Non-Mendelian inheritance during inbreeding of Ca\textsubscript{v}3.2 and Ca\textsubscript{v}2.3 deficient mice

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The mating of 77 heterozygous pairs (Ca\textsubscript{v}3.2[+\textpipe−] x Ca\textsubscript{v}3.2[+\textpipe−]) revealed a significant deviation of genotype distribution from Mendelian inheritance in weaned pups. The mating of 14 pairs (Ca\textsubscript{v}3.2[−\textpipe−] female x Ca\textsubscript{v}3.2[+\textpipe−] male) and 8 pairs (Ca\textsubscript{v}3.2[+\textpipe−] female x Ca\textsubscript{v}3.2[−\textpipe−] male) confirmed the significant reduction of deficient homozygous Ca\textsubscript{v}3.2[−\textpipe−] pups, leading to the conclusion that prenatal lethality may occur, when one or both alleles, encoding the Ca\textsubscript{v}3.2T-type Ca\textsuperscript{2+} channel, are missing. Also, the mating of 63 heterozygous pairs (Ca\textsubscript{v}2.3[+\textpipe−] x Ca\textsubscript{v}2.3[+\textpipe−]) revealed a significant deviation of genotype distribution from Mendelian inheritance in weaned pups, but only for heterozygous male mice, leading to the conclusion that compensation may only occur for Ca\textsubscript{v}2.3[−\textpipe−] male mice lacking both alleles of the R-type Ca\textsuperscript{2+} channel. During the mating of heterozygous parents, the number of female mice within the weaned population does not deviate from the expected Mendelian inheritance. During prenatal development, both, T- and R-type Ca\textsuperscript{2+} currents are higher expressed in some tissues than postnatally. It will be discussed that the function of voltage-gated Ca\textsuperscript{2+} channels during prenatal development must be investigated in more detail, not least to understand devastative diseases like developmental epileptic encephalopathies (DEE).

Calcium ions are crucial for reproduction and development\textsuperscript{1}. Changes of cytosolic Ca\textsuperscript{2+} concentrations translate a diverse set of signals into specific cellular responses. More than 100 Ca\textsuperscript{2+} channels, pumps, exchangers, sensors and buffers contribute to the fundamental processes involved in development and propagation of living cells\textsuperscript{2}. Voltage-gated Ca\textsuperscript{2+} channels (VGCCs) are a key mediator of Ca\textsuperscript{2+} entry from the extracellular space and enable Ca\textsuperscript{2+} signaling in a dual manner, electrophysiologically, via Ca\textsuperscript{2+}-induced changes in membrane potential, and biochemically, through the activation of Ca\textsuperscript{2+} dependent enzymes and other proteins affecting cellular regulation\textsuperscript{3}.

Ten mammalian genes are known to encode different ion conducting Ca\textsubscript{a1} subunits of these VGCCs, which have been subdivided into 7 high- and 3 low-voltage activated Ca\textsuperscript{2+} channels (for details,\textsuperscript{4}). In vivo, they are assembled with additional auxiliary subunits\textsuperscript{4}, for which the complete setup of components is only partially known. Additional structural variation arises from alternative splicing, which increases structural and functional variability\textsuperscript{5,6}.

The ion conducting Ca\textsubscript{a1} subunits of VGCCs have been inactivated in mice to deduce their individual functions\textsuperscript{8}. Some of the resulting mouse models are related to human diseases (for a summary see\textsuperscript{9}).

Several voltage-gated Ca\textsuperscript{2+} channels play a role in rodent models of acquired epilepsy, including the Ca\textsubscript{v}2.3 / R-type\textsuperscript{10,11} and the Ca\textsubscript{v}3.2/T-type channel\textsuperscript{12,13}, both of which are highly sensitive towards divalent trace metal cations\textsuperscript{14,15,16}. Mouse models lacking both Ca\textsuperscript{2+} channel types were investigated in previous studies to describe in detail their phenotypes and sensitivities towards divalent metal cations when co-injected with kainate. During the breeding of these mice the number of weaned pups did not correspond the expected ratios for a Mendelian inheritance, pointing to a possible prenatal lethality.

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Materials and Methods

Material and reagents. 

Unless noted otherwise, all reagents were obtained from Sigma-Aldrich and used without further purification (Sigma-Aldrich Chemie GmbH, Schnelldorf, Germany). Solutions were prepared with deionized, double-distilled or type I ultrapure water dispensed from an ELGA LabWater (Purelab Flex 2, United Kingdom) system respectively.

Animals.

Mice were housed at a constant temperature (20–22 °C) in makrolon type II cages, with light on from 7 a.m. to 7 p.m. (light intensity at the surface of the animal cages was between 5 and 10 lx) and ad libitum access to food and water.

The cacna1h gene encoding Ca,3.2 was disrupted in mice by homologous recombination17. These mice were inbred in C57Bl/6 background for more than 10 generations. C57Bl/6 were used as Ca,3.2-competent control mice. The strain abbreviation for the mouse line is C57Bl/6-cacna1h+/− for the mice lacking one Ca,3.2-allele (heterozygous mice) and C57Bl/6-cacna1h−/− for the mice lacking both Ca,3.2-alleles (homozygous mice deficient of Ca,3.2). Ca,3.2-deficient mice are available from Mutant Mouse Resource & Research Centers (MMRRC) with the strain name B6.129-Cacna1hm1Kcam/Mmjh.

The cacna1e gene encoding Ca,2.3 was disrupted in vivo by agouti-colored Ca,2.3[fl +] and deleter mice expressing Cre-recombinase constitutively18. Thus, exon 2 was deleted by Cre-mediated recombination. Ca,2.3-deficient mice were fertile, exhibited no obvious behavioral abnormalities and the born Ca,2.3-deficient mice had the same lifespan as control mice. Parallel breeding of parental inbred mouse lines of Ca,2.3-deficient and control mice ensured their identical background. The strain abbreviation for the mouse line is C57Bl/6.129SvJ-cacna1e+/- for the mice lacking one Ca,2.3-allele (heterozygous mice) and C57Bl/6-cacna1e−/− for the mice lacking both Ca,2.3-alleles (homozygous mice deficient of Ca,2.3). Ca,2.3-deficient mice are available from MMRRC with the strain name B6.129P2(4-Cg)-Cacna1e(-/-)Tsch/Mmjax.

The animal experimentation described in the text was approved by the institutional committee on animal care (Landesamt für Natur, Umwelt und Verbraucherschutz North Rhine Westfalia, Az number 84-02.04.2013. A186 and 81-02.04.2018.A176) and conducted in accordance with accepted standards of humane animal care, as described in the UFAB handbook on the care and management of laboratory animals.

Genotyping of mice. 

Tail biopsies from 21 day old mice were used for the extraction of genomic DNA. Contaminating protein and RNA were enzymatically digested by protease and RNAses, respectively.

For the PCR amplification of indicative Ca,3.2 DNA-fragments, about 1 µg DNA was introduced and amplified with the WT-forward primer 5′-ATT CAA GGG CTT CCA CAG GGT A-3′ and the WT-reverse / KO-forward primers 5′-CAT CTC AGG GCC TCT GGA CCA C-3′ and the WT-reverse / KO-reverse primer 5′-GCT AAA GCG TCT GCT GAG CAT GCC CCA CAC TG-3′17. The sizes of DNA fragments expected are 480 bp for the WT and 330 for the Ca,3.2-KOs.

For the PCR amplification of indicative Ca,2.3 DNA-fragments, about 1 µg DNA was introduced and amplified with the forward primer (B45Hilx1) 5′-AAA AAC AGC CGG GGA AAG CTT AT-3′ and the reverse primer (a1eb45r) 5′-CTG CCC TTT CTT CTT GCC TGA C-3′. The sizes of DNA fragments expected are 1047 bp for the WT and 86 bp for the Ca,2.3-KOs.

PCRs for both genotypings were performed using a DNAEngine Peltier thermal cycler (BioRad, Germany) or a PTC-200 Peltier thermal cycler (MJ Research, Biozym Diagnostik, Germany) with the initial denaturation (94 °C for 10 min) followed by 34 cycles (denaturation at 94 °C for 60 s, annealing at 60 °C for 90 s, extension at 72 °C 4 min) and final extension at 72 °C for 10 min. The PCR products were separated by agarose gel electrophoresis and fluorescent bands were detected on a Herolab UVT-28 M transilluminator by UV irradiation (312 nm excitation wavelength) (Fig. 1).

Data analysis and statistics. 

The assumption of normal distribution of data was tested by the Kolmogorov–Smirnov test. The Student’s t-test was used for the comparison of two experimental groups. Data were analyzed by one-way ANOVA for multiple comparisons. Statistical analysis was performed with the GraphPad Prism software (version 8). The Mendelian genotype distributions were tested by a chi-square test for Mendelian ratios by the use of the algorithm on the web page https://www.ihh.kvl.dk/htm/kc/popgen/genetik/apple_tw/k1.htm. The calculated chi-square values were evaluated and converted into a probability (p-)value by using tables 4-119.

Ethical approval.

All applicable international, national and institutional guidelines for the care and use of animals were followed.

Results.

During the routine breeding for Ca,3.2-deficient mice over a time period of 12 years, the number of genotyped knockout mice (Fig. 1) was severely under represented (the distribution for the genotypes within each group of born mice is summarized in Supplement-table S1 to S3). The consecutive systematic evaluation of wild type, heterozygous and Ca,3.2-deficient pups from 99 breeding pairs (Table 1) revealed a highly significant reduction of heterozygous and even more significant reduction of homozygous Ca,3.2-deficient mice. For comparison, the breeding history was also analyzed for the Ca,2.3-deficient mouse lines.

Analysis of the genotypes for Ca,3.2/T-type mice. 

Ca,3.2 channels mediating T-type Ca2+ currents have been inactivated in mice by homologous recombination17. The deletion of exon 6 of the murine cacna1h
gene was designed to delete the IS5 region in the channel protein and to impair the synthesis of a functional full-length protein. It caused a severe reduction of Cav3.2 mRNA in heterozygous mice as quantified by Northern blot analysis and a complete loss of Cav3.2 mRNA in Cav3.2-deficient mice. In differentiated myotubes from the individual genotypes, the transcript for Cav3.2 identified by RT-PCR was well detected in Cav3.2-competent and completely lost in Cav3.2-deficient mice. Simultaneously, the amount of transcript for Cav3.1 was strongly increased in these myotubes\(^1\), suggesting a compensatory upregulation of these channels.

**PCR-genotyping results.** The genotyping of the litter was performed postnatally by PCR on total DNA isolated from tail biopsies (Fig. 1C,D). The amplified DNA fragments were clearly separated from each other by agarose gel electrophoresis to ensure exact genotype identification (Fig. 1A). The oligonucleotide primers were designed to detect easily and precisely DNA fragments from wild type and Cav3.2-deficient mice. The intensities of DNA fragments were strong enough to identify wild type (3 mice plus 1 reference DNA), heterozygous (11 mice plus 1 reference DNA) and Cav3.2-deficient candidates (3 candidates plus 1 reference DNA) (Fig. 1A). The negative control (no tail DNA included) did not contain DNA fragments of the references sizes (480 bp for wt or 330 bp for KO).

**Distribution of individual genotypes in the mouse lines for the Cav3.2 gene inactivation.** During 77 breeding events from heterozygous parents, in total 83 male Cav3.2(+) (+) mice were born (Suppl.-Tab. 1). As null hypothesis and the low number of events (line in the middle of the table), * = For the summarized data in this table line, no significant difference was observed. However, in 8 matings with a homozygous male and a heterozygous female partner the number of female pups was significantly reduced (p = 0.023, Students t-test).

| Genotype of parents | Statistics of pups and genotype distribution |
|---------------------|---------------------------------------------|
| Female(s) Male       | Mating pairs (n) | Mean litter size Female pups | Chi-squared p | Male pups | Chi-squared p | Sex ratio F / M |
| Cav3.2(+) (+)        | 77              | 2.95 ± 0.25 | < 0.01 | 2.97 ± 0.22 | < 0.001 | 1.28 ± 0.17 |
| Cav3.2(+) (+)        | 14              | 1.68 ± 0.22 | > 0.05 | 2.41 ± 0.33 | < 0.01 | 0.71 ± 0.13 (p = 0.07 *) |
| Cav3.2(−) (−)        | 8               | 3.29 ± 0.21 | > 0.1 | 3.59 ± 0.21 | < 0.005 | 1.05 ± 0.07 |

Table 1. Litter sizes for male and female weaned pups and deviations of genotype distribution from Mendelian inheritance. Mating pairs with a homozygous Cav3.2(−) (−) partner are summarized because of the low number of events (line in the middle of the table). * = For the summarized data in this table line, no significant difference was observed. However, in 8 matings with a homozygous male and a heterozygous female partner the number of female pups was significantly reduced (p = 0.023, Students t-test).

Analysis of the genotypes for Cav2.3/R-type mice. Next, we were interested in the breeding results for Cav2.3-deficient mice, which are known to exhibit a deficit in the flagellar speed of moving sperms as well as in the acrosome reaction\(^2\,\,^22\). The Cav2.3 channels mediating R-type Ca\(^{2+}\) currents have been inactivated in mice by homologous recombination and by successive breeding with cre-deleter mice\(^23\). The deletion of exon 2 of the murine cacna1e gene was designed to delete the IS1 region in the channel protein and to impair the synthesis of a functional full length channel transcript. It caused the complete loss of Cav2.3 channel protein as proven...
by Western blotting using Cav2.3-selective antibodies. In heterozygous mice, the brain Cav2.3 protein level was about half of the amount detected in Cav2.3-competent mice. Cav2.3 channels are inactivated by deleting exon 2 introducing an early stop codon. Exon 2 encoding the first transmembrane segment of the Cav2.3 Ca2+ channel was deleted by Cre-mediated recombination. After inbreeding of heterozygous parents, the genotyping of the litter was performed postnatally by PCR on total DNA isolated from tail biopsies (Fig. 1B,C). The oligonucleotide primers were designed to detect reliably DNA fragments from all genotypes. The intensities of DNA fragments were sufficiently strong to identify wild type (6 mice plus 1 reference DNA), heterozygous (3 mice plus 1 reference DNA) and Cav2.3-deficient candidates (7 mice plus 1 reference DNA), heterozygous (3 mice plus 1 reference DNA) and Cav2.3-deficient candidates (7 mice plus 1 reference DNA).
Distribution of individual genotypes in the mouse lines for the Cav2.3 gene inactivation. During 63 breeding events from heterozygous parents, the mean litter size did not differ between male (3.6 ± 0.2) and female pups (3.3 ± 0.2) (Table 1). In total, 68 male Cav2.3(+|−) mice were born (Suppl.-Tab. 3). As null hypothesis and according to Gregor Mendel one would expect 136 heterozygous and 68 homozygous male Cav2.3-deficient pups. But only 86 heterozygous mice were born. Different results were achieved for the females. During the same 63 breedings, in total 58 Cav2.3(+|−) female mice were born, 105 heterozygous and 44 Cav2.3-deficient pups were counted (Suppl.-Tab. 3). So far, only the number of heterozygous male mice was significantly different from the expected number (CHISQ p < 0.005) (Table 1). No deviation was observed for female pups (CHISQ p > 0.1) (Table 1 and Fig. 3), leading to the conclusion that in male mice the null hypothesis has to be rejected and that as an alternative hypothesis the inactivation of one allele of Cav2.3 must cause developmental problems, leading to a clear reduction of heterozygous male pups, which may only be compensated when both Cav2.3 alleles are missing.

Discussion
Our most important findings are related to deviations of genotype distributions from the expected normal Mendelian inheritance among weaned pups. For both Ca2+ channel types, the genotypes of the weaned offspring were significantly different from the expected Mendelian ratios.

To demonstrate that one may exclude a false genotyping, examples for the determination by PCR were included showing that indicative DNA-fragments could reliably be amplified. Further, an erroneous determination of sex can be excluded, because the sex determination was performed by an experienced coworker. If a continuous miss-determination of male heterozygous pups would have occurred, the number of heterozygous female pups must have been significantly elevated from the expected Mendelian ratio, which is not the case.

Another mistake, which could explain the “non-Mendelian ratios”, would be if unintentionally the breeding pairs would not have been all heterozygous. We can exclude it, as all parents were re-genotyped when the breeding was started. Further, we checked the data for the Ca3.2-breeding lines (see Suppl.-Tab. 1) only a single breeding pair with no Cav3.2 null pups. For the Ca2.3-breeding data (see Suppl.-Tab. 3) we revealed in the 17 breeding lines only 2 of them, which did not have a Cav2.3 null pups.

So far, only for the ion conducting Cavα1 subunit of the cardiac L-type Ca2+ channel a prenatal lethality is known. No viable Cav1.2(−|−) mice were born, but the number of heterozygous pups was normal, corresponding to the expected Mendelian ratio. The developing Cav1.2-deficient pups died before day 14.5 postcoitum (p.c.) but up to day 12.5 p.c., the embryonic hearts contracted with identical frequency in wild type, heterozygous and homozygous Cav1.2 deficient mice. So far, it has remained unclear, which unidentified L-type like Ca2+ current may enable the normal prenatal beating between day 12.5 and 14.5 p.c. in Cav1.2-deficient mice.
For the Ca,3.2-deficient matings, a continuous reduction in the offspring number was observed for both sexes, when one or both alleles were inactivated in the pups. It is currently unknown and would be interesting to investigate, why the lack of the Ca,3.2 allele causes prenatal lethality in some but not in all cases.

Ca,3.2 belongs to the subfamily of low-voltage activated T-type Ca²⁺ channels. They are expressed in many developing tissues and involved in regulating cell proliferation, differentiation, growth and death²⁷. Both, the development of T-type channel isotypes and the development of electrophysiologically defined T-type currents reveals higher levels during embryonic states compared to the postnatal development (see Fig. 1 in²⁷). There is sufficient evidence for a high expression of T-type Ca²⁺ channels in embryonic tissues at the molecular level²⁸, which appears to be especially important for cardiac²⁹,³⁰ and neuronal development³¹,³².

Using information from the gnomAD data base, which quantifies the functional constraints for human genes, CACNA1H is not under significant functional constraint in the human population, though no individuals with homozygous loss of function alleles have been observed. Its o/e number with 0.38 (CI 0.28–0.5) is high, illustrating that the number of observed per expected (o/e) nucleotide variants found indicates a much higher functional constraint for CACNA1E, which is among the most constrained genes in the human genome with an o/e value of only 0.07 (CI 0.04–0.12).

While for the inherited mutations in humans, the functional constraints for the CACNA1E are much higher than for the CACNA1H gene, it does not seem to be the case for the investigated mouse models in the present study. The investigation of the role of the cacna1e gene in a neurotoxin Parkinson's mouse model revealed that the Cav2.3 knockout even reduced activity-associated nigral somatic Ca²⁺ signals and Ca²⁺-dependent afterhypermperpolarizations, leading to full protection from degeneration in vivo [2a].

On the other side, the o/e evaluation for the CACNA1E gene in the gnomAD data base fits well with the observation that de novo mutations in CACNA1E are critical. Recently, for Ca,2.3 in 30 children de novo gain-of-function mutations were identified, which cause developmental and epileptic encephalopathy with contractures, macrocephaly and dyskinesias³³. These disturbances in addition cause early death in the young patients.

The ion conducting subunit Ca,2.3 forms the central pore of the pharmacoresistant R-type Ca²⁺ channels, which also exhibit higher expression levels during prenatal development than postnatally³⁴,³⁵. The sex specific effect of one allele loss in heterozygotes may relate to the function of Cav2.3 during acrosome formation²¹,²². Sperms lacking Ca,2.3 show altered Ca²⁺ responses, a reduced acrosome reaction and a strong subfertility phenotype²⁶. If the loss of one Ca,2.3 allele affects the acrosome reaction substantially, the loss of both alleles in homozygous KO mice could have triggered a corresponding compensation reaction, e.g. by upregulation of another voltage-gated Ca²⁺ channel.

Probably the sex-selective deviation from the Mendelian ratio may include sex-specific hormonal effects, similar as it was reported for effects of Zn²⁺ ions on glucose homeostasis³⁷.

In the literature, paradoxical inheritance with heterozygosity has been listed as one out of ten different non-Mendelian inheritance patterns³⁸. Such rare cases of unusual segregation patterns are found in some specific diseases, as for example for glaucoma involving the K423E allele of TIGR (trabecular meshwork-inducible glucocorticoid response) gene, which is only seen in heterozygotes³⁹. Obviously, the mutated proteins in homozygotes may still form functional response elements that interact with other proteins. Two additional examples
are reported in the same review, which are related to a defect in the ephrin-B1 gene and to the craniofrontonasal syndrome, for which even heterozygous females are more severely affected than hemizygous mutant males\(^6\)\(^{31}\).

**Conclusion and future perspectives**

Our findings show that in depth investigations are needed to understand the prenatal developmental role of voltage-gated Ca\(^{2+}\) channels. For mutations of the human Ca\(_{2.3}\) R-type Ca\(^{2+}\) channel several gain-of-function mutations have been reported and they severely change the juvenile development during the mentioned developmental and epileptic encephalopathy\(^{32}\). A better understanding of this complex disease would help to find a better therapy for treating the children, which have a low life time expectancy.

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Author contributions
T.S. contributed to conception and design of the work, acquisition, analysis and interpretation of the data for the work, drafting the work and revising it critically for important intellectual content. S.A., R.C. and F.N. contributed to conception and design of the work and revising it critically for important intellectual content. J.H. contributed to conception and design of the work. All authors approved the final version of the manuscript and agree to be accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. All persons designated as authors qualify for authorship, and all those who qualify for authorship are listed.

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Competing interests
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