Sleep in Populations of *Drosophila Melanogaster*¹,²,³

Chang Liu, Paula R. Haynes, Nathan C. Donelson, Shani Aharon, and Leslie C. Griffith

DOI: http://dx.doi.org/10.1523/ENEURO.0071-15.2015

Department of Biology, National Center for Behavioral Genomics and Volen Center for Complex Systems, Brandeis University, Waltham, Massachusetts 02454-9110

Abstract

The fruit fly *Drosophila melanogaster* is a diurnal insect active during the day with consolidated sleep at night. Social interactions between pairs of flies have been shown to affect locomotor activity patterns, but effects on locomotion and sleep patterns have not been assessed for larger populations. Here, we use a commercially available locomotor activity monitor (LAM25H) system to record and analyze sleep behavior. Surprisingly, we find that same-sex populations of flies synchronize their sleep/wake activity, resulting in a population sleep pattern, which is similar but not identical to that of isolated individuals. Like individual flies, groups of flies show circadian and homeostatic regulation of sleep, as well as sexual dimorphism in sleep pattern and sensitivity to starvation and a known sleep-disrupting mutation (*amnesiac*). Populations of flies, however, exhibit distinct sleep characteristics from individuals. Differences in sleep appear to be due to olfaction-dependent social interactions and change with population size and sex ratio. These data support the idea that it is possible to investigate neural mechanisms underlying the effects of population behaviors on sleep by directly looking at a large number of animals in laboratory conditions.

Key words: *Drosophila*; population; sleep

Significance Statement

Most species live in an interactive environment in their natural habitats, and sleep can be affected by social cues. Although flies have been widely used to understand the mechanisms of sleep in recent years, sleep in large populations has not been systematically studied. Here we report both similarities and differences between sleep in populations of flies compared with individuals, as well as provide a new method for the study of social behavior under constant environmental conditions.
suggested that flies have a genuine sleep state. Flies have therefore been increasingly used for the neurogenetic dissection of sleep/wakefulness behavior and the circuits that produce it.

In Drosophila, sleep is defined as quiescence for longer than 5 min (Hendricks et al., 2000; Shaw et al., 2000). This criterion was established by examination of the timing of changes in arousal threshold after the onset of quiescence and allows measurement of locomotor activity with the standard Drosophila activity monitor (DAM2) system to be used to assess the amount and structure of sleep. Most previous studies of social behaviors focused on sleep by recording a pair of flies’ locomotor activity (Fujii et al., 2007; Lone and Sharma, 2012; Hanafusa et al., 2013); however, as social animals (Hay, 1973; Schneider et al., 2012), whether flies in a group sleep in a similar manner as isolated individual flies remains largely unknown. In the present study, we observed activity in larger groups in order to characterize sleep in populations of flies and compare it to individual fly sleep. We show that the commercially available Drosophila population monitor (LAM25H) system can be used to analyze population sleep/activity patterns and investigate the neural mechanisms of population behavior.

Materials and Methods

Animals

Flies were raised in a 12 h light/dark cycle on modified Brent and Oster cornmeal-dextrose-yeast agar food (Brent and Oster, 1974). Per batch: 60 L H2O, 600 g Agar, 1950 g flaked yeast, 1,451 g cornmeal, 6300 g dextrose, 480 g NaKt, 60 g CaCl2, and 169 g Lexgard dissolved in ethanol. Canton S, amn1, w;Orco2 (also known as Or83b2; Larsson et al., 2004) and wCG flies were raised at 25°C in an incubator after eclosion. Males and mated females were used for all experiments.

Behavioral analysis

Newly enclosed flies were raised in standard bottles and transferred to new food bottles every 2–3 d. Mating was allowed to happen freely before sorting into storage vials. Flies were sorted into small vials (50 flies per vial) 1 d prior to the loading day. Flies were 2- to 7-d-old at the start of each experiment. Individuals were placed into 65 × 5 mm glass tubes and populations of 50 flies into 95 × 25 mm glass-like vials. All sleep tubes/vials contained 2% agarose with 5% sucrose food. Flies were entrained in 12 h light/dark (LD) conditions for 2–3 d. Activity was then recorded for 2 d in LD then switched to constant darkness for another 2 d (data not shown).

For sleep deprivation (SD) experiments, a mechanical stimulus was applied using a Trikinetics plate attached to a VWR vortex mixer with a shaking frequency of 2 s of every 10 s for the entire 12 h dark period. Populations of flies housed in vials were removed from the LAM25H system onto the shaker 10 min before ZT12, and placed back immediately in the LAM25H after the 12 h of sleep deprivation to record their sleep rebound. Because data could not be collected during this period, no data are shown for the period of deprivation for the SD group. For starvation experiments, flies were transferred to 2% agarose vials at ZT0 for 24 h, and put back onto sucrose-agarose food for recovery.

Calculation of relative sleep changes and statistical analyses

The behavioral patterns of individuals and groups of flies were monitored using the DAM2 and LAM25H systems (Trikinetics, respectively). Diagrams of the apparatus are shown in Figure 1. Sleep parameters were analyzed using an in-house MATLAB program described previously (Donelson et al., 2012) from averages of 2 d of LD data in most experiments. All sleep manipulations (sleep deprivation and starvation) were performed for 1 d. Total sleep, number of sleep episodes, mean episode length, activity while awake, and sleep latency were analyzed for 24 h and/or 12 h light and dark periods (LP and DP). Sleep data were analyzed using Prism 6 software (GraphPad). For experiments that had multiple variables, a two-way ANOVA was performed (Table 1). Multiple comparisons after two-way ANOVA were used for each analysis period (24 h, LP and DP), and were performed to determine which pairs were significantly different and if major effects are significantly different. Holm–Sidak’s/Dunn’s test were used according to the distribution of datasets (Table 2). Datasets are marked with letters (A, B, C, or D) for statistical equivalence groups; i.e., data that are significantly different are indicated by different letters. To evaluate the sleep changes (∆Sleep) during and/or after manipulations, we subtracted the sleep during manipulation days and the sleep after manipulations from its baseline day sleep. The sleep change of the experimental group was compared with the control groups using an unpaired t test if it passed a normality test or Mann–Whitney test if it did not pass a normality test (Table 3). For experiments with different ratio of males in the population, datasets that did not have a normal distribution, nonparametric statistics (Kruskal–Wallis test followed by Dunn’s multiple-pairwise-comparison test) were applied. Otherwise, a one-way ANOVA followed by Holm–Sidak’s test was applied (Tables 4, 5). Figures are all presented as mean ± SEM in a uniform figure style for clarity. For single comparisons, asterisk (*) indicates a significant difference between the experimental group and the control group. The significance level of statistical tests was set to 0.05.

Results

Population sleep patterns differ from those of isolated individuals

Drosophila are normally social animals (Hay, 1973), and their behavior and daily activity patterns can be changed by interactions with other individuals in a population (Levine et al., 2002; Krupp et al., 2008; Schneider et al., 2012). To address whether the features of sleep in populations of flies are similar to those observed for individual flies, we compared sleep patterns of isolated individual Canton S wild-type flies to those from groups of Canton S flies using data collected with the DAM2 and LAM25H systems. DAM2 records the activity of 32 individual animals using two infrared beams across a 5 mm tube (Fig.
LAM25H records 32 vials (25 mm diameter) with high resolution using nine infrared beams across the center of each vial to detect activity (Fig. 1B).

As shown in Figure 2A and C, individual males slept more than individual females during the day, consistent with previous reports of sexual dimorphism (Huber et al., 2004; Andretic and Shaw, 2005; Isaac et al., 2010). When kept in same-sex groups, daytime sleep was still greater in males than in females, but the total amount of sleep for both sexes was lower (Fig. 2B,C) suggesting that perhaps the presence of other flies and the attendant sensory stimulation may decrease sleep.

A more interesting difference between individuals and groups was seen in nighttime sleep. When flies were alone, males slept more than females at night (Fig. 2A). In same-sex populations, however, males slept significantly less than females and this difference was mostly due to a decrease in sleep in the last half of the night (Fig. 2B). The total amount of nighttime sleep compared between individuals and same-sex groups was the same for females, but significantly less for males (Fig. 2C). These data imply that male–male interactions either increase late at night or are more arousing in that time window than female–female interactions.

The differences in sleep were not due to sex-dependent differences in locomotor activity. Activity levels during waking periods were higher in males than in females for both isolated individuals and groups of flies, and activity counts during waking periods were higher for groups than for isolated flies for both sexes (Fig. 2D). Because an activity count (beam break) can be generated by a single fly, groups will naturally have more beam breaks during active periods and this will lead to an increase in overall “activity”. Activity during wake periods in the population datasets is therefore not a measure of locomotor activity or speed of individuals in the population, it is an aggregate measure that reflects both individual locomotor activity level and the number of flies that are active in a time window. If the presence of other flies stimulates locomotion differentially at different times of day, this would manifest as differences in activity but would not necessarily mean that individual flies were moving faster/slower at that time of day. The difference between male and female groups suggests that there are sex-specific increases in population activity that might be caused by interactions between males. We speculate that this might reflect increased aggression, but without direct observation it is difficult to know. In any case, this difference in basal activity likely contributes to the differences in nighttime sleep between same-sex groups.

Sleep architecture metrics were also affected by sex and group interactions. Individual male flies exhibit more consolidated sleep than individual female flies; i.e., fewer episodes but longer episode duration. In populations of flies, however, there was no difference between male and female groups (Fig. 2E,F). The increase in number of episodes and the decrease in their duration for groups compared with individuals is likely a reflection of the fact that during a population sleep episode all flies in the group must necessarily be immobile, but the activity of a single fly in the group can terminate a population sleep bout. The calculated average population sleep bout duration therefore reflects the minimum sleep bout duration for individuals in the group rather than an actual average length, which is an important distinction in interpreting population data. In general, sleep structure parameters for individuals cannot be extrapolated from population data in a quantitative manner.

Figure 1. Diagrams of DAM2 and LAM25H systems. A, DAM2 apparatus. Left, Side view of DAM2 sleep tube (5 × 65 mm) for individual fly recording showing location of infrared beams and food. Right, Cross-section of the tube with the orientation of the two infrared beams. B, LAM25H apparatus. Left, Side view of LAM25H vial (25 × 95 mm) for population recording showing location of infrared beams and food. Right, Cross-section of the vial with the orientation of the nine infrared beams. Dark blue bars and light blue bars indicate transmitters and receivers. Red arrow lines indicate how pairs of infrared beam sensors work, as well as the coverage of the cross-sectional area.
In contrast to total sleep and sleep structure parameters, latency to sleep appeared to scale similarly between isolated flies and groups. Isolated males had shorter latency to sleep onset than isolated females, but there were no significant differences between grouped males and females (Fig. 2G). The absolute latency to sleep onset was much higher in groups during the day, whereas nighttime latencies were of similar magnitude for both individuals and groups. This may reflect a difference in sleep drive during the day and the night. At night, sleep drive is strong enough to overcome the sensory stimulation provided by other individuals in the group, but during the day these sensory inputs are disruptive in groups.

These results suggest that quiescence in populations of flies shares similarities in overall presentation with sleep that has been characterized in individual animals in terms of day/night distribution and sexual dimorphisms in the amount of daytime sleep. The fact that same-sex groups

Table 1. Two-way ANOVA

| Source of variation | Gender (female vs male) | Interaction |
|---------------------|-------------------------|-------------|
|                    | F          | p     | F          | p     | F          | p     |
| Group (individual vs population) | 123.7 | <0.0001 | 86.91 | <0.0001 | 0.02634 | 0.8715 |
| Gender (female vs population) | 136.5 | <0.0001 | 37.59 | <0.0001 | 18.87 | <0.0001 |
| Genotype (CS vs amn) | 12.8 | <0.0001 | 2.376 | 0.1274 |
| Source of variation | Food (complete vs sucrose agar) | Interaction |
|---------------------|-------------------------|-------------|
|                    | F          | p     | F          | p     | F          | p     |
| Group (individual vs population) | 123.7 | <0.0001 | 86.91 | <0.0001 | 0.02634 | 0.8715 |
| Gender (female vs mixed) | 12.8 | <0.0001 | 2.376 | 0.1274 |
| Genotype (w vs Orco2) | 20.44 | <0.0001 | 24.5 | <0.0001 | 3.26 | 0.0721 |
| Source of variation | Size (10 vs 50 vs 100) | Interaction |
|---------------------|-------------------------|-------------|
|                    | F          | p     | F          | p     | F          | p     |
| Group (individual vs population) | 123.7 | <0.0001 | 86.91 | <0.0001 | 0.02634 | 0.8715 |
| Gender (female vs male) | 12.8 | <0.0001 | 2.376 | 0.1274 |
| Genotype (w vs Orco2) | 20.44 | <0.0001 | 24.5 | <0.0001 | 3.26 | 0.0721 |
| Source of variation | genotype (w vs Orco2) | Interaction |
|---------------------|-------------------------|-------------|
|                    | F          | p     | F          | p     | F          | p     |
| Group (individual vs population) | 123.7 | <0.0001 | 86.91 | <0.0001 | 0.02634 | 0.8715 |
| Gender (female vs male) | 12.8 | <0.0001 | 2.376 | 0.1274 |
| Genotype (w vs Orco2) | 20.44 | <0.0001 | 24.5 | <0.0001 | 3.26 | 0.0721 |

July/August 2015, 2(4) e0071-15.2015 eNeuro.sfn.org
show qualitative differences in sleep patterns and activity during the night, however, implies that the presence of other animals affects sleep in ways that are not simply due to changes in the number of flies in the apparatus. Sex-specific social interactions appear to modulate the amount and pattern of nighttime sleep.

**Homeostatic sleep regulation in populations of flies**

Although populations demonstrate quiescence periods with the same basic structure and many of the properties of sleep that have been characterized in individual flies, to be considered true sleep, this quiescence has to be homeostatically regulated. To examine this issue we used two methods for disrupting sleep. First, we mechanically deprived female flies of sleep for 12 h overnight and measured the amount of excess sleep that was produced over the following 24 h. Both isolated females and groups of females had a significant amount of homeostatic rebound sleep during the day following sleep deprivation (Fig. 3). As a second method of sleep deprivation, we used starvation (Keene et al., 2010) to deprive both males and females. We used a 3
Population sleep is disrupted by mutation of amnesiac

It is clear that quantitative parameters, such as bout length and number in population sleep data, cannot quantitatively reflect the architecture of sleep of individuals in a group due to the manner in which the locomotor data are acquired (see above). To determine whether these measurements can qualitatively inform our understanding of sleep architecture, we compared population sleep in the amnesiac mutant, which is known to have disrupted sleep structure (Liu et al., 2008), with Canton S wild-type to see if population measurements would be able to capture the previously characterized defects.

Similar to what had been reported for amnXa mutants (Liu et al., 2008), we observed loss of sleep both during the day and night, and significant sleep fragmentation in isolated amnX females. amnX Mutant female populations also showed significantly lower sleep during the day and night, and a significantly increased number of sleep episodes at night compared with the wild-type control (Fig. 5). No difference was found between populations of amnXa mutants and wild-type flies in episode length, but absolute episode length of populations is very short compared with that of individuals (Fig. 5E), perhaps reflecting a floor effect. Surprisingly, the number of sleep episodes of populations of amnXa flies during the day time actually decreased compared with wild-type, going in the opposite direction from individual fly measurements (Fig. 5F).

This difference in the number of episodes may be due to changes in locomotor activity. Individual amnXa mutants did not differ in locomotor activity level during wake periods from wild-type flies; however, populations of amnXa mutants exhibited increased activity during the daytime while they were awake and decreased activity at night during wake periods (Fig. 5D). As discussed above this could reflect either changes in individual fly locomotion or could reflect changes in the number of flies active during these time windows. In either case it suggests that the amn gene might have a specific role in the responses to social situations that is not seen in isolated animals. One possibility is that interactions with other flies cause amn mutants to become hyperaroused during the day. amn Mutants lack ability to focus selective attention on visual stimuli (Wu et al., 2000) and have an exaggerated locomotor response to ethanol (Wolf et al., 2002), consistent with altered regulation of arousal (Chi et al., 2014). Until this is experimentally addressed, however, this conclusion remains speculative.

In contrast to a previous study (Liu et al., 2008), we found that amnX mutants appeared to have no change in nighttime sleep latency compared with wild-type flies in either individual or population measurements (Fig. 5G).
The differences between our study and that of Liu et al. (2008) may arise from a number of factors. We used an amn1 stock which was outcrossed to w+ Canton S, whereas Liu et al. (2008) used amn2/+ on a w background. The differences in latency effects may be allele- or genetic background-specific. Our latency results with amn1 are qualitatively similar to what we observed with wild-type flies (Fig. 2), which supports the idea that very high sleep drive can overcome differences in arousal state caused by genotype and the presence of other individuals.

Food quality alters sleep patterns in populations of flies

Although many mutants, such as amn, have been shown to affect sleep, environmental factors such as food quality can also have a profound influence (Zimmerman et al., 2012). To examine whether food quality influences population sleep parameters, we examined sleep patterns on either standard complete food or sucrose-agar food (see Materials and Methods for food details) and compared wild-type Canton S females and males. Both female and male groups slept significantly longer on standard food compared with sucrose-only during the day (Fig. 6A–C). Interestingly, standard food resulted in a significantly more consolidated nighttime sleep pattern (fewer sleep episodes but longer episode duration) in groups of females but in not groups of males (Fig. 6E,F). With standard food, activity during wake periods was significantly elevated in males during the day and in females at night (Fig. 6D), suggesting an interaction of food quality and sex on locomotor activity. Populations of females synchronized their sleep on standard food faster than on sucrose-agar food after lights on/off, but no significant difference was detected in males (Fig. 6G). Altogether these results suggested that food quality has a sex-specific impact on population sleep parameters.

Social interactions alter population sleep

All the data shown in previous figures used a population size of 50 individuals of the same sex. To determine whether population sleep parameters are correlated with population size and to test the effect of mixing the sexes, we measured sleep in all-female and 1:1 male–female mixed populations of 10, 50, and 100 individuals (Fig. 7). Increasing the number of flies dramatically decreased total sleep (Fig. 7A–C), and also increased locomotor activity during wake periods (Fig. 7D) for both female and mixed groups. This is consistent with increased population density providing a higher level of sensory input and consequent arousal. Interestingly, daily sleep episodes exhibited opposite trends in female and mixed populations. Increasing the number of females in a group increased the number of episodes, whereas in mixed populations the number of episodes decreased with increasing population size (Fig. 7E). Nighttime sleep latency was neither significantly affected by the number of flies nor by gender of the population (two-way ANOVA, gender: $F_{(1,26)} = 3.294, p = 0.0550$; size: $F_{(2,26)} = 3.807, p = 0.0619$; interaction: $F_{(2,26)} = 2.859, p = 0.0754$; Fig. 7F), suggesting sleep drive can overcome the arousing effects of increased population size.

A previous study looking at pairs of flies showed that the locomotor activity pattern is driven dominantly by males (Fuji et al., 2007). To further address whether the ratio of males to females in a group has an impact on population sleep, we compared female and males with a ratio of 1:1 (50% male), 1:2 (67% male), and 2:1 (33% male) to same-sex male and female populations. Total daily sleep in all three mixed groups were equivalent, about one-half of that of the single-sex groups (Fig. 8A,B), indicating that total sleep was not sensitive to changes in sex ratio around equivalence. Interestingly, the pattern of

### Table 5. Multiple-comparisons following one-way ANOVA and nonparametric test

| Data | n1 | n2 | 24 h Mean (rank) differences | Adjusted p value | LP Mean rank differences | Adjusted p value | DP Mean rank differences | Adjusted p value |
|------|----|----|-----------------------------|-----------------|------------------------|-----------------|-------------------------|-----------------|
| Fig. 8B Female vs male 100% | 6 | 6 | -92.42 | 0.3864 | -6.833 | >0.9999 | 6 | >0.9999 |
| Female vs male 50% | 6 | 7 | 388.2 | <0.0001 | 11.31 | 0.3021 | 17.79 | 0.0065 |
| Female vs male 77% | 6 | 7 | 427.3 | <0.0001 | 12.6 | 0.1579 | 19.5 | 0.0019 |
| Female vs male 33% | 6 | 6 | 435.2 | <0.0001 | 11.83 | 0.2887 | 19.83 | 0.0025 |
| Male 100% vs male 50% | 6 | 7 | 480.6 | <0.0001 | 18.14 | 0.0051 | 11.79 | 0.2393 |
| Male 100% vs male 77% | 6 | 7 | 519.7 | <0.0001 | 19.43 | 0.002 | 13.5 | 0.0069 |
| Male100% vs male 33% | 6 | 6 | 527.6 | <0.0001 | 18.67 | 0.0057 | 18.33 | 0.1064 |
| Male 50% vs male 77% | 7 | 7 | 39.07 | 0.7816 | 1.286 | >0.9999 | 1.714 | >0.9999 |
| Male 50% vs male 33% | 7 | 6 | 46.98 | 0.8781 | 0.5238 | >0.9999 | 2.048 | >0.9999 |
| Male 77% vs male 33% | 7 | 6 | 7.905 | 0.8861 | -0.7619 | >0.9999 | 0.3333 | >0.9999 |
| Male 100% vs male 100% | 33% | 6 | 6 | 45.67 | >0.9999 | 519.7 | >0.9999 | 6 |
| Fig. 8D Female vs 100% male | 8 | 8 | -24 | 0.1486 | -44.5 | <0.0001 | 23 | 0.2018 |
| Female vs 4% male | 8 | 12 | 5.667 | >0.9999 | -5.5 | >0.9999 | 8.167 | >0.9999 |
| Female vs 10% male | 8 | 12 | 22 | 0.1441 | 1.167 | >0.9999 | 39.17 | >0.9999 |
| Female vs 90% male | 8 | 12 | 3.333 | >0.9999 | -22.17 | 0.1361 | 33.5 | 0.0012 |
| Female vs 96% male | 8 | 12 | -9.667 | >0.9999 | -31.83 | 0.0027 | 23.83 | 0.0754 |
| 100% male vs 4% male | 8 | 12 | 29.67 | 0.0072 | 39 | <0.0001 | -14.83 | >0.9999 |
| 100% male vs 10% male | 8 | 12 | 46 | <0.0001 | 45.67 | <0.0001 | 16.17 | 0.8556 |
| 100% male vs 90% male | 8 | 12 | 27.33 | 0.0194 | 22.33 | 0.1285 | 10.5 | >0.9999 |
| 100% male vs 96% male | 8 | 12 | 14.33 | >0.9999 | 12.67 | >0.9999 | 0.8333 | >0.9999 |
| 4% male vs 10% male | 12 | 12 | 16.33 | 0.4738 | 6.667 | >0.9999 | 31 | 0.0007 |
| 4% male vs 90% male | 12 | 12 | -2.333 | <0.0001 | -16.67 | <0.0001 | 25.33 | 0.0128 |
| 4% male vs 96% male | 12 | 12 | -15.33 | 0.6539 | -26.33 | 0.0079 | 15.67 | 0.5883 |
| 10% male vs 90% male | 12 | 12 | -18.67 | 0.2104 | -23.33 | 0.032 | -5.667 | >0.9999 |
| 10% male vs 96% male | 12 | 12 | -31.67 | 0.0005 | -33 | 0.0002 | -15.33 | 0.6539 |
| 90% male vs 96% male | 12 | 12 | -13 | >0.9999 | -9.667 | >0.9999 | -9.667 | >0.9999 |
Nighttime sleep in all these groups looked very much like that of males, with a sharp drop in sleep during the last one-half of the night (Fig. 8A). To look at more skewed ratios farther from equivalence we monitored sleep in groups of 50 flies that were 4, 10, 90, and 96% male. Interestingly, 10% males in a female-dominant group could have a significant effect on sleep, driving it into a male-like pattern at the end of the night (Fig. 8C,D). These
results suggested that the sexual interactions play a role in regulating population sleep.

Previous studies on mixed-sex pairs of flies have shown that courtship, a behavior that can be driven by olfactory cues, likely plays an important role in increased nighttime locomotor activity (Fujii et al., 2007; Hanafusa et al., 2013). To determine whether the social interaction influences on sleep we have seen were mediated via olfactory input, we compared isolated animal and population sleep for the olfactory receptor mutant Orco2 (also known as Or83b2) with a w genetic background control line in normal LD. For individual flies (Fig. 9A–C), olfaction affects sleep similarly in males and females. During the day, flies with compromised olfaction slept less than w control flies. At night, the main effect of genotype (Orco2 mutant, w control) on total sleep was not significant (two-way ANOVA, $F_{(1,21)} = 0.4748, p = 0.4921$). Overall, these results suggest a mild sleep-promoting effect of olfaction on isolated flies.

In populations (Fig. 9D–G), Orco2 mutant generally slept more at night, suggesting a role for olfactory input in the suppression of sleep in populations. The biggest effects were seen in mixed-sex populations. Orco2 mutants failed to decrease nighttime sleep (Fig. 8F,G), suggesting the change in sleep in mixed-sex groups reflects olfaction-dependent, perhaps sexual or aggression-related, behavior. Interestingly, Orco2 mutants also lacked the late-night male-specific decrease in sleep seen (with varying magnitude; Zimmerman et al. 2012) in Canton S male popu-
lations (Fig. 2B) and w control male populations (Fig. 9E) supporting the idea that this might reflect an olfactory-driven male–male interaction like aggression (Liu et al., 2011).

Discussion
Mechanisms of sleep have been studied widely; however the effects of social context on the characteristics of sleep have not been systematically evaluated. In the studies we report here, we provide evidence that sleep occurs in populations of flies through assessment of sleep patterns and homeostasis. We find that sleep patterns in populations of flies are distinct from those of individual flies and that many of these differences are likely to be rooted in the effects of social interactions on sleep.

The utility of the fruit fly in the study of population sleep depends on whether a group of flies can synchronize their sleep/wake activity. In this respect we find that populations of flies behave in the same manner as individuals. They exhibit the same morning and evening activity peaks, as well as having a siesta during the light period and more sleep during the dark period. In addition, and consistent with previous observations (Hendricks et al., 2000), we found that populations of flies fall asleep quickly, within the first hour after lights off with both female and male populations having similar sleep onset (Fig. 2G). These phenomena are unaffected by population size or gender. Importantly, we also find that the quiescence we see in populations is under homeostatic control,
an important criterion for inactivity to be called “sleep” (Huber et al., 2004; Cirelli and Bushey, 2008). Sleep deprivation generated by mechanical shaking or by starvation induced recovery sleep in populations of flies (Figs. 3, 4).

A critical question for these studies is whether the population activity measurements we present quantitatively reflect sleep in individuals within the population. Population sleep has a number of characteristics which...
suggest that what we are measuring is rooted in the sleep behavior of the individuals in the group; i.e., it is packaged as circadian clock-regulated periods of immobility and it is homeostatically regulated. But the quantitative relationship of population and individual measurements is more complicated. All the metrics previously used to characterize sleep are based on measurements of the behavior of individuals and we are measuring the activity of the population as a whole. What we can say about LAM25H data is that when the population is "asleep", every individual by

Figure 6. Populations of flies sleep better with complete food. Sleep profiles for populations of females (A) and males (B) on different food. C, Quantification of total sleep. Both female and male populations of flies slept significantly longer when on standard fly food compared with sucrose during the day, but there was no statistically significant difference during the night. D, Activity while awake. Complete food significantly increased activity levels during daytime wake periods in males, and at night in females. E, Number of sleep episodes. F, Sleep bout length. Females, but not males, had significantly consolidated sleep at night, i.e., fewer but longer sleep episodes. G, Latency. Females fell asleep faster on the complete food than on the sucrose agar food, whereas males exhibited similar latency on both food media. n = 8 groups for all conditions. ZT, Zeitgeber time; F, female; M, male.
definition must also be because there are no beam breaks. What we cannot determine is what waking activity in the population means: is one fly active or all the flies active? Video recording of populations did not reveal any obvious cases of single flies “driving” wake activity in wild-type populations (data not shown) but this would have to be more rigorously examined for other genotypes to rule it out completely.

There is also a strong caveat to interpretation of sleep architecture data from populations of flies. Sleep fragmentation is usually assessed in individual animals by looking at the number and length of sleep bouts. In population data, a sleep episode is a period during which all the flies are inactive, but an episode can be terminated by the activity of a single fly. This means that the bout duration in population data reflects a minimum value for the sleep of individuals in that population. Together, this implies that population measurements are likely to be underestimates of the sleep of individuals in that population and that the interpretation of sleep structure measurements is necessarily different.

To really look quantitatively at the effects of being in a group on an individual’s sleep pattern, one would need a system where the activity of individuals could be monitored in the context of the population, but for the densities of flies we are looking at here and the geometry of the arena (a vial), that is likely to be quite difficult even with currently available tracking software (Branson et al., 2009; Swierczek et al., 2011; Ardekani et al., 2012). Most of these tracking systems are based on capture of two-dimensional images which distinguish subjects from the background and determine their path. This would be dif-

Figure 7. Sleep is affected by population size and social behavior. A, Sleep profiles for female populations of different sizes. B, Sleep profiles for populations of males and females (1:1 ratio of sexes). C, Quantification of total sleep. Total sleep was decreased significantly with increasing number of flies and mixed populations with the same number of total flies exhibit lower sleep than populations of female flies. D, Activity while awake. Increasing the number of flies increases population activity. E, Number of sleep episodes. The number of episodes scales with population size in opposite directions for female only and mixed populations. F, Sleep latency does not change significantly with population size. n = 5–6 groups for all conditions. ZT, Zeitgeber time.
developed a 3D movement-tracking system by Wu et al., 2008), which found increased total sleep. In a previous study (Keene et al., 2010), we observed that populations of flies with reports of hyperarousability to other stimuli (Cirelli et al., 2004; Wu et al., 2008). A previous study suggested amn plays a major role on sleep architecture (Liu et al., 2008). In our study, populations of amn7 mutants exhibited fragmented sleep at night with significantly increased number of sleep episodes and an overall reduced amount of sleep, similar to individual flies (Fig. 5). In addition, we observed populations of amn7 flies had a novel locomotor phenotype with hyperactivity during the day and hypoactivity at night. We speculate that this may reflect a role for amn in social interactions that has not been previously reported, but is consistent with reports of hyperarousability to other stimuli (Wolf et al., 2002).

Population sleep measurements also are able to detect the suppressing effects of starvation on sleep similar to those that have been previously reported. Starvation has been shown to induce sleep loss during both the day and night in individual females but only at night in individual male flies (Keene et al., 2010). Interestingly, in our study, male flies housed in populations had an immediate decrease in sleep (Fig. 6B), whereas female populations only suppressed sleep after 12 h (Fig. 6A), suggesting that sleep suppression is sexually dimorphic, perhaps dependent on metabolic effects of starvation, social interaction or possibly survival competition within a group. In contrast to a previous study which found no effects of starvation on sleep homeostasis in individual flies (Thimgan et al., 2010), we found a robust sleep rebound in populations after starvation-induced sleep loss, providing a potential method to investigate homeostatic regulation by feeding state. We can also capture effects of food quality on sleep in populations. Although a previous study suggested sufficient caloric intake with no amino acids was able to support normal levels of sleep in individuals (Keene et al., 2010), we observed that populations of flies housed with sucrose food had less sleep than those given complete food. Housing in populations may exaggerate an effect that was undetectable in individuals, or increased activity of populations may generate more metabolic need. In contrast to a previous study (Zimmerman et al., 2012), which found increased total sleep/consolidated sleep when flies were switched to sucrose-based food from molasses- or dextrose-based food, we observed the opposite phenotypes in female populations. This might be due to differences in genetic background, mating status, or social context. Interestingly, we observed that sucrose food did not influence sleep architecture in male populations, whereas females robustly increased the number of sleep episodes and reduced sleep episode length during the dark period. In a previous study (Linford et al., 2012), diet was
shown to alter sleep architecture such that both individual male and female flies exhibited an increasing number of shorter sleep episodes with a low dietary sugar (5%) compared to higher sugar medium. Our results support the idea that the nutritional environment has an impact on sleep behavior, but suggest that these effects are sex-specific in a group context. The ability of population sleep measurements to find similar effects of mutations and nutritional state on sleep as have been reported for individuals supports the idea that they represent sleep at least at a qualitative level.

The quantitative differences between population and individual sleep are likely a function of social behaviors. Social interactions have been demonstrated to influence rhythmicity in humans (Stern and McClintock, 1998), rodents (Mrosovsky, 1988), bees (Toma et al., 2000), as well as in flies (Levine et al., 2002; Schneider et al., 2012). In Drosophila, it has been shown that the clocks of group-housed individuals are more synchronized than animals that have been isolated (Levine et al., 2002). In our study, we also observed that populations of flies (regardless of whether they were male, female, or mixed) synchronized sleep onset very quickly, within 1.5 h of lights off (Fig. 7F). The synchronized sleep/wake pattern in populations of flies may be due to the same volatile chemical signals which synchronize locomotor activity (Levine et al., 2002; Lone and Sharma, 2011), though the ability of Orco2 mutant flies to show similar synchronization suggests that these volatile signals may be detected by ORCO-independent olfactory pathways or that other cues can also be used to synchronize behavior.

There are also likely to be direct courtship- and mating-related effects on sleep. Courtship activity has been shown to be higher during the night and morning (Sakai and Ishida, 2001; Tauber et al., 2003; Hanafusa et al., 2013) and male flies play a dominant role in nocturnal locomotor activity in pairs of flies (Fujii et al., 2007). Male sex peptide, which is transferred to females when they mate, can inhibit female daytime sleep (Isaac et al., 2010).

Figure 9. Olfactory input modulates sleep amount by influencing social interactions. Sleep profiles of individual female (A) or male (B) flies. C, Quantification of data. Orco2 mutants slept significantly less than w controls during the day, and male Orco2 mutants slept longer than w males at night. No significant difference was detected between individual female Orco2 mutants and w. n = 30–32 for all genotypes. Population sleep profiles for female (D), male (E), and 1:1 mixed-sex populations (F). G, Quantification of data. Total sleep in populations of Orco2 mutant flies was similar to w controls within the male and female groups during the day and night. However, mixed female and male populations of Orco2 mutants exhibited drastically elevated sleep compared with w controls during the night. n = 5–6 groups for all genotypes. ZT, Zeitgeber time; F, female; M, male.
Reproduction-specific roles in regulating group dynamics could be a critical biological function in ecological contexts. Our sleep data showing drastically decreased sleep in the last half of the night in mixed populations also suggest the possibility that this is a time window during which mating occurs.

Many studies have shown that olfaction is a major driver of social interactions in flies (Levine et al., 2002; Fujii et al., 2007; Lone and Sharma, 2012). In this study, we find that populations of Orco2 mutant flies have no difference in total sleep compared with wild-type populations during the day, suggesting that other sensory input can compensate for the loss of olfaction for the synchronization of daytime activity. At night, however, loss of olfactory input, especially in male–female mixed populations, increases sleep time. This suggests that social interactions regulated by olfaction, such as courtship and aggression (Wang and Anderson, 2010; Liu et al., 2011; Dweck et al., 2015) specifically affect nighttime sleep. Although other social stimuli (sight, sound, mechanosensation, etc.) could also influence nighttime behavior, olfaction appears to have an important role.

In summary, we find that sleep can be measured in populations of flies, but the characteristics of population sleep vary from those of individual sleep. Some of the differences are due to technical considerations, e.g., the interpretation of sleep structure metrics from populations, but other reflect major effects of social interactions on sleep. Our demonstration of sex-specific and olfaction-related changes in sleep will provide interesting new avenues for understanding social behavior.

References
Andretic R, Shaw PJ (2005) Essentials of sleep recordings in Drosophila: moving beyond sleep time. Methods Enzymol 393:759-772. CrossRef Medline
Ardekani R, Huang YM, Sanchetti P, Stanciuskaus R, Tavaré S, Tower J (2012) Using GFP video to track 3D movement and conditional gene expression in free-moving flies. PLoS One 7:e40506. CrossRef Medline
Branson K, Robie AA, Bender J, Perona P, Dickinson MH (2009) High-throughput ethomics in large groups of Drosophila. Nat Methods 6: 451-457. CrossRef Medline
Brent MM, Oster II (1974) Nutritional substitution: a new approach to microbial control for Drosophila cultures. Dro Inf Ser 155-157.
Chi M W, Griffith LC, Vecsey CG (2014) Larval population density alters adult sleep in wild-type Drosophila melanogaster but not in amnesiac mutant flies. Brain Sci 4: 453-470. CrossRef Medline
Cirelli C, Bushey D (2008) Sleep and wakefulness in Drosophila melanogaster. Ann N Y Acad Sci 1129:323-329. CrossRef Medline
Cirelli C, Gutierrez CM, Tononi G (2004) Extensive and divergent sleep in Drosophila. Curr Biol 14:896-902. CrossRef Medline
Donelson NC, Donelson N, Kim EZ, Slawson JB, Vecsey CG, Huber R, Griffith LC (2012) High-resolution positional tracking for long-term analysis of Drosophila sleep and locomotion using the “tracker” program. PLoS One 7:e37250. CrossRef Medline
Dweck HK, Brahman SA, Thoma M, Mohamed AA, Keesy IW, Trona F, Lavista-Llanos S, Svatko A, Sachse S, Knaden M, Hansson BS(2015) Pheromones mediating copulation and attraction in Drosophila. Proc Natl Acad Sci U S A 112:E2829-35. CrossRef Medline
Fujii S, Krishnan P, Hardin P, Amrein H (2007) Nocturnal male sex drive in Drosophila. Curr Biol 17:244-251. CrossRef Medline
Hanafusa S, Kawaguchi T, Umezaki Y, Tomioka K, Yoshii T (2013) Sexual interactions influence the molecular oscillations in DN1 pacemaker neurons in Drosophila melanogaster. PLoS One 8:e84495. CrossRef Medline
Hay DA (1973) Effects of genetic variation and culture conditions on the social behavior of Drosophila melanogaster. Behav Genet 3:107-119. Medline
Hendricks JC, Finn SM, Panckera KA, Chavkin J, Williams JA, Sehgal A, Pack AI (2000) Rest in Drosophila is a sleep-like state. Neuron 25:129-138. Medline
Huber R, Hill SL, Holloway C, Biesiadecki M, Tononi G, Cirelli C (2004) Sleep homeostasis in Drosophila melanogaster. Sleep 27: 628-639. Medline
Isaac RE, Li C, Leedale AE, Shirras AD (2010) Drosophila male sex peptide inhibits siesta sleep and promotes locomotor activity in the post-mated female. Proc Biol Sci 277:65-70. CrossRef Medline
Keene AC, Duboué ER, McDonald DM, Dus M, Suh GS, Waddell S, Blau J (2010) Clock and cycle limit starvation-induced sleep loss in Drosophila. Curr Biol 20:1209-1215. CrossRef Medline
Krupp JJ, Kent C, Billette JC, Azanchi R, So AK, Schonfeld JA, Smith BP, Lucas C, Levine JD (2008) Social experience modifies pheromone expression and mating behavior in male Drosophila melanogaster. Curr Biol 18:1373-1383. CrossRef Medline
Larsson MC, Domingos AI, Jones WD, Chiappe ME, Amrein H, Vosshall LB (2004) Or83b encodes a broadly expressed odorant receptor essential for Drosophila olfaction. Neuron 43:703-714. CrossRef Medline
Levine JD, Funes P, Dowse HB, Hall JC (2002) Resetting the circadian clock by social experience in Drosophila melanogaster. Science 298:2010-2012. CrossRef Medline
Linford NJ, Chan TP, Fletcher SD (2012) Re-patterning sleep architecture in Drosophila through gustatory perception and nutritional quality. PLoS Genet 8:e1002668. CrossRef Medline
Liu W, Guo F, Lu B, Guo A (2008) Amnesiac regulates sleep onset and maintenance in Drosophila melanogaster. Biochem Biophys Res Commun 372:798-803. CrossRef Medline
Liu W, Liang X, Gong J, Yang Z, Zhang YH, Zhang JX, Rao Y (2011) Social regulation of aggression by pheromonal activation of Or65a olfactory neurons in Drosophila. Nat Neurosci 14:896-902. CrossRef Medline
Lone SR, Sharma VK (2011) Social synchronization of circadian locomotor activity rhythm in the fruit fly Drosophila melanogaster. J Exp Biol 214:3742-3750. CrossRef Medline
Lone SR, Sharma VK (2012) Or47b receptor neurons mediate sociosexual interactions in the fruit fly Drosophila melanogaster. J Biol Rhythms 27:107-116. CrossRef Medline
Mrosovsky N (1988) Phase response curves for social entrainment. J Comp Physiol A 162:35-46. Medline
Sakai T, Ishida N (2001) Circadian rhythms of female mating activity governed by clock genes in Drosophila. Proc Natl Acad Sci U S A 98:9221-9225. CrossRef Medline
Schneider J, Dickinson MH, Levine JD (2012) Social structures depend on innate determinants and chemosensory processing in Drosophila. Proc Natl Acad Sci U S A 109:17174-17179. CrossRef Medline
Shaw J, Cirelli C, Greenspan RJ, Tononi G (2000) Correlates of sleep and waking in Drosophila melanogaster. Science 287:1834-1837. Medline
Stern K, McClintock MK (1998) Regulation of ovulation by human pheromones. Nature 392:177-179. CrossRef Medline
Swierzczek NA, Giles AC, Rankin CH, Ken RA (2011) High-throughput behavioral analysis in C. elegans: Nat Methods 8:592-598. CrossRef Medline
Tauber E, Roe H, Costa R, Hennessey JM, Kyriacou CP (2003) Temporal mating isolation driven by a behavioral gene in Drosophila. Curr Biol 13:140-145. Medline
Thimgan MS, Suzuki Y, Seugnet L, Gottschalk L, Shaw PJ (2010) The periplin homologue, lipid storage droplet 2, regulates sleep ho-
meostasis and prevents learning impairments following sleep loss. PLoS Biol 8:e1000466. CrossRef
Toma DP, Bloch G, Moore D, Robinson GE (2000) Changes in period mRNA levels in the brain and division of labor in honey bee colonies. Proc Natl Acad Sci U S A 97:6914-6919. Medline
Wang L, Anderson DJ (2010) Identification of an aggression-promoting pheromone and its receptor neurons in Drosophila. Nature 463:227-231. CrossRef Medline
Wolf FW, Rodan AR, Tsai LT, Heberlein U (2002) High-resolution analysis of ethanol-induced locomotor stimulation in Drosophila. J Neurosci 22:11035-44. Medline
Wu MN, Koh K, Yue Z, Joiner WJ, Sehgal A (2008) A genetic screen for sleep and circadian mutants reveals mechanisms underlying regulation of sleep in Drosophila. Sleep 31:465-472. Medline
Wu Z, Gong Z, Feng C, Guo A (2000) An emergent mechanism of selective visual attention in Drosophila. Biol Cybern 82:61-68. Medline
Zimmerman JE, Chan MT, Jackson N, Maislin G, Pack AI (2012) Genetic background has a major impact on differences in sleep resulting from environmental influences in Drosophila. Sleep 35: 545-557. CrossRef Medline