Neurophysiological and behavioral synchronization in group-living and sleeping mice

Graphical abstract

Mice seek physical contact prior to sleep initiation and sleep in close proximity to conspecifics. Somatolonging—a motivation for prolonged physical contact—drives huddling behavior. Co-sleeping individuals show coordination in multiple neurophysiological features, including REMS timing synchronizes among male siblings, but not female or unfamiliar mice.

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In brief
Sotelo et al. demonstrate that mice seek to huddle prior to sleep initiation and that sleep huddling is a motivated behavior. They further reveal social-context-dependent synchronization in multiple neurophysiological features among co-sleeping individuals, including sleep and wakefulness onset times, NREMS intensity, and REMS timing.

Highlights
- Mice seek physical contact before sleep initiation and huddle during their sleep.
- Somatolonging—a motivation for prolonged physical contact—drives huddling behavior.
- Co-sleeping individuals show coordination in multiple neurophysiological features.
- REMS timing synchronizes among male siblings, but not female or unfamiliar mice.
Neurophysiological and behavioral synchronization in group-living and sleeping mice

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SUMMARY

Social interactions profoundly influence animal development, physiology, and behavior. Yet, how sleep—a central behavioral and neurophysiological process—is modulated by social interactions is poorly understood. Here, we characterized sleep behavior and neurophysiology in freely moving and co-living mice under different social conditions. We utilized wireless neurophysiological devices to simultaneously record multiple individuals within a group for 24 h, alongside video acquisition. We first demonstrated that mice seek physical contact before sleep initiation and sleep while in close proximity to each other (hereafter, “huddling”). To determine whether huddling during sleep is a motivated behavior, we devised a novel behavioral apparatus allowing mice to choose whether to sleep in close proximity to a conspecific or in solitude, under different environmental conditions. We also applied a deep-learning-based approach to classify huddling behavior. We demonstrate that mice are willing to forgo their preferred sleep location, even under thermoneutral conditions, to gain access to social contact during sleep. This strongly suggests that the motivation for prolonged physical contact—which we term somatolonging—drives huddling behavior. We then characterized sleep architecture under different social conditions and uncovered a social-dependent modulation of sleep. We also revealed coordination in multiple neurophysiological features among co-sleeping individuals, including in the timing of falling asleep and waking up and non-rapid eye movement sleep (NREMS) intensity. Notably, the timing of rapid eye movement sleep (REMS) was synchronized among co-sleeping male siblings but not co-sleeping female or unfamiliar mice. Our findings provide novel insights into the motivation for physical contact and the extent of social-dependent plasticity in sleep.

INTRODUCTION

Social interactions play a crucial role in animal life, significantly affecting the development, physiology, and behavior of individuals.1 Consequently, social isolation in various species is associated with negative health outcomes, including compromised immunity and mood dysregulation.2–5 However, the extent to which social contact modulates sleep—a central behavioral and neurophysiological process6—remains unclear. This is mainly because sleep has traditionally been studied in experimental subjects that are individually isolated and deprived of their natural ecological and social milieu.7,8

Studies have suggested that sleep quality and need are socially modulated in diverse species.9,10 For instance, honeybees and fruit flies sleep more following an awake period with social interactions than following an awake period without social interactions.11 In baboons, 3D acceleration tracking suggests that

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in biologging technologies, which permit continuous neurophysiological monitoring across days from multiple untethered individuals in semi-natural and wild conditions, have emerged as a breakthrough in the capacity to study sleep in naturalistic environments. Indeed, the recent use of these technologies has revealed that sleep neurophysiology and behavior are plastic and modulated by internal and ecological demands.

Various species, including humans, baboons, gulls, and laboratory mice, exhibit coordination in the timing of sleep and wake episodes between co-living individuals. Recent studies have also uncovered coordination in brain oscillatory patterns between awake interacting individuals in humans, bats, and mice. However, there is still a significant knowledge gap regarding the extent of this coordination to brain oscillatory patterns during sleep.

Here, we utilized innovative wireless neurophysiological recording devices to investigate the sleep behavior and neurophysiology of freely moving mice living together under various social conditions. We reveal that, prior to sleep initiation, mice actively seek physical contact and sleep while in close proximity to each other, a behavior we refer to as huddling. Employing a novel preference assay and deep-learning-based huddling classification, we demonstrate that huddling during sleep is a motivated behavior. We also uncover social-dependent changes in sleep architecture and coordination in sleep oscillatory patterns among co-sleeping individuals. We reveal that, although social sleeping fragments non-rapid eye movement sleep (NREMS), it synchronizes rapid eye movement sleep (REMS) episodes. Notably, REMS synchronization was specific to male sibling mice and not observed in female or unfamiliar male mice, suggesting a dependence on both sex and social familiarity. Our findings further suggest that the internal state of an animal plays a crucial role in coordinating REMS and oscillatory neurophysiological activity and expand current understanding of the extent to which social factors can modulate sleep.

RESULTS

Mice huddle prior to sleep initiation

To examine the influence of the presence of conspecifics on behavioral preparation for sleep, we recorded same-sex triplet sibling adult wild-type (CD-1) mice in their home-cage environment before and after transitioning into the light phase (Figure 1A). We identified eight behavioral states (“in-motion,” “transiently socially interacting,” “eating/drinking,” “allo-grooming,” “self-grooming,” “nesting,” “resting,” and “sleeping”) and noted whether these behaviors were displayed while in close physical proximity to conspecifics—hereafter referred to as huddling—or not (Figure 1B). We compared the amount of time spent in each behavior within two time windows: 40–20 and 20–0 min before sleep onset (Figures 1C and 1D), informed by our recent identification of the 20-min period preceding sleep initiation as a pre-sleep transitional phase. We performed a second-by-second annotation of the videos for one individual from eight triplets (n = 4 females and 4 males). Four behavioral states (self-grooming, nesting, resting, and sleeping) were displayed both while huddling and not-huddling during the 40-min period prior to sleep onset. While not-huddling, mice spent significantly less time in-motion and transiently socially interacting as sleep approached yet did not alter time spent eating/drinking, self-grooming, nesting, and resting (Figure 1C; data not shown for eating/drinking). Conversely, while huddling, mice spent significantly more time self-grooming, nesting, and resting as sleep approached, but not allo-grooming (Figure 1D). Overall, the proportion of behaviors performed while not-huddling decreased, and those performed while huddling increased, as sleep onset approached (Figure 1E). These findings imply that mice seek physical contact in the pre-sleep phase and engage in pre-sleep behaviors such as nesting and grooming, while huddling.

Huddling before and during sleep is driven by somatolonging—A motivation for physical contact

We next aimed to examine whether the observed increase in huddling prior to sleep was driven by a motivation for physical contact or due to difficulty sleeping alone in the shared home-cage environment. To facilitate a high throughput analysis of huddling behavior, we employed a deep-learning-based approach for huddling classification (Figures 2A and S1A–S1G). First, we performed pose estimation via DeepLabCut (DLC) to identify and track mice. We annotated two back keypoints for each mouse, which remained predominately visible even when most of the mouse’s body was occluded by a conspecific or nesting material (Figures 2A and S1A). Next, we computed the minimum Euclidean distance between the mouse’s keypoints (Figure 2A). Finally, we classified huddling instances when the minimum distance fell below a pre-set threshold indicating physical contact (5 cm; Figure S1F; Table 1) and which lasted over a pre-set duration threshold for huddling (1 s, excluding transient social interactions) (Figures 2A and S1G).

Subsequently, we devised a novel behavioral assay, which we named the “immersive social interactions assay (ISIA),” to investigate whether the increase in huddling prior to sleep signifies a motivation for physical contact. The ISIA apparatus consists of two home-cage chambers connected by a plastic tube, whose inner diameter can be reduced by the insertion of a plastic ring (Figure S2A). We studied same-sex pairs of adult sibling mice (n = 8 pairs). In each pair, one mouse was implanted with a 6-pin connector (hereafter referred to as “head-bar”), while the other underwent a sham operation. On the first day of the experiment, both mice were allowed free movement between chambers, and they consistently spent most of the ensuing light phase in one chamber, where they built their nest and slept (Figures 2B, 2C, S2B, and S2C; Video S1). On the second day, at the end of the dark/active phase, a plastic ring was inserted into the tube connecting the chambers to restrict the head-bar mouse to the least preferred chamber, while the test mouse remained free to move between the chambers. During the following light phase, test mice spent significantly less time in their preferred chamber and more time in their least preferred chamber (Figure 2C; Video S1), indicating that mice are willing to sacrifice their preferred sleep location for physical contact during sleep. Additionally, the proportion of time spent huddling during the light phase did not decrease under the restriction (Figure 2D), suggesting that the increase in huddling prior to sleep is driven by a motivation for physical contact rather than by difficulty sleeping alone in the shared home-cage environment.

Given that the laboratory environment falls below the thermoneutral zone for mice, we wondered whether the observed...
increase in huddling behavior might be due to mice seeking physical contact as a means to mitigate the cold. We thus repeated the previous experiment, but in this instance, we modified the ISIA setup to utilize two different chambers: one kept at room temperature, henceforth referred to as the “cold chamber” and another maintained at the thermoneutral zone for mice (30°C), referred to as the “warm chamber” (n = 8 pairs of sibling mice; Figures 2E, S2D, and S2E). Under baseline conditions, all mice spent significantly more time in the warm chamber during the light phase, where they constructed their nests and slept (Figures 2F and S2E; Video S2). However, when the head-bar mouse was confined to the cold chamber, the test mice spent significantly less time in the warm chamber and more time in the cold chamber (Figure 2F; Video S2). Intriguingly, the time spent huddling did not decrease but rather slightly increased when the head-bar mouse was restricted to the cold chamber.
Figure 2. Huddling behavior is an expression of a motivation for physical contact

(A) Schematics illustrating the pipeline for pose estimation, coordinate data filtering, and huddling classification.

(B) Schematics detailing the 2-day ISIA procedure conducted under room temperature conditions, accompanied by representative data. The top row illustrates day 1 under freely moving conditions, whereas the bottom row details day 2 under sibling restriction conditions. Left: schematic illustrating the ISIA procedure. Middle: representative heatmaps depicting the locations of a head-bar fitted mouse and a test mouse during the 12-h light cycle in the ISIA apparatus (log transformed histogram counts). Right: snapshots from the DLC tracking output depicting predicted keypoints, the mice, and their nest. Note that both the heatmaps and snapshots originate from the same recording sessions.

(C) Percent time (mean ± SE) spent in each of the ISIA chambers during the light phase, measured under room temperature conditions, across the 2 days of the experiment. n = 8 pairs of female and male mice, repeated-measures (RM) two-way ANOVA (p(day) = 0.03, p(chamber) = 0.57, p(interaction) < 0.0001) followed by Sidak’s multiple comparisons test (****p < 0.0001).

(D) Percent time (mean ± SE) spent huddling during the light phase, measured under room temperature conditions, across the 2 days of the experiment. n = 8 pairs of female and male mice, paired t test, two-tailed, t = 2.017, df = 7, p = 0.0836.

(E) Similar to that in (B), but with the ISIA procedure conducted under optional thermoneutrality conditions. This was achieved by maintaining the ambient temperature in one of the chambers at the mice’s thermoneutral zone (30°C).

(F) Percent time spent in each of the ISIA chambers during the light phase, measured under optional thermoneutrality conditions, across the 2 days of the experiment. n = 8 pairs of female and male mice, RM two-way ANOVA (p(day) = 0.16, p(chamber) = 0.9, p(interaction) < 0.0001), followed by Sidak’s multiple comparisons test (****p < 0.0001).

(G) Percent time spent huddling during the light phase, measured under optional thermoneutrality conditions, across the 2 days of the experiment. n = 8 pairs of female and male mice, paired t test, two-tailed, t = 2.281, df = 7, p = 0.0565. See also Figures S1 and S2, Table 1, and Videos S1 and S2.
Table 1. Classifier performance and evaluation

<table>
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<th>Distance (cm)</th>
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<th>Predictive values</th>
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<tr>
<td></td>
<td></td>
<td></td>
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<tr>
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<td></td>
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</tr>
<tr>
<td>6 cm</td>
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<td></td>
<td>4 cm</td>
<td>5 cm</td>
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<tr>
<td>F1 score</td>
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</table>

The table illustrates the confusion matrices generated for the three evaluation sets using three distinct thresholds for physical contact (4, 5, and 6 cm). It also presents corresponding metrics for accuracy, precision, recall, specificity, and F1 score for each of these distance thresholds.

See also Figures 2 and S1 and Videos S1 and S2.

(Sibling presence fragments NREMS) Next, we investigated the influence of sibling presence on the sleep/wake architecture in mice, employing wireless devices capable of monitoring neurophysiological signals in groups over extended periods (Neurologger 2A; Figure 3A). We examined the sleep/wake architecture of same-sex sibling adult mice over a 24-h period, both in groups and isolated settings (n = 9 males and 9 females from 3 triplets per sex; Figures S3B and 3C). Importantly, sibling mice were recorded simultaneously, either while cohabiting in the same home-cage or when housed in separate cages within the same experimental room. Sibling presence significantly altered the proportion of time spent in the different sleep/wake states in female, but not in male, mice (Figures 3D and 3E). Specifically, while with their siblings, females spent less time awake and more time asleep during the dark phase, compared with when individually isolated (Figure 3E). Notably, the presence of siblings significantly fragmented NREMS in both male and female mice, as evidenced by an increase in the number of NREMS episodes and a decrease in the average NREMS episode duration throughout most of the light and dark phases (Figures 3F–3I and S3A–S3D). Additionally, delta power during NREMS was significantly lower in group-housed mice compared with when individually isolated (Figure S3E). Collectively, these findings suggest that the presence of siblings adversely affects the quality of NREMS in mice.

Considering that our mice were fitted with wireless devices, we wondered whether the observed sleep fragmentation is due to the devices themselves rather than a genuine fragmentation induced by conspecifics. We thus compared the average duration of sleep bouts in mice with and without wireless devices. We first validated that video-based sleep scoring yielded a similar average duration of sleep episodes as EEG-EMG based scoring in mice equipped with wireless devices, recorded under both social and individual housing conditions (n = 6 mice; Figure S3F). Subsequently, employing video-based scoring, we compared the average sleep episode in mice with and without wireless devices under both individual and social conditions (Figure S3G). We found that the social context (isolation vs. social housing), but not the presence of wireless devices, significantly influenced sleep episode duration (n = 6 mice per group; Figure S3G). This finding further supports our premise that the presence of conspecifics, rather than the wireless devices, negatively affects sleep quality.

To quantify the proportion of huddling during sleep under group-housed conditions, we employed the deep-learning-based approach for huddling classification described above (Figures 2A, S1, and S3H) and aligned the huddling data with the sleep-state data. Both female and male mice huddled with their siblings during the majority of their NREMS and REMS epochs during the light phase (n = 9 mice per sex; Figure S3I). These findings suggest that while mice actively seek physical contact prior to sleep and continue huddling throughout, such close physical proximity to conspecifics leads to a fragmentation of NREMS.

Co-sleeping siblings show coordination of multiple neurophysiological features

To investigate the effects of group housing on brain states and oscillatory brain activity synchronization between animals (n = 18 male pairs and 14 female pairs), we first assessed the synchrony of sleep and wake onset times between pairs of mice, either cohabiting in the same home-cage or housed separately within the same experimental room. Both female and male mice displayed a significant degree of synchronization in the timing of sleep/wake onsets in group-housed conditions as compared with isolated settings (Figures 4A–4I). Notably, we found a positive correlation between the degree of synchronization in the timing of falling asleep and the degree of synchronization in the timing of waking up across mouse pairs, suggesting that pairs who typically wake-up together also tend to fall-asleep together (Figure 4J). Further, we detected a strong synchrony in the overall sleep and wake times when mice were cohabiting in the same home-cage compared with isolated settings and shuffled data (Figures 4K and 4M). Our cross-correlation analysis further revealed a significant coordination in sleep/wake state timing with a peak at zero lag time in group-housed but not in isolated settings (Figures 4L and 4N). Together, these findings suggest that sibling mice demonstrate strong coordination, with high temporal precision, of sleep and wake timing. We then examined whether, beyond the overall sleep/wake synchrony, co-sleeping mice showed synchrony in their sleep states (Figure 4O). Notably, male mice demonstrated significant
synchrony in the timing of REMS (Figures 4P and 4Q), whereas female mice exhibited minimal to no coordination (Figures 4R and 4S). Together, these findings suggest that social living promotes the coordination of sleep/wake states, with REMS coordination particularly pronounced in male sibling mice.

Lastly, we examined the synchronization of oscillatory brain activity between pairs of mice. Co-habiting male and female mice exhibited synchrony in the theta band power, but not the delta band, of the EEG during wakefulness (Figures 4T and 4U). Notably, during NREMS, only male mice demonstrated...
Figure 4. Sibling mice show coordination of multiple neurophysiological features while co-sleeping

(A) On the left, binary vectors representing wake (depicted in blue) and sleep (depicted in purple) states over a 24-h period for a pair of sibling mice. On the right, a magnified section of the vectors reveals the exact moments of wake onset (depicted in green) and sleep onset (depicted in red) for the two mice.
Synchronization in the power of the delta band, whereas female mice did not (Figure 4V); however, both males and females showed synchronization in the power of the theta band during NREMS (Figure 4W). Collectively, our data unraveled a significant degree of coordination in multiple neurophysiological features among co-sleeping individuals, with the extent of coordination being particularly pronounced in male sibling mice.

Social factors modulate sleep/wake architecture
We examined the impact of non-familial social interactions on sleep/wake architecture by introducing intruder mice (female or male) into a resident male’s territory, studying both male-female and male-male pairings (Figure 5A). Anticipating significant immediate behavioral changes in both resident and intruder mice, we conducted neurophysiological analysis on the fourth day post-introduction—when sexual and agonistic behaviors had significantly subsided (n = 8 male-female and 8 male-male dyads; Figures 5B, 5A, and 5B). We first investigated the propensity of non-familial mice to huddle during sleep. Not only male-female dyads but also male-male dyads huddled for most of their sleep epochs on the fourth day of cohabitation (Figure 5C), even when the home-cages contained two shelters (Figure S4C). When analyzing the sleep/wake architecture of resident and intruder mice (n = 9 male-female and 9 male-male dyads; Figures 5D–5I and S5A–S5F), we compared, for each mouse, data from non-familial dyadic experiments with data collected when the mouse was with its siblings. This comparison enabled us to pinpoint changes attributable specifically to non-familial interactions rather than to the mere presence of conspecifics. Male residents, when in the presence of a male intruder, displayed minimal alterations in their sleep/wake architecture (Figures 5D–5I and S5A–S5F). The primary change observed was a consolidation of NREMS during the light phase, marked by fewer yet lengthier NREMS bouts compared with when they were with their siblings (Figure S5B). Conversely, when a female intruder was present, male residents exhibited an increase in wakefulness and a decrease in REMS during the light phase compared with when with their siblings (Figures 5D and 5F). Notably, the proportion of REMS in male residents while in the presence of a female matched that of their female counterparts (Figure S5G). The surge in wakefulness in male residents was accompanied with its consolidation, characterized by fewer but longer episodes (Figure S5A), while the decrease in REMS was primarily attributed to fewer episodes rather than alteration in the duration of REMS bouts (Figure S5C). However, during the dark phase when in the presence of a female intruder, the male residents showed no significant alterations in their sleep/wake architecture (Figures 5G–5I and S5D–S5F).

Unlike resident mice, male intruders displayed significant sleep/wake architecture changes during the dark, but not the light, phase (Figures 5D–5I and S5A–S5F). Male intruders spent more time awake and less time asleep during the dark phase compared with when with siblings (Figures 5G–5I). Wake episodes were more consolidated (Figure S5D), while sleep bouts were not shorter but less frequent (Figures S5E and S5F). Notably, female intruders showed minimal to no alterations in their sleep/wake architecture when housed with a male resident, compared with when with siblings (Figures 5D–5I and S5A–S5F). Together, our results suggest that, for females, the presence of a social partner influences sleep more than its familiarity or sex. In contrast, for males, the prevailing social context plays a pivotal role in their sleep/wake architecture.
Figure 5. Sex and social-context dependent modulation of sleep
(A) Schematic illustrating experimental paradigm.
(B) Total time dedicated to sexual behaviors in male-female dyads (left; n = 8 mice) and agonistic behaviors in male-male dyads (right; n = 8 mice) during zeitgeber time (ZT) 23–0 immediately following the placement of an intruder mouse into the home-cage of a resident mouse on days 1 and 4 later. Sexual behaviors include “seeking,” “mounting without lordosis,” and “mounting with lordosis.” Agonistic behaviors include “mounting,” “attacking,” “chasing,” “aggressive grooming,” and “fleeing.” Behavioral data were obtained from resident mice. Paired t tests, two-tailed; p(sexual behaviors) = 0.0003; p(agonistic behaviors) = 0.048.
(C) Percentage of time spent huddling during NREMS and REMS in the light phase on the fourth day following cohabitation for male-female dyads (left) and male-male dyads (right). n = 9 mice per group.
(D–I) Sleep/wake architecture across the light (left) and dark (right) cycle in resident and intruder mice compared with the architecture of the same mice while cohabiting with their siblings.
role. Our findings highlight the nuanced interplay between sex and context in social modulations of sleep.

**Social context modulates the degree of neurophysiological coordination among mice**

Finally, we explored the impact of non-familial social interactions on the synchronization of brain states and oscillatory brain activity among mouse pairs (n = 9 male-female and 10 male-male dyads). We compared synchrony in mice cohabiting the same home-cage with that of the same mice when housed separately (Figure 6). We discovered a significant sleep and wake onset synchronization among both male-female and male-male dyads, compared with individually housed (Figures 6A–6H). Moreover, dyads waking up together often fell asleep together (Figure 6I). Notably, the degree of synchronization in waking up and falling asleep times among dyads surpassed that observed in sibling mice, particularly among males (Figures 6J and 6K). Furthermore, we identified robust synchrony in overall sleep and wake times among dyads compared with solitary housing and shuffled data (Figures 6L and 6N). Our cross-correlation analysis further revealed a significant coordination in sleep/wake state timing with a peak at zero lag time among dyads, a finding not observed in solitary housing (Figures 6M and 6O). Collectively, these findings provide compelling evidence of robust coordination in the timing of sleep and wakefulness among non-familial mouse dyads.

Surprisingly, when we examined the synchrony in sleep states, we found no coordination in REMS epochs for either male-female or male-male dyads (Figures 6P–6S). We also observed minimal synchronization of oscillatory brain activity between male-female and male-male dyads (Figures 6T–6W). Only male-male dyads demonstrated significant synchronization in the delta power band during wakefulness (Figure 6T) and theta power band during NREMS (Figure 6W), relative to solitary housing. Taken together, these findings strongly suggest that neurophysiological coordination among mice is modulated by social context and sex.

**DISCUSSION**

To assess the impact of the social environment on sleep, we employed state-of-the-art wireless neurophysiological devices for extended monitoring of multiple freely moving individuals within a group, alongside detailed behavioral recordings and deep-learning-based behavioral classification. We reveal that mice seek physical contact prior to sleep initiation and opt to sleep in close proximity to each other. Using a novel behavioral assay, we establish huddling during sleep as a motivated behavior. We also find that conspecific presence fragments NREMS, whereas co-sleeping animals show neurophysiological coordination across various measures. Notably, we found a synchronization in REMS timing among co-sleeping male siblings but not female or unfamiliar mice. Lastly, we highlight a social context-dependent modulation of sleep/wake architecture. Our research offers significant insights into the ethological context of sleep and further our understanding of sleep physiology. Considering the prevalence of sleep disturbances and social stress, including loneliness, understanding the impact of social factors on sleep dynamics is essential. Gaining insights into these behavioral and neurophysiological interactions could inform strategies to mitigate sleep disturbances related to social stress, potentially enhancing human well-being.

Social factors influence both the quality and quantity of sleep across diverse species. For instance, in insects, social interactions increase subsequent sleep need (based on behavioral indices), while in baboons, prior social interactions and social status modulate sleep duration and intensity (based on 3D acceleration data). Laboratory studies utilizing brief recordings or focusing on single individuals within a group further substantiate the premise that context and social status influence sleep. However, until now, much remained unknown about how round-the-clock social dynamics and mating-related interactions affect sleep, as well as the extent to which social factors impact neuronal oscillatory activity during sleep. Our study provides novel insights into the complex interplay between sleep and social dynamics, through 24-h-long neurophysiological recordings from multiple subjects while monitoring the same individuals in diverse social contexts. These contexts span from cohabitation with siblings to interactions with non-familial individuals, including mates and same-sex conspecifics.

Our study reveals that sleep/wake patterns are modulated by both sex and social context. Female mice shift their sleep timing when cohabiting with their siblings, sleeping more during the active/dark phase compared with when alone. The presence of a male mouse does not further alter the sleep/wake states of female mice. This suggests that they are more sensitive to the presence or absence of a sleep partner, rather than its familiarity or sex. In contrast to females, male mice display sleep/wake modulation that is dependent on social context. The presence of siblings or a male intruder does not significantly alter sleep/wake duration or timing in males compared with when isolated. However, their social status plays a pivotal role. Intruding into...
Figure 6. Social context modulates the degree and nature of neurophysiological coordination among mice

(A–H) Synchrony of sleep and wake onset times between pairs of mice, either cohabiting in the same home-cage or housed separately.

(A) Heatmap representation of the cross-correlation of sleep onset times for each male-female dyad, whether housed together (right) or separately (left).
another male’s territory increases wakefulness and reduces NREMS, especially during the dark/active phase. Mating-related interactions also alter sleep-wake states in male mice but mainly during the light/inactive phase. In the presence of a female, males stay awake longer and spend less time in REMS during the light phase, aligning their REMS duration with their female counterparts. Although increased wakefulness could be due to greater opportunities to engage in motivated behaviors, the specific mechanisms and causes behind the decrease in REMS, but not NREMS, remain to be elucidated. In summary, our findings underscore the complex relationship between social factors and sleep regulation, highlighting the importance of considering both social context and sex when investigating sleep dynamics.

Our study provides novel insights into the coordination of neurobehavioral states and neurophysiological oscillatory activity during sleep among co-sleeping individuals. We identified synchronization in sleep/wake onset times, duration of sleep and wakefulness, and REMS timing. The strong synchronization in sleep/wake onset times, coupled with shorter and more frequent NREMS bouts, suggests that co-sleeping mice disturb each other’s sleep. Conspecific presence has been reported to disrupt sleep in several species. For instance, insomnia can be transferred between human bed partners,29 sleep duration in baboons is reduced when in close proximity to group-mates,13 and collective waves of awakenings have been observed in gulls.20 Given the well-established relationship between the consolidation of sleep and its health benefits,30,31 it is somewhat perplexing that animals willingly choose to sleep in situations that appear to compromise their sleep. Social living is recognized as beneficial in numerous ways,1,32–35 suggesting that this type of sleep fragmentation may not be as detrimental as other forms, such as those triggered by stress. Alternatively, the benefits of group sleeping may outweigh its costs. These unresolved possibilities provide promising avenues for future research, particularly concerning any protective mechanisms that might be involved.

As opposed to the synchronization in sleep/wake onset times, the mechanisms underlying REMS synchronization remain elusive. Our data do not suggest a simple correlation between synchronized sleep initiation and REMS synchronization because both female and male dyads exhibited strong synchronization in sleep/wake onsets but not in REMS timing. These findings imply that, while REMS timing can be synchronized among co-sleepers, this process is gated by an individual’s internal state.

Although the adaptive value of REMS synchronization remains unclear, several potential benefits can be postulated. The synchronization might provide thermal benefits as thermoregulation is suppressed during REMS3 and could help mitigate the fragmentation of REMS caused by conspecific movements. Furthermore, by synchronizing a period of reduced responsiveness, animals might decrease the risk of conspecifics accessing shared resources or posing harm. Further research is needed to elucidate these potential benefits and reveal why females and unfamiliar mice do not exhibit REMS synchronization.

We also discovered synchronized oscillatory brain activity in social-living mice that was gated by the social context. Specifically, sibling mice exhibited strong theta power synchronization during both wakefulness and NREMS, whereas non-sibling mice showed weak to no synchronization. Synchronization in oscillatory activity during wakefulness has been reported in various species, including humans, bats, and mice.21–23 These studies propose that such coordination is not reliant on shared task engagement or physical contact but is rather linked to a shared cognitive space, facilitating social communication.36 Given the fundamental differences in cognitive processes between wakefulness and sleep, it is likely that diverse mechanisms oversee synchronization during these states.

Various physiological processes, including cardiac and respiratory activities, could underlie the synchronized timing of REMS and oscillatory brain activity during sleep. Although responsiveness to

<table>
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<tr>
<th>(A)</th>
<th>Average cross-correlations of sleep onset times from all male-female dyads, either housed together or separately (mean ± SE). Shuffle test, p &lt; 0.0001.</th>
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<td>(B)</td>
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<td>(K)</td>
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<td>Average cross-correlations of REMS (male-female) either housed together or separately (mean ± SE). Wilcoxon rank-sum test, p = 0.0019.</td>
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ties, 53–57 enhancing understanding of the motivation for physical contact. Nonetheless, much remains to be revealed about the decision-making processes that initiate and maintain huddling, as well as the sensory modalities and circuitry triggering the behavior. Deepening our understanding of the behavioral-, neurophysiological-, and circuit-level facets of the motivation for social contact could provide novel insights into the regulation of natural, ethologically relevant behaviors. Such understanding could also shed light on how social interactions influence health and well-being in humans and other species.

We have devised a novel behavioral apparatus, the ISIA, that presents several key features significantly enhancing current capacity to conduct comprehensive, dynamic, and ethologically relevant assessment of animal behavior within a neuroscience context (whether in its current two-chamber design or a modified N-chamber version). First, our apparatus permits dynamic interactions between freely moving individuals, enabling a full spectrum display of natural behaviors. This facet is typically missing in commonly used tests, such as the three-chamber social interaction test, which simplifies social behaviors to parameters such as proximity to other individuals and sniffing time. 58 Second, it offers animals the autonomy to either engage in or avoid social activities by choosing whether to enter or avoid a chamber. This feature introduces a new layer of complexity to the study of social behaviors such as aggression and sexual behaviors, where traditionally, animals are “forced” into a situation rather than given a choice to participate or not. Third, in addition to allowing the placement of animals in novel testing cages, our apparatus enables the use of the animal’s home-cage. This feature may reveal behaviors not typically exhibited in a new environment, it grants animals the choice to leave their territory or not, and it eliminates the need to manipulate animals to participate in the test—eliminating the confounding effect of handling stress. Lastly, because our apparatus employs modified standard housing cages, it is accessible and cost effective for widespread use. The integration of our apparatus with deep-learning methodologies for the detection and classification of behaviors in a social context will further support the high throughput, yet ethologically relevant and nuanced, study of animal behavior in the lab.

STAR METHODS

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SUPPLEMENTAL INFORMATION

Supplemental information can be found online at https://doi.org/10.1016/j.cub.2023.11.065.

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AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

INCLUSION AND DIVERSITY

We support inclusive, diverse, and equitable conduct of research.

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KEY RESOURCES TABLE

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Experimental models: Organisms/strains

Mouse: CD-1 Charles River Laboratories 022

Software and algorithms

Adobe Illustrator/ Premiere Pro Adobe Creative Cloud https://www.adobe.com/ |
BORIS Friard and Gamba59 https://www.boris.unito.it/ |
iSpy iSpy https://www.ispyconnect.com/ |
Prism 9.5 Graphpad software https://www.graphpad.com/ |
SleepSign for Animal 3.0 Kissei Comtec http://www.sleepsign.com/ |
BioRender BioRender https://biorender.com |

RESOURCE AVAILABILITY

Lead contact
Further information and requests for resources should be directed to and will be fulfilled by the lead contact, Dr. Ada Eban-Rothschild (adae@umich.edu).

Materials availability
This study did not generate new unique reagents.

Data and code availability

- All original neurophysiological data reported in this paper will be available via Zenodo. 3D printer files will be available via github. DOIs are listed in the key resources table.
- All codes will be publicly available via github.
- Any additional information required to reanalyze the data reported in this paper is available from the lead contact, Ada Eban-Rothschild adae@umich.edu, upon request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

We utilized CD-1 mice (Strain #: 022, Charles River Laboratories) of both sexes that were reproductively inexperienced unless otherwise specified. The mice were bred in-house and were between 7–10 weeks old at the beginning of the experimental procedures. They were housed in a controlled environment with a 12-hour light/dark cycle at a temperature of 22 ± 1°C, and were provided with nesting material consisting of 4-6 grams of compressed cotton ("Nestlet"; Ancare, Bellmore, NY, U.S.A) and 1 gram of shredded paper ("Enviro-Dri"; Shepherd Specialty Papers, Watertown, TN, U.S.A) as well as ad libitum access to food and water. All experiments were conducted in...
METHOD DETAILS

Surgery
The mice were anesthetized with a ketamine-xylazine mixture (100 and 10 mg kg\(^{-1}\), respectively; administered via intraperitoneal injection, IP) and received lidocaine and carprofen (4 mg kg\(^{-1}\) and 5 mg kg\(^{-1}\), respectively) prior to being placed in a stereotaxic frame (David Kopf Instruments, Tujunga, CA, U.S.A.). During the surgery, the mice were kept under isoflurane anesthesia (~1% in O\(_2\)). To enable individual identification and tracking, all mice, except for two male triplets (see below), were dyed with either Electric Tiger Lily (orange), Blue Moon (blue), or Electric Lizard (green) dyes from Tish & Snooky’s Manic Panic (manicpanic.com), as in Shemesh et al.\(^{60}\)

EEG-EMG implantation
Mice were surgically fitted with five miniature screws. Two screws serving as reference electrodes and placed on the right hemisphere (frontal: \(\text{AP} = 1.5\ mm\), \(\text{ML} = -1.5\ mm\), parietal: \(\text{AP} = -3.5\ mm\), \(\text{ML} = -2.8\ mm\)) and one screw serving as a ground electrode (\(\text{AP} = -5.9\ mm\), \(\text{ML} = 0\)) and two additional screws serving as support and were placed over the left hemisphere. Two EMG electrodes were inserted between the trapezius muscles. All electrodes were pre-soldered to a 7-pin connector and secured to the skull using either C&B Metabond (Parkell) and dental cement or OptiBond Universal adhesive and Revolution flowable light-cured composite (Kerr). Mice were given a minimum of 10 days for recovery before initiating experimental habituation.

Head-bar implantation
Mice underwent surgery to secure a 6-pin connector (10 mm (L) x 2 mm (W) x 10 mm (H), referred to as ‘head-bar’) to the skull using two support screws (\(\text{AP} = 1.5\ mm\), \(\text{ML} = 1.5\ mm\) and \(\text{AP} = -3.5\ mm\), \(\text{ML} = -2.8\ mm\)) and either C&B Metabond (Parkell) and dental cement or OptiBond Universal adhesive and Revolution flowable light-cured composite (Kerr). Sibling mice of head-bar implanted mice underwent a sham operation in which their skin was opened at the top of the skull and closed with surgical sutures. Mice were allowed at least 5 days of recovery before experimental habituation.

Video and neurophysiological acquisition
Triplet same-sex sibling mice were implanted with EEG-EMG electrodes and placed in custom plexiglass chambers (50 cm x 25 cm x 40 cm). Mice were habituated to dummy wireless neurophysiological devices, which were of similar size and weight as the genuine devices, for over a week. Video data was collected via USB cameras (either Logitech C920 or Angetube 1080p webcams) mounted above the chambers, using iSpy software (ISpyConnect.com). Neurophysiological data was collected at 400 Hz by Neurologger 2A devices (Evolocus, Tarrytown, NY, USA) powered by two 1.45 V batteries (#312, Renata Batteries). The loggers were equipped with a 3D accelerometer and an infrared (IR)-receiver micro-board, and IR pulses (Neurologger Synchronizer, Evolocus) were used to synchronize recordings from multiple devices mounted onto different mice. To protect the loggers from damage, they were covered with parafilm and a custom 3D printed perforated plastic enclosure (Figure 2A), resulting in a final combined weight of 3 gr (which accounts for approximately 8% and 9% of the body weight for CD-1 adult male and female mice, respectively). The loggers were mounted onto the experimental mice 2-4 hrs prior to the initiation of recordings. The neurophysiological recordings were performed in mice that had inhabited their chambers for at least two days, and simultaneous video and neurophysiological data were collected for 24 consecutive hours, starting from Zeitgeber time (ZT) 0 (‘Lights on’).

We recorded mice under four different sequential conditions: 1) same-sex triplet sibling, 2) individual isolation, 3) non-sibling stranger male dyads, and 4) opposite-sex dyads. Conditions 1-3 were carried out while sexually inexperienced mice were used. For Condition 2, mice were individually housed for three days before the experiment. Simultaneous 24-hour long EEG-EMG and video recordings were carried out for mice from the same triplet while they were individually housed in adjacent cages in the same room. For Conditions 3 and 4, mice were individually housed for approximately one week prior to the experiment. In Condition 3, experiments were conducted in the home-cage of a “resident” mouse, into which a male “intruder” was placed at the end of the dark phase (ZT 23). In a subset of experiments (6/9), the “resident”’ home-cage contained two shelters (one: 8 x 5 x 5 cm and the second: 5 x 5 x 5 cm; Figure S4C). In Condition 4, experiments were conducted in the home-cage of a male “resident” mouse into which a female “intruder” mouse at the proestrus/estrus phase of the reproduction cycle was introduced at the end of the dark phase (ZT 23). Only video data was collected on Day 1 for Conditions 3 and 4, while both video and neurophysiological data were collected on Day 4.

Estrus cycle determination
To determine the estrus cycle phase, we conducted vaginal cytology by flushing vaginal cells with saline using a pipette, collecting the liquid onto a glass slide, and examining it under an epifluorescence microscope (Zeiss Axio Imager.M2). Females were tested once a day at ZT 22 for up to 5 days until nucleated epithelial cells and/or cornified cells were identified, indicating the proestrus/estrus phase.

Immersive social interactions assay (ISIA)
The ISIA apparatus consisted of two chambers (each 27 (L) x 16 (W) x 25 (H) cm), which were equipped with video cameras and connected by a perforated plastic tube (Length: 13.5 mm; Inner diameter: 3.7 mm; Figure S2A). To restrict the movement of mice with head-bars, a
plastic ring (Inner diameter: 2.5 cm; Length: 1.5 cm) was inserted into the tube, reducing its diameter. This modification ensured that mice with head-bars were confined to one of the chambers, while mice without head-bars could move freely between the two chambers. Ad libitum food and water, and nesting material were provided in both chambers of the apparatus. Two successive experiments were conducted, each lasting two days. In Experiment 1, both chambers were maintained at room temperature (21.4 ± 0.01°C). In Experiment 2, one chamber was kept at room temperature (23.8 ± 0.01°C) while the other was at the thermoneutral zone for mice (29.9 ± 0.01°C). To maintain a temperature of 30°C in the latter chamber, an infrared ceramic heat lamp (Wuhostamp, Black bulb, 100W) was placed above it. Temperature was continuously monitored via iButton devices (one measurement / 10 min; DS1925, Maxim Integrated) placed on the inner walls of the chambers (14.5 cm above chamber floor) during the experiments.

Pairs of head-bar and sham-operated sibling WT CD-1 mice (4 pairs of females and 4 pairs of males) were placed in the apparatus one day before the experiments began. One hour prior to the recordings (at ZT 23) on each experimental day, nesting material was divided between the two chambers. On Day 1, both mice were free to move between chambers. During the following light phase, mice built their nests and slept mostly in one chamber (Figures 2B and 2C). On Day 2, one hour prior to the recording (at ZT 23), the plastic ring was inserted into the tube, restricting the movement of mice with head-bars to the least preferred chamber defined as the one where mice spent less than 50% of the time during the light phase on Day 1. Video recordings were made during the light phase (ZT 0–12) on all experimental days. It is important to note that only mice from litters with fewer than 9 pups were used for the experiments.

To generate a unified video that incorporated the two chambers of the ISIA, each originally captured by a separate video camera, we merged the individual recordings using Adobe Premiere Pro (Adobe Creative Cloud). The original videos were resized to standardize the chambers dimensions, temporally synchronized, and subsequently converted to a .3gp file format, at a resolution of 352 x 288 pixels.

**Polysomnographic analysis**

We down-sampled the neurophysiological data to 256 Hz using custom MATLAB code. We digitally filtered the EEG/EMG signal and then performed a fast Fourier transformation analysis using SleepSign 3.0 for Animal software (Kissei Comtec America). The EEG was filtered between 0.3–25 Hz and the EMG was filtered between 25–50 Hz. We classified arousal states (wakefulness, NREMS, and REMS) during 4-sec time windows. An automatic analysis was first performed based on EEG power spectra, EEG integral, and EMG integral, followed by visual inspection and correction when necessary. Wakefulness was defined as desynchronized, low-amplitude EEG signal occurring concomitantly with high tonic EMG signal with phasic bursts. NREMS was defined as synchronized, high-amplitude, low-frequency (0.5–4 Hz; delta band) EEG signal with reduced EMG activity compared to wakefulness. REMS was defined as EEG signal with a dominant theta band (6–9 Hz) with very low EMG activity. The scoring was performed by investigators (M.I.S. and T.K.) who were blind to the experimental manipulation. EEG power spectrum data were expressed as values relative to the average total power during wakefulness at ZT 12–15 of each recording. Due to EEG artifacts, we excluded the data of two, five, and four mice, respectively, from the analyses presented in Figures 4, 6, and S3H.

**Sleep/wake synchronization**

To quantify sleep/wake synchronization between pairs of mice, we transformed their sleep scoring vectors into binary vectors. These vectors indicated whether each mouse was awake or awake in each time bin over a 24-hour period. We then counted the number of time bins where both mice were asleep. To evaluate the extent of sleep/wake synchronization, we employed a shuffle test. In this procedure, the binary sleep/wake vectors of two mice underwent circular shifting 500 times by a random quantity. With each iteration, we counted the number of time bins where both shuffled vectors indicated sleep. This process allowed us to derive a distribution of sleep/wake synchronization values based on random alterations of the original data. The synchronization ratio between two mice (Figures 4K, 4M, 6L, and 6N) was calculated by dividing the sleep/wake synchronization value from the original data by the average sleep/wake synchronization value derived from the shuffled data.

**Cross-correlation analyses**

To further explore the temporal profile of sleep/wake synchronization, we implemented a complementary approach to the one outlined above. We computed the cross-correlation of binary sleep/wake vectors for each pair of animals, normalizing the cross-correlation vectors so that autocorrelations equaled one at zero lag (Figures 4L, 4N, 6M, and 6O). Similarly, cross-correlations were derived for shuffled vectors as well. To determine the significance of the cross-correlation, we derived a p-value by computing the proportion of shuffled cross-correlations that exhibited a peak surpassing the peak of the original data. To test whether the peak cross-correlation of group-housed mice was higher than that of isolated mice, we utilized a shuffling procedure. We calculated the difference between the peaks of the average cross-correlations for the two groups. Next, we randomly mixed cross-correlation vectors from the group-housed and isolated pairs 100,000 times, and computed the differences between the average groups of cross-correlations. The p-value for the difference between the original mean cross-correlations was obtained by calculating the proportion of differences from the shuffled data that exceeded the difference of the original data.

To compute the cross-correlation of ‘falling asleep’ and ‘waking up’ (Figures 4C, 4E, 4G, 4I, 6B, 6D, 6F, and 6H), we utilized a procedure akin to that detailed above for sleep/wake cross-correlation. However, in this case, the cross-correlated vectors included ‘1’s in bins where an animal transitioned from wakefulness to sleep (for the ‘falling asleep’ analysis) and from sleep to wakefulness (for the ‘waking up’ analysis), while ‘0’s were placed in all other time bins.

For the computation of REMS synchronization and cross-correlation (Figures 4P–4S), we utilized a similar method to the one outlined above for sleep/wake synchronization. The key difference was that the vectors incorporated ‘1’s for bins during which the animal was in the REMS, and ‘0’s for bins when the animal was in NREMS. Notably, bins in which either animal was awake were omitted. 

Please cite this article in press as: Sotelo et al., Neurophysiological and behavioral synchronization in group-living and sleeping mice, Current Biology (2023), https://doi.org/10.1016/j.cub.2023.11.065
from both vectors prior to the REMS synchronization quantification. Thus, this analysis specifically quantified REMS synchronization within the sleep period.

**EEG power synchronization**

The power of the EEG at the delta and theta bands (detailed above) was computed by filtering the EEG within the specific bands and taking the envelope of the squared filtered signal. To assess the synchronization of the EEG power in the delta and theta bands within a specific arousal state (Figures 4T–4W and 6T–6W), we first identified and concatenated all time bins during which both animals were in a specific arousal state, excluding any time bins in which either animal was not in that state. Subsequently, we calculated the Pearson correlation of the EEG power within the specific band across these time periods.

**Manual behavioral analysis**

We annotated mice behavior in detail using the Behavioral Observation Research Interactive Software (BORIS)

**Pre-sleep behavior**

We began by identifying the sleep initiation time for each mouse closest to ZT 0 and preceded by a 1-hour period of wakefulness. We then analyzed 50 minutes of data for each mouse, starting 40 minutes prior to sleep initiation and ending 10 minutes after. We annotated nine different behaviors: 1) "In-motion," where the mouse was actively moving, including walking, running, rearing, or digging; 2) "Eating/Drinking," where the mouse was holding or consuming food pellets or feces with its forepaws or where the mouse was licking the nozzle of the water bottle spout; 3) "Transient social interactions," where the mouse was touching another mouse with its snout or chasing/fleeing from another mouse; 4) "Self grooming," where the mouse was licking its fur, stroking its body with its forepaws, or scratching itself with any limb; 5) "Allo-grooming," where the mouse was grooming the body or head of another mouse or being groomed by another mouse; 6) "Nesting," where the mouse was engaged in pulling, carrying, fraying, push-digging, sorting or fluffing of nesting material; 7) "Resting," where the mouse was awake and alert, either lying or sitting inside the nest with its head lifted. Occasional movements were present, and it was distinguished from sleeping by the lifted head; 8) "Sleeping," where the mouse was motionless in a sleeping posture (lying curled up or sitting with its face tucked into its body) for ≥ 8 consecutive seconds. Brief twitches (<1 second long) may be present. We also recorded whether behaviors were performed in close physical proximity to a conspecific (‘huddling’) or otherwise (‘non-huddling’). Huddling was classified for incidents during which the bodies of two mice were in contact, resulting in the lack of visible background between the two bodies, for > 1 second. Mice bodies included the abdomen and thorax but not the head, legs, paws, and tail.

**Sleep/wake states**

We manually classified sleep/wake states during the ZT 5–6 time window—a period during which mice are predominantly asleep—in 3 female and 3 male WT CD-1 mice per group (with and without wireless devices). We assigned the label "Wake" when a mouse was engaged in active behaviors such as eating, drinking, grooming, walking, nesting or digging, or when it was lying down with its head lifted or displaying body movement for ≥ 1 second. "Sleep" was assigned as described in the Pre-sleep behavior section.

**Sexual behavior**

We manually classified the behavior of resident male mice during ZT 23-0, immediately following the placement of a female intruder into their home-cage, and four days later. We annotated four different behaviors: 1) "Seeking": Mouse is approaching or sniffing conspecific; 2) "Mounting with lordosis": Mouse lays over the hind quarter of a conspecific with his two front paws over their back, while conspecific displays a downward arching of the back; 3) "Mounting without lordosis": Mouse lays over the hind quarter of a conspecific with his two front paws over their back, while conspecific displays a straightened back; 4) "Other": Mouse is not engaged in the aforementioned behaviors but may display non-sexual-related behaviors, including eating, drinking, grooming, nesting, resting and sleeping.

**Agonistic behavior**

We manually classified the behavior of resident male mice during ZT 23-0, immediately following the placement of a male intruder into their home-cage, and four days later. We annotated six different behaviors: 1) ‘Mounting’: mouse lays over the hind quarter of a conspecific with his two front paws over their back; 2) ‘Attacking’: mouse is running/leaping at a conspecific, and may bite; 3) ‘Chasing’: mouse is running towards a conspecific, while conspecific is running away, but not biting; 4) ‘Aggressive grooming’: mouse is allo-grooming a conspecific while the conspecific is rigid and motionless; 5) ‘Fleeing’: mouse is running away from a conspecific while the conspecific is running toward it, with no bites present; 6) ‘Other’: mouse is not engaged in the aforementioned behaviors, but may display non-aggressive-related behaviors, including eating, drinking, grooming, nesting, and sleeping.

**Markerless pose estimation**

We utilized DeepLabCut software (version 2.3.1) to identify and track individual mice body parts. For each mouse, we annotated two back keypoints. The first was located midway between the tail base and the ears, while the second was situated midway between the first keypoint and the ears. We selected these keypoints as they remained predominantly visible even when the majority of the mouse’s body was occluded by nesting material or a conspecific (Figure S1A). We generated and validated separate models for each of our datasets: (a) two-chambers containing two mice (ISIA data), (b) a single chamber containing three mice with wireless devices, and (c) a single chamber containing two mice with wireless devices.

For the ISIA video dataset, we annotated a total of 811 frames. Approximately 65% of these frames were randomly selected from the dataset, consisting of 32 videos across 16 mice from 8 different experiments. The remaining frames were manually chosen to

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represent instances that posed a tracking challenge. This includes frames depicting one or more mice in the connecting plastic tube, frames where the mice were entirely obscured, frames containing visually confusing objects outside the behavioral apparatus, and frames from video sections where a prior model underperformed (see below). For the datasets involving either two or three mice with wireless devices in a single chamber, we applied a similar strategy, utilizing both randomly selected images from the datasets and frames selected to represent challenging tracking scenarios. We annotated a total of 986 frames for the dataset with three mice and 841 frames for the dataset with two mice.

We employed a ResNet-50-based neural network with default parameters and trained each model for 250,000 iterations. To test the performance of our models on novel data, we evaluated each model using an evaluation set, which we generated by randomly sampling 20% of each dataset. This evaluation procedure was repeated five times for each model, with resampled evaluation sets for each run. We computed the mean average Euclidian error (MAE, which is proportional to the average root mean square error) between manual labels and the ones predicted by DLC. For the ISIA dataset, we recorded an average (± SE) training error of 3.934 ± 0.2 pixels and an average (± SE) testing error of 5.864 ± 0.19 pixels (the size of most images was 352 x 288 pixels). The finalized models, which were used to perform markerless pose estimation, incorporated all annotated frames from each dataset. An experimenter validated the accurate identification and tracking of keypoints across all videos at 2 to 64 times the normal playback speed. We progressively increased the size of the training set, specifically incorporating instances that posed tracking challenges, until no identification switches or tracking errors were detected.

For each recording, we also annotated the edge points of the chambers, which allowed us to standardize the coordinate data across all trials (Figure S1B). This step was necessary due to varying distances between the cameras and the cages in different recordings. Accordingly, we transformed the DLC output coordinates so that a consistent real-world distance (1 mm) corresponded to the same pixel distance (1 unit) both within and across datasets (Figure S1C).

Huddling classification
We started by filtering the coordinate data, excluding frames where the coordinates either fell outside the edge points of the chambers or had a likelihood value equal to or below 0.1 (Figures 2A and S1C). The latter typically corresponded to instances where the mice were visually undetectable, due to being completely occluded by nesting material, conspecifics, or tucked away inside the tube (Figures S1D and S1E).

We then computed, for every frame, the closest proximity (minimum Euclidian distance) between each of the two keypoints of each mouse. We classified framed by comparing this distance to a pre-set hyper-parameter threshold value for physical contact (5 cm; see below). We first classified frames into two categories: those in which the minimum Euclidian distance was equal to or below the preset threshold, designated as ‘physical contact,’ and those in which the distance exceeded the threshold, designated as ‘no physical contact.’ If the ‘physical contact’ state persisted for more than a second, we classified the frames as instances of ‘huddling.’ If not, they were considered ‘non-huddling’ (Figure S1G).

To establish the hyper-parameter threshold value for physical contact, we initially generated a validation set by randomly sampling frames from our entire dataset. We manually excluded frames where the outlines of the mice bodies were occluded, which hindered a clear determination of whether the mice were touching or not. Subsequently, we manually classified the images as depicting either ‘physical contact’ or ‘no physical contact,’ using the previously described criteria (see Pre-sleep behavior section), resulting in 935 ‘physical contact’ frames and 1178 ‘no physical contact’ frames. We plotted a histogram of the minimum Euclidean distances for the two classified states (Figure S1F). This visualization highlighted 5 cm as an optimal threshold value for physical contact. To assess the chosen hyper-parameter threshold, we generated three test sets by randomly re-sampling our dataset (excluding images used for training), testing three distinct threshold values for physical contact: 4 cm, 5 cm, and 6 cm. Once again, we excluded frames where the outlines of the mice were obscured, yielding 1988, 2037, and 2020 frames for the 4 cm, 5 cm, and 6 cm thresholds, respectively. We computed false and true positives, along with Accuracy (number of correct predictions divided by dataset size), Precision (True Positives divided by the sum of True Positives and False Positives), Recall (True Positives divided by the sum of True Positives and False Negatives), Specificity (True Negatives divided by the sum of True Negatives and False Positives), and F1 score (2 multiplied by ([Precision multiplied by Recall] divided by the sum of Precision and Recall)) for the three distance values (Table 1). This evaluation procedure further affirmed our selection of 5 cm as the threshold value for physical contact.

QUANTIFICATION AND STATISTICAL ANALYSIS
Data and statistical analyses were conducted using SleepSign for Animals (3.0), MATLAB (2020a), and Prism 9.5.1 (GraphPad Software). The sample sizes were analogous to those reported in previous studies.61 The data analysis was carried out blind to the experimental conditions. While we assumed data distribution was normal, this wasn’t formally tested in every instance. For comparing the data of two paired groups, we employed Wilcoxon signed-rank tests or paired t-tests. In cases involving more than two groups, we utilized the Friedman test or one-way repeated-measures (RM) ANOVA. For multiple comparisons, we used RM two-way ANOVA or, for datasets with missing values (e.g., absence of a particular sleep state at a specific time point), we utilized mixed-effects model ANOVA. Post-hoc tests were conducted using Sidak’s multiple comparisons tests. The exact P values and n values for the statistical tests are detailed in the figure legends. Adobe Illustrator 2023 (Adobe Creative Cloud) was used for figure preparation.