Altered Anesthetic Sensitivity of Mice Lacking Ndufs4, a Subunit of Mitochondrial Complex I

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Abstract

Anesthetics are in routine use, yet the mechanisms underlying their function are incompletely understood. Studies in vitro demonstrate that both GABA_A and NMDA receptors are modulated by anesthetics, but whole animal models have not supported the role of these receptors as sole effectors of general anesthesia. Findings in C. elegans and in children reveal that defects in mitochondrial complex I can cause hypersensitivity to volatile anesthetics. Here, we tested a knockout (KO) mouse with reduced complex I function due to inactivation of the Ndufs4 gene, which encodes one of the subunits of complex I. We tested these KO mice with two volatile and two non-volatile anesthetics. KO and wild-type (WT) mice were anesthetized with isoflurane, halothane, propofol or ketamine at post-natal (PN) days 23 to 27, and tested for loss of response to tail clamp (isoflurane and halothane) or loss of righting reflex (propofol and ketamine). KO mice were 2.5- to 3-fold more sensitive to isoflurane and halothane than WT mice. KO mice were 2-fold more sensitive to propofol but resistant to ketamine. These changes in anesthetic sensitivity are the largest recorded in a mammal.

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Introduction

The molecular mechanisms responsible for the effects of volatile anesthetics are far from clear. Although volatile anesthetics inhibit excitatory synaptic transmission and enhance inhibitory signaling, there is little agreement as to how this phenomenon occurs [1–3]. Ligand-gated ion channels initially emerged as the leading candidates to mediate these effects. Both GABA_A and NMDA receptors were initially viewed as likely volatile anesthetic targets, by virtue of their physiologic functions and anatomic locations within the central nervous system (CNS) [3]. A large number of compelling in vitro studies substantiated these hypotheses, since volatile anesthetics could potentiate inhibitory currents through GABA_A channels, or inhibit excitatory transmission in glutamatergic neurons [1,4]. However, for a number of different possible reasons, whole animal models have not supported the hypothesis that NMDA and GABA_A receptors mediate all aspects of general anesthesia produced by volatile anesthetics [5,6]. To date the largest change in a mammal to a volatile anesthetic is a 40% decrease in sensitivity to halothane in a mouse that lacks a 2-pore potassium channel, TREK-1 [7].

In a forward genetic screen in the nematode, C. elegans, we identified a mutation, gas-1(fc21), that caused a very significant hypersensitivity to all volatile anesthetics [8]. The gas-1 gene encodes a highly conserved subunit of complex I of the electron transport chain (83% similar to the human orthologue NDUFS2) [9,10]. RNAi inhibition of most complex I subunits also increased volatile anesthetic sensitivity [11]. Interestingly, mutations in subunits of complex II, III, or IV did not change sensitivity of C. elegans to volatile anesthetics, even though animals carrying these mutations share many other phenotypes with gas-1 [12,13]. Children with defects in complex I function were hypersensitive to sevoflurane, whereas children with defects in other steps of electron transport within the mitochondrion were not, even though they were indistinguishable in symptoms of mitochondrial disease from the complex I-deficient patients [14]. Although the data obtained from patients predicted enhanced sensitivity of other mammals with complex I dysfunction, the sample size was low, the genes involved were unknown, and controls were a mixed population.

A mouse model with complex I deficiency was developed by conditional inactivation of the Ndufs4 gene that encodes an 18 kD subunit of complex I. This subunit is not directly involved in electron transport, but appears to play a role in assembly or stability of the complex [15,16]. Homozygous Ndufs4-null mice appear neurologically normal at weaning, but post-natal day 35 (PN35) the KO mice manifest increasing ataxia, failure to thrive, and usually die by PN55. This strain has been established as a model for Leigh syndrome, the most common infantile mitochondrial disorder. Mice with selective loss of Ndufs4 function within the CNS have the same behavioral phenotype as the total KO mice [16].
Initial attempts to anesthetize KO mice using standard conditions were often fatal. Those observations along with the knowledge of sensitivity of worms and children with complex I deficiencies prompted us to hypothesize that the KO animals would be hypersensitive to volatile anesthetics. We determined the sensitivity to anesthetics shortly after weaning (PN23 to 27), when the animals are still behaviorally normal and before there is any evidence of neuronal degeneration in the CNS [16]. Remarkably, KO mice displayed the greatest hypersensitivity to volatile anesthetics ever recorded for a mammal. This sensitivity extended to the non-volatile GABA<sub>A</sub> facilitator and agonist, propofol [17], but not to the NMDA antagonist, ketamine [18,19]. The differential sensitivity to different classes of anesthetics may provide a clue to the role of complex I in mediating anesthetic action.

**Methods**

**Ethics Statement**

This study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. All animal experiments were performed with the approval of the Animal Care and Use Committee of the University of Washington (IACUC #2183-02). No surgery was performed and all efforts were made to minimize suffering.

**Anesthetic sensitivity**

Mice were maintained with rodent diet (5053, Picolab, Hubbard, OR) and water available ad libitum in a vivarium with a 12-h light-dark cycle at 22°C. The KO mice were generated by crosses of heterozygotes on a C57Bl/6 genetic background, genotyped by polymerase chain reaction at PN22; KO and WT littermate controls were tested for anesthetic sensitivity during the next week.

Mice were anesthetized with halothane or isoflurane and their temperature was maintained by radiant heat according to the techniques of Sonner [20,21]. Animals were between 23–27 days old at the time of exposure to anesthetic. Failure to respond to a non-damaging tail clamp was recorded. Responses of the same mouse to different doses of the volatile anesthetics were measured after 15 min of equilibration between doses. All animals were exposed to a single anesthetic drug. Samples of isoflurane or halothane were taken at different delivery settings and measured by gas chromatography as described [8]. The non-volatile anesthetics: propofol (Diprivan®, AstraZeneca, Wilmington DE, USA) or ketamine (Ketaset®, Fort Dodge, IA, USA) were injected intra-peritoneally with drug at a concentration of 10 mcg/ul. Animals were tested for loss of righting reflex (LORR) at 5-min intervals following injection. Animals were kept warm on a heating pad throughout. Animals were allowed to recover for at least 24 h before testing again at a different dose of the same anesthetic. No animal received more than four test doses. Animals that did not lose righting reflex within 30 min of injection were denoted as a no LORR.

**Statistics**

The effective concentration for 50% of the animals tested (EC<sub>50</sub>) for volatile anesthetics was determined as described by Sonner et al., using an up and down method [20]. The effective dose for 50% of maximum effect (ED<sub>50</sub>) values for propofol and ketamine were determined by constructing a dose-response curve for each drug and taking the midpoint of the curve. Values for EC<sub>50</sub>s and ED<sub>50</sub>s were compared between the WT and KO strains using a Students t-test. Significance was defined as a p<0.01. Error bars in Figure 1 represent the standard deviations of the mean. Errors for propofol and ketamine represent Standard Errors of the mean.

**Results**

**Ndufs4** KO mice were extremely hypersensitive to isoflurane (Figure 1), with an EC<sub>50</sub> that was about one third that of their WT littermates (KO EC<sub>50</sub> = 0.44±0.07%; WT EC<sub>50</sub> = 1.23±0.13%). KO mice were also hypersensitive to halothane (KO EC<sub>50</sub> = 0.52±0.11%; WT EC<sub>50</sub> = 1.28±0.07%, WT) as shown in Figure 1. No animals displayed any seizure-like activity with exposure to the volatile anesthetics. Animals reached steady state for their response within 5 min of volatile anesthetic exposure and they recovered from exposure to the gases within 15 min of breathing room air. KO and WT mice displayed vigorous responses to tail pinch in air and at sub-anesthetics doses of volatile anesthetics. The EC<sub>50</sub> values of WT mice were similar to that previously reported for the C57Bl/6 strain [20]. Animals lost righting reflex at concentrations too low to be delivered with standard vaporizers. KO mice were also hypersensitive to propofol, although the shift was not as extreme as that for the volatile agents (Figure 2). The dose of propofol that produced LORR in the KO mice was about one half that of their WT littermates (KO ED<sub>50</sub> = 58±5 mg/kg; WT ED<sub>50</sub> = 67±6 mg/kg). The maximum effect of propofol was observed within 5 min of injection in both the WT and KO mice and all animals recovered righting reflex within 15 min of injection. The ED<sub>50</sub> for propofol in WT mice agrees with published data [23,24].

In contrast to both the previous results, the KO animals were strikingly resistant to the effects of ketamine (Figure 3), and were significantly resistant to the LORR (KO ED<sub>50</sub> = 106±5 mg/kg; WT ED<sub>50</sub> = 69±4 mg/kg). The maximum effect of ketamine on LORR was seen within 5 min of injection for both WT and KO animals and all animals recovered by 15 min after injection. The ketamine data for the WT animals agree with a published value of

![Figure 1. EC<sub>50</sub> as for isoflurane (ISO) and halothane (HAL) to cause immobility in response to tail pinch.](10.1371/journal.pone.0042904.g001)
65 mg/kg [23,24]. Recovery times for all drugs were similar between WT and KO animals.

Discussion

We report here the largest change in sensitivity to volatile anesthetics recorded for a mammal. The ability of complex I mutations to change response to volatile anesthetics transcends many phyla, which implies an ancient, common mechanism of action. The KO mice were equally hypersensitive to two volatile anesthetics that are quite different in structure. These results are in contrast to our results with mutations in transmembrane leak channels that result in differential sensitivity to isoflurane and halothane in C. elegans [23]. As we have noted previously, we believe that mitochondrial defects affect a downstream target relative to the leak channels, such that sensitivities to all volatile anesthetics are affected [26].

Isoflurane and halothane both enhance GABA\textsubscript{A} receptor signaling while antagonizing NMDA receptor signaling [17] although it is not clear that these effects cause the anesthetized state. Numerous structure-function experiments have shown that GABA\textsubscript{A} receptors are targets of volatile anesthetics, using in vivo assays. However, when putative targets that were thought to be resistant to anesthetic action were tested in genetically engineered mice, the responses to volatile anesthetics were insufficiently affected [5,27–29]. Thus, the true target(s) of volatile anesthetics remain enigmatic.

The degree/direction of changes in sensitivity of Ndufs4(KO) is not uniform across different classes of anesthetic drugs. The KO mice were also hypersensitive to propofol, which is known to act primarily on GABA\textsubscript{A} receptors [17]. However, the hypersensitivity was not as great as with the volatile anesthetics. In addition, any explanation for the hypersensitivity of the KO animals to volatile agents and to propofol must also account for the surprising resistance of these animals to an NMDA antagonist, ketamine [18,19]. Since times of onset and recovery for propofol and ketamine were the same for WT and KO animals, pharmacokinetesc did not play a major role in these responses; the responses represent changes in pharmacodynamics. The observation that the KO mice are resistant to ketamine argues against the possibility that the KO mice manifest a general neuronal dysfunction at the time of testing that makes them hypersensitive to all neuronal depressants. This is in agreement with studies in a mouse model of Alzheimer’s disease, which also demonstrated no increase in anesthetic sensitivity despite generalized CNS depression. [30,31]. It also indicates that the targets that produce the anesthetic state are not identical between ketamine and the other anesthetics tested here. Ketamine anesthetic action may be unique, as it has been suggested to involve increased activation and cortical synchronization rather than neuronal inactivation [32]. Inhibition of HCN1 channels has also been recently suggested as a contributing factor in the hypnocic actions of ketamine further indicating that ketamine function is more complicated than usually thought [33]. The resistance to ketamine in these mice raises an intriguing question as to whether similar changes might be present in humans and may suggest future studies.

How can mitochondrial dysfunction cause extreme hypersensitivity to volatile anesthetics and propofol, and why are defects in complex I function so important? Complex I is responsible for over half of the electron transport necessary to generate the mitochondrial membrane potential and drive ATP synthesis [34]. The mitochondrial TCA cycle generates glutamate and the precursors of GABA and, while a small part of total energy requirements, the glutamate/glutamine cycle between neurons and glia is dependent on glycolysis and oxidative phosphorylation [35]. Complex I also has the potential to generate reactive oxygen species (ROS), which can result in deleterious oxidation events and/or serve as a critical signaling molecule, when not functioning optimally [34]. Thus, there are many possible ways that loss of complex I might cause hypersensitivity to volatile anesthetics. Most notable is the finding that presynaptic function in glutamatergic neurons is extremely sensitive to complex I function [36].

It is possible that anesthetic sensitivity of the KO mice (as well as C. elegans and children) with complex I deficiency is due to the direct actions of these compounds on defective complex I [37],
further inhibiting its activity and resulting in the inability to maintain ATP production and/or essential signaling necessary to maintain neuronal activity. Xi et al. [37] noted that several mitochondrial proteins bind halothane, including three from complex I, consistent with the possibility that complex I may be a direct target. In both worms and in mammals, movement of electrons through complex I is clearly the most sensitive step within the mitochondrial respiratory chain to disruption by volatile anesthetics [9,38,39] whereas defects in respiratory complexes II, III, and IV do not affect anesthetic sensitivity [12–14,40].

An alternative idea is that anesthetics act primarily on ion channels as is generally hypothesized [6,17,35], but select populations of neurons within the CNS may depend on optimal complex I function to maintain neuronal activity [34]. Assuming that volatile anesthetics work by altering synaptic transmission in some specific areas of the CNS, it may be that the animal is able to match ATP supply to demand in most of the CNS, but has insufficient ATP to support synaptic transmission by some crucial neurons. Thus, if those neurons were already compromised due to complex I mutations and consequently functioning at maximum capacity, then modulation of ion currents by anesthetics could selectively compromise their ability to function adequately.

Although many authors hypothesize that anesthetics act diffusely throughout the CNS, others attribute their actions to specific brain regions; for example, a portion of the rat brainstem produces anesthesia [41–44]. The central medial thalamic region of the rat brain has also been shown to be as crucial to maintaining consciousness [44]. However, considerable controversy still surrounds the putative location in the brain for producing the anesthetic state. The KO mice display progressive glosis and eventually neurodegeneration in specific brain regions; primarily the olfactory bulb, vestibular nucleus, posterior lobes of the cerebellum and deep cerebellar nuclei. However, some other brain regions are without obvious glosis, for example, the pre-Bötzing complex, yet are also affected by the mitochondrial defect [16,43]. Consequently, we assume that Ndufs4 deficiency does not affect all neurons equally. The differences in sensitivity to loss of Ndufs4 could be attributable to differences in (a) intrinsic activity of the neurons, (b) extent of activation in response to changing conditions, or (c) regulation of complex I, for example, by phosphorylation.

It may be possible to identify brain region(s) and neuronal type(s) where Ndufs4 functions to maintain anesthetic sensitivity. Because the Ndufs4 allele in our KO mice can be inactivated by Cre recombinase, it is possible to use Cre-expressing viruses or specific Cre-driver lines of mice to selectively inactivate Ndufs4 in specific cell types or brain regions. Alternatively, it is possible to restore Ndufs4 function to specific cells or brain regions in KO mice. These approaches were used to demonstrate that the vestibular nucleus of KO mice is selectively compromised leading to fatal breathing abnormalities [45]. An additional challenge will be to ascertain whether complex I dysfunction indirectly facilitates anesthetic sensitivity or whether volatile anesthetics interfere with complex I function to directly control sensitivity.

Author Contributions
Conceived and designed the experiments: AQ PGM SEK RDP MMS. Performed the experiments: AQ PGM MMS. Analyzed the data: AQ PGM RDP MMS. Contributed reagents/materials/analysis tools: AQ PGM RDP MMS. Wrote the paper: AQ PGM SEK RDP MMS.

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