Biogenic action of *Lactobacillus plantarum* SBT2227 promotes sleep in *Drosophila melanogaster*

**Highlights**
- *Lactobacillus plantarum* SBT2227 promotes sleep at the onset of nighttime.
- Existing intestinal microbes do not affect the SBT2227 sleep effect.
- Heat-stable intracellular/intramembrane components are candidates for active substances.
- Neuropeptide F is required for the sleep-promoting effect of SBT2227.

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SUMMARY
Lactic acid bacteria (LAB) influence multiple aspects of host brain function via the production of active metabolites in the gut, which is known as the pre/probiotic action. However, little is known about the biogenic effects of LAB on host brain function. Here, we reported that the Lactobacillus plantarum SBT2227 promoted sleep in Drosophila melanogaster. Administration of SBT2227 primarily increased the amount of sleep and decreased sleep latency at the beginning of night-time. The sleep-promoting effects of SBT2227 were independent of the existing gut flora. Furthermore, heat treatment or mechanical crushing of SBT2227 did not suppress the sleep-promoting effects, indicative of biogenic action. Transcriptome analysis and RNAi mini-screening for gut-derived peptide hormones revealed the requirement of neuropeptide F, a homolog of the mammalian neuropeptide Y, for the action of SBT2227. These biogenic effects of SBT2227 on the host sleep provide new insights into the interaction between the brain and gut bacteria.

INTRODUCTION
Nutrients obtained from food sources are not only necessary for growth and sustenance but also for various vital functions and, in some cases, for improving adaptability (Milton, 2003; Watanabe et al., 2019). In general, fermentation increases the rate of nutrient absorption and improves the nutritional value of foods (Kärland et al., 2020; Sharma et al., 2020). Animals prefer fermented food and consume the bacterial population responsible for fermentation along with it. Therefore, part of the intestinal microbiota is constructed by bacteria consumed (Pasolli et al., 2020).

From studies in mammals such as rodents and humans, intestinal bacteria are known to improve host health via their probiotic functions such as the activation of immunity (Shi et al., 2017) and anti-obesity action (Barthikannan et al., 2019), and by supplying nutrients (Yadav et al., 2018), as well as via prebiotic functions that regulate the balance of intestinal bacteria (Gibson et al., 2017; Hill et al., 2014). In addition to these various biological functions, studies in rodent models have shown that intestinal bacteria can affect higher-order brain functions, such as anxiety (Peric and Alvina, 2019), depression (Peric and Alvina, 2019), and cognitive functions (Chu et al., 2019), via communication with the brain-gut axis (Grenham et al., 2011; Mayer et al., 2014; Peric and Alviña, 2019). In the past decade, the probiotic and prebiotic functions of these intestinal bacteria have attracted considerable attention, and the causative factors and mechanisms have been elucidated (Erny et al., 2015; Lyte, 2014; Strandwitz, 2018; van de Wouw et al., 2018). Furthermore, functional substrates produced by bacteria may direct biological effects irrespective of the presence of intestinal flora, a concept known as biogenics (Mitsuoka, 2000). For example, biogenic actions of bacterial substrates have been shown to reduce body fat in humans (Nakamura et al., 2016) and, in rodents, to prevent cancer (Badr El-Din et al., 2020; Zhang et al., 2013), hypertension (Chanson-Rolle et al., 2015; Cheng and Pan, 2017) and inflammation (Yang et al., 2015, 2017). Furthermore, in vitro model systems have suggested that lipid metabolism (Monagas et al., 2010) and the immune system (Kobatke and Kabuki, 2019) are modulated by bacterial substrates. Compared to probiotics and prebiotics, knowledge regarding the effects of biogenics on higher-brain functions is limited.

Recently, the fruit fly Drosophila melanogaster, a model organism with a simple brain and nervous system, is being increasingly used for studying the brain-gut-microbiome axis (Broderick and Lemaitre, 2012; Wong et al., 2011). The intestinal flora of the fly is a miniature version of that of mammals, with Lactobacillus and Acetobacter being the main bacterial species (Erkosar et al., 2013). Furthermore, similarities between flies...
and mammals have already been shown in host-microbe interactions, such as in the immune system and metabolic pathways (Buchon et al., 2014; Wong et al., 2016). Indeed, studies on the gut microbiota of D. melanogaster have revealed probiotic effects on brain functions such as locomotion (Schretter et al., 2018), foraging (Leitão-Gonçalves et al., 2017; Wong et al., 2017), odor-guided egg laying (Qiao et al., 2019), and food preference (Venu et al., 2014). These evidence indicated that D. melanogaster, with its compact brain-gut axis structure, can be used as an experimental system for studying the biogenic functions of digested bacteria using the many sophisticated molecular genetic tools that are currently available.

Sleep is one of the behaviors regulated by higher-order brain functions. Behavioral genetics of D. melanogaster have revealed genes and neural mechanisms that regulate sleep, most of which are shared with mammals (Hendricks et al., 2000; Ly et al., 2018; Sehgal and Mignot, 2011; Shaw et al., 2000). A recent study showed that the absence of commensal bacteria or axenic conditions affects sleep in flies negligibly (Jia et al., 2021; Selkrig et al., 2018). This indicates that probiotics and prebiotics in the gut microbiota do not significantly affect fly sleep, although evidence for biogenics is inconclusive so far. This gap in our knowledge of the biogenic effects could be filled by a detailed analysis of the sleep behavior of flies following the oral administration of a particular lactic acid bacteria (LAB) strain. To achieve this, we used an L. plantarum strain, SBT2227, and demonstrated for the first time that the biogenic effect of the LAB can facilitate fly sleep. Furthermore, transcriptome analysis suggested that the endocrine system is a potential target for the action of the biogenic actions. The results of a mini-screening using flies with gene knockdowns suggested the involvement of a specific peptide hormone, neuropeptide F, in the biogenic action of SBT2227. These findings provide a novel perspective on the brain-gut-microbe interaction and open up new avenues for the functional use of LAB.

RESULTS
Effects of oral administration of SBT2227 on fly sleep
L. plantarum is a Gram-positive LAB commonly found in plants and fermented food (Guidone et al., 2014; Mundt and Hammer, 1968) and is also known to be one of the major intestinal bacteria of D. melanogaster (Chandler et al., 2011; Cox and Gilmore, 2007). As fruit flies may ingest L. plantarum from fermented fruits in the field, it is highly possible that this lactobacillus species influences the biological activities of flies. Therefore, we decided to investigate the biogenic effects of SBT2227, a strain of L. plantarum, on the sleep of fruit flies.

The sleep pattern of flies orally administered SBT2227 (1% w/v in food) was compared to that of untreated control flies (Figure 1A). There was no significant difference in fly consumption between the SBT2227 food and the control food (Figure S1). Flies in both groups showed typical sleep patterns reported previously: napping in the middle of the subjective daytime and sleeping mostly during subjective night-time (Andretic and Shaw, 2005; Hendricks et al., 2000; Shaw et al., 2000). Daytime sleep was significantly reduced on the first day of SBT2227 treatment, although this was not sustained after the second day (Figure 1B). By contrast, obvious change was not detected in night-time sleep on the first day, although it increased significantly on the second and third day in SBT2227-fed flies (Figure 1C).

Based on the night-time sleep pattern on day 3 (Figure 1A), the sleep-promoting effect was preferentially detected at a specific time window (ZT12-15, zeitgeber time (ZT), where ZT0 = lights on and ZT12 = lights off, Figure 1D). By contrast, no obvious sleep effect was detected during ZT15-24 (Figure 1E). The sleep effect appeared in early night-time, presumably because of the shorter latency of sleep. Sleep latency was significantly reduced (Figure 1F). On the other hand, the sleep bout length and number of sleep bouts were not altered on day3 (Figures 1G and 1H). These effects of SBT2227 (increase in sleep amount during ZT12-15 and shortened sleep latency on the third day) were also conserved in another fly strain with a different genetic background (Figure S2). Furthermore, these sleep phenotypes were dose-dependent for SBT2227 (Figure S3). Therefore, we decided to use these two characteristics, sleep amount of ZT12-15 and sleep latency on the third day, as the main parameters for further analyses, as they are considered to show the typical effects of SBT2227.

Effects of oral administration of SBT2227 on the awakening state
Sleep and wakefulness are two sides of the same coin; as sleep increases, wakefulness decreases, and vice versa. However, this reciprocal relationship is not always true. For example, Fernandez-Chiappe et al. reported a case where flies did not increase their sleep when they reduced the activity level (Fernandez-Chiappe et al., 2020). On the other hand, Potdar and Sheeba reported another case where flies increased...
sleep without changing their activity level (Potdar and Sheeba, 2018). Therefore, we investigated the effects of SBT2227 on wakefulness. The activity of SBT2227-fed flies was examined over a 3-day period. The graph showing the activity patterns showed that the flies fed SBT2227 showed a clear decrease in their night-time activity after the second day (Figure 2A). Separate quantitative analysis of daytime and night-time activity counts showed the absence of any obvious change in the daytime; however, a significant decrease was detected at night-time in flies fed SBT2227 (Figures 2B and 2C). Reduction in activity counts may be attributed to the decrease in motor control. To assess whether SBT2227 disrupts motor ability, we examined the activity index, which is the number of activity counts per waking time. We did not detect any obvious change during the daytime, which is the main waking period of flies (Figure 2D). By contrast, we found a significant reduction in the activity index at night on day 3 (Figure 2E). As daytime locomotion was normal in flies fed SBT2227, the night-time arousal level on the third day was lower than that of the control; in other words, sleep was strongly promoted.

Commensal bacteria did not alter the sleep-promoting effect of SBT2227

To investigate whether the sleep-promoting effect of SBT2227 requires interaction with the commensal bacterial flora, the existing intestinal bacteria were disturbed with antibiotics (Storelli et al., 2018).

Figure 1. Oral administration of SBT2227 increased sleep at the onset of night-time and decreased sleep latency
(A) Sleep patterns of flies fed control food (red) or SBT2227 food (green). Sleep traces are presented as mean ± SEM. (B–H) Amount of sleep during daytime (ZT0-12); (C) amount of sleep during night-time (ZT12-24), amount of sleep at specific timing; (D) ZT12-15 and (E) ZT15-24. (F) Sleep latency, (G) sleep bout length, and (H) number of sleep bouts during night-time. n = 96 for each group. Wilcoxon-Mann-Whitney test was used for statistical analysis, adjusted with the Bonferroni correction. *p < 0.05, **p < 0.01, ***p < 0.001, n.s., not significant. See Table S2 for detailed statistics.
behavior was evaluated with or without SBT2227 treatment (Figure 3A). The number of aerobic bacteria in the antibiotic-treated flies (ABT flies) was reduced to approximately 1/100 compared to that in flies fed the control conventional food (CC flies) (Figure 3B). LAB, including *L. plantarum*, are anaerobic bacteria. Therefore, we determined the number of anaerobic bacteria present in the gut and found that anaerobic bacteria were almost eliminated in ABT flies (Figure 3B). When these post-ABT flies were fed SBT2227 for 2 days, the median number of anaerobic bacteria in the gut was $1.7 \times 10^3$ cfu/fly (Figure S4).

Significant differences were not observed between the sleeping behaviors of ABT and CC flies (Figures 3C and 3D). By contrast, the sleep-promoting effect of SBT2227 was observed in both ABT and CC flies (Figures 3E and 3F). The effect sizes of SBT2227 treatment on the sleep amount (ZT12-15) and sleep latency in CC and ABT flies were similar (Figures 3E and 3F, sleep amount effect size ($r$) = 0.35 and 0.29, and sleep latency effect size ($r$) = 0.31 and 0.34, respectively). These results suggested that intestinal commensal bacterial flora is not required for the action of SBT2227 on fly sleep.

The sleep-promoting effect of SBT2227 was sustained after protein denaturation

In order to clarify whether the SBT2227 needs to be alive to have the sleep-promoting effect, heat-killed SBT2227 was administrated to flies and evaluated its effect on sleep. SBT2227 was treated at 65°C for 1 h or autoclaved at 121°C for 15 min and then administered to flies. Compared to that in the control, the amount of sleep during ZT12-15 on day 3 increased significantly in all the SBT2227-fed groups (unheated, 65°C, or autoclaved) (Figure 4A). Similarly, compared to that of the control group, sleep latency decreased significantly in all SBT2227 treated groups (Figure 4B). These results suggested that SBT2227 does not need to be alive for the sleep-promoting effect. Moreover, protein denaturation in SBT2227
did not alter its sleep-promoting effect. The heat-stable substances in SBT2227 were possibly the biogenic factors required for the sleep-promoting effect.

Supernatant of crushed SBT2227 cells may contain active substances required for the sleep-promoting effect

To identify the SBT2227-derived components that exert biogenic effects on sleep, we crushed the bacteria and separated it into a supernatant rich in intracellular/membrane components and a precipitate rich in cell wall components. To verify the efficiency of crushing the bacteria, we performed Gram staining of the bacterial samples before and after crushing. Most of the bacteria were decolorized after crushing, and the cell walls were destroyed (Figure 5A). We then examined the sleep effects of SBT2227 under four different conditions (uncrushed, crushed, crushed supernatant, and crushed precipitate). The results showed that irrespective of the conditions of SBT2227, the amount of sleep in ZT12-15 on day 3 increased significantly, except in the case of the crushed precipitate (Figure 5B). Consistently, sleep latency decreased significantly in flies treated with uncrushed, crushed, or crushed supernatants of SBT2227, except for those treated with the precipitated fraction (Figure 5C). These results suggested that the intracellular/membrane components of SBT2227 may contain active substances required for the sleep-promoting effects.

Biological processes altered by SBT2227 administration

The gut is possibly a primary target of the action of the sleep-promoting substances of SBT2227, as it has abundant immune-responsive cells and enteroendocrine cells (EECs), and the nerve fibers connecting the brain or ventral nerve cord to the gut have been identified (Miguel-Alia et al., 2018). Hence, it is possible that the substances in SBT2227 are digested and absorbed and directly affect the brain, but it is also possible that the active substances or digested and absorbed substances act on the gut, and changes that occur in the gut affect the brain via immune, endocrine, and neural pathways. Therefore, we investigated the changes in gene expression in the gut with or without SBT2227 administration using RNA sequencing-based transcriptome analysis.

The intestines containing fore-, mid-, and hindgut of flies were harvested at ZT12-14, which acts as the main time window for the sleep-promoting effect, and the mRNA extracted was subjected to RNA sequencing.
The number of genes with more than one read detected in either the SBT2227-fed or SBT2227-unfed (control) group was 12,015 genes, of which 787 genes were identified as differentially expressed genes (DE-Gs) (Figure 6A, see also Table S1). Next, we conducted Gene Ontology (GO) enrichment analysis to identify the categories of molecular functions associated with these 787 genes. Multiple categories of significantly enriched DE-Gs were detected, including those associated with nucleosome/DNA binding, sodium ion transporter activity, neuropeptide receptor activity, and peptidoglycan muralytic activity (Figure 6B). Studies have shown that peptide hormones secreted by endocrine cells in the gut, such as CCHamide-1, act on brain neurons and regulate fly behaviors such as wakefulness (Titos and Rogulja, 2020). Therefore, among these GO categories, we focused on neuropeptide receptor activity and hypothesized that peptide hormones produced by EECs may mediate the sleep-promoting effects of SBT2227 and tested this possibility in subsequent experiments.

Neuropeptide F was required for the sleep-promoting effect of SBT2227

In total, 16 different peptide hormones have been reported to be expressed in EECs of D. melanogaster (Chen et al., 2016a). The expression of seven of these peptide hormones in the gut EECs, namely, allatostatin A (AstA), myoinhibiting peptide precursor (Mip), allatostatin C (AstC), CCHamide-1 (CCHA1), CCHamide-2 (CCHA2), diuretic hormone 31 (Dh31), and neuropeptide F (NPF), has been confirmed using in situ hybridization and immunostaining (Chen et al., 2016a). These hormones have also been found to be related to sleep, wakefulness, and circadian rhythms (Chen et al., 2016b; Chung et al., 2017; Diaz et al., 2019; Fujitaka et al., 2018; He et al., 2013; Hermann et al., 2012; Kunst et al., 2014; Oh et al., 2014; Ren et al., 2015). Therefore, we focused on these seven peptide hormones and evaluated their involvement in the sleep effects of SBT2227.

First, we combined Act-Gal425FO1 with UAS-RNAi against each target gene to ubiquitously knockdown their expression and analyzed the sleep of flies with or without the administration of SBT2227. As Act-Gal425FO1 > AstARNAi flies showed severe lethality, we could not evaluate them under these conditions. Quantitative comparison of the amount of sleep during the ZT12-15 period on the third day showed a significant increase in sleep in the following order: Act-Gal425FO1 > AstCRNAi, Act-Gal425FO1 > CCHa2RNAi, or Act-Gal425FO1 > Dh31RNAi flies fed SBT2227 (Figure 7A). By contrast, in Act-Gal425FO1 > MipRNAi, Act-Gal425FO1 > CCHA1RNAi, or Act-Gal425FO1 > NFRNAi flies, the amount of sleep did not increase after SBT2227 feeding (Figure 7A). Furthermore, the sleep latency decreased significantly after SBT2227 feeding in AstC or CCHA2 knockdown flies, but not in Mip, CCHA1, Dh31, or NPF knockdown flies (Figure 7B). In contrast, all parental control flies showed increased the sleep amount and shortened the sleep latency after SBT2227 feeding (Figure S5). These results indicate that Mip, CCHA1, and NPF are required for both sleep effects (increasing the amount and decreasing the latency) of SBT2227. Thus, they are the putative candidate mediators.

Continuous repression of AstA expression using Act-Gal425FO1 may affect developmental processes. To exclude the possibility of a developmental abnormality, we performed molecular genetic manipulations to temporally suppress the expression of AstA, Mip, CCHA1, or NPF using the GeneSwitch system (Osterwalder et al., 2001; Roman et al., 2001). To suppress the expression of these genes ubiquitously at the adult stage, we used tub-GeneSwitch-Gal4 in combination with UAS-RNAi corresponding to each gene. A significant reduction in gene expression was observed in flies after RNAi induction with RU486 feeding (Figure S6). The amount of sleep during ZT12-15 on day 3 increased significantly in flies with temporary
knockdown of AstA, Mip, or CCHA1, but not in flies with NPF knockdown (Figure 7C). In addition, the effect of SBT2227 administration on sleep latency was not observed in flies with NPF knockdown (Figure 7D). On the other hand, the effect of SBT2227 on sleep was similar to that of the wild type in tub-GeneSwitch-Gal4 >NPFRNAi flies treated with the vehicle control (Figure S7). These results suggested that NPF was the most potent mediator of the sleep-promoting effects of SBT2227.

**DISCUSSION**

We investigated the biological effects of LAB feeding using the fruit fly *D. melanogaster* and found the sleep-promoting effects of the LAB strain *L. plantarum* SBT2227. SBT2227 retained its sleep-promoting effects even after heat denaturation. Therefore, SBT2227 did not act on the host organism as a probiotic. Furthermore, SBT2227 promoted sleep in flies even when commensal intestinal bacteria were removed, suggesting that it did not act as a prebiotic. This finding is consistent with that of a previous study showing that removal of the *Drosophila* intestinal flora negligibly affects sleep (Jia et al., 2021; Storelli et al., 2018). These results suggested that the biological action of SBT2227 is mostly biogenic.

In biogenic action, substances present in LAB induce biological effects (Mitsuoka, 2000). The supernatant of the cytoreductive fluid of SBT2227 promoted sleep in flies, suggesting that the active substances were present in the supernatant fraction. The fact that autoclaved SBT2227 promoted sleep in flies suggested that heat-denatured proteins and hydrolyzed DNA/RNA can be excluded from the list of potential active substances. Molecules such as peptidoglycans and cell wall-associated components in the precipitated fraction of crushed SBT2227 may also be excluded as candidates, as they did not exert any sleep-promoting effect under our study conditions. The active substances can be identified by comprehensively identifying the substances in the supernatant fraction and administering the identified substances to the flies. Our preliminary investigations showed that different strains of SBT2227 of *L. plantarum* retain a similar sleep effect on flies, while LABs of the genus *Streptococcus* had no detectable sleep effect in our experimental condition. Metabolomic comparisons between these different species of bacteria will help to identify the causative substances responsible for the sleep effects of SBT2227.

RNA-seq-based gut transcriptome analysis in this study revealed multiple biological pathways that were enriched by SBT2227 administration, which may contain molecules responsible for SBT2227’s action on...
fly sleep. Neuroactive peptide hormones are involved in these candidate biological pathways. The first candidate for the biogenic action of LAB in the target tissue is the cell population comprising the intestine, as it is the site of absorption of the active substance of LAB. EECs are one such group of cells (Chen et al., 2016a; Guo et al., 2019; Hung et al., 2020), from which peptide hormones are secreted into the hemolymph, which possibly transfers these hormones to brain regions to regulate fly sleep. Some EEC-derived peptide hormones, such as AstA, Mip, AstC, CCHA1, CCHA2, Dh31, and NPF, have been reported as sleep/wake-affective or circadian rhythm-affective hormones (Chen et al., 2016b; Chung et al., 2017; Diaz et al., 2019; Fujiwara et al., 2018; He et al., 2013; Hermann et al., 2012; Kunst et al., 2014; Oh et al., 2014; Ren et al., 2015).

To identify the molecules responsible for the sleep-promoting effect of SBT2227 from among these peptide hormones, we performed mini-screening for suppression of gene expression by combining RNAi (Fire et al., 1998) with Gal4/UAS method (Fischer et al., 1988). NPF was found to be an endocrine factor necessary for the sleep-promoting effect of SBT2227.

NPF is a peptide hormone composed of 36 amino acid residues, the vertebrate homolog of which is neuropeptide Y (NPY) (Brown et al., 1999). NPFR, the receptor for NPF, encodes a G-protein-coupled receptor (GPCR)-type receptor that activates inhibitory G proteins (Garczynski et al., 2002). The biological functions

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**Figure 6.** RNA-seq-based transcriptome analysis indicated multiple biological pathways altered by the administration of SBT2227

(A) A flow chart of transcriptome analysis is shown.

(B) Statistically significant GO terms in the category of "Molecular Function" are shown with p value and enrichment score. Enrichment score was calculated as (number of genes associated with the GO term)/(number of all genes in the GO term).
of NPF include the regulation of feeding (Wu et al., 2005), courtship behavior (Liu et al., 2019), alcohol sensitivity (Wen et al., 2005), aggression (Dierick and Greenspan, 2007), and learning and memory (Krashes et al., 2009). It is also involved in the regulation of circadian rhythms, especially in the evening (Hermann et al., 2012). In zebrafish, NPY promotes sleep by inhibiting the wake-promoting noradrenergic system (Singh et al., 2017). By contrast, the sleep-promoting effect of NPY in mammals is still controversial; intravenous injection of NPY promotes sleep in humans (Antonijevic et al., 2000; Held et al., 2006). Furthermore, in rodents, injection of NPY into the brain has been reported to promote sleep and decrease locomotion (Akanmu et al., 2006; Jolicoeur et al., 1991); however, the opposite effect has also been reported (Szentirmai and Krueger, 2006; Ushimura et al., 2015).

**Drosophila** NPF exerts a similar effect on sleep: overexpression of NPF promotes sleep but is restricted to males (He et al., 2013); conversely, activation of NPF-producing cells promote wakefulness (Chung et al., 2017). One possible reason for such contradictory effects is that the directions of sleep regulation by NPF and NPY may differ with the target neurons (Singh et al., 2017).

NPF is expressed in the brain and the midgut. NPF produced from the midgut is known to regulate germ-line stem cell proliferation, although its effect on sleep is unknown (Ameku et al., 2018). Furthermore, the relationship between NPF in the brain and sleep is not understood. *Drosophila* sleep is controlled by two oscillators: a morning oscillator and an evening oscillator. Dorsal lateral neurons (LNds) and fifth small ventral lateral neurons (sLNvs) expressing NPF are known to be part of the evening oscillator, and their neural activity increases in the evening (Liang et al., 2016). LNds are composed of six neurons, three of which are NPF-positive neurons and one among which is a cryptochrome (CRY)-positive neuron (Yoshi et al., 2008). According to Chung et al., CRY-positive NPF-producing cells promote wakefulness (Chung et al., 2017).

**Figure 7. Mini-screening of flies with gene knockdown of the peptide hormones expressed in the gut**

(A and B) Knockdown was performed ubiquitously using Act-Gal4. (A) Amount of sleep during ZT12-15 on day 3. (B) Sleep latency on day 3.

(C and D) Knockdown was performed ubiquitously and temporally using tub5-GS-Gal4. (C) Amount of sleep during ZT12-15 on day 3 and (D) sleep latency on day 3. The sample size is shown below each graph. The Wilcoxon-Mann-Whitney test was used for statistical analysis of the control group vs SBT2227 administered group in each genotype. *p < 0.05, **p < 0.01, ***p < 0.001, n.s. means not significant at p < 0.05. See Table S2 for detailed statistics.
However, the target neurons expressing NPFR are still not clear. Identification of NPF-producing cells that are controlled by the active substances of SBT2227 and the NPFR-expressing cells that receive its signal should be identified in the future to understand the mechanism of action of SBT2227. This may also clarify the relationship between NPF/NPFR signals and sleep.

In our RNA-seq analysis, peptidoglycan muralytic activity, a GO category associated with bacterial recognition, was detected as an enriched category in the gut of flies treated with SBT2227. The immune-active state of animals enhances sleep, which has been reported in both mammals (Krueger et al., 1982) and flies (Kuo et al., 2010). SBT2227 harbors peptidoglycan on the cell wall surface, which is recognized by the Toll signaling pathway for gut immune responses (Valanne et al., 2011). Therefore, the cell wall fraction should be a potential substance for sleep promotion via immune responses. However, the precipitated fraction of crushed-SBT2227, which could be rich in peptidoglycan components, did not significantly affect fly sleep (Figure 5). Hence, the immune system may not be directly regulating SBT2227’s effects on sleep.

In conclusion, we found a novel biogenic action of L. plantarum strain SBT2227 as sleep-promoting effects on D. melanogaster. The active substance derived from cellular components of SBT2227 would be ingested, digested, and absorbed via the intestinal tract. The active substance could then act on the responsible brain circuits via the neuropeptide NPF to exert the sleep-promoting effect (Figure 8). Mammals also harbor the NPF homolog, NPY, which is reported to control sleep behavior. Therefore, we expected SBT2227 to exert sleep-promoting effects in mammals. Currently, the mainstream approach to improving sleep involves the use of pharmaceuticals. In the near future, elucidation of the mechanism of action of functional LABs that act on brain functions, such as SBT2227, will improve our understanding of new aspects of brain-gut-bacteria interactions and also assist in the development of functional fermented foods.

Limitations of the study
A contract with Megmilk Snow Brand Co., Ltd. is required for the use of L. plantarum SBT2227.

Figure 8. Model showing the biogenic action of the SBT2227 on fly sleep
The active substance of SBT2227 is released from the cells, or the cells are digested and flow out into the gut tract. The active substance acts directly or indirectly on NPF-producing cells and finally acts on the brain neuronal circuits to exert the sleep-promoting effect.
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SUPPLEMENTAL INFORMATION
Supplemental information can be found online at https://doi.org/10.1016/j.isci.2022.104626.

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AUTHOR CONTRIBUTIONS
T.K., H.M., and H.I. designed the research; T.K., H.M., and H.I. performed the research; H.I. contributed new reagents/analytic tools; T.K., H.M., and H.I. analyzed the data; T.K. and H.I. wrote the article; and A.K. and H.I. reviewed and edit the article. All authors read and approved the article.

DECLARATION OF INTERESTS
T.K. and H.M. are employees of Megmilk Snow Brand Co., Ltd. The other authors declare no competing interest.

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# STAR METHODS

## Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and virus strains** | | |
| *Lactobacillus plantarum* SBT2227 | This paper | RRID: N/A |
| **Chemicals, peptides, and recombinant proteins** | | |
| Tetracycline Hydrochloride | FUJIFILM Wako Pure Chemical | Cat# 205-08591 |
| Ampicillin Sodium | FUJIFILM Wako Pure Chemical | Cat# 016-23301 |
| Kanamycin Sulfate | FUJIFILM Wako Pure Chemical | Cat# 117-00341 |
| Erythromycin | FUJIFILM Wako Pure Chemical | Cat# 057-07151 |
| RNAiso Plus | Takara Bio | Cat# 9109 |
| RNA using ReverTra Ace qPCR RT master mix with gDNA Remover | TOYOBO | Cat# FSQ-301 |
| **Critical commercial assays** | | |
| THUNDERBIRD SYBR qPCR Mix | TOYOBO | Cat# QPS-201 |
| **Experimental models: Organisms/strains** | | |
| *D. melanogaster*: Canton-S2202u | Dr. Sakai, Tokyo Metropolitan Univ. Japan | RRID: N/A |
| *D. melanogaster*: Amherst_3 | Bloomington Drosophila Stock Center | RRID: BDSC_4265 |
| *D. melanogaster*: Act-Gal425FO1 / CyO, Act-GFP: y[1] w[+]; P[014C-GAL425FO1/CyO, y[+] | Dr. Awasaki, Kyorin Univ. Japan, available from the BDSC | RRID: BDSC_4414 |
| *D. melanogaster*: tub5-GS-Gal4 | Dr. Scott D. Pletcher, Baylor College of Medicine, Houston, TX, USA | RRID: N/A |
| *D. melanogaster*: UAS-AstA-RNAi: y[1] v[1]; P[014C-GAL425FO1/CyO, y[+] | Dr. Sakai, Tokyo Metropolitan Univ. Japan | RRID: BDSC_25866 |
| *D. melanogaster*: UAS-Mip-RNAi: y[1] sc[+] v[1] sev[21]; P[0777 v[1] v[1] 1.8]=TRIP:HS02244attP2 | Bloomington Drosophila Stock Center | RRID: BDSC_41680 |
| *D. melanogaster*: UAS-AstC-RNAi: y[1] v[1]; P[014C-GAL425FO1/CyO, y[+] | Dr. Awasaki, Kyorin Univ. Japan, available from the BDSC | RRID: BDSC_25868 |
| *D. melanogaster*: UAS-CHa1-RNAi: y[1] sc[+] v[1] sev[21]; P[0777 v[1] v[1] 1.8]=TRIP:HS04879attP2 | Bloomington Drosophila Stock Center | RRID: BDSC_57562 |
| *D. melanogaster*: UAS-CCha2-RNAi: y[1] sc[+] v[1] sev[21]; P[0777 v[1] v[1] 1.8]=TRIP:HS04565attP2 | Bloomington Drosophila Stock Center | RRID: BDSC_41957 |
| *D. melanogaster*: UAS-Dh31-RNAi: y[1] sc[+] v[1] sev[21]; P[0777 v[1] v[1] 1.8]=TRIP:HS02354attP2/TM3, Sb[1] | Bloomington Drosophila Stock Center | RRID: BDSC_57183 |
| *D. melanogaster*: UAS-NPF-RNAi: P[011184]VIE-260B | Vienna Drosophila Resource Center | VDRC ID: 108772 |
| **Oligonucleotides** | | |
| Primer: rp49-f; AGTATCTGATGCCCAACATCG | This paper | RRID: N/A |
| Primer: rp49-r; CAATCTCCTTGCCCATACATCG | This paper | RRID: N/A |
| Primer: AstA-f; TTGCACCGCGCCTCCGCTT | This paper | RRID: N/A |
| Primer: AstA-r; ATGCTATGGGCACGGGATGG | This paper | RRID: N/A |
| Primer: CChA1-f; CCAAAAATCAGTGGCGCAATG | This paper | RRID: N/A |
| Primer: CChA1-r; GCAATTGGCCTCGGAATGTT | This paper | RRID: N/A |
| Primer: Mip-f; CTCTAGCACCTAGTCTCCACG | This paper | RRID: N/A |

(Continued on next page)
Continued

**REAGENT or RESOURCE** | **SOURCE** | **IDENTIFIER**
--- | --- | ---
Primer: Mip-r, GTTGCCATTITGGTATGTATTGATGT | This paper | RRID: N/A
Primer: NPF-t, TCCGCGAAAGAGACGATGTC | FlyPrimerBank | Cat# PD44548, RRID:N/A
Primer: NPF-r, CTCTCATTAAACCCGCAG | FlyPrimerBank | Cat# PD44548, RRID:N/A

**Software and algorithms**

R Project for Statistical Computing 4.0.0 | R Core Team | https://www.r-project.org; RRID: SCR_001905
R Studio 1.4.1106 | RStudio PBC | https://www.rstudio.com/; RRID: SCR_000432
Rethomics | Geissmann et al. 2019 | https://rethomics.github.io/; RRID: N/A
StringTie | Pertea et al. 2015 | https://ccb.jhu.edu/software/stringtie/; RRID: SCR_016323
edgeR | Robinson et al. 2010 | http://bioconductor.org/packages/edgeR/; RRID: SCR_012802
gProfiler | Reimand et al., 2007 | https://bit.cs.ut.ee/gprofiler/gost; RRID: SCR_006809
G*power | Faul et al., 2007 | http://www.gpower.hhu.de/; RRID:SCR_013726

**Other**

TriKinetics Drosophila Activity Monitoring System, DAM2 | Trikinetics | https://trikinetics.com; RRID: SCR_021798
Bead cell disrupter | TOMY SEIKO | Micro Smash,MS-100R
5.0 mL tube (cell disrupter) | TOMY SEIKO | Cat# TM-657
0.1 mm glass beads (cell disrupter) | TOMY SEIKO | Cat# GB-01
Conical tube (food consumption) | Greiner | Cat# 227261
Feeder cup (food consumption) | Molecular BioProducts | Cat# 3500
Spectrophotometer | Thermo Fisher Scientific | Varioskan Flash

**RESOURCE AVAILABILITY**

**Lead contact**
Further information and requests for resources should be directed to and will be fulfilled by the lead contact Hiroshi Ishimoto (hishimoto@bio.nagoya-u.ac.jp).

**Materials availability**
*L. plantarum* SBT2227 was deposited in the International Patent Organism Depositary, National Institute of Technology and Evaluation (Chiba, Japan) under the accession number FERM BP-03104.

**Data and code availability**
The original/source data are available from the lead contact on request. This paper does not report original code. Any additional information required to reanalyze the data reported in this paper is available from the lead contact upon request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Fly strains and rearing conditions**
All fly strains used in this study were reared and maintained on standard cornmeal yeast (50 g/L glucose, 45 g/L yeast, 40 g/L corn flour, 8 g/L agar, 4 ml/L propionic acid, and 3 ml/L methyl 4-hydroxybenzoate) food at 24 ± 1°C and 60 ± 3% humidity, with a 12 h light/12 h dark (12 h L/D) cycle. The Canton-S2202u strain was used as the wild-type strain. Amherst_3 (BDSC_4265), another wild-type strain, was used to confirm the sleep phenotypes obtained using Canton-S2202u. To suppress gene expression, the following Gal4 and UAS strains were used: Act-Gal4^{SigFO1} / CyO, Act-GFP (Dr. Asakawa, Kyorin Univ. Japan, BDSC_4414), and tub5-GS-Gal4 (Dr. Scott D. Pletcher, Baylor College of Medicine, Houston, TX, USA). UAS-AstARNAi (BDSC_25866), UAS-Mip-RNAi (BDSC_41680), UAS-AstC-RNAi (BDSC_25868), UAS-CCHa1-RNAi
(BDSC_57562), UAS-CCHA2-RNAi (BDSC_57183), and UAS-Dh31-RNAi (BDSC_41957) were obtained from BDRC. UAS-NPF-RNAi (VDRC ID: 108772) was obtained from the Vienna Drosophila Resource Center. For all experiments, virgin female flies were collected under CO2 anesthesia and used for experiments. Flies were maintained in food vials (15 - 20 flies per vial) until the experiments.

Preparation of LAB samples

*L. plantarum* SBT2227 was obtained from Megmilk Snow Brand Co., Ltd. (Tokyo, Japan). SBT2227 was grown in Man, Rogosa, and Sharpe (MRS) broth (BD Biosciences, CA, USA) at 37°C for 16 h. The cultured bacteria were collected via centrifugation (7,000 × g, 20 min, 4°C) and then washed twice with sterilized saline (0.9% NaCl). The bacterial pellet was then suspended in a trehalose solution (12.5%) and concentrated to final volume of one-tenth of the culture medium. The bacterial suspension was incubated at 4°C for 2–5 h and then rapidly frozen in liquid nitrogen for storage at –80°C until further use. The frozen stock contained at least 5.2 × 10¹⁰ cfu/mL viable cells. For heat denaturation, SBT2227 was incubated at 65°C for 1 h using an incubator HB-80 (TAITEC Co., Saitama, Japan) or autoclaved at 121°C for 15 min using Panasonic MLS-3751 (PHC Corporation, Tokyo, Japan). After heat denaturation, the number of viable cells was <3,000 cfu/mL (65°C for 1 h) or <100 cfu/mL (autoclave). To crush SBT2227, 60.3 mg cells were suspended in 3 mL ultrapure water and placed in a 5.0 mL tube (TOMY SEIKO Co., Tokyo, Japan, Cat#TM-657) along with 2 g of 0.1 mm glass beads (TOMY SEIKO Co., Tokyo, Japan, Cat#GB-01). Cells were crushed at 4,200 rpm for 12 cycles at intervals of 30 s using a bead cell disrupter (MS-100R, TOMY SEIKO Co., Tokyo, Japan). After crushing, the glass beads were removed using a cell strainer (mesh size 40 μm, Thermo Fisher Scientific, MA, USA).

METHODS DETAILS

Measuring food consumption

Food consumption was measured as previously reported with some modifications (Shell et al., 2018; Wu et al., 2020). Briefly, the control and SBT2227 foods are colored blue (1% w/v, Blue No. 1, Fujifilm Wako Pure Chemicals Corporation, Osaka, Japan) and filled into feeder cups (10 μL tip, Molecular BioProducts, CA, USA, Cat# 3500), respectively. Drill a hole in the center of the lid of conical tube (50 ml conical tube, Greiner, Kremsmünster, Austria, Cat# 227261), and insert the feeder cup inside that lid. Ten flies were placed in each conical tube and capped with a lid with a feeder cup containing colored control or colored SBT2227 food for 3 h. The flies were then collected and ground in 1 mL of water to extract the blue dye from their bodies. In addition, another 1 mL of water was added to the conical tube from which the flies were removed, and the dye was extracted from the excretions on the tube wall. These dye extracts were mixed in equal amounts. The absorbance of the mixture was then measured at 630 nm in a spectrophotometer (Varioskan Flash, Thermo Fisher Scientific, MA, USA).

Antibiotic treatment

The antibiotic-supplemented food contained 50 μg/mL tetracycline (Fujifilm Wako Pure Chemicals Corporation, Osaka, Japan, Cat#025-08591), 50 μg/mL ampicillin (Fujifilm Wako Pure Chemicals Corporation, Osaka, Japan, Cat#016-23301), 50 μg/mL kanamycin (Fujifilm Wako Pure Chemicals Corporation, Osaka, Japan, Cat#117-00341), and 15 μg/mL erythromycin (Fujifilm Wako Pure Chemicals Corporation, Osaka, Japan, Cat#057-07151) in standard fly food. Newly enclosed flies were grown on antibiotic-supplemented food for four days. On the fourth day, the internal bacterial load was counted.

Quantification of bacterial load in flies

Flies were washed in 70% ethanol for 15 s and then rinsed twice with sterile phosphate-buffered saline (PBS) for 15 s. Three flies were placed in 100 μL sterile PBS in a 1.5 mL tube and then homogenized using a sterilized pestle. The homogenates were diluted and plated on MRS agar plates. After 3 days of incubation under either aerobic or anaerobic conditions at 37°C, the number of colonies on the plate was counted. Anaerobic conditions were generated using AnaeroPack-Anaero (Mitsubishi Gas Chemical, Tokyo, Japan).

Gram staining of bacteria

The bacterial suspension was spread on a glass slide, fixed with methanol, primary stained with Victoria blue, decolorized with picric acid in ethanol, and counterstained with fuchsin (Muto Pure Chemicals Co., Ltd. Tokyo, Japan).
Sleep analysis

Fly sleep was measured using the Drosophila activity monitoring system (Trikinetics, MA, USA). Four-day old virgin female flies were transferred into a glass tube (5 x 65 mm) with control food or test food on one end of the tube, which was sealed with a cotton plug on the other end after loading the fly. The flies were allowed to acclimate to the new environment overnight, and locomotor activity was measured for 3–4 days at 25 ± 1°C in a 12 h light/dark cycle.

The control food contained trehalose (6.25%), sucrose (5.0%), and bacto-agar (1.0%). The LAB food contained approximately 1% of SBT2227 (10.1 mg dry weight per mL). For evaluating crushed SBT2227 (10.1 mg/mL final concentration) and its centrifuged supernatant (5.4 mg/mL final concentration) and precipitate (3.0 mg/mL final concentration), each of them were mixed with the control food lacking trehalose. For the GeneSwitch system, RU486 (500 μM final conc., Mifepristone, Sigma-Aldrich, MO, USA) was mixed with SBT2227 food.

Locomotor data were collected in 1-min bins, and inactivity for at least 5 min was defined as a sleep bout. All behavior parameters were calculated and statistically analyzed for individual flies using R package Re-thomics (Geissmann et al., 2019). Dead animals were excluded from analysis. The sleep amount was calculated by summing the durations of all sleep events within the time of interest. Sleep latency was defined as the time from ZT12 (start of nighttime) to the detection of the first sleep event. The duration of a sleep event detected during the time period of interest was used as the sleep bout length. The number of sleep events detected during the time of interest was defined as the number of sleep bouts. The number of times a fly crossed the infrared beam during the time of interest was used as the activity counts. The daytime or nighttime activity index was calculated by dividing the number of activity counts by the total awake time (minutes) for each time period.

RNA isolation

Total RNA was extracted using RNAiso Plus (Takara Bio, Shiga, Japan) according to the manufacturer’s protocol. Five flies were placed in 500 μL RNAiso Plus reagent and thoroughly homogenized with a plastic pestle. Total RNA was dissolved in 100 μL RNase-free water.

For RNA sequencing, total RNA was extracted from the fly gut using the RNeasy mini kit (Qiagen, Hilden, Germany) according to the manufacturer’s protocol. The guts were collected between ZT12-14 4 days after the start of the sleep experiment. The guts (from the lower part of the crop to the upper part of the Malpighian tubules) were dissected and immediately immersed in 2-mercaptoethanol with Buffer RLT of the RNeasy mini kit. This procedure was repeated thrice independently.

Real-time polymerase chain reaction (PCR)

First-strand cDNA was synthesized from total RNA using ReverTra Ace qPCR RT master mix with gDNA Remover (TOYOBO, Osaka, Japan). Real-time PCR was performed using Thunderbird SYBR qPCR Mix (TOYOBO, Osaka, Japan). All steps were performed according to the manufacturer’s protocol. PCR amplification was performed using the following primers: NPF primers were designed using the FlyPrimerBank (Hu et al., 2013). Ribosomal protein 49 (rp49) was used as the internal control.

rp49-f; 5’-AGTATCTGATGCCAACATCG-3’
rp49-r; 5’-CAATCTCCTTGCGCTTCTT-3’
AstA-f; 5’-TTGCACCGGTATCCTGTCT-3’
AstA-r; 5’-ATGCTATGCGCAGGGATGG-3’
CCHA1-f; 5’-CCCAAATCGATGCGACAATG3’
CCHA1-r; 5’-GCAATTGGCCTCGGAATGGT-3’
Mip-f; 5’-CTCTAGCACCTAGTCCACG-3’
RNA sequencing analysis

RNA sequencing libraries were constructed and sequenced by Macrogen Inc. using the Illumina NovaSeq6000 system. Read fragments were mapped to the *D. melanogaster* genome NCBI GCF_000001215.4_Release_6_plus_ISO1_MT, and assembled using StringTie (Pertea et al., 2015). edgeR (Robinson et al., 2010) was used for comparison of RNA expression levels and the exact test for negative binomial distribution was applied. Among the genes expressed in both the control group and the SBT2227 administered group, the expression level of the genes was more than 2-fold, and \( p < 0.05 \) was considered for DEGs. For genes expressed in only one of the groups, those with \( p < 0.05 \), were included in DEGs. We performed GO enrichment analysis of these DEGs using g:Profiler (https://biit.cs.ut.ee/gprofiler).

QUANTIFICATION AND STATISTICAL ANALYSIS

All statistical analyses were performed using the R Studio version 1.0.136. (www.r-project.org). Sample size was estimated by G*Power software (Faul et al., 2007). All data were checked for normality with the Shapiro-Wilk test and appropriate statistical methods were applied. The statistical methods and statistics are summarized in detail in Table S2. When the Welch’s t-test or Wilcoxon-Mann-Whitney tests were repeated, the \( p \) values were adjusted using the Bonferroni method. Statistical significance was set at \( p < 0.05 \). The boxes in box and whisker plots represent the median and interquartile range (the distance between the first and third quartiles), and whiskers represent the highest and lowest data points, excluding any outliers. To evaluate the sleep-promoting effect, the effect size \( r \) was calculated from the \( z \) value of the Wilcoxon-Mann-Whitney test using the following formula: \( r = z / \sqrt{n} \).