Mice with Mutations in Trpm1, a Gene within the locus of 15q13.3 Microdeletion Syndrome, Display Pronounced Hyperactivity and Decreased Anxiety-Like Behavior.

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Research

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Abstract

15q13.3 microdeletion syndrome is a genetic disorder caused by a deletion of a region containing seven genes on chromosome 15, MTMR10, FAN1, TRPM1, MIR211, KLF13, OTUD7A, and CHRNA7, and characterized by a wide spectrum of psychiatric disorders. The contribution of each gene in this syndrome has been studied using mutant mouse models, but the phenotypes of these mice do not account for human phenotypes and the results are still controversial. The behavior of Trpm1−/− mice with relation to 15q13.3 microdeletion syndrome has not been investigated due to the visual impairment in these mice, which may confound the results of behavior tests that involve vision. We have now applied a comprehensive behavioral test battery to examine the relationship of TRPM1 and 15q13.3 microdeletion syndrome by using Trpm1 null mutant mice. Our data indicate abnormal behavior of Trpm1−/− mice which may explain some phenotypes of 15q13.3 microdeletion syndrome, including reduction of anxiety behavior, abnormality of social interaction, attenuation in fear memory, and hyperactivity, which is the most prominent phenotype of Trpm1 mutant mice. While the ON visual transduction pathway is impaired in Trpm1−/− mice, we did not detect compensatory high sensitivities for other sensory modalities. Although Trpm1−/− mice share the same pathway for visual impairment with mGluR6−/− mice, hyperlocomotion activity has not been reported in mGluR6−/− mice. These data suggest that the phenotype of Trpm1−/− mice extends beyond that expected from visual impairment alone. This is the first evidence to associate TRPM1 with impairment of cognitive function similar to that found in the phenotypes of 15q13.3 microdeletion syndrome.

Introduction

TRPM1, the first member of the melanoma-related transient receptor potential (TRPM) subfamily to be discovered, is the visual transduction channel downstream of metabotropic glutamate receptor 6 (mGluR6) in retinal ON bipolar cells (BCs) [1, 2]. Mutations in TRPM1 are found in humans with an autosomal recessive form of complete congenital stationary night blindness (CSNB) and Trpm1 mutant mice lack a b-wave in the electroretinogram (ERG) and have no light responses in ON BCs [3]. TRPM1 is located in human chromosome 15q13.3, a region that is associated with 15q13.3 microdeletion syndrome, a genetic disorder caused by a deletion encompassing a region of ~ 1.5 megabase (MB) from break-point (BP) 4 to BP5, with seven genes; MTMR10; FAN1; TRPM1; MIR211; KLF13; OTUD7A; and CHRNA7 (OMIM #612001) [4]. Individuals with 15q13.3 microdeletion syndrome have mild to moderate intellectual disorder, mild learning delays, autism spectrum disorder, epilepsy (recurring seizures), attention deficit hyperactivity disorder (ADHD) and mental illnesses including schizophrenia and bipolar disorder [5]. Deletion of CHRNA7 is thought to account for the neuropsychiatric disorder in 15q13.3 microdeletion syndrome, although the phenotype of Chrna7 deficient mice is controversial [6]. Otud7a mutant mice also show many features, including neurological features, in common with those in 15q13.3 microdeletion syndrome, including reduced body weight, developmental delay, abnormal electroencephalogram (EEG) patterns and seizures, reduced ultrasonic vocalizations, decreased grip strength, impaired motor learning/motor coordination, and reduced acoustic startle [7].
The role of TRPM1 in behavioral disorders has not been studied, probably because of its strong relationship with vision. TRPM1 and its regulator, mGluR6, both cause CSNB. In mutant retinas, both cause ON but not OFF pathway to fail to respond to light stimuli [1, 8, 9]. We previously reported an unexpected difference between \( Trpm1^{-/-} \) and \( mGluR6^{-/-} \) mouse retinas. Spontaneous oscillations are observed in \( Trpm1^{-/-} \) retinas, but not in \( mGluR6^{-/-} \) retinas by recording spiking in retinal ganglion cells (RGCs) using a multielectrode array (MEA) [10]. We also previously reported that rod ON BC terminals were significantly smaller in \( Trpm1^{-/-} \) retinas than those of \( mGluR6^{-/-} \)[10]. These data suggest that a deficiency of TRPM1, but not of mGluR6, in rod ON BCs may effect synaptic terminal maturation and could underlie the observed differences in oscillatory response. Spurred on by the observation of \( Trpm1^{-/-} \) deletion specific RGC oscillations and a gene location in the targeted region of 15q13.3 microdeletion syndrome, we searched for central and behavior changes that might attribute to a persistent, rhythmic visual outflow.

In this study, we thoroughly examined \( Trpm1^{-/-} \) mice with a battery of behavioral tests [11]. We also examined structural and functional change in \( Trpm1^{-/-} \) mouse brain, which may explain the abnormal behaviors of this mutant as a model of 15q13.3 microdeletion syndrome.

**Methods**

**Animals and Experimental Design**

\( Trpm1^{-/-} \) mice were generated as described previously [1]. In this study, we analyzed \( Trpm1^{-/-} \) mice with their wild type (WT) littersmates on the 129 Sv/Ev Taconic background. All behavioral tests were performed with male that were 11–12 weeks old at the starting of the testing (\( Trpm1^{-/-} \) mice, \( n=24 \); WT littersmates, \( n=24 \)). Mice were housed as 2 pairs of \( Trpm1^{-/-} \) and WT mice in a cage with a 12-hour light/dark cycle (light on 7:00 a.m. and off 7:00 p.m.). All mice had access to food and water *ad libitum*. Behavioral testing was performed between 8:30 A.M. and 6:30 P.M. unless otherwise noted. Our behavioral test battery consists of the tests listed in Table 1. After the tests, all the testing apparatus were cleaned with diluted hypochlorous solution or 70% ethanol to prevent a bias due to olfactory cues.

Brain weight measurement and monoamine quantification in brain tissues were performed with 129 Sv/Ev male at 4 months (\( Trpm1^{-/-} \) mice, \( n=24 \); WT littersmates, \( n=24 \)) or 1 months (\( Trpm1^{-/-} \) mice, \( n=4 \); WT littersmates, \( n=5 \)). Gene expression analysis was performed with 129 Sv/Ev male at 1 months (WT, \( n=5 \)). Mice used for monoamine quantification were housed as 2 pairs of \( Trpm1^{-/-} \) mice and WT mice in a cage with a 12-hour light/dark cycle (light on 8:00 a.m. and off 8:00 p.m.), and tissue dissection was performed at the same time point (1:00 p.m.). All mice had access to food and water *ad libitum*. The experimental procedures and housing conditions for animals were approved by Institutional Animal Care and Use Committee of National Institute for Physiological Sciences, Fujita Health University and Ritsumeikan University.

**General health and neurological screen**
A general health and neurological screen examined the body weight, rectal temperature, whisker, coat, simple reflexes such as righting, whisker touch, eye blink, ear twitch reflexes and reaching behavior using 11 ~ 12 weeks old, as described previously [12]. Grip strength test and wire hang test were conducted to measure muscle strength. Grip strength was measured by using a grip strength meter (O’Hara & Co., Japan). In the wire hang test, the mouse was placed on a wire cage lid that was then inverted, so that the subject gripped the wire. Latency to fall onto the bedding was recorded, with a 60 sec cutoff time.

**Light/dark transition test**

Light/dark transition test was performed as described previously [13–15]. The apparatus used for light/dark transition test consisted of a cage (21 × 41.5 × 25 cm) divided into two sections of equal size by a partition with a door (O’Hara & Co., Japan). One section was brightly illuminated (390 ± 20 lux), whereas the other section was dark (< 2 lux). Mice were placed into the dark side of the apparatus, and allowed to move freely between the two sections for 10 min, while the door remained open. In the same way, mice (34–35 weeks old) were placed into the light side of the apparatus allowed to move freely between the two sections for 10 min. The total number of transitions, time spent in each section, initial latency to the light section, and distance traveled were recorded automatically using Image LD software.

**Open field test**

Open field test was performed as described previously [14, 15]. Mice was allowed to move freely in an open field apparatus (40 × 40 × 30 cm; Accuscan Instruments, U.S.A.), which was illuminated at 10.0 lx for 120 min. Each subject was placed in the corner of the apparatus. The total distance, vertical activity (rearing measured by counting the number of photobeam interruptions), time spent in the center area, and stereotypic behaviors were recorded.

**Elevated plus maze test**

Elevated plus maze test was performed as described previously [14, 16]. The apparatus (O’Hara & Co., Japan) consisted of two open arms (25 × 5 cm) and two enclosed arms of the same size, with central square (5 × 5 cm). The closed arms have 16 cm high transparent walls. To minimize the likelihood of animals falling from the apparatus, 3-mm-high Plexiglas ledges were provided for the open arms. The arms and were made of white plastic plates and were elevated to a height of 50 cm above the floor. Arms of the same type were arranged at opposite sides to each other. Mice was placed in the central square of the maze, facing one of the closed arms and behavior was recorded during a 10-min test period. The percentage of open arm entries, the percentage of time spent on the open arms, the total number of arm entries, and total distance traveled were measured automatically using Image EP software.

**Hot plate test**

Hot plate test was performed as described previously [16]. Mice were placed on a 55.0 °C hot plate (Columbus Instruments, U.S.A.), and latency to the first hind paw response was recorded. The hind paw response was either a foot shake or a paw lick.

**Social interaction test**
Social interaction test was performed as described previously [16]. A pair of mice (12–13 weeks old) was placed simultaneously at opposing corners in the open field apparatus (40 × 40 × 30 cm; O’Hara & Co., Japan), whose illumination level was 10.0 lx at the center of the floor, and allowed to explore freely for 10 min. The pair of mice had been housed in different cages. The number of active contacts, the number of contacts, mean duration per contact, total duration of contact, and total distance traveled were measured. Analysis was performed automatically using Image SI software.

**Rota-rod test**

Motor coordination and balance were tested with the rota-rod test old as described previously [16]. The rota-rod test using an accelerating rota-rod (UGO Basile, Italy) was performed by placing a mouse on a rotating drum (3 cm diameter) and measuring the time each animal was able to maintain its balance on the rod. The speed of the rota-rod accelerated from 4 to 40 rpm over a 5-min period.

**Social approach and novelty preference test**

Social approach and preference for social novelty were tested with the The three-chamber social approach test as described previously [14, 16]. The apparatus comprised a rectangular, three-chambered box and a lid with a video camera (O’Hara& Co., Japan). Each chamber was 20 cm × 40 cm × 22 cm and the dividing walls had small openings (5 cm × 3 cm) to allow exploration into each chamber. The day before testing, the mice were individually placed in the middle chamber and allowed to freely explore the entire apparatus for 10 min. During the test session, the amount of time spent in each chamber and time spent around each cage were recorded and analyzed automatically using Image CSI.

**Acoustic Startle response/prepulse inhibition tests**

Acoustic Startle response/prepulse inhibition tests were performed as described previously [16] (O’Hara & Co., Japan). A test session began by placing a mouse in a Plexiglas cylinder where it was left undisturbed for 10 min. The duration of white noise that was used as the startle stimulus was 40 msec for all trial types. A test session consisted of six trial types (i.e., two types for startle stimulus-only trials and four types for prepulse inhibition trials). The intensity of startle stimulus was 110 or 120 dB. The prepulse was presented 10.0 msec before the startle stimulus, and its intensity was 74 or 78 dB. Four combinations of prepulse and startle stimuli were used (74–110, 78–110, 74–120, and 78–120). Six blocks of the six trial types were presented in pseudorandom order such that each trial type was presented once within a block. The average intertrial interval was 15 sec (range, 10–20 sec).

**Porsolt Forced Swimming test**

Depression-related behavior was assessed, using the forced swimming test as described previously [12]. The apparatus consisted of Plexiglas cylinders (22 cm height x 12 cm diameter). The cylinders were filled with water (room temperature, 23 ± 2 °C), up to a height of 7.5 cm. Mice were placed into the cylinders, and their behavior was recorded over a 10-min test period. Immobility and distance traveled were measured analyzed automatically using Image PS software.

**Gait analysis**
The gait during walk/trot locomotion was assessed using DigiGait Imaging System (Mouse Specifics, U.S.A.) as described previously [17]. Digital video images of the underside of mice were collected at 150 frames per second. We placed the mice on a treadmill belt that moves at a speed of 24.7 cm/s. The percent of the time of stride or stance duration, stride length, stance width, step angle and paw angle were calculated.

**Barnes maze**

The Barnes maze test was performed as described previously [12]. The circular open field (O’Hara & Co., Japan) was elevated 97 cm from the floor. Training session were conducted one to three per day. After 24 hours after 15th training session, a probe test was conducted without the escape box, to confirm that this spatial task was acquired based on navigation by distal environmental room cues. One month after last (16th) training session, probe trial tests were conducted again to evaluate memory retention. After five additional training sessions conducted after the memory retention test, the escape box was moved to a new position opposite to the original (reversal learning). Mice were then trained with 8 sessions to locate the new position of the escape hole using the same procedure as described above. Latency to reach the target hole, distance to reach the target hole, number of errors and time spent around each hole were recorded automatically using Image BM software.

**T-maze Spontaneous Alteration**

T-maze spontaneous alteration test was performed as described previously [15] using the automatic modified T-maze apparatus (O’Hara & Co., Japan). Mice were subjected to a the spontaneous alteration protocol for 5 sessions. One session consists of 10 choices with a 50-min cutoff time. Mice were first placed in the start compartment of the T-Maze. Mice chose entering either the left or the right arm and could return to the start compartment. The mice was then given a 3-sec delay followed by a free choice between both T arms. A correct choice was made if the mouse entered the arm which was not visited in the previous choice. The percentage of correct response, latency (sec) to complete a session, distance traveled during the session. Data acquisition was performed automatically using Image TM software.

**Tail suspension test**

Depression-related behavior was assessed by the tail suspension test as described previously [17]. Mice were suspended 30 cm above the floor in a visually isolated area by adhesive tape placed ,1 cm from the base of the tail, and their behavior was recorded over a 10-min test period. Data acquisition and analysis were performed automatically using Image TS software.

**Contextual and cued fear conditioning**

The ability of mice to learn and remember an association between environmental cues and aversive experiences was assessed by fear conditioning test as described previously [15, 16]. Each mouse was placed in a test chamber (26 × 34 × 33 cm, O’Hara & Co., Japan) and allowed to explore freely for 2 min. A 55 dB white noise, which served as the conditioned stimulus (CS), was presented for 30 sec. Next, a mild (2 sec, 0.3 mA) foot shock, which served as the unconditioned stimulus (US), was presented immediately
after the CS. Two more CS-US pairings were presented with a 2-min interstimulus interval. Context testing was conducted 1 day after conditioning in the same chamber for 30.0 sec without CS and US presentations.

Cued testing with altered context was conducted after conditioning using a triangular box (33 × 33 × 33 cm) made of white opaque Plexiglas, which was located in a different room. Mice are allowed to explore the chamber for 360 sec. In the first 3 min, neither a CS nor US is presented, then a CS (a 55dB white noise) is presented for the last 3 min. Freezing and distance traveled were recorded. Data acquisition, control of stimuli (i.e. tones and shocks), and data analysis were performed automatically using Image FZ software.

**Twenty-four hour home cage monitoring test**

The 24-hour home cage test was performed as described previously [15]. The system for monitoring social interaction comprised a home cage (19 × 29 × 13 cm) and a filtered cage top with an infrared video camera (O'Hara & Co., Japan). Two mice with the same genotype that had been housed separately were placed together in a home cage. To evaluate their locomotor activity and social interaction, their behavior was monitored with a video camera for a week. Distance traveled was measured automatically using ImageHA software. The occurrence of social interaction was detected by counting the number of particles consisting of the mice as follows: 2 particles indicated that the mice were not in contact whereas 1 particle indicated that 2 mice were in contact. Locomotor activity of the mice was also measured.

**Methylphenidate administration in the open field**

After the behavioral test battery, the behavioral response to methylphenidate (MPH) was assessed in the open filed. A quarter of the area of the open field apparatus (20 × 20 × 30 cm) was used by installing a divider. Other conditions were the same as for the open filed test. The mice of each genotype were randomly divided into two groups for treatment with MPH and saline. The experiment was repeated twice with varying drug doses. Locomotor activity was recorded continuously during the 60-min habituation period and for 120 min after injection of saline or MPH (3 mg/kg or 10 mg/kg).

**Monoamine quantification in brain tissues**

Monoamine transmitter quantification was performed as described previously [18]. Tissue concentrations of biogenic monoamines were analyzed after dissection in various brain regions; prefrontal cortex, hippocampus, striatum, cerebral cortex, olfactory bulb, cerebellum, midbrain, pons and medulla, thalamus, hypothalamus. The weight of the brain tissue was measured, and homogenized in 0.2 M ice-cold perchloric acid (including 10.0 µM EDTA·2Na) and the homogenates were cooled on ice for 30 min to deproteinize. The homogenates were centrifuged at 20,000 G for 15 min at 0 °C. Then, the pH of the supernatant was adjusted to approximately 3.0 by adding 1 M sodium acetate. The samples were filtered through a 0.45 mm filter (Millipore, Billerica, USA). Next, 10 µL of filtrate was loaded into a high performance liquid chromatography (HPLC) system (Eicom, Japan). The HPLC system had a ø3.0 mm x 150 mm octadecyl silane column (SC-50DS, Eicom), and an electrochemical detector (ECD) (HTEC-50.0;
Eicom, Japan) set to an applied potential of +750 mV versus an Ag/AgCl reference analytical electrode. The change in electric current (nA) was recorded using a computer interface at 25 °C. The mobile phase was composed of 0.1 M aceto-citric acid buffer (pH 3.5), methanol, sodium-1-octane sulfonate (0.46 M), and EDTA2Na (0.015 mM) [830: 170: 1.9: 1]. The flow rate was 0.5 mL/min.

**Gene expression analysis in the brain**

Total RNA was isolated from each brain part using Biomasher (nippi, Japan) and ISOGEN (NIPPON GENE, Japan). For complementary DNA (cDNA) synthesis, 1 µg of total RNA was reverse transcribed (RT) into cDNA using the SuperScript (TaKaRa, Japan) according to the manufacturer's instructions. For quantitative polymerase chain reaction (qPCR) was conducted on a Thermal Cycler Dice® Real Time System (TaKaRa, Japan) using TB Green® *Premix Ex Taq™* (Tli RNaseH Plus) (TaKaRa, Japan) according to the manufacturer's instructions. Primers for mouse *Trpm1*: forward, 5'-GAGATGCAGCCAAACTGAAGC-3'; reverse, 5'-TGACGACACCAGTGCTCACA-3'. Primers for mouse *b-actin*: forward, 5'-CTCTGGCTCCTAGCACCATGAAGA−3'; reverse, 5'-GTAAAACGCAGCTCAGTAACAGTCCG−3'.

**Corticosterone measurement**

Blood was collected from mice at 4 months old by Cardiac puncture immediately after cervical dislocation. A serum was separated by centrifuging at 2,000 g for 20 min, and stored at -80 ºC until use. Corticosterone measurements were performed on serum by enzyme-linked immuno sorbent assay (ELISA) using a Corticosterone immunoassay (R&D Systems, U.S.A.) according to the manufacturer's instructions.

**Image analysis**

Behavioral data were obtained automatically by customized applications based on a public domain ImageJ program (Image LD, Image EP, Image SI, Image CSI, Image PS, Image BM, Image TM, Image TS, Image FZ, Image HA). The ImageJ plugins, and the precompiled plugins for light/dark transition test (Image LD), elevated plus maze (Image EP), open field test (Image OF), fear conditioning test (Image FZ), and T-maze (Image TM) are freely available on the website of “Mouse Phenotype Database” (http://www.mouse-phenotype.org/software.html).

**Data analysis**

All statistical analyses were performed using Graph Pad Prism7. Statistical methods are indicated in the figure legends. Data are presented as mean ± SEM. Unpaired 2-tailed Student’s *t* test or Welch’s *t* test were used for 2-group comparisons. Two-way analysis of variance (ANOVA) or repeated-measures two-way ANOVA following Tukey’s test and one-way ANOVA following Dunnett’s test was used for multiple comparison. Unless otherwise noted, the *p* values are for the genotype effect.

**Data repository**

The raw data of the behavioral tests and the information about each mouse are accessible on the public database “Mouse Phenotype Database” (http://www.mouse-phenotype.org/).
Results

Trpm1−/− mice show significantly high daily locomotor activity.

We performed a battery of more than 20 behavioral tests (Table 1). There was almost no significant difference for general physical characteristics, such as body weight, body temperature, grip strength, and motor coordination between Trpm1−/− and wild type mice (Fig. S1A-K). Trpm1−/− did not show depression-like behaviors in Porsolt forced swim test and tail suspension test (Fig. S1L-N). Intriguingly, Trpm1−/− mice showed significantly high daily locomotor activity (Fig. 1A) in contrast to mGluR6−/− mice [19]. It has been reported that the absence of vision in human and animals enhances auditory, haptic, and pain sensitivities [20–29], and causes structural changes in the visually deprived cortex and in other areas [23, 30, 31]. We examined sensory responses in Trpm1−/− mice, but we did not find any significance difference between Trpm1−/− mice and wild type for the hot plate test, the acoustic startle response, and prepulse inhibition (Fig. 1B-D). Although Trpm1−/− mice lack a functional ON visual pathway and thus have limited light responses, Trpm1−/− mice showed hyper- locomotor activity but did not show sensory hypersensitivities that was reported in other visually impaired animals.

Hyperactivity and reduction of anxiety-like behavior in Trpm1−/− mice

To assess anxiety-like behavior, we performed the light/dark transition test, the open-field test, and the elevated plus maze test (Fig. 2). In the light/dark transition test, distance traveled in the light and dark chamber were significantly increased in Trpm1−/− mice suggesting a reduction in anxiety-like behavior (Fig. 2A). However, the defect in the ON visual pathway may have affected the longer stay time in light and increased transition time and shorter latency to light for tests started at dark (Fig. 2B-D). In the open field test, which measures voluntary locomotor activity in a novel environment, Trpm1−/− mice exhibited a significant increase in total distance, vertical activity, center time and stereotypic behavior relative to WT mice (Fig. 2E-H), suggesting strong hyperactivity, which also explains longer distance traveled in the light/dark transition test. To implicate the hyperactivity in Trpm1−/− mice with ADHD, we performed the open field test after administration of MPH, which is often used as a first choice for treatment of ADHD in human (Fig. 2I) [32]. 120 min after administration of MPH, both WT and Trpm1−/− mice showed prominent hyperactivity, especially in mice which were injected at 10 mg/kg. As a result, MPH administration did not support the idea that the ADHD behavior displayed in Trpm1−/− mice can be reduced by MPH administration [33].

Additionally, in the elevated plus maze test, Trpm1−/− mice exhibited significantly increased numbers of entries and longer traveled distance, which are also explained by hyperactivity (Fig. 2J, M). Although visually impaired, Trpm1−/− mice did not show differences for entries to open arms, but stayed longer time in open arms suggesting a reduction of anxiety-like behavior (Fig. 2K, L).
To examine what causes a reduction in anxiety-like behavior in \textit{Trpm1}^{−/−} mice, we examined serum corticosterone levels in \textit{Trpm1}^{−/−} mice by ELISA [34, 35]. The idea is that an authentic reduction in anxiety should be correlated with a decrease in serum corticosterone levels, while a reduction in anxiety-like behavior in the absence of a decrease may have some other cause. The serum levels of corticosterone were not significantly different between \textit{Trpm1}^{−/−} mice and WT mice (Fig. S1O). This observation suggests that the hyperactivity of \textit{Trpm1}^{−/−} mice simulates a reduction of anxiety in our tests.

Abnormal social interaction in \textit{Trpm1}^{−/−} mice

Four kinds of social interaction tests (novel environment, sociability, novelty preference, and home cage test) were performed to evaluate social behaviors in the \textit{Trpm1}^{−/−} mice (Fig. 3). The novel environment test revealed significant differences between \textit{Trpm1}^{−/−} and WT mice, including a shorter duration per contact, increased contact number, and total traveled distances, which may be explained by the hyperactivity in \textit{Trpm1}^{−/−} mice (Fig. 3A, D, E). Although total duration of contact was shortened, but not significantly, active contacts were longer in \textit{Trpm1}^{−/−} mice (Fig. 3B, C). Both Crawley’s sociability and social novelty preference test and the test in the home cage did not reveal significant differences between WT and mutant mice (Fig. 3F-M, S1P). These results suggest that social interactions were attenuated in \textit{Trpm1}^{−/−} mice, which may be partially explained by hyperlocomotion behavior.

Attenuation of fear and spatial memories in \textit{Trpm1}^{−/−} mice

The contextual and cued fear conditioning test is used to assess fear memory (Fig. 4). In the conditioning phase, \textit{Trpm1}^{−/−} mice showed a lower level of freezing and traveled longer distances during sessions (Fig. 4A, B). The mutant mice traveled longer immediately after foot shock, an index of pain sensitivity (Fig. 4C). 24 hours after conditioning, \textit{Trpm1}^{−/−} mice showed decreased freezing and increased distance traveled. Similar significant differences were observed in tests 28 days after conditioning (Fig. 4D, E). These data suggest that attenuation of fear memory in \textit{Trpm1}^{−/−} mice could be related to hyperactivity.

We performed the Barnes maze test to determine whether the deficit in fear memory of \textit{Trpm1}^{−/−} contributes to hyper-locomotion or results from a deficit of memory. In both training sessions and reversal task tests, the distance to the escape box (Fig. 4F) and the number of errors to reach the escape box were significantly higher in \textit{Trpm1}^{−/−} mice (Fig. 4G), but latency to first reach the escape box was equivalent or shorter in \textit{Trpm1}^{−/−} mice than in WT mice (Fig. 4H), which may be related to hyper-locomotion activity. The probe tests were performed 24 hours and 1 month after the final training sessions. In these tests, \textit{Trpm1}^{−/−} and WT mice exhibited a significant effect of hole location target against the rest holes: 24 hour, WT \( p < 0.0001, \) \textit{Trpm1}^{−/−} \( p < 0.0001; \) 1 month, WT \( p < 0.0001, \) \textit{Trpm1}^{−/−} \( p < 0.0001; \) one-way ANOVA followed by Dunnett’s multiple comparison test), indicating that both genotypes were able to distinguish the location of the target. Time spent around the correct hole did not differ significantly between both genotypes at 24 hours after training, but was significantly shorter in \textit{Trpm1}^{−/−} mice 1 month later (Fig. 4I, J). These results suggest that \textit{Trpm1}^{−/−} mice have a deficit in long-term memory. In the reversal probe
test, although both genotypes distinguished the location of the target (WT $p<0.0001$, $Trpm1^{-/-} p<0.0001$; one-way ANOVA followed by Dunnett’s multiple comparison test), there was no significant difference in time spent around the correct hole between both genotypes (Fig. 4K). This result indicates that there was no deficit in behavioral flexibility in $Trpm1^{-/-}$ mice. We also performed a T-maze test to examine the working memory of $Trpm1^{-/-}$ mice. Although the shorter latency and longer distance traveled in $Trpm1^{-/-}$ mice were both significantly different, the correct responses at each trial were not (Fig. 4L-N). Taken together, $Trpm1^{-/-}$ mice showed attenuation for fear and long term memories, but no obvious deviation for flexibility and working memory.

Abnormal Structural and biochemical changes in the brains of $Trpm1^{-/-}$ mice

We have shown differences in the behavioral phenotype in $Trpm1^{-/-}$ mice relative to WT. However, $Trpm1$ functions predominantly as a component of the retinal ON bipolar transduction cascade and its expression is quite minor in the brain. To determine whether there are central structural changes, we compared brain regions between $Trpm1^{-/-}$ and WT mice. The cerebral cortex, olfactory bulb, and pons and medulla of $Trpm1^{-/-}$ mice were significantly heavier than in WT mice at 1 month old (Fig. 5A). In addition, the cerebral cortex, hippocampus, midbrain and cerebellum of $Trpm1^{-/-}$ mice were significantly heavier than those of WT mice at 4 months old (Fig. 5B).

We detected a subtle expression of $Trpm1$ mRNA throughout the WT mouse brain with the exception of the cerebellum (Fig. 5C). We also quantified levels of biogenic monoamines ex vivo, including dopamine (DA), noradrenaline (NA), serotonin (5-HT), and their major metabolites using HPLC-ECD in several adult brain regions. Levels of DA, NA, and NM were significantly decreased in the cerebellum (Fig. 5D-F). There was no significant change in the levels of the other monoamines and their metabolites in any other brain region (Fig. S2). The decreased levels of monoamines in the cerebellum of $Trpm1^{-/-}$ mice could influence hyper-locomotory activity, locomotion is regulated by the cerebellum. The lack of overlap between the $Trpm1$ expression pattern and the change of monoamine distribution in the brain is consistent with the idea that $Trpm1$ expresses in monoaminergic neurons that project to the cerebellum.

Discussion

15q13.3 microdeletion syndrome shows a spectrum of neurobehavioral phenotypes in human patients. So far, several investigations have suggested that mutants that abolish the functions of OTUD7A and CHRNA7 can partially explain the phenotypes of 15q13.3 microdeletion syndrome. However, we still lack a full accounting for the microdeletion phenotypes, especially those related to hyperactivity. In this paper, we examined the behavior of $Trpm1$ deficient mice with a comprehensive behavioral test battery. Our data show abnormal behaviors in $Trpm1$ deficient mice including a reduction of anxiety behavior, an abnormality of social interaction, the attenuation of fear and spatial memories, and hyper-locomotion activity, which is the most prominent phenotype of $Trpm1$ mutant mice and may also underlie or contribute to other phenotypes of $Trpm1^{-/-}$ mice (Fig. 1–3). Hyperactivity is one of the features of ADHD,
and a patient with 15q13.3 deletion, who had a relative lack of expression of genes including TRPM1, showed ADHD behavior [36–43]. We examined the effect of MPH which is often used for treatment of ADHD, however, MPH significantly increase the locomotion activity of Trpm1−/− mice (Fig. 2I). Intriguingly, MPH-like compounds were ineffective in approximately 35% of patients with ADHD [44, 45]. There are several mouse models of hyperactivity which were also insensitive to MPH. In Ndrg2 deficient mice, their ADHD-like hyperactivity was not rescued by MPH [46]. Shank2 and Fmr1 mutant mice, both are the model of autism, display hyperactivity, but the injection of MPH increased their locomotion activity [47, 48]. Relevant to the effect of MPH in Shank2 and Fmr1 deficient mice, hyperactivity in Trpm1 deficient mice may not be related to ADHD, but instead autism which is also one of the phenotypes in 15q13.3 microdeletion syndrome (Table S1) [5, 49].

Trpm1−/− mice showed prominent locomotion activities (Fig. 1A, 2E) that are not observed in mGluR6−/− mice [19]. Both mice lack of a functional ON visual transduction pathway and lack a b-wave in the ERG [3, 50] as well as having no ON response [1, 8, 9]. Additional evidence for visual impairment in Trpm1−/− mice comes from measurements of the spatial frequency and contrast sensitivity thresholds of the optokinetic response (OKR) was reported. Thresholds were reduced by approximately 10% and 30%, respectively, compared to WT [51]. While both mGluR6 and Trpm1 deficient mice lack ON BCs responses, spontaneous oscillatory firing in RGCs, the output cells of the retina, was only detected in Trpm1−/− mice [10]. An attractive idea is that these retinal oscillations might be communicated along the optic nerve to higher visual centers and result in hyperlocomotion in Trpm1−/− mice.

Visual impairment can cause behavioral abnormalities in human and mice related to hypersensitivity to sensory stimuli. Moreover, the visual cortex receives feedback projections from auditory and somatosensory cortex as well as from motor and multisensory cortices [22, 52–56]. However, Trpm1−/− mice did not show hypersensitivity to sensory stimuli, which is observed in visually impaired animals as well as in human patients [20–29], at least for thermal perception and auditory responses (Fig. 1B-D). Thus is it unlikely that the behavioral changes in Trpm1−/− mice are secondary to changes in non-visual sensory perception.

There is another possible explanation for the behavioral phenotypes in Trpm1−/− mice. The expression of mGluR6 is known to be restricted to retinal ON BCs, whereas TRPM1 is expressed in the retina and skin in humans [57–59], and a short form of TRPM1, which does not have channel function, is expressed in embryonic retinal pigment epithelial and skin in mouse [1, 2]. Thus, there is a precedent for the expression of TRPM1 outside of the retina, including alternate splice forms. We analyzed the expression of Trpm1 in the brain and detected a faint expression by qPCR thoughout most of the brain with the exception of the cerebellum (Fig. 5C). Thus, Trpm1 may be expressed in some parts of brain and the presence or lack of Trpm1 in a particular region may effect behavior. A link between TRPM1 and brain function was previously suggested by the demonstration that capsaicin-induced activation of TRPM1 channels contributed to the induction of long-term depression (LTD) in the lateral amygdala, which was specifically mediated by group I mGluRs and via interactions with another member of the TRP family, TRPC5 [60].
Deficiency of Trpm1 expression in the brain including the amygdala may lead to a neurochemical attenuation in brain function that may cause behavioral phenotypes in Trpm1 deficient mice similar to those demonstrated here.

In summary, our results are consistent with the idea that spontaneous oscillatory firing in the retina may be transmitted to the higher visual system through the optical nerve and more central projections during development and after, and as a result may modify the function and structure of the brain leading to the observed behavior changes. An alternate but not mutually exclusive view is that the lack of expression of Trpm1 in the brain may change the distribution of biogenic monoamines and behaviors in Trpm1−/− mice. Irrespective of the mechanism, this is the first report to implicate TRPM1 loss in 15q13.3 microdeletion syndrome. Further experiments will be required to determine if retinal dysfunction causes the alteration in the brain, or whether there is a specific contribution of Trpm1 in certain brain regions.

**Abbreviations**

TRPM1  
melanoma-related transient receptor potential,  
mGluR6  
metabotropic glutamate receptor 6  
BC  
bipolar cell  
CSNB  
congenital stationary night blindness  
ERG  
electroretinogram  
MB  
megabase  
BP  
break-point  
MTMR10  
myotubularin related protein 10  
FAN1  
FANCD2 And FANCl associated nuclease 1  
KLF13  
Kruppel-like factor 13  
OTUD7A  
OTU deubiquitinase 7A  
CHRNA7  
cholinergic Receptor Nicotinic Alpha 7 Subunit  
ADHD
attention-deficit hyperactivity disorder
EEG
electroencephalogram
RGC
retinal ganglion cell
MEA
multielectrode array
WT
wild-type
CS
conditioned stimulus
US
unconditioned stimulus
MPH
methylphenidate
HPLC
high performance liquid chromatography
ECD
electrochemical detector
cDNA
complementary DNA
RT
reverse transcribed
qPCR
quantitative polymerase chain reaction
ELISA
enzyme-linked immuno sorbent assay
ANOVA
analysis of variance
DA
dopamine
NA
noradrenaline
5-HT
5-hydroxytryptamine (serotonin)
OKR
optokinetic response
LTD
long-term depression
DOPAC
3,4-dihydroxyphenylacetic acid
3-MT
3-methoxytyramine
HVA
omovanillic acid
Ad
adrenaline
NM
normetanephrine
MHPG
3-methoxy-4-hydroxyphenylglycol
5-HIAA
5-hydroxyindoleacetic acid

Declarations

Ethics approval and consent to participate

All experimental procedures and housing conditions for animals were approved by Institutional Animal Care and Use Committee of National Institute for Physiological Sciences, Fujita Health University and Ritsumeikan University.

Consent for publication

Not applicable.

Availability of data and materials

The datasets generated and/or analysed during the current study are available in the [Mouse Phenotype Database] repository, [http://www.mouse-phenotype.org/]

Competing interests

The authors declare that they have no competing interests.

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Authors' contributions

TH contributed to data acquisition and analysis of brain extraction, HPLC-ECD and RT-qPCR and ELISA, and co-wrote the manuscript. SI and TY contributed to data acquisition of behavior test battery. SH, KT and TM supervised behavior test battery and performed part of them. KT and CK designed this projects and write this manuscript. All authors read and approved the final manuscript.

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Tables

Due to technical limitations, table 1 is only available as a download in the Supplemental Files section.

Figures

**Figure 1**

Physical characteristics of Trpm1−/− mice (A) Daily activity averaging 6 days in the home cage; n = 22 for both genotypes. (B) latency to the first response in the hot plate test. n = 24 for both genotypes. (C, D)
Acoustic startle response/prepulse inhibition tests; (C) the amplitude of the startle response to the 110 and 120 dB acoustic stimuli, (D) the percentage of prepulse inhibition at the 74 and 78 dB prepulse sound level. n = 24 for both genotypes. *P < 0.05; repeated measures two-way ANOVA (A), Student t test (B-D).

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Figure 2

Locomotor activity and anxiety-like behavior of Trpm1−/− mice (A-D) Light/Dark transition test; (A) the total distance traveled, (B) time spent in light, (C) the number of transitions, (D) the latency of opposite side. n = 24 for both genotypes. (E–H) Open field test; (E) the total distance traveled, (F) time spent in center of the field, (G) The number of vertical activities, (H) The number of stereotypies. n = 24 for both genotypes. (I) The total distance traveled with treatment of MPH. n = 9 for WT + Saline, n = 9 for WT + 3 mg/kg MPH.
mg/mL MPH, n = 12 for WT + 10 mg/mL MPH, n = 11 for Trpm1−/− + saline, n = 12 for Trpm1−/− + 3 mg/mL MPH, n = 12 for Trpm1−/− + 10 mg/mL MPH. (J–M) Elevated plus maze test; (J) the total distance traveled, (K) time spent on open arms, (L) number of entries into open arms, (M) and number of entries. n = 24 for both genotypes. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; two-way ANOVA followed by Tukey’s multi comparison test (A–D), repeated measures two-way ANOVA (E–H), repeated measures two-way ANOVA followed by Tukey’s multi comparison test (I), Student’s t test (J, L, M), Welch’s t test (K)
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Figure 3

Social interaction of Trpm1−/− mice (A-E) Social interaction in novel environment; (A) the total distance traveled, (B) total duration of contact, (C) total duration of active contact, (D) the number of contacts, (E) mean duration per contact. n = 12 for both genotypes. (F-I) Social preference; (F) the total distance traveled, (G) average speed, (H) time spent in each chamber, (I) social preference (calculated as the ratio of time spent in stranger chamber to all chamber). n = 24 for both genotypes. (J-M) Social novel preference; (J) the total distance traveled, (K) average speed, (L) time spent in each chamber, (M) novel preference (calculated as the ratio of time spent in stranger chamber to all chambers; n = 24 for both
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**Social interaction test (novel environment)**

- **A** Total distance traveled (cm)
- **B** Total duration of contact (sec)
- **C** Total duration of active contact (sec)
- **D** Mean duration per contact (sec)
- **E** Counts of contact

**Social interaction test (social preference)**

- **F** Total distance traveled (cm)
- **G** Average speed (cm/s)
- **H** Time spent around cage (sec)
- **I** Social preference index (%)

**Social interaction test (Novel preference)**

- **J** Total distance traveled (cm)
- **K** Average speed (cm/s)
- **L** Time spent around cage (sec)
- **M** Social novelty index (%)

**Figure 3**

Social interaction of Trpm1−/− mice (A-E) Social interaction in novel environment; (A) the total distance traveled, (B) total duration of contact, (C) total duration of active contact, (D) the number of contacts, (E) mean duration per contact. n = 12 for both genotypes. (F-I) Social preference; (F) the total distance traveled, (G) average speed, (H) time spent in each chamber, (I) social preference (calculated as the ratio of time spent in stranger chamber to all chamber). n = 24 for both genotypes. (J-M) Social novel
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**Fear conditioning test**

**Barnes maze test**

**T-maze spontaneous alteration test**

Figure 4
Cognitive function of Trpm1−/− mice (A-E) Fear conditioning test; (A) distance traveled in the conditioning phase, (B) the percentage of the freezing time in the conditioning phase. Conditioned stimulus (CS: white noise) and unconditioned stimulus (US: foot shock) were presented, (C) distance traveled during and after foot shocks, (D) the percentage of the freezing time in the context tests or cued tests one day and 30 days after conditioning, (E) distance traveled in the context tests or cued tests one day and 30 days after conditioning. n = 24 for both genotypes. (F-K) Barnes maze test; (F) the distance, (G) the count of the error, (H) the latency to first reach the correct hole above the escape box in the training, acquisition and reversal sessions, (I-K) Time spent around each hole in the probe trial conducted 24 hours (I), 1 month (J) after the last training session and 24 hours after last reversal training session (K). n = 24 for both genotypes. (L-N) T-maze forced alternation task test; (L) the percentage of correct response, (M) the latency, (N) distance traveled. n = 24 for both genotypes. *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001; repeated measures two-way ANOVA (A-H, L-N), Student t-test (I, K), Welch t-test (J).
Figure 4

Cognitive function of Trpm1−/− mice (A-E) Fear conditioning test; (A) distance traveled in the conditioning phase, (B) the percentage of the freezing time in the conditioning phase. Conditioned stimulus (CS: white noise) and unconditioned stimulus (US: foot shock) were presented, (C) distance traveled during and after foot shocks, (D) the percentage of the freezing time in the context tests or cued tests one day and 30 days after conditioning, (E) distance traveled in the context tests or cued tests one day and 30 days after conditioning.
day and 30 days after conditioning. \( n = 24 \) for both genotypes. (F-K) Barnes maze test; (F) the distance, (G) the count of the error, (H) the latency to first reach the correct hole above the escape box in the training, acquisition and reversal sessions, (I-K) Time spent around each hole in the probe trial conducted 24 hours (I), 1 month (J) after the last training session and 24 hours after last reversal training session (K). \( n = 24 \) for both genotypes. (L-N) T-maze forced alternation task test; (L) the percentage of correct response, (M) the latency, (N) distance traveled. \( n = 24 \) for both genotypes. *\( P < 0.05 \), **\( P < 0.01 \), ***\( P < 0.001 \), ****\( P < 0.0001 \); repeated measures two-way ANOVA (A-H, L-N), Student t-test (I, K), Welch t-test (J).

**Figure 5**

Abnormal Structural and biochemical change of Trpm1 −/− mice (A, B) The Brain weight at 1 month old (A), \( n = 5 \) WT, 4 Trpm1 −/− and 4 months old (B), \( n = 24 \) for both genotypes. (C) The expression of Trpm1 gene in WT brain. \( n = 4 \) for both genotypes. All amplification data were normalized with mean cycle threshold (Ct) value of WT whole brain group for \( \Delta \)Ct and normalized with \( \beta \)-actin for \( \Delta \Delta \)Ct. Trpm1 mRNA was not detectable in whole brain of Trpm1−/− mice. (D-F) Quantification of monoamine neurotransmitters in cerebellum at 4 months old; (D) dopaminergic, (E) noradrenergic, (F) serotonergic neurotransmitter and their metabolites. DOPAC: 3,4-dihydroxyphenylacetic acid, 3-MT: 3-methoxytyramine, HVA: homovanillic acid, Ad: adrenaline, NM: normetanephrine, MHPG: 3-methoxy-4-hydroxyphenylglycol, 5-HIAA: 5-hydroxyindoleacetic acid. \( n = 24 \) for both genotypes. *\( P < 0.05 \), **\( P < 0.01 \), ***\( P < 0.001 \), ****\( P < 0.0001 \); Student’s t test or Welch’s t test.
Abnormal Structural and biochemical change of Trpm1 –/– mice (A, B) The Brain weight at 1 month old (A), n = 5 WT, 4 Trpm1 –/– and 4 months old (B), n = 24 for both genotypes. (C) The expression of Trpm1 gene in WT brain. n = 4 for both genotypes. All amplification data were normalized with mean cycle threshold (Ct) value of WT whole brain group for ΔCt and normalized with β-actin for ΔΔCt. Trpm1 mRNA was not detectable in whole brain of Trpm1–/– mice. (D-F) Quantification of monoamine neurotransmitters in cerebellum at 4 months old; (D) dopaminergic, (E) noradrenergic, (F) serotonergic neurotransmitter and their metabolites. DOPAC: 3,4-dihydroxyphenylacetic acid, 3-MT: 3-methoxytyramine, HVA: homovanillic acid, Ad: adrenaline, NM: normetanephrine, MHPG: 3-methoxy-4-hydroxyphenylglycol, 5-HIAA: 5-hydroxyindoleacetic acid. n =24 for both genotypes. *P <0.05, **P <0.01, ***P <0.001, ****P <0.0001; Student’s t test or Welch’s t test.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- HorietalTable1.pdf
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