

Empirically simulated spatial sorting points at fast epigenetic changes in dispersal behaviour

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1 **Empirically simulated spatial sorting points at fast epigenetic**
2 **changes in dispersal behaviour**

3

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16

1 **Abstract**

2

3 During range expansion, the most dispersive individuals make up the range front, and
4 assortative mating between these dispersive phenotypes leads to increased dispersiveness (*i.e.*
5 spatial sorting). The precise inheritance of dispersal, however, is to date largely unknown in
6 many organisms, thereby hampering any progress in evaluating the adaptive potential of
7 species during range expansion.

8 Using the spider mite *Tetranychus urticae*, we therefore empirically simulated spatial
9 sorting by means of artificial selection on a unique pre-dispersal behaviour, tightly related to
10 emigration. To separate directionality of the response from potential drift, we mimicked a
11 recurrent low number of founders in replicated selection regimes. Afterwards, we inferred the
12 mode of inheritance of the pre-dispersal behaviour by performing reciprocal crosses between
13 selected (*i.e.* dispersive) and non-selected (*i.e.* non-dispersive) mites and by screening for
14 endosymbionts known to be associated with changes in dispersal behaviour.

15 Despite the recurrent low number of founders, the aerial dispersal behaviour
16 responded strongly to the imposed selection pressure. The behaviour furthermore showed a
17 maternal inheritance, though independent of any known dispersal-related endosymbionts.
18 Though cytoplasmic inheritance cannot fully be excluded, we attribute the observed strong
19 and rapid, maternally influenced response in dispersal to transgenerational epigenetic effects.
20 Consequently, we can expect fast evolutionary dynamics during range expansion in the
21 species.

22

1 Introduction

2

3 Dispersal is the main mechanism maintaining gene flow among populations, and of
4 primary importance for population regulation (Krebs et al. 1969), metapopulation dynamics
5 (Hanski 1999; Hanski and Gaggiotti 2004; Levins 1969) and the spatial distribution of
6 populations (Holt 1985; Kot et al. 1996). Crucially, dispersal is the key to the dynamics of
7 genetic diversity in space and time (Gaggiotti and Couvet 2004), allowing species to respond
8 to environmental challenges. Global warming, for example, is currently creating exciting
9 evolutionary experiments where species expanding their ranges experience a strong selection
10 for dispersal (Perkins et al. 2013; Shine et al. 2011). Indeed, during range expansion, the most
11 dispersive individuals make up the range front, and assortative mating between these
12 individuals (*i.e.* ‘the Olympic Village effect’) results in increased dispersal abilities (Phillips
13 et al. 2010). This process is referred to as spatial sorting (or spatial selection) (Shine et al.
14 2011).

15

16 Range expansion is typically achieved by only those few individuals that reach and
17 successfully settle in a previously unoccupied environment (*i.e.* not occupied by
18 conspecifics). These individuals thus represent only a limited subset of the original genetic
19 variation (*i.e.* founder effect) (Mayr 1963). Moreover, because of the low number of
20 individuals, the new population will be highly susceptible to inbreeding and genetic drift,
21 resulting in a clear genetic signature of range expansion, even in species with high dispersal
22 capacity (Pierce et al. 2014). Repeated founder effects may furthermore result in gene surfing:
23 a process where neutral alleles hitchhike with pioneer genotypes, thereby attaining high
24 frequencies over large areas while not being under direct selection (Klopfstein et al. 2006).
25 Life history-traits in founder populations will thus be shaped through the combined effect of

1 natural selection (*i.e.* adaptation to the new local environment), spatial sorting (*i.e.* selection
2 for dispersal and correlated traits) and genetic drift.

3

4 While selection on dispersal is expected to be strong during spatial sorting, the rate of
5 evolutionary change will largely depend on the genetic architecture of the trait, the trait
6 heritability and, equally, on the degree of adaptive plasticity potentially imposed by ‘non-
7 classical’ inheritance (see Moran and Alexander 2014 for a complete overview of limits to
8 evolutionary responses to environmental change). The latter mechanism might be especially
9 relevant in cases where standing genetic variation is low, like in case of subsequent founder
10 effects during range expansion. Indeed, there is mounting evidence that parental effects and
11 cytoplasmic inheritance might be alternative pathways for species to respond fast and
12 adaptively to environmental perturbations (Bonduriansky et al. 2012; Bossdorf et al. 2008;
13 Ledon-Rettig et al. 2013). Parental effects on dispersal, described for both plants (reviewed in
14 Roach and Wulff 1987) and animals (*e.g.* Bitume et al. 2011; Diss et al. 1996; Li and
15 Margolies 1994; Sinervo et al. 2006), as well as cytoplasmic inheritance of, for example,
16 mitochondrial DNA (see Van Leeuwen et al. 2008) or endosymbiotic bacteria (see Goodacre
17 et al. 2009), are therefore expected to induce fast phenotypic changes that might affect future
18 ecological dynamics in a profound manner (but see Donohue 1999).

19

20 The two-spotted spider mite (*Tetranychus urticae* Koch - Acari, Tetranychidae) is a
21 very useful model species for testing these dynamics, as previous research demonstrates a
22 significant and high broad-sense genetic component underlying ambulatory ($h^2=0.52$ in
23 Bitume et al. 2011) and aerial ($h^2=0.22$ in Li and Margolies 1993) dispersal behaviour. These
24 behaviours can easily be quantified under standardised laboratory conditions. The species is
25 considered as one of the most damaging pest species and is expanding its natural range

1 towards higher latitudes (see Carbonnelle et al. 2007). This range expansion is assumed to be
2 due to long distance dispersal events through passive aerial dispersal, where new populations
3 are founded by a limited number of individuals. In contrast, local dynamics are merely
4 mediated by short distance ambulatory dispersal in response to elevated densities and kin
5 competition (Bitume et al. 2013).

6

7 We simulated the spatial sorting process, as is expected to occur during range
8 expansion, by means of artificial selection on a unique pre-dispersal behaviour, tightly related
9 to emigration: the aerial take-off posture, which has a very high probability of resulting in
10 mites becoming airborne (see Smitley and Kennedy 1985). We used *T. urticae* as a model
11 species and mimicked a recurrent low number of founders in replicated selection regimes, to
12 separate directionality of the response from potential drift. Several other studies already
13 performed artificial selection on dispersal in *T. urticae*, but these focused on ambulatory
14 dispersal (Bitume et al. 2011; Tien et al. 2011; Yano and Takafuji 2002) or used a very high
15 number of individuals (Li and Margolies 1994) and can therefore not be interpreted in the
16 context of spatial sorting during range expansion.

17 Our aims were twofold. Firstly, we assessed the evolutionary response of the aerial
18 take-off behaviour to a strong selection pressure, given only minimal genetic variation.
19 Secondly, we assessed the mode of inheritance of the aerial dispersal trait by crossing
20 dispersers and non-dispersers and by screening for five endosymbiotic bacteria known to be
21 associated with changes in dispersal behaviour (Goodacre et al. 2009). We hypothesised that a
22 fast evolutionary response in aerial dispersal behaviour in *T. urticae* could be the result of
23 ‘non-classical’ inheritance, as indicated by the slight maternal influence in the study of Li and
24 Margolies (1994).

25

1 **Materials and methods**

2

3 **Study species**

4 The two-spotted spider mites (*Tetranychus urticae* Koch - Acari, Tetranychidae) used
5 in this study originated from a laboratory strain (“LS-VL”) that was originally collected in
6 October 2000 from roses in a garden near Ghent, Belgium. Since then, this strain was
7 maintained on potted beans (*Phaseolus vulgaris* L. cv. ‘Prelude’) in a climate-controlled room
8 at 26±0.5°C, 60%RH and 16/8 h (L/D) photoperiod (Van Leeuwen et al. 2006).

9

10 **Artificial selection**

11 We largely based our experimental procedure on Li and Margolies (1993, 1994). The
12 propensity to disperse by air was assessed by counting the percentage of female mites
13 showing the aerial take-off posture (*i.e.* upraised first pair of legs and cephalothorax, to
14 increase drag). Dispersal latency was furthermore assessed by counting the number of minutes
15 between the start of the one hour observation and the moment the focal female showed the
16 aerial take-off posture. Only one-day-old, freshly mated females were considered, since they
17 are the main dispersing stage (see Li and Margolies 1993, 1994 for more details).

18 To ensure having a sufficient number of dispersive individuals to start the selection
19 experiment (dispersal propensity was relatively low in the base population), an initial
20 selection procedure was first run. For this initial generation of selection, 20 females were put
21 on each of three separate arenas -black painted squares of hard plastic on wet cotton. They
22 were then subjected to a light and wind regime known to elicit dispersal initiation behaviour
23 (*i.e.* the aerial take-off posture) in *T. urticae* (see Li and Margolies 1993). During one hour,
24 the females were observed and screened for the aerial take-off posture in a climate room at
25 20°C. From one of the three arenas, the first five females showing the aerial take-off posture

1 were collected and put separately on a fresh bean leaf. These would make up the high
2 dispersal regime (HD). On a second arena, all females showing the posture were killed, and
3 afterwards, five females were randomly chosen from the remaining ones. These females were
4 then put on a fresh bean leaf and made up the low dispersal regime (LD). For the control
5 regime (C), five females were randomly chosen out of all original 20 females from the third
6 arena and put on yet another fresh bean leaf. All of these females (HD, LD, C) were
7 subsequently allowed to lay eggs in a breeding room at 20°C with a light-regime of 16:8 LD.
8 Their female offspring, once adult and freshly mated, were then used for the next generation
9 of selection.

10 From this point onwards, the selection experiment was performed, using three
11 replicates per selection regime (thus nine lines, hence nine arenas to be observed: HD1, HD2,
12 HD3, LD1, LD2, LD3, C1, C2, C3). Using these replicates allowed us to differentiate
13 between random genetic drift and evolutionary changes caused by the selective pressure. The
14 selection criteria remained the same as for the initial selection procedure (see above). In cases
15 where less than five females showed the aerial take-off posture in a HD line, we just
16 continued with this lower number (*i.e.* we did not increase the number to five by adding
17 randomly chosen females as this would weaken the selection pressure).

18 In order to create sufficient divergence between the selection regimes, the selection
19 procedure was repeated for 13 generations (Li and Margolies 1994). Dispersal latency was
20 only assessed for the HD regime, as too few females showed the dispersal behaviour in the C
21 and LD regime.

22

23 **Crosses between selection regimes**

24 The adult HD and LD females resulting from the 13th generation of selection were put
25 on fresh bean leaves and allowed to lay eggs. The freshly adult male and female offspring of

1 the six lines (2 selection regimes x 3 replicates) were then used for the crosses. This
2 experiment consisted of females and males from HD and LD selection regimes mating
3 amongst each other according to a 2x2-factorial design. (To ensure a sufficient number of
4 female offspring to later conduct an aerial dispersal test, five females were each time put
5 together with five males and allowed to mate for four days.) The crosses (HD_{fem} x HD_{male},
6 HD_{fem} x LD_{male}, LD_{fem} x HD_{male}, LD_{fem} x LD_{male}) were made in each of the three original
7 replicates of the HD and LD selection regimes, resulting in a total of twelve (3x4) crosses.
8 Finally, the one-day-old female offspring from these crosses were all tested for their aerial
9 dispersal propensity. This was done in a similar setup as described earlier. Twelve (3x4)
10 arenas were observed for one hour, during which all females showing the aerial take-off
11 posture were immediately removed from their arena. Such a removal was each time recorded.
12 As such, the proportion of females showing the posture could afterwards be calculated for
13 each arena by dividing the number of removals by the original number of females on that
14 arena.

15

16 **Endosymbionts**

17 Four hundred adult females were pooled from the parental spider mite strain prior to
18 selection and from the HD, LD and C selection regimes after the 13th generation of selection.
19 Genomic DNA was collected following a phenol-chloroform extraction method (Van
20 Leeuwen et al. 2008). Additionally, 20 DNA samples, each originating from a single adult
21 female mite, were extracted from the parental strain and from the selection regimes, following
22 the procedure described in Van Leeuwen et al. (2008). In all DNA samples, the presence of
23 five endosymbionts known to be associated with changes in dispersal behaviour (*Cardinium*,
24 *Caulobacter*, *Rickettsia*, *Spiroplasma* and *Wolbachia* -see Goodacre et al. 2009) was tested by

1 PCR, using the primers listed in Table 1. Amplified PCR products were sequenced by LGC
2 Genomics (Berlin, Germany) after purification with Cycle-Pure Kit (EZNA™).

3

4 **Statistics**

5 All analyses were performed with SAS 9.4 (SAS Institute Inc. 2013), using the
6 glimmix procedure for generalised linear mixed models (GLMM). We analysed aerial
7 dispersal propensity with selection regime (HD, LD, C), generation and the interaction
8 between selection regime and generation as the independent variables. For the analysis of
9 aerial dispersal latency, only generation was used as an independent variable, as only the HD
10 regime was involved. The analysis of the regime crosses was performed with the dispersal
11 type of the mother and the father (*i.e.* originating from the HD selection regime opposed to
12 the LD selection regime) and their interaction term as the independent variables. According to
13 the dependent variable, a binomial (dispersal propensity) or Poisson (dispersal latency) error
14 structure was modelled with the proper link function (logit/log). Replicate was always
15 modeled as a random effect. This was done to control for dependency among the replicates of
16 each selection regime. By modeling residual variation as an additional random factor, we
17 corrected for potential overdispersion (Verbeke and Molenberghs 2000). The denominator
18 degrees of freedom for the tests of fixed effects were computed according to a general
19 Satterthwaite approximation. All non-significant contributions ($p > 0.05$) were omitted by a
20 backwards selection procedure. Finally, *post-hoc* Tukey tests were used to obtain the pair-
21 wise differences among treatments (only for the regime crosses).

1 **Results**

2

3 **Artificial selection**

4 Aerial dispersal propensity responded well to the artificial selection: selection regime
5 significantly interacted with generation ($F_{2,1673}=4.72$; $p=0.0091$) (Fig. 1). The slopes of the
6 HD and LD regime differed significantly from zero: HD: 0.0696 ± 0.0210 ($t_{1835}=3.32$;
7 $p=0.0009$), LD: -0.1024 ± 0.0370 ($t_{1835}=-2.77$; $p=0.0056$), C: -0.0509 ± 0.0307 ($t_{1835}=-1.66$;
8 $p=0.0977$). These slopes imply that the odds of successful dispersal multiply by a factor of
9 1.07, 0.90 and 0.95 per generation, in the HD, LD and C regime, respectively.

10 Furthermore, latency (which was only measured for the HD regime) significantly
11 decreased with generation ($F_{1,215}=112.81$; $p<0.0001$) (Fig. 2).

12

13 **Crosses between selection regimes**

14 The proportion of female offspring displaying the aerial take-off posture differed
15 significantly among some of the crosses (Fig.3). Only the dispersal type of the mother had a
16 significant effect ($F_{1,8}=19.40$; $p=0.0023$): the proportion of females showing the aerial take-
17 off posture was higher for HD mothers (0.4054 ± 0.0496 SE) than for LD mothers ($0.0636 \pm$
18 0.0287 SE) ($t_8=4.40$; $p=0.0023$). The dispersal type of the father did not affect the dispersal
19 propensity of the offspring ($F_{1,8}=2.44$; $p=0.1566$). There was also no significant interaction
20 between the dispersal type of the mother and the father ($F_{1,8}=0.37$; $p=0.5587$).

21

22 **Endosymbionts**

23 *Wolbachia* and *Rickettsia* were detected by species-specific primers in the pooled
24 sample of the non-selected parental strain, and infection showed a prevalence of 0.15 and 0.9
25 respectively, as determined by single-mite PCR. BLASTn-searches, using the sequenced PCR

1 products as queries against the NCBI database, confirmed the correct identification of
2 endosymbionts. No endosymbionts were found in the replicated HD, C and LD selection
3 regimes in both the pooled samples of 400 mites and the single mite DNA samples.

1 Discussion

2

3 We empirically simulated the process of spatial sorting by artificially selecting on
4 aerial dispersal behaviour (more specifically, the aerial take-off posture), using a recurrent
5 low number of founders, and demonstrate a fast response to selection and a maternal
6 inheritance of this dispersal behaviour in *T. urticae*.

7

8 As in comparable studies (Li and Margolies 1993, 1994; Yano and Takafuji 2002),
9 ‘highly dispersive mites’ (HD selection regime) were selected for showing a specific
10 behaviour, while ‘low dispersive mites’ (LD selection regime) were selected for not showing
11 this behaviour. Possibly, the mites from the LD regime might therefore not have been selected
12 for a low propensity to disperse, but for a general low viability, hence incapability of showing
13 the aerial take-off posture (see Tien et al. 2011). However, even with our very strong selection
14 criteria (the number of founders was each time restricted to only five mites), the aerial
15 dispersal behaviour never really disappeared in the LD regime. This suggests that the
16 capability for the aerial take-off posture remained present throughout the selection procedure.
17 All mites were moreover maintained at similar, low densities, resulting in overall fit females
18 in all selection treatments; *i.e.* the selection regimes did not differ in daily fecundity or
19 longevity (Pétilion et al., unpublished data).

20

21 Despite the recurrent low number of founders used in our artificial selection
22 experiment, the aerial dispersal behaviour responded equally strong to the imposed selection
23 pressure as in the study of Li and Margolies (1994). Aerial dispersal propensity gradually
24 increased in the HD regime while it showed a steady decrease in the LD regime. The strong
25 response across all replicates, in a minimal number of generations, is unlikely to have resulted

1 from new mutations or from standing genetic variation. Standing genetic variation was most
2 probably low due to the low number of initial founders that were used to start the replicated
3 selection regimes. Instead, in accordance with a growing amount of studies in lizards (*e.g.*
4 Massot et al. 2002; Meylan et al. 2002; Sinervo et al. 2006), birds (*e.g.* Tschirren et al. 2007)
5 and arthropods (*e.g.* Bitume et al. 2011, 2014; Bonte et al. 2007; Li and Margolies 1994;
6 Mestre and Bonte 2012), we attribute the strong response in our experiment to
7 intergenerational plasticity of dispersal through a maternal influence on offspring phenotype.
8 Indeed, the crosses between our selection regimes indicated a strong maternal inheritance.
9 Several mechanisms could be responsible for this, including cytoplasmic inheritance, classical
10 Mendelian inheritance through sex specific chromosomes and maternal effects.

11

12 Cytoplasmic inheritance includes, amongst others, the vertical transmission of
13 bacterial endosymbionts. Goodacre et al. (2009) demonstrated an effect of such cytoplasmic
14 endosymbionts on the tendency for long-distance movement in the spider *Erigone atra*. We
15 thus investigated whether the maternal inheritance in our study could have been mediated by
16 such maternally transmitted endosymbiotic bacteria. The absence of all endosymbionts known
17 to be associated with changes in dispersal behaviour (*Cardinium*, *Caulobacter*, *Rickettsia*,
18 *Spiroplasma* and *Wolbachia*) in our selection regimes, however, makes this mechanism
19 unlikely. Surprisingly, where *Rickettsia* was absent in the selection regimes, it showed a high
20 prevalence in the non-selected parental stock population. This apparent discrepancy may be
21 ascribed to the density of *Rickettsia* bacteria within individual mites of the parental
22 population. Although we measured the prevalence of endosymbionts in the mite populations,
23 we did not evaluate infection density (*i.e.* the number of *Rickettsia* within each single
24 individual). It is well established that this infection density affects many aspects of the
25 endosymbiont-host interaction, including the vertical transmission of these endosymbionts

1 (Noda et al. 2001; Perrot-Minnot and Werren 1999). Low infection densities can cause
2 imperfect vertical transmission, and over multiple generations, this can lead to an eventual
3 overall loss of the endosymbiont. Our results suggest that the *Rickettsia* bacteria suffered from
4 imperfect vertical transmission, potentially caused by low bacterial densities within single
5 mites. *Rickettsia* can therefore not have been responsible for the maternal inheritance of aerial
6 dispersal behaviour in our experiment. Nonetheless, there could always be some other
7 endosymbiotic bacteria contributing to this observation.

8 Another form of cytoplasmic inheritance is the vertical transmission of mitochondrial
9 DNA. When individuals possess more than one mtDNA haplotype (*i.e.* mitochondrial
10 heteroplasmy), genotype frequencies can shift greatly in a single transmission from mother to
11 offspring, as for instance found for loci related to pesticide resistance (Van Leeuwen et al.
12 2008). Though mitochondrial heteroplasmy is generally rare (Jenuth et al. 1996), we should
13 thus keep in mind that loci on the mitochondrial genome may also contribute to strong and
14 rapid phenotypic effects.

15
16 Maternal inheritance of a trait can in some cases also be the consequence of classical
17 Mendelian inheritance of the mother's nuclear genome. If dispersal-related traits are
18 controlled by loci on the female sex-chromosome (as is, for example, present in butterflies,
19 where females are the heterogametic sex), Mendelian inheritance could be enough to explain
20 mother to daughter similarities. This, however, does not apply for *T. urticae*. In tetranychid
21 species, sexual reproduction occurs through arrhenotokous parthenogenesis, in which
22 unfertilised eggs develop into males and fertilised eggs into females. *T. urticae* females do not
23 possess a 'female-specific chromosome' that is passed on from mother to daughter only.
24 Mendelian inheritance of nuclear genomes can therefore not explain the result of our cross-
25 mating experiment.

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A third potential mechanism causing the observed maternal inheritance could be maternal effects. Maternal effects are defined as any maternal influence on offspring phenotype that cannot be attributed solely to offspring genotype, the direct action of the non-maternal components of the offspring environment, or to their combination (Lacey 1998). According to Lacey (1998), there are three types: (i) genetically based maternal effects (*e.g.* genomic imprinting), (ii) indirect genetic maternal effects (*e.g.* genetically determined maternal care) and (iii) environmentally induced maternal effects (*i.e.* all effects induced by the maternal environment, independent of maternal genotype). Since *T. urticae* does not provide any form of maternal care and since all individuals were raised in identical circumstances for several generations, we can rule out indirect genetic and environmentally induced maternal effects. Though we did not specifically assess any genetically based maternal effect, we consider it most likely that such transgenerational epigenetic effects caused the maternal inheritance of aerial dispersal behaviour observed in our experiment.

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In the context of the current global change, the potential of species to colonise new habitats may be essential for their persistence. Since shifts at the leading edge of expanding ranges are often achieved by a small number of long-distance dispersing individuals, newly established populations may suffer from founder bottlenecks (reviewed in Hill et al. 2011). Not only does this lead to a reduced genetic diversity, with all its possible consequences, it also implies that the individuals at expanding ranges may differ substantially from the ones that stay behind. Individuals at the shifting edge may carry a suite of typical characteristics, shaped through selection on dispersal and other potentially correlated traits that may eventually constrain genetic adaptation. In the presence of such genetic constraints, transgenerational epigenetic inheritance may provide alternative pathways for species to

1 respond fast and adaptively to environmental perturbations like global change (Bossdorf et al.
2 2008; Ledon-Rettig et al. 2013; Tuomainen and Candolin 2011). The ability to epigenetically
3 control dispersal can be considered as a form of phenotypic plasticity that evolved in response
4 to the ephemeral nature of the species' habitat (*T. urticae* is characterised by a rapid
5 development and a high fecundity, leading to exponential growth and systematic depletion of
6 its host plant). Indeed, spatiotemporal variation in habitat quality is known to select for
7 plasticity as an adaptive trait on its own (Via and Lande 1985).

8

9 In conclusion, despite the recurrent low number of founders used to simulate the
10 process of spatial sorting during range expansion, the aerial dispersal behaviour in *T. urticae*
11 responded strongly to the imposed artificial selection pressure. We attribute this strong and
12 rapid response to transgenerational epigenetic mechanisms, though we should acknowledge
13 that cytoplasmic inheritance cannot fully be excluded as a potential (co-)affecting factor. As a
14 consequence of this epigenetic inheritance, we can expect fast, non-classical evolutionary
15 dynamics during range expansion in the species.

16

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2

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10

1 **Figure legends**

2

3 **Figure 1: Evolution of aerial dispersal propensity.** For each generation, the mean aerial
4 dispersal propensity in the different selection regimes is represented by a symbol. Standard
5 errors are represented by bars. Aerial dispersal propensity is averaged over replicates and
6 females within replicates. Closed circles: high dispersal selection regime (HD), open circles:
7 control regime (C), rectangles: low dispersal selection regime (LD). (Note: Due to
8 circumstances, the selection procedure did not last a full hour in generation 8.)

9

10 **Figure 2: Evolution of aerial dispersal latency.** For each generation, the mean aerial
11 dispersal latency in the high dispersal regime (HD) is represented by a dot. Standard errors are
12 represented by bars. Aerial dispersal latency is averaged over replicates and females within a
13 replicate. The aerial dispersal latency of a female mite is the number of minutes between the
14 start of the one hour observation of the aerial dispersal propensity of this female and the
15 moment she shows the aerial take-off posture. (Note: Due to circumstances, the selection
16 procedure did not last a full hour in generation 8.)

17

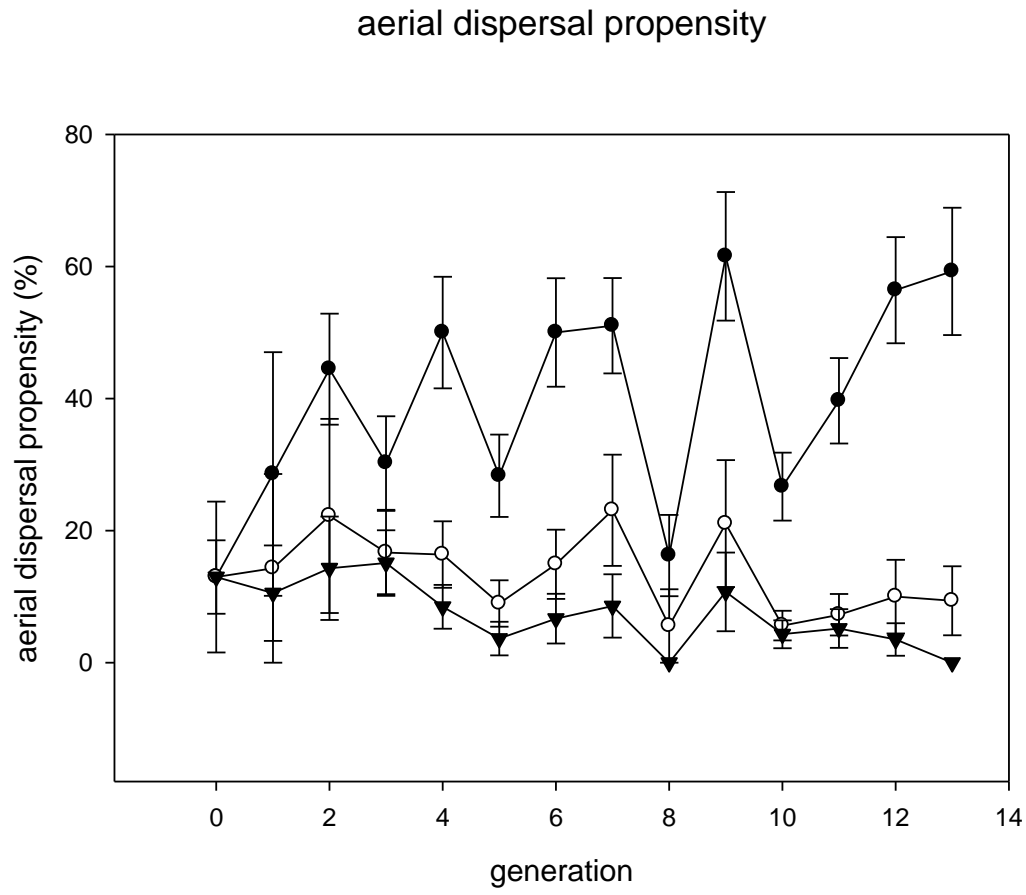
18 **Figure 3: Aerial dispersal propensity in the offspring of the different regime crosses.** The
19 mean aerial dispersal propensity of the female offspring from the four different regime crosses
20 is represented by columns. Standard errors are represented by bars. Different letters above the
21 error bars indicate significantly different outcomes according to the used statistical test. Aerial
22 dispersal propensity is averaged over the replicates within each of the regime crosses and over
23 female offspring within a replicate. HD_{m/f}: high dispersal male/female, LD_{m/f}: low dispersal
24 male/female.

25

1 **Figures**

2

3 **Figure 1**

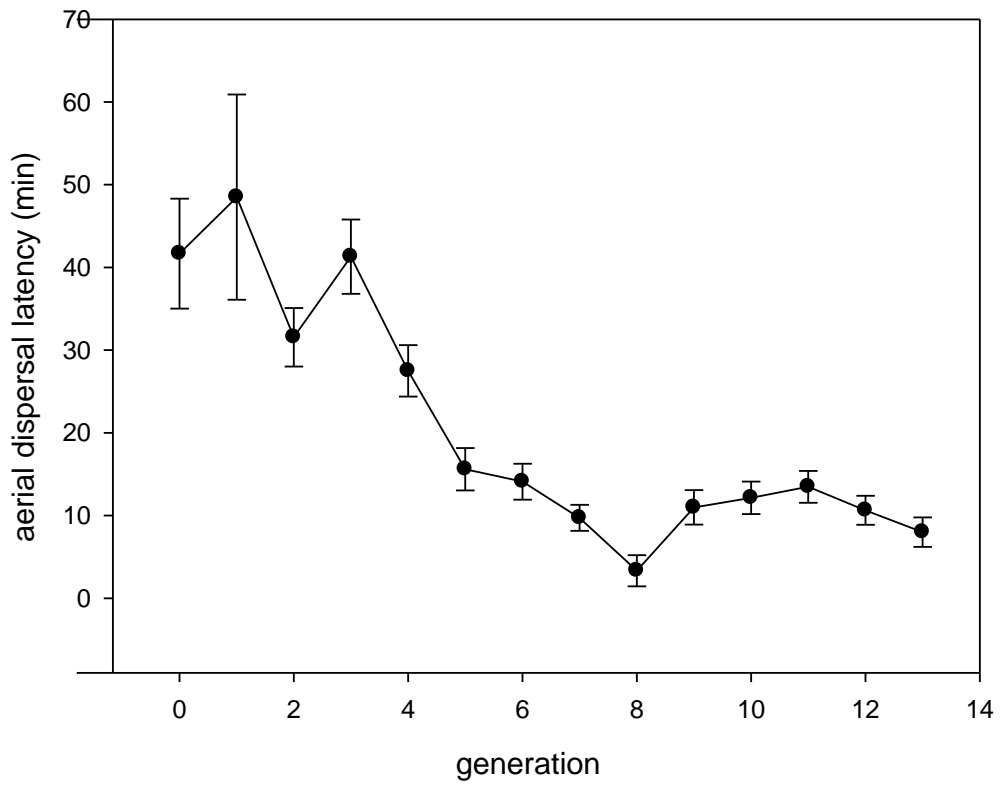


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5

1 **Figure 2**

aerial dispersal latency

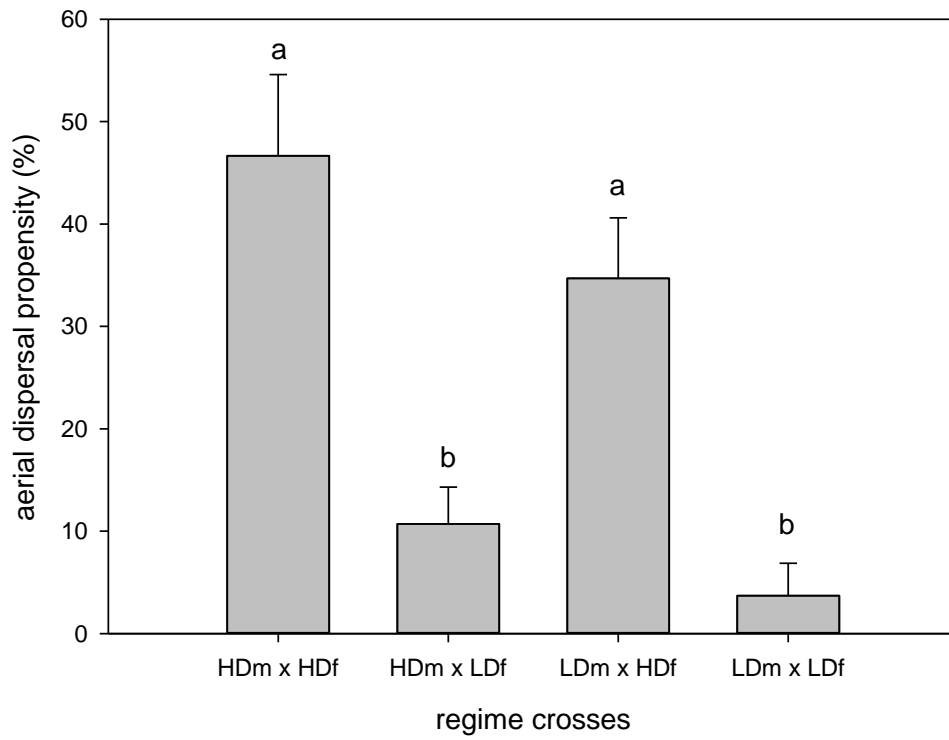


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3

1 **Figure 3**

inheritance of aerial dispersal



2

3

1 **Tables**

2

3 **Table 1**

4 Primers used in this study for the detection of endosymbionts in *T. urticae*. For each primer pair, sequences are given for the forward (F) and the
5 reverse (R) primer, with the corresponding reference. The presence of *Cardinium* was investigated using three different primer pairs.

6 Bacteria	Primer sequence [5' – 3']	Reference
7 <i>Cardinium</i>	F1: GCGGTGTAAAATGACGTG	(Weeks et al. 2003)
	R1: ACCTMTTCTTAACTCAAGCCT	
	F2: GGAACCTTACCTGGGCTAGAATGTATT	(Gotoh et al. 2007)
	R2: GCCACTGTCTTCAAGCTC TACCAA	
	F3: AGAAGAGCCTTGACCCGCAC	own design using Primer-BLAST
	R3: TCTGTGCCCATGGTTACGCC	
15 <i>Caulobacter</i>	F: GAGACCTTCGGGTCTAGTG	(Hoy and Jeyaprakash 2005)

1 R: CTTGCGAGTTAGCGCAACG

2 *Wolbachia* F: TGGTCCAATAAGTGATGAAGAACTAGCTA (Jeyaprakash and Hoy 2000)

3 R: AAAAATTAAACGCTACTCCAGCTCTGCAC

4 *Rickettsia* F: GGACTAATTGGGGTTTGCTC (Hoy and Jeyaprakash 2005)

5 R: GCTGCCTCTTGCGTTAGCT

6 *Spiroplasma* F: GAGAGTTTGATCCTGGCTCAG (Weisburg et al. 1991)

7 R: TAGCCGTGGCTTTCTGGTAA (Fukatsu and Nikoh 2000)

8

9