RESEARCH ARTICLE

Longitudinal analysis of ultrasonic vocalizations in mice from infancy to adolescence: Insights into the vocal repertoire of three wild-type strains in two different social contexts

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Abstract

Ultrasonic vocalizations (USV) are emitted by mice under certain developmental, social and behavioral conditions. The analysis of USV can be used as a reliable measure of the general affective state, for testing the efficacy of pharmacological compounds and for investigating communication in mutant mice with predicted social or communication deficits. Social and communication studies in mice have focused mainly on the investigation of USV emitted by neonatal pups after separation from the dam and during social interaction between adult males and females. Longitudinal USV analysis among the different developmental states remained uninvestigated. In our study, we first recorded USV from three inbred mouse strains C57BL/6N, DBA/2 and FVB/N during the neonatal stages after separation from the littermates and then during a reunion with one littermate. Our results revealed significant strain-specific differences in the numbers and categories of USV calls. In addition, the USV profiles seemed to be sensitive to small developmental progress during infancy. By following these mice to the adolescent stage and measuring USV in the three-chamber social test, we found that USV profiles still showed significant differences between these strains in the different trials of the test. To study the effects of social context on USV characteristics, we measured USV emitted by another cohort of adolescent mice during the direct social interaction test. To this end, this study provides a strategy for evaluating novel mouse mutants in behavioral questions relevant to disorders with deficits in communication and sociability and emphasizes the important contribution of genetics and experimental contexts on the behavioral outcome.
Introduction

Wild-type mice emit ultrasonic vocalizations (USV) with frequencies higher than 20 kHz, which cannot be detected by human ears. USV are produced in different situations including aversive affective states and negative situations [1–3], social interaction to enhance social bonding in juvenile [4] and adult mice [3, 5–7] and appetitive situations such as rough-and-tumble play in juvenile and courtship in adult mice [8, 9]. Moreover, pups separated from dam and littermates during their first two weeks of life emit a large number of USV to trigger the mother’s retrieval behavior [10–12] and in response to body temperature dropping, handling and specific smells. In particular, recording USV in mice has great potential to provide a model system for the investigation of communication in wild-type and mutant mice with neuropsychiatric-like phenotypes [13, 14]. Alterations of USV have been reported in several mouse models of neuropsychiatric disorders including autism [15–24] and schizophrenia [13, 25]. These studies have proved that studying USV can help to understand the mechanisms of human communication and associated disorders. Moreover, recording USV is a useful tool for testing the efficacy of pharmacological compounds [26–28] and for monitoring the general health of mice [5].

USV can be classified into ten different patterns with distinctive temporal-structural characteristics designated as frequency steps, harmonics, composite, short, complex, chevron, flat, downward, upward, two-syllable [4, 6, 29]. The specific sequential pattern of calls is suggested to convey specific information; e.g., female mice preferentially approach specific patterns of USV emitted by males [30]. However, other important factors including number, rate, duration and peak frequency of USV are still important cues used by mice to interpret the nature of the social interaction [31] and these factors are dependent on age and genetic background [32–35]. The detailed characterization of USV properties requires many steps including recording, call detection, call type classification and clustering [36].

In the present study, we compared three inbred mouse strains, C57BL/6N, DBA/2 and FVB/N, in respect to the basic USV parameters including the number of calls, frequency, duration and amplitude as well as the call structure based on a previously published USV category scheme [6]. We have selected these strains since they are typically used as background strains for breeding genetically modified mice and are able to show robust social interaction ability [37]. Although they represent the most used standard controls for many behavioral studies, each strain can display different characteristics under certain behavior paradigms [38–42]. For example, DBA/2 mice perform poorly in several hippocampus-dependent contextual and spatial learning tasks compared to C57BL/6 mice [43]. Moreover, DBA/2 mice have a decreased locomotor activity in the open field task compared to FVB/N mice [40]. On one hand, the FVB/N strain is preferable for transgenic analysis because its fertilized eggs contain large and prominent pronuclei, which facilitates the microinjection of DNA [44]. On the other hand, this strain shows high anxiety, aggressive behavior and disturbed circadian rhythm [45]. To this end, we first assessed possible differences in the USV properties between neonatal pups of the selected mouse strains after their separation from the dam and littermates. Moreover, we checked the effects of small developmental progress during infancy as well as the reunion with a littermate on the patterns of the USV emissions. Furthermore, we measured USV from the same mice at the adolescent stage in the three-chamber social test [46]. Additionally, we tested the effects of social context on the USV patterns by measuring USV of the three selected mouse strains in the direct social interaction test.

This study offers a deeper insight into the discrete functions, individual calls and calls sequences within the diversity of the vocal repertoire, which can be used for further understanding of the acoustic signals associated with social interactions in mice and in testing the
role of motivational states in the emission of USV. Moreover, it can be used as a reference for testing mice with predicted communication deficits or for testing the efficacy of drugs treating social deficits by considering the crucial importance of the developmental state, social context and genetic background in the regulation of different behaviors.

Materials and methods

Animals and housing conditions

C57BL/6N, DBA/2 and FVB/N female and male mice were purchased from Janvier Laboratories (Le Genest-Saint-Isle, France) at the age of 8 weeks. After ten days, female and male mice were paired. Then, females were isolated from males after ten days of pairing, housed individually and inspected daily for pregnancy and delivery. The day of birth was considered as postnatal day (P) 0. Pups were marked by paw tattoo at P3 using a non-toxic animal tattoo ink (Ketchum permanent tattoo green paste, Ketchum Manufacturing, Inc., Brockville ON Canada). The ink was injected subcutaneously through a 25 gauge hypodermic needle tip into a toe of a paw. At P20, mice were weaned, and the littermates from the same sex were grouped in the same cage (three to four per cage) (Fig 1A). All animals had ad libitum access to food and water under a standard 12 h light/dark cycle (7:00 pm—7:00 am) with a regulated ambient temperature of 22 °C +/- 1 and at a relative humidity of 40–50%. All experiments were conducted in strict compliance with the National Institutes of Health Guidelines for the Care and Use of Laboratory Animals and approved by the Regional Council in Karlsruhe (Regierungspräsidium Karlsruhe, Germany; Animal Ethic Protocol 35–9185.81/G-129/15).

USV analysis during infancy after separation and reunion with a littermate

USV were recorded in infant mice at P4, P8 and P12. At each developmental stage, the testing pup was separated from the littermates and placed in an empty glass container (6 x 9.5 x 7.5 cm) with the floor covered with bedding material. The container was placed inside a box with the walls covered with Styrofoam to attenuate surrounding noise. USV were recorded for five min (separation trial 1) which was immediately followed by placing one littermate next to the testing pup for an additional five min (reunion trial 2) (Fig 1B). Before placing the pup in the glass container, it was weighed, and the body temperature was measured by gently applying a thermal probe on the belly (digital thermometer Bosotherm medical, Boso, Jungingen, Germany). After the USV measurement, the pup was immediately returned to its home cage. The same paired pups were used among the 3 days of recordings. The numbers of pups used in this study were 7 for C57BL/6N (2 males and 5 females), 8 for DBA/2 (4 males and 4 females) and 11 for FVB/N (6 males and 5 females).

USV analysis in adolescent mice during the three-chamber social test

USV were recorded among adolescent mice between P24 and P29. The pairs of tested mice were from the same sex and litter. In our adapted protocol, the three-chamber social test was designed to assess the social ability with a familiar conspecific and with a stranger adult mouse. USV were measured from all chambers in parallel using one microphone placed 30 cm above the arena floor of the middle chamber. The three-chamber social apparatus was made up of a transparent plexiglas box (20 x 61 x 40 cm) with three compartments including sliding doors (5 x 8 cm) between the compartments. The top of each compartment was closed with a plexiglas top cover. A wire mesh cylinder was placed in the left and right compartments. The test was subdivided into three trials, 10 minutes each, with a 15 min intertrial interval. In trial 1 (habituation), the testing mouse, isolated from the home cage for 24 h, was introduced into the
apparatus with free access to all compartments (Fig 3A). In trial 2 (social recognition), a littermate mouse was placed into the right wire mesh cylinder and the subject mouse was allowed to explore all compartments. In trial 3 (social discrimination), an unfamiliar adult mouse from a different strain was placed into the left mesh wire cylinder: one DBA/2 mouse for C57BL/6N strain; one FVB/N for the DBA/2 strain and one C57BL/6N mouse for the FVB/N strain, and the subject mouse was again allowed to explore freely all compartments. The numbers of adolescent mice used were 7 for C57BL/6N (2 males and 5 females), 8 for DBA/2 (4 males and 4 females) and 11 for FVB/N (6 males and 5 females).

**USV analysis in adolescent mice during the direct social interaction test**

USV were recorded among adolescent mice between P24 and P29. The pairs of tested mice were from the same sex and the same litter. One adolescent mouse was isolated from the home cage for 24 h in a separate colony housing room. The social test included 2 trials with trial 1 serving as a habituation phase by placing one mouse from the colony into a white acryl open-
field box (40 × 40 × 40 cm) for 2 min. In trial 2, the sibling mouse was placed gently next to the isolated mouse for 5 min (social interaction trial). USV were recorded along both trials using a microphone placed 30 cm above the arena floor (Fig 4A). The numbers of tested adolescent mice were 25 for C57BL/6N (14 males and 11 females), 14 for DBA/2 (7 males and 7 females) and 18 for FVB/N (7 males and 11 females).

The USV recording and analysis

The USV recording and analysis were conducted with an equipment and a software from Avisoft Bioacoustics (Berlin, Germany). The acoustic signals were recorded, amplified and digitized at 250 kHz with a 16-bit resolution by the Ultrasound Gate 416 Hb USB audio device and ultrasonic condenser microphones CM16/CMPQ were placed 30 cm above the floor of the testing area. For acoustical analysis, signals saved as wave-files were transferred to the Avisoft SLabPro version 5.2.07 (Avisoft Bioacoustics, Berlin, Germany), and a Fast Fourier Transformation was conducted (512 FFT-length, 100% frame Hamming window and 75% time window overlap). Spectrograms displaying frequency versus time were produced with a 488 Hz frequency resolution and 0.512 ms time resolution. The number of calls and a wide range of acoustic parameters were automatically measured using the automatic whistle tracking algorithm with a minimum duration of 3 ms and a hold time of 10 ms. A cut-off filter of 15 kHz and a post filter (minimum duration of 1 ms and entropy of 0.5) were applied. Accuracy of the number of calls was verified manually and existing extraneous noise was removed from the sonograms. USV categories were defined manually according to a previously published classification scheme [6].

Statistical analysis

Prism 6 software (GraphPad Software) and Microsoft Office Excel software were used for data analysis. For all measurements, data were calculated and presented as mean ± SEM. The two-way ANOVA test with strain and sex as the two factors was performed to assess statistical significance. According to Bonferroni correction for multiple testing, P-values were corrected by multiplying the original p-values by the number of hypotheses (6 hypotheses for pups and adolescent mice in the direct social interaction test; 9 hypotheses for adolescent mice in the three-chamber social test). Corrected P-values ≤ 0.05 were considered significant.

Results

Body weight and temperature during infancy

In order to monitor the developmental progress in infant mice, the body weight was measured at postnatal day (P) 4, 8 and 12. All strains gained weight during the first 12 days after birth. However, DBA/2 pups were significantly smaller compared to C57BL/6N and FVB/N pups at the three investigated developmental stages (S1A Fig). Infant mice are unable to control their body temperature during the first days of life and therefore tend to suffer from hypothermia which is a major cause of infant death in nature. In order to control body temperature and ensure the standardization of the testing procedures, the body temperature was measured pre- and post-experimentally at all measurement days. At P4, C57BL/6N pups lost on average $3.46 \pm 0.64$ °C, DBA/2 pups 1.75 ± 2.04 °C and FVB/N pups 2.89 ± 0.55 °C after testing (S1B Fig). The temperature loss diminished for C57BL/6N mice at P8 indicating a fast improvement in the thermoregulation of the developing pups. However, for DBA/2 and FVB/N, the temperature loss diminished later at P12 (S1B Fig).
Ultrasonic vocalization in pups

Effect of separation (trial 1) on the number of calls. Separated pups tend to emit USV which are suggestive for mother seeking behavior. At P4, there was no difference in the number of calls between C57BL/6N, DBA/2 and FVB/N pups (Fig 1C). At P8, the number of calls emitted by FVB/N pups was significantly lower than DBA/2 (p = 0.034) and lower than both C57BL/6N (p = 0.032) and DBA/2 (p = 0.015) at P12 (Fig 1C).

By comparing the three developmental states during infancy, we found that FVB/N pups emitted more calls at P4 than at P8 and P12 (p = 0.0084 and 0.0074, respectively) (S2 Fig). In contrast, C57BL/6N and DBA/2 pups did not show any difference in the number of calls between P4, P8 and P12 (S2 Fig).

Effect of reunion with a littermate (trial 2) on the number of calls. After a reunion with a littermate, the number of USV calls emitted by C57BL/6N pups slightly increased at P4 and P8. In contrast, the number of calls did not change at these stages for the DBA/2 strain. For the FVB/N strain, there was a slight increase in the number of calls at P4, but not at P8 or P12. Reunion with a second pup did not lead to a diminution of calls in all strains except for DBA/2 at P12 (p = 0.024) (Fig 1C).

Effect of separation and reunion on the call characteristics. Additionally to the number of calls, the acoustic characteristics like duration, frequency as well as the amplitude of the emitted USV were analyzed. The duration of calls ranged from 30 to 80 ms and varied across strains and developmental states in the separation trial 1 (Fig 1D). In all strains, the duration of calls declined significantly during infant development from P4 to P12 in both the separation and reunion trials (S2 Fig). The C57BL/6N pups showed the lowest duration across all developmental states in both trials (P4: trial 1: p = 0.008 vs DBA/2 and 0.007 vs FVB/N; trial 2: p = 0.0094 vs DBA/2 and 0.0068 vs FVB/N) (P8: trial 1: p = 0.08 vs DBA/2 and 0.1 vs FVB/N; trial 2: p = 0.12 vs DBA/2 and 0.14 vs FVB/N) (P12: trial 1: p = 0.02 vs DBA/2 and 0.009 vs FVB/N; trial 2: p = 0.037 vs DBA/2 and 0.054 vs FVB/N), while the FVB/N pups showed the longest duration at P4 during both the separation and reunion trials (Fig 1D). By comparing trial 2 to trial 1, no gross changes in calls durations were present in any of the three tested strains (Fig 1D).

The frequency of USV calls ranged from 65 to 80 kHz throughout all tested postnatal stages and was not affected by the presence or absence of a littermate in all strains (Fig 1E). The DBA/2 pups emitted calls at a lower frequency than that emitted from the C57BL/6N and FVB/N pups with significance observed at P12 in both trials (Trial 1: p = 0.016 vs C57BL/6N and 0.017 vs FVB/N; trial 2: p = 0.013 vs C57BL/6N and 0.009 vs FVB/N) (Fig 1E). FVB/N pups showed an increasing call frequency between P4 and P12 in trial 2 (S2 Fig).

Measuring the amplitude of USV calls revealed that C57BL/6N pups emitted softer calls than DBA/2 and FVB/N pups at P12 in both trials (Trial 1: p = 0.005 vs DBA/2 and 0.02 vs FVB/N; trial 2: p = 0.007 vs DBA/2 and 0.014 vs FVB/N) (Fig 1F). These pups also displayed a significant upward trend of amplitude across the developmental states with calls becoming softer with age (S2 Fig). In contrast, the USV calls emitted by DBA/2 pups at P4 were softer than at P8 and P12 (S2 Fig). For all strains, the amplitude of USV did not change when a second littermate joined the first pup (Fig 1F).

Call categories in pups. According to their spectrographic shape, frequency and duration, USV were classified into ten categories [6]. These ten categories have been detected in the three strains and are presented with their relative frequencies and durations in Fig 2A. Our dissection of USV showed that the call categories were unique for each strain at specific developmental stages. The pattern emitted the most by C57BL/6N pups at P4 and P8 was the two-syllable pattern (25–30% of all patterns) (Fig 2B). At P12, C57BL/6N mice emitted mostly the
Fig 2. Analysis of USV categories of C57BL/6N, DBA/2 and FVB/N pups. (A) Representative sonograms of different call categories with time in milliseconds (ms) represented on the X-axis and frequency in kHz on the Y-axis. (B1-10) Percentages of different call categories emitted by each strain at P4, P8 and P12 during separation (trial 1) and reunion with a littermate (trial 2); n = 7 for C57BL/6N (2 males and 5 females), n = 8 for DBA/2 (4 males and 4 females) and n = 11 for FVB/N (6 males and 5 females), all data as mean ± SEM (two-way ANOVA, *p< 0.05, **p< 0.01, ***p< 0.001).

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Fig 3. USV analysis of adolescent mice from C57BL/6N, DBA/2 and FVB/N strains during the three-chamber social test. (A) A schematic drawing of the experimental setup for USV recording in adolescent mice. USV were recorded for 10 min in the apparatus as a habituation phase (trial 1), followed by a social contact phase for 10 min with a same-sex littermate on the right compartment (trial 2) followed by another trial for 10 min with an adult unfamiliar mouse from another strain (trial 3). The intertrial interval between trials was 15 min. (B-E) USV analysis including number (B), duration (C), frequency (D) and amplitude (E) of calls during the three trials. (F) Percentages of different call categories emitted by each strain during each trial; n = 7 for C57BL/6N (2 males and 5 females), n = 8 for DBA/2 (4 males and 4 females) and n = 11 for FVB/N (6 males and 5 females), all data as mean ± SEM (two-way ANOVA, *p< 0.05, **p< 0.01).
Fig 4. USV analysis of adolescent mice from C57BL/6N, DBA/2 and FVB/N strains during the direct social interaction test. (A) A schematic drawing of the experimental setup for USV recording in adolescent mice. USV were recorded for 2 min in a new arena as a habituation phase (trial 1), followed by a social contact phase for 5 min with a same-sex littermate (trial 2). (B-E) USV analysis including number (B), duration (C), frequency (D) and amplitude (E) of calls at P24 during the habituation and social contact phases. (F) Percentages of different calls categories emitted by each strain during social contact phase; n = 25 for C57BL/6N (14 males and 11 females), n = 14 for DBA/2 (7 males and 7 females) and n = 18 for FVB/N (7 males and 11 females), all data as mean ± SEM (two-way ANOVA, *p<0.05, **p<0.01, ***p<0.001).

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upward pattern (≈ 25%) (Fig 2B). For DBA/2 pups, most of the calls produced at P4 and P8 were chevron (30–50%) (Fig 2B10), but upward at P12 (≈ 35%) (Fig 2B4). The most emitted patterns by FVB/N pups were two-syllable at P4 (≈ 30%) (Fig 2B1), frequency steps at P8 (≈ 25%) (Fig 2B5) and flat at P12 (≈ 30%) (Fig 2B9). The comparison of the individual call categories used by separated (trial 1) and reunited (trial 2) pups did not reveal a socially-induced alteration of the call categories with the exception of a significant decrease in harmonic calls by DBA/2 P4 pups in trial 2 and its increase by FVB/N pups at P12 (Fig 2B6).

**USV in adolescent mice in the three-chamber social test**

The USV profiles of adolescent mice were also investigated using the three-chamber social interaction test (Fig 3A). Analyzing the number of calls within the 10 min habituation phase (trial 1) of the three chamber box revealed no differences between the three strains (Fig 3B). By presenting a familiar littermate in the right compartment (social recognition trial 2), the number of calls did not change significantly. However, the additional presence of an adult unfamiliar mouse (social discrimination trial 3) in the left compartment elicited an increase in the number of calls for all strains without reaching significance (Fig 3B). In all trials, FVB/N mice showed the longest calls durations with significant results in trial 1 (p = 0.008 vs C57BL/6N and 0.014 vs DBA/2 and trial 2 (p = 0.0074 vs C57BL/6N and 0.009 vs DBA/2) (Fig 3C). Interestingly, the frequency of the USV calls was significantly lower in all strains in the presence of the adult stranger mouse in trial 3 (C57BL/6N: p = 0.031 vs trial 1 and 0.054 vs trial 2; DBA/2: p = 0.037 vs trial 1 and 0.22 vs trial 2; FVB/N: p = 0.02 vs trial 1 and 0.034 vs trial 2) (Fig 3D). The amplitude of USV calls varied across trials and strains and tended to decrease with the introduction of the stranger mouse (Fig 3E). When considering the gender, no differences in the number, duration, frequency or amplitude of the calls were detected between male and female adolescent mice from all tested strains during all trials (S3 Fig).

The comparison between the USV calls emitted by adolescent mice in trials 1 and 2 of the three-chamber social test with the calls emitted in trials 1 and 2 during infancy revealed a significant decrease in the number of calls and their average durations in all strains during adolescence (Trials 1 and 2 of the three-chamber social test during adolescence vs trials 1 and 2 at P4: p<0.001 for all tested strains for both number and duration of calls). In contrast, the frequency of calls at the adolescent stage was significantly higher compared with infancy at P4 (Trial 1: p<0.001 for all tested strains; trial 2: C57BL/6N: p = 0.0096, DBA/2: p = 0.041 and FVB/N: p = 0.0092). Because of the difference of the arenas sizes and shapes, amplitudes of calls cannot be directly compared between adolescent and infant stages.

By analyzing the USV patterns, DBA/2 mice presented less heterogeneity in the USV categories than both FVB/N and C57BL/6N mice. DBA/2 mice emitted mainly downward calls during the habituation (trial 1) and social recognition (trial 2) (50%) and flat calls (40%) when the unfamiliar mouse was presented (trial 3) (Fig 3F). In all trials, the calls with a flat shape were the most frequent calls emitted by C57BL/6N mice (40%) (Fig 3F). For FVB/N mice, the USV patterns of trial 1 and trial 2 were more similar compared to trial 3 and represented by complex (> 35%), downwards (> 25%), flat (> 10%) and composites (> 10%) calls (Fig 3F). Interestingly, the harmonic calls (5%) were emitted only by FVB/N mice in trial 3. Moreover, the short calls were emitted only by C57BL/6N mice in trial 3.

**Ultrasonic vocalizations in adolescent mice in the direct social interaction test**

**Number of calls and calls characteristics.** In this task, the USV profiles were measured during a direct social approach of 2 adolescent littermates of the same gender. An adolescent
mouse was isolated from the litter in a home cage. After 24 h, the isolated mouse was put in a new arena for 2 min for habituation (habituation trial 1) followed by a social interaction for 5 min with a same-sex littermate (social interaction trial 2) (Fig 4A). In trial 1, C57BL/6N marginally emitted USV in contrast to DBA/2 and FVB/N (Fig 4B). In trial 2, adolescent mice from the three tested strains emitted USV when they were reunited with a littermate. Noticeably, the numbers of calls produced by the FVB/N mice were significantly more than the calls emitted by the C57BL/6N and DBA/2 mice (p = 0.003 vs C57BL/6N and 0.0007 vs DBA/2) (Fig 4B). In addition, the duration of calls was significantly longer in FVB/N compared with C57BL/6N mice (p = 0.0094 vs C57BL/6N and 0.012 vs DBA/2) (Fig 4C), but the peak frequencies were similar between all three strains (Fig 4D). FVB/N mice emitted louder calls than C57BL/6N and DBA/2 mice (Fig 4E). When considering the gender, no difference in the number, duration, frequency or amplitude of the USV calls was detected between male and female adolescent mice from the three tested strains (S4A Fig).

**Call categories.** The patterns of USV calls produced by the three tested strains during trial 2 of the direct social interaction test were investigated. The highest percentage of USV pattern emitted by C57BL/6N mice was short (≈ 30%) (Fig 4F). For DBA/2 and FVB/N adolescent mice, the highest percentages of calls patterns were flat (≈ 40%) and upward (≈ 30%), respectively (Fig 4F). The heterogeneity of the calls patterns in DBA/2 mice was less compared with the other two strains, similarly to the results obtained from the three-chamber social test. Moreover, DBA/2 mice did not emit any harmonic or frequency steps calls (Fig 4F). In addition, there was no gross difference in call categories between males and females mice except in DBA/2 mice (S4B Fig).

**Discussion**

Because USV are extensively used nowadays in mouse models of neurological diseases including verbal dyspraxia, depression, autism and stuttering [47], their detailed analysis in wild-type mice compared to different strains at different developmental states is a must in order to better understand their basic characteristics under different conditions. This, in turn, can direct the choice of the best strain during a specific developmental state for the investigation of neuropsychiatric disorders. Although many studies have already shown that USV properties vary across strains [4, 30, 48, 49], social contexts [50–52] and between different ages especially pups and adults [53, 54], all of these parameters were not thoroughly compared in one single study. In the present comprehensive study, we analyzed many USV characteristics including the number, duration, frequency and amplitude in addition to the categories of calls in three wild-type mouse strains C57BL/6N, DBA/2 and FVB/N under different contexts in a longitudinal manner following the same mice from infancy to adolescence, in order to provide a more unified data set.

During infancy, when pups were separated from their dam and littermates, the USV properties varied across postnatal days and strains indicating an influence of developmental state and genetic background on the ultrasound emissions. When comparing the number of calls at P4, no difference was found between the strains. In contrast, differences in all USV characteristics can be observed at P8 and P12, suggesting that the differentiation in communication behavior might occur after P4. FVB/N pups vocalized longer USV, and the number of their calls decreased with age compared to C57BL/6N and DBA/2. Although the literature suggests a strain-specific peak in vocalization rate at around P8 [13, 48], this was not observed in the current study. The duration of calls decreased from P4 to P12 in all tested strains, which is consistent with previous studies that showed that the duration of calls decreases with ages [47, 54, 55]. Compared with other tested strains, The DBA/2 pups consistently displayed lower frequency and higher intensity calls.
The USV properties did not grossly differ between the contexts of pups separation and reunion with a littermate, which may indicate that the presence of a littermate is not sufficient to change the affective state of the separated pup. However, a current limitation of the spectrograms is the inability to distinguish which of the two pups vocalized more or which remained silent. Interestingly, studies in rats with anesthetized littermates introduced to the separated pups illustrated a reduction in vocalization rate [56]. Considering the fact that neonate mice are deaf till P10 [11], two assumptions can be made: (i) Both pups were unable to interact and hence both produced distress calls resulting in significantly increased number of USV or (ii) pups were able to establish contact (tactile or olfactory perception) leading to a reduction in ultrasound emissions. However, none of these hypotheses could be confirmed by the observed results leaving the open question whether contact between two conscious littermates changes the ultrasound emission.

The body weights and temperatures of pups can have an effect on the USV characteristics [57]. Therefore, these parameters were checked prior to experimentation and revealed reduced body weights of DBA/2 pups compared with C57BL/6N and FVB/N pups. However, the influence of smaller body weight on temperature control and ultrasound emission was excluded since all strains lost body temperature during separation at P4 and P8. This result was also confirmed by Scattoni et al. excluding the influence of body size on the vocalization rate [6].

Following these mice to the adolescent stage, the USV category profiles illustrated similarities across three different trials in the three-chamber social test. However, the first two trials displayed more similarities compared to the third one, especially with the number of calls in the three tested strains. Although we cannot exclude that some of the recorded calls were obtained from the littermate in trial 2 or the adult unfamiliar mouse in trial 3, we can reasonably consider that most of USV calls were emitted by the tested mouse since we did not observe overlapping vocalization streams, consistent with a single vocalizer and as the host mouse has been shown to be the main emitter of USV [58]. As the unfamiliar adult mice in trial 3 were from different strains: DBA/2 for the tested C57BL/6N strain; FVB/N for the tested DBA/2 strain and C57BL/6N for the tested FVB/N strain, the consistent differences in USV characteristics between trial 3 and both trial 1 and 2 in all tested strains may not be a consequence of individual variability and likely indicates an influence of the territorial interaction with an unfamiliar mouse versus a normal social interaction with a littermate. This confirms that mice are able to adapt their vocal behaviors in a context-dependent manner in order to convey different information [59]. The frequency of calls from all tested strains was very high during the habituation trial and decreased in the social recognition trial when there was a mouse in the arena, especially when it was an unfamiliar one from a different strain. The low-frequency calls during the exposure to an adult stranger mouse might be similar to the 22 kHz alarm calls emitted by rats in an aversive context [60, 61] [62]. Moreover, it has been shown that high-frequency calls were recorded during social interaction tasks, while the lowest frequency calls were recorded under stressful situations [3]. The duration of the USV calls in trial 2 and trial 3 (social trials, littermate or unfamiliar mouse, respectively) was similar to trial 1 (non-social trial, habituation) although it has been shown that the calls emitted in a social interaction task were longer in comparison with the calls emitted in other non-social situations [3].

A comparison of the USV characteristics between the adolescent and the infant stages revealed a decrease in the number of calls and their durations in all strains during adolescence, despite the longer duration of the experiment (5 min in the infant stage and 10 min in the adolescent stage). This suggests that the role of USV in survival as shown by calling the dam after separation is more vital than their role in social or territorial interactions. Additionally, adolescent mice from all tested strains emitted higher frequency calls in the habituation phase (trial 1) and social contact phase (trial 2) in the three-chamber social test than while they were
infants after separation from the dam (trial 1) and reunion with a littermate (trial 2). This also confirms that the low-frequency calls indicate highly stressful situations which may present during infancy, in particular during the risk of hypothermia. However, apart from stress, other factors including the social context and normal development might explain the high-frequency calls during adolescence compared with the low-frequency calls during infancy.

To test the effects of the social context on the USV characteristics, we also recorded USV from another cohort of mice during the direct social interaction test. We did not perform this test on the same mice used during the three-chamber social test to avoid a previous social experience which can have a major effect on the pattern of the USV emission. In trial 1, when the mice were alone in an open arena, they hardly emitted any USV. In contrast, they emitted many USV calls in trial 2, when they were socially active with same-sex littermates, indicating a positive correlation between USV and the social approach. Compared with the three-chamber social test, the three tested strains emitted more ultrasound signals in the direct social test during the social recognition trial, although the test lasted shorter. The low vocalization rates observed in the three-chamber social test can be explained by the limited social interaction, given the presence of a barrier with the wire mesh cylinder and the bigger arena area, which would leave the tested mouse with less option to socialize directly. In contrast, a comparison of the duration and frequency of calls between the social contact phases (trial 2) in both the direct social interaction test and the three-chamber social test revealed no differences. In the direct social interaction test, the USV properties were different between different strains, with the FVB/N mice showing the highest number and duration of calls with reduced frequency and amplitude. Moreover, call categories were unique for each strain which confirms that the USV calls emitted by mice are a communication tool during different types of situations or contexts. Therefore, the USV patterns of adolescent mice can serve as indicators for strain-specific social approaches.

During both social interaction experiments, DBA/2 adolescent mice showed less USV calls, which can be partially explained by their hearing disability. This strain shows a high-frequency hearing loss at as early as 3 weeks of age caused by $Cdhn_{23753G>A}$ [63] and $Fscn_{2326G>A}$ alleles [64] genetic variants. However, the significant role of optimal hearing ability on the emission of USV is still controversial. Adult mice from several strains have been shown to emit USV during courtship without an auditory feedback [65]. Moreover, previous studies have revealed that adult DBA/2 mice can emit USV to the same degree as other strains including CD1 and 129SV [58]. Therefore, other factors may have caused the reduced number of USV in juvenile DBA/2 mice. Therefore, C57BL/6N, DBA/2 and FVB/N should be further investigated in various social contexts during adulthood to check whether the reduced number of calls in DBA/2 mice is constant at different developmental stages.

In all experiments performed on the three tested strains during infancy and adolescence, we did not observe any significant differences between males and females in the number of calls or other USV characteristics consistent with studies performed on adult male and female mice [32] [66] [67] [68], indicating that the social communication between mice of the same sex could be considered as different from the reproductive context between opposite sex as shown previously [69].

Mice did not show clear visual cues of their vocal behavior during social interactions. Therefore, we were not able to confirm that the produced USV calls were mainly emitted by the tested mice during infancy or adolescence, which was the main limitation of our study. However, as the tested mice were the host mice that were habituated in the experimental devices before the introduction of a littermate or a stranger mouse, these mice were considered the main emitters of USV, as shown previously [58]. Additionally, in most of our spectrograms, we did not observe overlapping vocalization patterns, suggesting a single vocalizer. To
avoid this dilemma in the future, the USV measurement should be accompanied by an accurate social behavior analysis using tracking software for better synchronization of the calls and confirmation of the emitter mouse. A recent study has proposed a new approach for a high-precision spatial localization of the mouse USV during social interaction test, which would allow a proper understanding of social vocalizations [66]. A detailed examination of the temporal and sequencing aspects of vocalizations bouts could further provide an insight into how particular patterns of vocalizations can confer different meanings.

The long-term study following the same individuals from the little to the juvenile age should reduce the variability within the study groups. Therefore, it could also be conceivable to measure USV every single day from P4 to the late adolescence at P40 to give more precise insights over the entire developmental progression. However, this may represent a high burden on mice, and the habituation effects due to the daily manipulation might additionally influence the results. To this end, we propose, in addition to the paradigm of pup isolation test and reunion, additional measurements of USV during adolescence which can be performed weekly until P40.

Here, we report for the first time a comprehensive analysis of USV by providing all calls characteristics from three different wild-type mouse strains in a longitudinal manner starting from infancy to adolescence. As observed for other behaviors with these three inbred mice, differences in USV characteristics can also be detected at different ages and social contexts using our paradigms. Together, this study gives a more complex picture of all variables that can modify USV. These variables need to be considered during the design and analysis of mouse models of mental disorders and/or communication deficits.

Outlook

This study focused on a detailed USV analysis of different strains during isolation and reunion of pups and two social interaction contexts in adolescent mice. Other commonly used social interaction tests include social conditioned place preference, partition and operant condition tests [70]. The effects of these tests on the USV characteristics should be further investigated along with other kinds of behaviors in which mice emit USV like nursing, play and aggression. Moreover, the synchronization of different call characteristics with a specific type of social interactions is of great importance, in order to assign calls sequencing patterns with specific types of behaviors. This, in turn, will facilitate the examination of acoustic features that are related to specific biological states and social outcomes. In addition, it can determine the type of information encoded in the different call types and their functions [4, 29, 71, 72]. This can be carried out with the recent advancement of USV analysis from a manual to a total automated analysis using the new Mouse Ultrasonic Profile ExTraction (MUPET) technique [73]. With this technique, the USV repertoire can now be measured and compared, while providing an automated time-stamp of individual USV events. Such analysis tools will accelerate the progress of social communication studies in rodents [73].

Supporting information

S1 Fig. Body weight (A) and temperature (B) of pups during separation from the home cage at P4, P8 and P12. Body temperature was measured before separation (pre-experiment) and immediately after testing (post-experiment); n = 7 for C57BL/6N, n = 8 for DBA/2 and n = 11 for FVB/N, all data as mean ± SEM (two-way ANOVA, *p < 0.05, **p < 0.01). (TIF)
S2 Fig. Comparison of USV in infancy including the number, duration, frequency and amplitude of calls across P4, P8 and P12 during separation (trial 1) and reunion with a littermate (trial 2); n = 7 for C57BL/6N, n = 8 for DBA/2 and n = 11 for FVB/N, all data as mean ± SEM (two-way ANOVA, *p < 0.05, **p < 0.01, ***p < 0.001).

(TIF)

S3 Fig. Comparison of USV between male and female juvenile mice from the C57BL/6N, DBA/2 and FVB/N strains during the three-chamber social test. USV analysis includes the number, duration, frequency and amplitude of calls during three trials (habituation trial 1, social recognition trial 2 and social discrimination trial 3); n = 2 males and 5 females for C57BL/6N, n = 4 males and 4 females for DBA/2 and n = 6 males and 5 females for FVB/N, all data as mean ± SEM (two-way ANOVA).

(TIF)

S4 Fig. Comparison of USV between male and female juvenile mice from the C57BL/6N, DBA/2 and FVB/N strains during the direct social interaction with a littermate after 24 h of isolation. (A) USV analysis including the number, duration, frequency and amplitude of calls at P24 during the social interaction phase. (B) Percentages of call categories emitted by males and females from each strain during the social interaction phase; n = 14 males and 11 females for C57BL/6N, n = 7 males and 7 females for DBA/2 and n = 7 males and 11 females for FVB/N, all data as mean ± SEM (two-way ANOVA, *p < 0.05, **p < 0.01, ***p < 0.001).

(TIF)

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