Intergenerational paternal effect of adult density in *Drosophila melanogaster*

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**Abstract**

1. Notwithstanding recent evidences, paternal environment is thought to be a potential but unlikely source of fitness variation that can affect trait evolution. Here we studied intergenerational effects of males’ exposure to varying adult density in *Drosophila melanogaster* laboratory populations.

2. We held sires at normal (N), medium (M) and high (H) adult densities for 2 days before allowing them to mate with virgin females. This treatment did not introduce selection through differential mortality. Further, we randomly paired males and females and allowed a single round of mating between the sires and the dams. We then collected eggs from the dams and measured the egg size. Finally, we investigated the effect of the paternal treatment on juvenile and adult (male) fitness components.

3. We found a significant treatment effect on juvenile competitive ability where the progeny sired by the H-males had higher competitive ability. Since we did not find the treatment to affect egg size, this effect is unlikely to be mediated through variation in female provisioning.

4. Male fitness components were also found to have a significant treatment effect: M-sons had lower dry weight at eclosion, higher mating latency, and lower competitive mating success.

5. While being the first study to show both adaptive and non-adaptive effect of the paternal density in *Drosophila*, our results highlight the importance of considering paternal environment as important source of fitness variation.

**KEYWORDS**
crowding adaptation, juvenile competitive fitness, male reproductive success, mating latency, sire effect
Parental environment has the potential to influence offspring traits and fitness through intergenerational effects (and more stable transgenerational effects, see Dias & Ressler, 2014 for the distinction between trans and intergenerational effects). While it can potentially pass on deleterious effects of different components of the environment to the following generation (Yehuda et al., 2000), intergenerational effect can also be adaptive, especially under fluctuating environment (Bonduriansky & Day, 2009). Among the myriad components of an organism's ecology, few factors are as variable as density and nutritional availability. Both have been recently found to have intergenerational effects, especially through the maternal route (i.e., maternal effect), in a wide variety of organisms (Mousseau & Fox, 1998a, 1998b). There is a growing body of evidence showing the importance of the intergenerational effect of paternal nutrition, social experience and density on fitness related traits of the offspring (Adler & Bonduriansky, 2013; Crean, Dwyer, & Marshall, 2013; Dasgupta, Halder, & Nandy, 2016; Friberg, Stewart, & Rice, 2012). However, the prevalence and adaptive significance of such paternal effect is yet to be ascertained.

There are many reports of environment dependent maternal effect mediated through variation in maternal provisioning in egg/offspring (Mousseau & Fox, 1998a, 1998b; Rossiter, 1996). For example, females living under high density may suffer from adverse effects of crowding (such as, malnutrition) and may therefore struggle to allocate resources in maternal provisioning—either in the form of stored resources in egg or lactation, which in turn may lead to poor quality progeny (Christian & Lemunyan, 1958). Alternatively, females raised in high density may strategically produce fewer eggs/progeny while investing more resources (e.g., yolk) in each of them—thereby giving the progeny a better start for the impending challenges of crowding (Holbrook & Schal, 2004; Mitchell & Read, 2005; Prasad, Shakarad, Rajamani, & Joshi, 2003; Vijendravarma, Narasimha, & Kawecki, 2010). Generally, under fluctuating environmental conditions, such parental ability to optimize offspring phenotype has been conjectured to be adaptive (Bonduriansky & Day, 2009; Kuijper & Hoyle, 2015). For example, Guppy (Poecilia reticulata) females were found to produce larger offspring (a) under food limitation (Reznick & Reznick, 1993) and (b) when they experienced high level of competition—priming the offspring for better competitive ability (Bashey, 2006). The larger eggs produced by Drosophilid melanogaster females that grew in nutritionally impoverished food, survive (egg-to-adult survivorship) better in impoverished food and give rise to smaller adults (Vijendravarma et al., 2010). In contrast, Valtonen, Kangassal, Pölkki, and Rantala (2012) found that D. melanogaster females grown on impoverished food produced larger offspring (adult) compared to those grown on nutritionally rich food. Note that many of the maternal effects discussed above are mediated through variation in resource provisioning by mothers.

Not surprisingly, most of the reports of environment dependent paternal effect (intergenerational and transgenerational) are from animals with paternal provisioning through nuptial gift transfer to the females (Dussourd et al., 1988; Gwynne, 1988; Smely & Eisner, 1996; Vahed, 1998; Zeh & Smith, 1995). However, it is only recently that studies have started addressing the presence of similar paternal effects in species without paternal provisioning. For example, in one of the first such explicit studies, female Neriid flies (Teleostylus angusticollis) raised on richer diet were found to produce larger eggs and offspring that developed faster, while males raised on richer diet sired larger offspring with better survival rate, especially under resource scarcity (Adler & Bonduriansky, 2013; Bonduriansky & Head, 2007). Further, in a solitary Ascidian, Styelapecta, males were found to produce offspring with phenotype corresponding to the population density experienced by the father (Crean et al., 2013). In fruit flies (D. melanogaster) Valtonen et al. (2012) reported that fathers fed on poor quality diet sire larger sons. Paternal experience of the intensity of competition (assessed by the number of co-inhabitant rival males) was found to affect reproductive behavior (duration of copulation) of male offspring in D. melanogaster (Dasgupta et al., 2016). In Desert locusts (Schistocerca gregaria), Islam, Roessingh, Simpson, and McCaffery (1994) showed paternal crowding to have a significant impact on hatching coloration and nymph behavioral traits. Paternal experience of ambient temperature was also found to affect offspring fecundity in D. melanogaster (Huey, Wakefield, Crill, & Gilchrist, 1995). Low temperature was found to affect offspring phenotype in two other species of Drosophila—D. simulans (Watson & Hoffmann, 1995) and D. serrata (Magiafoglou & Hoffmann, 2003). Thus, there is ample evidence showing environment dependent paternal effect. In addition to affecting viability, such paternal effect has been shown to affect progeny reproductive performance and hence is likely to be a key player in sexual selection (for example, see Bonduriansky & Head, 2007). However, such data are far from being plenty.

Here, we investigated the effect of paternal experience of population density on progeny fitness components, including male mating behavior in D. melanogaster laboratory adapted populations. As discussed previously, paternal effect has already been reported in these (Dasgupta et al., 2016) and other populations of D. melanogaster, establishing them as a relevant system to investigate the paternal effect and its consequences on Darwinian fitness (William et al., 2006). Further, laboratory adapted populations of D. melanogaster have been used to investigate the fitness consequence of a plethora of environmental parameters, including population density. Fruit flies naturally grow in ephemeral resource patches, such as rotting fruits and vegetables. Crowding in transiently available rich patches is expected to be a key component of their natural ecology. Density of adults in a resource patch not only determines the extent to which individuals must compete for food and limited space (e.g., oviposition substrate) but also for suitable mates. Increase in density also leads to an increase in the probability of disease transmission (Barnes & Siva, 2000). In essence, density often determines the nature and intensity of selection acting on a population and has been studied within the broader premises of density dependent selection (MacArthur...
& Wilson, 1967; Mueller, 1997; Prasad & Joshi, 2003). Much of the existing literature investigated adaptation to increased (but stable) juvenile or adult density, using experimental evolution on laboratory populations of *D. melanogaster* (Mueller, Guo, & Ayala, 1991; Mueller & Sweet, 1986; Nagarajan, Natarajan, Jayaram, & Joshi, 2016; Sarangi, Nagarajan, Dey, Bose, & Joshi, 2016; Shenoi et al., 2016; Shenoi & Prasad, 2016). However, little is known about adaptation to fluctuating density. Intergenerational and transgenerational effects, if used by the parents to optimize offspring phenotype, can be of adaptive value if density fluctuation across generation is, at least to some extent, predictable. Interestingly, these experimental evolution studies reported “rapid” adaptation to “crowding”. Though evidences unequivocally showed the genetic changes associated with such adaptation, non-genetic parental effects (trans and intergenerational) may, in addition, account for the “rapid” adaptation (Bonduriansky & Day, 2009). However, this idea has not been tested—an existing lacuna in the literature, which we intend to fill to some extent.

To investigate the paternally transmitted intergenerational effect of varying density, we subjected males to three adult density treatments and then allowed them to sire progeny by mating the treated males to untreated dams. We then assessed the effect of the paternal adult density (hereafter, referred to as paternal density) treatment on progeny fitness components in juvenile (juvenile competitive fitness) and adult stages (males: mating ability, mating latency, copulation duration, courtship frequency, competitive mating success). We found the paternal density treatment to have significant intergenerational effect on both juvenile and adult fitness components.

### 2 | MATERIALS AND METHODS

All the experiments were done using a set of laboratory adapted populations of *D. melanogaster*—BL. Full laboratory history of these populations can be found in (Nandy, Dasgupta, Halder, & Verma, 2016). Briefly, these are a set of five replicate populations (BL<sub>1–5</sub>) maintained on standard Banana–Jaggery–Yeast food, under 14-day discrete generation cycle at 25°C ambient temperature, 60%–80% relative humidity, with population size ~2,800. Larval density is maintained at ~70 per 6–8 ml food per vial (25 mm × 90 mm, diameter × height). Adult density is ~70 per vial for the first couple of days of their adult life and thereafter ~2,800 individuals in a ~6.4 L cage (19 cm × 14 cm × 24 cm). We also used a genetically marked population, BL<sub>st</sub> which was derived from BL<sub>1</sub> by introducing an autosomal recessive marker—scarlet eye, st (Dasgupta et al., 2016) through a series of six backcrosses. BL<sub>st</sub> population is maintained under a set of conditions identical to the other BL populations.

#### 2.1 | Paternal treatment

Sires and dams were generated from a BL population. The broad design of the protocol followed to generate the experimental flies

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**FIGURE 1** The design of the assay. The schematic diagram shows the design of the entire study, which spanned two generations. Treatment (Normal [N], Medium [M] and High [H] adult densities) was given in the paternal generation. Untreated dams were mated to the treated sires, followed by the collection of eggs from the dams. Assays were done with the eggs and the offspring emerging out of the eggs. Some eggs were subjected to mixed culture (along with competitor eggs) and juvenile competitive fitness vials were set up. Some eggs were cultured as monocultures (without any competitor eggs)—male progeny emerging from these vials were used for further assays, such as, mating ability, mating latency, copulation duration, competitive mating success and courtship frequency.

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is described in Figure 1. To generate the experimental sires and the
dams, eggs were collected from a BL population and cultured under
standard density (i.e., 70 per 6–8 ml food per vial). 100 such vials were
set up, of which 65 were used to collect the sires (=sire-vials) and the
remaining 35 for dams (dam vials). Dams were collected as virgins and
held in single sex vials at a density of 25 per vial with ad-lib food until
the day of the sire-dam mating (see below). In the sire-vials, all the
flies were allowed to eclose. These flies were used to set up three
density treatments—normal (N: 70 individuals per vial), medium
(M: 140 individuals per vial) and high (H: 210 individuals per vial; see
Figure 2). 10 vials were set up for each of the treatments, using flies
that were approximately 1-day old. These vials were left undisturbed
for 2 days, following which males from them were separated and used
as sires in the subsequent step. Here and elsewhere throughout the
study, all the fly sorting, including collection of virgins, were done
under light CO₂-anesthesia, unless mentioned otherwise.

2.2 Sire-dam mating

Following the 2-day-long conditioning, 25 males were randomly iso-
lated from each adult density treatment vials, to be used as sires.
They were then combined with dams (see previous section) in fresh
food vials (25 sires + 25 dams in a vial) and allowed to interact for
90 min, which is sufficient time for a single round of mating. This
method of ensuring single round of mating has been previously used
(Nandy, Joshi, Ali, Sen, & Prasad, 2012). In addition, mating was visu-
ally observed. Occasionally, in some vials, a small number of females
failed to mate within this time. We did not make any attempt to re-
move them. These un-mated females either mated with an already
mated male after a while (late mating) or remained un-mated. Most
males secured a single mating, while some very small number (those
which mated with the un-mated females mentioned earlier) may
have secured more. The number of such late matings (and hence,
male re-mating) was very small, and therefore very unlikely to have
any perceivable impact on the subsequent assays. Further, the fe-
males in this system usually do not re-mate within such short span
(i.e., 90 min) unless the first one was a failed mating, which is very
rare in our populations. Therefore, by following this protocol, we
generated singly inseminated females (average number per vial ~25).
10 mating vials were set up per density treatment. After mating, the
sires were discarded and the already inseminated dams from all 10
vials of a treatment (i.e., a total of 250 females) were transferred to a
2 L plastic cage with food smeared with ad-lib quantity of live yeast.
Three such cages were thus set up—one for each density treatment.
After 2 days, eggs were collected from these cages to set up the
remainder of the experiments. To collect the eggs, a fresh food plate
was introduced in the cage. The dams were allowed a short window
(2–3 hr) for oviposition. Using a fine brush, eggs were counted on to
a fine agar strip, which was then transferred to the culture vials (see
below). These eggs are hereafter referred to as treatment eggs.

2.3 Measurement of egg size

To test if the sires influenced the size of the eggs laid by the dams
(Pischedda, Stewart, Little, & Rice, 2010), a subset of these eggs
were frozen at −20°C and their size was measured. For this purpose,
egg were mounted on a glass slide on their dorsal side and photo-
graphed using Nikon Stereo-zoom trinocular microscope (SMZ745T)
and the area of the two-dimensional elliptical outline of the eggs
were measured in ImageJ, software. This area was taken as a proxy
for the size of each egg. A given egg was measured thrice and the av-
erage of these three measurements was taken as the unit of analysis.
50 eggs per treatment were measured for this purpose.

2.4 Experiment 1: Juvenile fitness assay

Egg-to-adult survivorship was taken as a measure of Juvenile fit-
ness. Survivorship of the treatment eggs were measured against a
back ground of a common competitor (BLsand) under two conditions—
crowded (C: 150 larvae per 1.5 ml food in each vial) and un-crowded
(UC: 70 larvae per 6 ml food in each vial). During the assay, treat-
ment eggs generated in the previous step were cultured with eggs
from common competitors in the ratio 1:4 (C: 30 targets, 120 com-
petitors; UC: 14 targets, 56 competitors). These common competi-
tors were collected from an untreated BLsand stock. On completion of
development, it was possible to identify the target progeny from the
competitor progeny based on eye color—progeny of the competitors
were scarlet-eyed, whereas the target progeny was red-eyed. 10 ju-
venile competition vials were set up for each of the three treatments
(viz., N, M and H) and two assay conditions (i.e., 10 as C and 10 as
UC for each treatment). These vials were left undisturbed until adult
emergence was complete (12th day post-egg deposition). The adults
were sorted based on eye color and counted. Juvenile fitness score
(w) was calculated for each vial following the formula:

\[ w = \frac{\text{number of red eyed progeny observed}}{\text{number of red eyed progeny expected}} \]
The number of red-eyed progeny expected was 14 and 30 for UC and C-assay conditions respectively.

2.5 | Experiment 2: Assay for behavior and fitness of the sons

To investigate the effect of the treatment on the male progeny, the treatment eggs were cultured in food vials in the usual density (i.e., 70 per 6 ml food in each vial) and the progeny were allowed to develop. Upon onset of eclosion, males were collected as virgins (<6 hr post-eclosion). Four assays were run with these males. (a) For each treatment, 50 males were immediately frozen at −20°C and were later dried at 60°C for 48 hr and weighed in groups of five using Shimadzu AUW220D to the nearest 0.01 mg. (b) A separate set of males were similarly collected and held in groups of 5 per vial for further assays. Ten such vials, for each treatment, were set up and left undisturbed till they were 3-days old. These males were then transferred to fresh food vials (hereafter referred to as mating vials) along with five age-matched, virgin females. Mating vials were set up without the use of anesthesia. The females used in this step came from the same replicate BL population and were generated under their standard maintenance conditions, collected as virgins and held in groups of five per vial with ample food until the day of the experiment. 10 mating vials were set up for each of the three treatments. They were observed (manually, without any video recording) continuously till all the flies finished mating. Every 2 min starting from the time when the females were introduced in these vials, the total number of mating pairs (nxy, n: number, x: time elapsed in minutes) was noted down at each time point (x = 0, 2, 4, 6...). Mean mating latency (ML, time taken by a virgin pair to start mating) and mean copulation duration (CD, duration for which a pair mated) were calculated following an algorithm mentioned below.

\[
ML = \frac{\sum (n_{x-2} - n_x) \times x}{N}
\]

For all values of x, until, \( n_{x-2} \leq n_x \)

\[
CD = \frac{\sum (n_{x-2} - n_x) \times x}{N} - ML
\]

For all values of x, until, \( n_{x-2} \geq n_x \)

Occasionally, some females did not mate within 1-hr long observation. These flies were excluded from the analysis. Similarly, some males also failed to secure mating. In vials having such an unsuccessful male, a mating was recorded much later—when one of the successful males finished its first mating and then initiated a second one with the un-copulated female. Such late copulations were also excluded from the analysis. Mating ability (MA) is measured as the proportion of the sons successfully copulated. MA was calculated for every single vial.

(c) Courtship frequency was quantified for the 3-day-old (post-eclosion) sons of the three paternal density treatments by setting up similar mating vials as described in the previous section. Ten vials were set up for each treatment. Therefore, a total of 30 vials were observed. After allowing the first mating, the courtship observation was initiated after a gap of approximately half an hour. Vials where all the flies did not mate were removed from the assay. Every 45 min, each vial was observed for 30 s, during which the total number of courtship bouts (male to female) was noted down. A total of 8 observations were taken. In Drosophila, courtship behavior includes chasing, tapping, courtship dance and song, genital licking and attempted mounting (Bastock & Manning, 1955; Sokolowski, 2010). Any of the above-mentioned courtship behaviors, displayed by the five males in each vial was counted as one. The total number of independent male to female courtship displays was counted within the observation window (Bedhomme, Prasad, Jiang, & Chippindale, 2008; Nandy et al., 2013). The treatment identities were unknown to the observers to avoid observer bias. (d) In the fourth assay, another set of males were similarly collected and held, to be used for quantifying their mating success under competitive condition (CMS, Competitive mating success). This was done by setting up mating vials with five 3-day-old target males, five competitor males (BLst) and five virgin females (BLst). Ten such mating vials were set up for each of the three treatments. After allowing a single round of mating (i.e., for 90 min) for all the females in a mating vial, the females were individually transferred to oviposition test tubes (12 mm diameter × 75 mm height) with ample food. The females were allowed to oviposit for 18 hr. Following oviposition, the females were discarded and the tubes were retained to allow the progeny to develop and eclose. For each female, the identity of their mate (whether target/competitor) was ascertained by observing the eye color of the progeny. Progeny sired by target males were red-eyed whereas those sired by competitors were scarlet-eyed. For a given vial, average CMS of the five target males in the vial was calculated as the proportion of the females mated to target males (i.e., produced red-eyed offspring).

2.6 | Experimental replications and data analyses

The entire study was carried out in three randomized blocks, using three different BL populations—BL1, BL5, and BL9. The blocks were handled on separate days. Number of replications within each block has been mentioned in the previous sections along with the assay design. Except for the egg size and dry body weight assay, all the experimental replication was done at the level of assay mating vials or juvenile competition vials. All the assays had 10 replicate vials. Vial means were used as the unit of analysis. For egg size assay, size of each egg was used as the unit of analysis. For dry body weight, weight of groups of five individuals was used as the unit of analysis. Data were analyzed using mixed-model analysis of variance (ANOVA). Block was treated as random factor, while paternal density treatment and assay density (wherever applicable) were treated as fixed factors. Multiple comparisons were done using Tukey’s HSD.
All the analyses were done in Statistica, version 10 (Statsoft, Tulsa, OK, USA).

### 3 | RESULTS

Variation in size of the eggs represents variation in maternal provisioning. The effect of the paternal density treatment on size of the eggs produced by the dams was not significant (Table 1, mean ± SE, μm²; N: 80,039.1 ± 387.5; M: 79,967.4 ± 415.3; H: 79,611.9 ± 411.4). The juvenile competitive fitness assay quantified overall egg-to-adult survival of the target juveniles compared to the same of juveniles from a common background (common competitors). While the data from un-crowded assay condition reflects the baseline survivorship, those from crowded assay condition represents difference in juvenile competitive ability across the three paternal density treatments. Paternal density treatment had a significant effect on Juvenile fitness (Table 1). While there was no significant difference between N and M-treatments, H-treatment had 8.9% higher juvenile fitness compared to that of the N-treatment. This relative advantage

| Trait                      | Effect          | SS       | DF  | MS       | DF Den | MS Den | F         | p       |
|----------------------------|-----------------|----------|-----|----------|--------|--------|-----------|---------|
| Egg size                   | Paternal density (PD) | 1.75 × 10⁷ | 2   | 8.75 × 10⁶ | 4.02   | 3.53 × 10⁶ | 2.48   | 0.20   |
|                            | Block           | 9.84 × 10⁸ | 2   | 4.92 × 10⁸ | 4.03   | 3.54 × 10⁶ | 139.11 | <0.01  |
|                            | PD × Block      | 1.41 × 10⁷ | 4   | 3.53 × 10⁶ | 438.00 | 2.25 × 10⁷ | 0.16   | 0.96   |
| Juvenile fitness           | Paternal density (PD) | 0.21 | 2   | 0.11   | 4.03   | 0.01   | 9.93   | 0.03   |
|                            | Assay density (AD) | 0.97 | 1   | 0.97   | 2.00   | 0.10   | 9.31   | 0.09   |
|                            | Block           | 0.40 | 2   | 0.20   | 1.58   | 0.09   | 2.15   | 0.35   |
|                            | PD × AD         | 0.23 | 2   | 0.11   | 4.01   | 0.02   | 5.42   | 0.07   |
|                            | PD × Block      | 0.04 | 4   | 0.01   | 4.00   | 0.02   | 0.51   | 0.74   |
|                            | AD × Block      | 0.21 | 2   | 0.10   | 4.00   | 0.02   | 4.96   | 0.08   |
|                            | PD × AD × Block | 0.08 | 4   | 0.02   | 148.00 | 0.02   | 1.24   | 0.30   |
| Dry body weight            | Paternal density (PD) | 2.67 × 10⁻⁷ | 2   | 1.34 × 10⁻⁷ | 4.00   | 6.22 × 10⁻⁹ | 21.49 | <0.01  |
|                            | Block           | 7.78 × 10⁻⁷ | 2   | 3.89 × 10⁻⁷ | 4.00   | 6.22 × 10⁻⁹ | 62.50 | <0.01  |
|                            | PD × Block      | 2.49 × 10⁻⁸ | 4   | 6.22 × 10⁻⁹ | 80    | 1.27 × 10⁻⁸ | 0.49   | 0.74   |
| Mating latency             | Paternal density (PD) | 41.810 | 2   | 20.90   | 4.00   | 1.12   | 18.59 | 0.01   |
|                            | Block           | 51.158 | 2   | 25.58   | 4.00   | 1.12   | 22.75 | 0.01   |
|                            | PD × Block      | 4.477 | 4   | 1.12   | 77.00  | 4.59   | 0.24   | 0.91   |
| Copulation duration        | Paternal density (PD) | 14.43 | 2   | 7.21    | 4.00   | 1.81   | 3.98   | 0.11   |
|                            | Block           | 80.87 | 2   | 40.44   | 4.00   | 1.81   | 22.32 | 0.01   |
|                            | PD × Block      | 7.22 | 4   | 1.81    | 77.00  | 5.51   | 0.33   | 0.86   |
| Mating ability             | Paternal density (PD) | 0.009259 | 2   | 0.00    | 4.00   | 0.04   | 0.10   | 0.90   |
|                            | Block           | 0.046678 | 2   | 0.02    | 4.00   | 0.04   | 0.52   | 0.63   |
|                            | PD × Block      | 0.179963 | 4   | 0.04    | 77.00  | 0.02   | 2.64   | 0.04   |
| Courtship frequency        | Paternal density (PD) | 13.28 | 2   | 6.64    | 4.03   | 10.79  | 0.62   | 0.58   |
|                            | Block           | 500.78 | 2   | 250.39  | 4.00   | 10.81  | 23.16  | 0.01   |
|                            | PD × Block      | 43.24 | 4   | 10.81   | 73.00  | 7.36   | 1.47   | 0.22   |
| Competitive mating success | Paternal density (PD) | 0.64 | 2   | 0.32    | 4.03   | 0.02   | 19.47  | 0.01   |
|                            | Block           | 0.09 | 2   | 0.05    | 4.06   | 0.02   | 2.86   | 0.17   |
|                            | PD × Block      | 0.07 | 4   | 0.02    | 75.00  | 0.04   | 0.41   | 0.80   |

Note. Paternal density and assay density (wherever applicable) were considered as fixed factor, while block as random factor. All tests were done considering α = 0.05 and significant p-values are mentioned in bold face.
Competitions. Multiple comparisons on the CMS results indicated that the M-sons had significantly lower CMS compared to H and N-treatments. CMS of the M-sons was approximately 34% less than that of the N-sons. This is however, not due to a reduced courtship performance by the M-sons as we found the effect of the treatment on CF (mean ± SE, N: 6.7 ± 0.6; M: 6.8 ± 0.6; H: 7.6 ± 0.8) to be non-significant. We also did not find any effect of the treatment on CD (mean ± SE, minutes, N: 18.6 ± 0.4; M: 17.6 ± 0.4; H: 17.9 ± 0.4), potentially indicating the lack of the treatment effect on post-copulatory traits of the sons (Table 1).

4 | DISCUSSION

Given that very few studies have shown the effect of paternal environment on offspring fitness components, there were two main objectives of the present study—(a) to assess if paternal exposure to varying population density affected progeny traits; if yes, then (b) to evaluate the adaptive significance of such effect. The results clearly showed that at sufficiently high density, males had an adaptive paternal effect on juvenile competitive fitness. As we did not find any effect of our treatment on size of the eggs produced by the dams, such paternal effect is unlikely to be mediated by variation in provisioning by the females. We further show that at intermediate density, males sire smaller sons which are inferior in acquiring mates. Interestingly, such maladaptive effect of paternal density on offspring adult fitness was not detected at high density.

In holometabolous insects like fruit flies, juvenile (larva and pupa) survival constitutes one of the most important components of fitness (Prasad & Joshi, 2003). In addition, juvenile ecology may also have a major effect on the life-history and fitness components of the adult stage (Heat shock: Khazaeli, Tatar, Pletcher, & Curtsinger, 1997; cold shock: Singh, Kochar, & Prasad, 2015; Singh & Prasad, 2016; crowding: Joshi & Mueller, 1988; Sarangi et al., 2016; Sheno et al., 2016). The observed paternal effect on juvenile competitive fitness therefore is extremely consequential. Some relatively recent studies have pointed out that evolving parental ability to optimize offspring fitness related traits can be an adaptation to ecological challenges (Galloway & Etterson, 2007), including crowding (Crean et al., 2013). Given that fruit fly natural ecology regularly involves adult and larval crowding, the observed paternal effect on juvenile competitive fitness can indicate males’ adaptation to crowding. Interestingly, we observed the paternal effect on juvenile competitive fitness, only at the highest density, which may indicate a certain threshold density beyond which such paternal effect starts affecting offspring traits. In addition, when assayed under un-crowded condition the progeny from the three sire treatments do not show any measurable difference in their egg-to-adult survival. This suggests that the juvenile competitive ability rather than baseline juvenile viability was affected by the treatment. Since a number of traits (e.g., feeding rate, waste tolerance, development time etc.) affect juvenile competitive ability in these flies, it will be interesting to...
find out the trait responsible for better competitive ability of the H-sons in our study.

In a wide range of species including D. melanogaster, maternal exposure to high density or poor nutrition has been found to affect offspring fitness components (Prasad & Joshi, 2003; Valtonen et al., 2012; Vijendravarma et al., 2010). Such effects can either be beneficial (Allen, Buckley, & Marshall, 2008; Bashey, 2006; Gorbí, Moroni, Sel, & Rossi, 2011; Mitchell & Read, 2005; Vijendravarma et al., 2010) or detrimental (Meylan, Clopert, & Sinervo, 2007) depending on the component of the fitness under consideration and the prevailing condition. As maternal provisioning and other maternal effects play vital roles in offspring survival and performance, such maternal density/nutrition effect is not surprising. However, what is not intuitive is the paternal density to have similar impact on offspring fitness, as our results suggest, given that Drosophila males do not pass on any nutrition to the offspring. It is well known in the Drosophila literature that even the laboratory populations harbor heritable genetic variation in survival under crowding both as adults and juveniles (see Sarangi et al., 2016 and the references therein for an updated review). Therefore, one possibility is that genetically superior males, which are better at surviving under high density, may produce offspring which are better both as juveniles, explaining at least part of our observations. Though larval competitive ability is known to respond to experimental evolution, indicating heritable genetic variation (Mueller, 1997; Prasad & Joshi, 2003), such heritable variation is very unlikely to have led to the observed treatment effect on juvenile competitive fitness. This is because (a) in our assay, we recorded very little mortality in males during the treatment, indicating negligible hard selection. In addition, we also ensured that there was no soft selection by randomly picking the set of males from the treatment vials to use them as sires. Further, we allowed the sires and the dams to mate only once by allowing them a limited window of time to interact after being put together in mating vials. (b) Even if there was selection in the current experimental design, the selection is likely to be weak (see Materials and Methods section). Such weak selection is unlikely to explain the observed differences in some of the traits (viz., 8.9% increase in juvenile competitive fitness, 35% increase in mating latency), especially within one generation. Alternatively, males may alter maternal provisioning and thereby indirectly affect offspring fitness components (Prasad et al., 2003; Vijendravarma et al., 2010). We, however, did not find any measurable difference in the size of the eggs produced by females mated to the males belonging to the three treatments, making variation in maternal provisioning an unlikely explanation. Therefore, although

**FIGURE 4** Effect of the paternal density treatment on male offspring. (a) Dry body weight at eclosion: five flies were weighted together to nearest 0.01 mg. This was then used as the unit of analysis; (b) Mean mating latency (time taken by a virgin male–female pair to start copulation): mean ML was calculated for five males in a vial following the algorithm given in the Materials and Methods section. This was done for all the mating vials in the assay. These values were then used as the unit of analysis; (c) Competitive mating success (CMS): CMS values were calculated for each vial having five target males as the proportion of females mated to target males in these assay vials. These values were then used as the unit of analysis. The blue broken line indicates the expected value of CMS if there is no mating bias. Black, blue and red color coding represent data from the progeny of N, M and H-males respectively. Treatments not sharing common alphabet were found to be significantly different from each other. The entire experiment was done following a randomized block design and the data were analyzed using three-factor mixed-model ANOVA with paternal treatment and assay condition as fixed factors and block as random factor.
sire effect on the quality of the eggs produced by the females cannot be completely ruled out without a more detailed qualitative analysis of the eggs, our results tentatively point at non-genetic paternal effect (Bonduriansky & Day, 2009) as the potential cause behind the observed effect of the treatment. Interestingly, a recent study on D. melanogaster has shown paternal effects to have important consequences on the expression of an array of genes in sons (Zajitschek, Zajitschek, & Manier, 2017, also see the corresponding correction). In addition, Garcia-Gonzalez and Dowling (2015) reported non-sire effect on daughters’ reproductive output in D. melanogaster, possibly caused by the seminal fluid proteins transferred by the males to their mates during copulation (Garcia-Gonzalez & Dowling, 2015).

While we found adaptive paternal effect on juvenile performance, adult performance however, was found to have a significant maladaptive effect of paternal density. Males that experienced intermediate density were found to sire sons which (a) are smaller, (b) take longer time to start mating and (c) have lower mating success. Since we did not find any effect of the treatment on courtship frequency, reduced mating success and increased mating latency was a likely outcome of females’ reluctance to accept relatively smaller males as their mates, a known fitness consequence of reduced size in Drosophila males (Jagadeeshan, Shah, Chakrabarti, & Singh, 2015; Partridge, Ewing, & Chandler, 1987). Body size has been reported to be affected by intergenerational paternal effect in another Dipteran—Telostylus angusticollis (Bonduriansky & Head, 2007). Unlike the maladaptive effect found in our study, the paternal effect on body size reported by Bonduriansky and Head (2007) was adaptive, especially under certain prevailing conditions. In D. melanogaster, the effect of body size on different fitness components has been found to be context specific (Lefranc & Bundgaard, 2000; Morimoto, Pizzari, & Wigby, 2016; Pitnick, 1991). Thus, although in the context of our experiment, the observed body size reduction in our study appears to be maladaptive, a detailed investigation where offspring fitness is measured under varying adult density is necessary to better understand the fitness consequences. An adaptive paternal effect theory would predict higher fitness of the offspring, particularly when offspring conditions match the paternal conditions. Interestingly in our study, the paternal (maladaptive) effect was found only at intermediate density and not in the high density treatment. However, at this point it is difficult to suggest any reason for such specific expression of the paternal effect at intermediate density.

As variation in population density and crowding related ecological challenges are common in almost all organisms, including fruit flies, paternal effect of the nature reported here is important to understand. Though paternal ability to optimize offspring traits is likely to be adaptive, especially under fluctuating environment, the results reported here show that paternal effect can be both adaptive and maladaptive. To the best of our knowledge this is the first evidence of the effect of paternal density on juvenile and adult fitness components in D. melanogaster. Importantly, our results emphasize the importance of considering paternal effect as a source of variation in fitness related traits. The full impact of such paternal effect in the evolution of life-history traits and the underlying mechanisms are emerging as an important topic of discussion, which is likely to see an increasing attention in years to come.

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CONFLICT OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS

BN, PD and SS conceived the ideas and designed the assays. PD, SS, AAD, TV, and to a lesser extent BN performed the assays and collected the data. Data analysis was primarily done by BN and to some extent, by PD. While BN and PD led the writing of the manuscript, SS and TV provided important assistance and inputs.

DATA ACCESSIBILITY

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