be hundreds or thousands of clones remaining at the end of the season. So while clonal erosion is very strong and clonal structure of the population is high, with many clones represented by thousands or even millions of individuals, it is unlikely that clonal diversity in such habitats would be reduced to the extent that only one or a few clones would remain. Moreover, while clonal selection is very effective as across generations both additive and interactive gene effects are inherited (Lynch and Walsh [1998\)](#page-13-0), it acts on the expressed phenotype, and to the extent that the same phenotypes can be coded by different gene combinations, this leads to the expression of ''hidden'' genetic variation upon recombination (Lynch [1983](#page-13-0)). This can be the case when different pathways or combinations of genes lead to the same phenotype, or when the clones that are selected for are heterozygous for specific loci. Several studies have reported an increase in heterozygosity during the course of a growth season or following a selection experiment (Hochmuth et al. [2015\)](#page-12-0), suggesting that clonal selection may even favour heterozygotes, leading to the release of genetic variation upon subsequent recombination. The high number of hatchlings at the start of the growth season in effect then results in a very broad sampling of the evolutionary potential offered by the allelic diversity present in the population (Orsini et al. [2016\)](#page-13-0). The huge clonal diversity at the beginning of the growth season thus likely has important consequences for evolutionary potential, population genetic structure, genetic drift and among-population genetic differentiation. In addition, the high number of hatchlings also has ecological consequences, as it results in a relatively high starting population, thus buffering for chance extinctions and reducing the length of the period where one would expect exponential growth in the absence of competition.

Hatching rates as quantified in the present study (on average 39.0 individuals m^{-2} day⁻¹ for OM2 and 38.4 m^{-2} day⁻¹ for OM3 across the growth season) were much higher than those measured in most other studies that monitored hatching in detail in the field. In the studies by Wolf & Carvalho [\(1989](#page-13-0)), Hairston et al. [\(2000](#page-12-0)) and Rother et al. ([2010\)](#page-13-0), the number of hatchlings ranged from 0.1 to 1 individuals m^{-2} day⁻¹. Similarly, Cáceres ([1998\)](#page-12-0) reported emergence rates for *D. pulicaria* of 0.1 to 0.75 individuals m^{-2} day^{-1} . The low hatching rates observed in some of these studies may be related to the fact that most of them involved relatively deep lakes that may provide a permanent habitat for the studied zooplankton species, so that populations invest less in the production of dormant eggs (Gilbert [2017](#page-12-0)). Moreover, the ponds studied by us are very productive and harbour a dense Daphnia population (often >50 and sometimes >200 individuals L^{-1} ; see Fig. [2](#page-5-0)). De Stasio ([1989\)](#page-12-0) observed hatching rates for Diaptomus sanguineus up to 200 individuals m^{-2} day⁻¹ in Bullhead Pond (NY). This population is eliminated each year between July and October due to intense sunfish predation.

Another example of a population that relies heavily on emergence to start up in the spring is Epischura nordenskioldi in the same pond, which shows hatching rates of 20 individuals m^{-2} day⁻¹ (De Stasio [1990\)](#page-12-0).

Hatching phenology

While hatching rates at the start of the growth season were high, hatching was not strongly synchronized and occurred, more than expected, in hatching bouts spread over the growth season. In 2002, for example, there were three approximately equally important hatching bouts in spring and summer, while in 2003 the hatching peak early in the growth season was the most pronounced, and the three hatching peaks occurred earlier in the season than in 2002. It is striking from Fig. [1](#page-4-0) that the two ponds show similar seasonality in hatching responses within a given year. This synchronicity among ponds suggests that hatching might among others be triggered by weather or other environmental conditions (e.g. food levels). The observation of several hatching peaks is in line with results on the patterns of genetic diversity observed in these populations. Multilocus genotype diversity of both OM2 and OM3 has been reported by De Meester et al. [\(2006](#page-12-0); their figure 5), and their results based on a survey carried out in 1996–1997 suggest that clonal diversity in these populations increased at different instances during the period from May to July. In addition, our observations are also consistent with the results of a hatching experiment starting from dormant eggs sampled in OM2, where high hatching rates were observed under both early spring and summer (Vanoverbeke and De Meester [2009\)](#page-13-0).

Hatching early in the season has important advantages as it gives a head start during the exponential growth phase (Vanoverbeke and De Meester [2009\)](#page-13-0). In addition, hatchlings from dormant eggs differ in life history traits from offspring that hatch from parthenogenetic eggs, as they are better adapted to the high food conditions that prevail during the spring algal bloom (Arbačiauskas and Lampert [2003\)](#page-12-0). On the other hand, there are also advantages to a more delayed hatching response that also allows hatching to occur later in the season. First, uncertainties on what is the optimal moment for early hatching, e.g. in the case of erratic fluctuations of the weather, may select for bethedging or the coexistence of alternative strategies (De Stasio and Hairston [1992](#page-12-0); Vanoverbeke and De Meester [2009](#page-13-0)). Second, if sedimentation rates are relatively high (which is the case in the OM ponds, with 1–2 cm per year Cousyn and De Meester [1998](#page-12-0)), then the reduced probability of being exposed to hatching conditions may select for a more generalistic response to hatching stimuli.

Ephippia production

Ephippia production of the D. magna populations in the two study ponds was very high, with the total number of ephippia produced yearly ranging from 1.2×10^4 to more than 17.3×10^4 m⁻². As each ephippium can harbour two eggs, the total number of dormant eggs produced yearly can mount up to $34.6 \times 10^{4} \text{ m}^{-2}$. Total dormant egg production in the copepod Diaptomus sanguineus in Bullhead Pond was similarly high, with total numbers varying between 8×10^4 and 4×10^5 individuals m⁻² per year (De Stasio [1989\)](#page-12-0). Cáceres ([1998\)](#page-12-0) found lower numbers in her study of dormant egg production by Daphnia species in Lake Oneida, with yearly totals ranging from 2.2 to 13.7 \times 10³ diapausing eggs m⁻² for *D. pulicaria* and from 0 to 1.2×10^3 diapausing eggs m^{-2} for *D. galeata mendotae.*

Phenology of ephippium production

The phenology of ephippia production is striking for both ponds, with the major peak at the end of May in 2001, in June and July in 2002 and in June in 2003. The peak in ephippium production in 2001 followed a population density peak (see Fig. [2a](#page-5-0)) and is therefore consistent with the expectation of sexual reproduction of dormant eggs induced by crowding (Hobæk and Larsson [1990;](#page-12-0) Kleiven et al. [1992\)](#page-12-0). This observation also corresponds with Alekseev and Lampert ([2001\)](#page-12-0) who revealed an association of ephippium production with maternal effects; Daphnia females switch to sexual reproduction if they are less well fed than their mothers. This in practice would often result in ephippium production after population density peaks and ensures proper timing of ephippium at the onset of less optimal conditions and at a moment that there are still many individuals in the population. The peaks in ephippium production in July and August are less clearly related to density peaks of the active population and may also be induced by the abiotic conditions in the ponds such as high water temperatures, or the occurrence of toxic algae or parasites (see also Gyllström 2004). The active *D. magna* populations inhabiting these ponds tend to disappear in late summer. The peaks in ephippium production in June and July may thus be seen as an insurance for survival. After this peak in production of sexual, dormant eggs, the females apparently go on reproducing parthenogenetically and try to survive, which sometimes results in overwintering populations (in practice, the studied populations are semi-permanent, see also De Meester et al. [2006\)](#page-12-0). We also noted that the peaks in ephippium production were determined by two factors: increases in the percentage of females that carry ephippia and increases in population size (see Fig. [2\)](#page-5-0).

Dormant egg budget

In both years, we observed a clear surplus of dormant eggs being produced compared to the number of hatchlings. Overall, only 3.7 to 9.5% of the eggs that are produced actually hatch at the next occasion (assuming on average only one dormant egg per ephippium). The high hatching percentage, 25%, of OM3 in 2002 was related to the low production of ephippia in 2001, especially when compared to the production in 2002 and 2003 (Table [1](#page-8-0)). The egg budget that we reconstructed did not cover the entire active period, as we stopped sampling in September. This is also apparent in Fig. [2](#page-5-0)c, where during our first sampling moment early spring an active population was already present, presumably consisting of individuals that had survived the winter (see also De Meester et al. [2006](#page-12-0)). In our egg budget, we only considered the main growth season from spring to late summer and did not monitor hatching and egg production in late autumn or winter. However, even considering the incomplete egg budget, our data do suggest that the dormant egg bank of the study ponds is increasing in size every year, with up to 31.2×10^8 ephippia (62.4 \times 10⁸ dormant eggs).

Our main conclusion is that even in moderately sized aquatic habitats, the amount of dormant eggs that are produced and hatch yearly can be very high. Moreover, the positive balance between eggs produced and eggs that hatch suggests a rapid build-up of a persistent dormant egg bank. We note that Daphnia densities are very high in our study systems. It might be that in ponds with lower productivity or higher predator densities, where Daphnia densities are lower, the ratio of egg production on the number of hatchlings is closer to one. Nevertheless, large egg banks as observed in our study systems have important consequences, among others because they ensure population persistence in the face of environmental change by buffering against occasional catastrophes (Rogalski et al. [2017](#page-13-0)). If, following unfavourable conditions, the habitat becomes favourable again, the population may rebound from the dormant egg bank even if the likelihood of an individual egg to hatch is very low. This is in line with the results of Mergeay et al. ([2007](#page-13-0)), who showed that a *Daphnia barbata* population that was absent from Lake Naivasha for more than 50 years was re-established by hatchlings from the local dormant egg bank. The large dormant egg bank also facilitates coexistence of alleles and species through the storage effect (Chesson [1994](#page-12-0); Chesson and Warner [1981\)](#page-12-0) and reduces the impact of genetic and ecological drift within populations through time. The increased genetic and species diversity may also favour environmental sorting and may thus lead to stronger patterns of isolation-by-environment in landscape genetic data (Orsini et al. [2013](#page-13-0)) and to stronger signals of species sorting in metacommunity data (Urban et al. [2008](#page-13-0)), although mass effects of dispersal in time may contribute to noise in the environmental signal in a metacommunity analysis (Leibold et al. [2004\)](#page-13-0) or to retarded adaptation under a scenario of directional selection (Hairston and De Stasio Jr [1988](#page-12-0)). Overall, the presence of massive dormant egg banks such as reported here and in other studies (García-Roger et al. [2006;](#page-12-0) Hairston et al. [1995;](#page-12-0) Thompson and Grime [1979](#page-13-0)) may strongly contribute to a dominance of local compared to regional dynamics in determining metapopulation and metacommunity structure, because of the combined effects of large numbers and increased genetic and species diversity.

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