Characterization of a novel model of chronic migraine

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ABSTRACT

Chronic migraine is a disabling condition that affects hundreds of millions of individuals worldwide. The development of novel migraine treatments has been slow, in part as a result of a lack of predicative animal models. We have developed a new model of chronic migraine involving the use of nitroglycerin (NTG), a known migraine trigger in humans. Chronic intermittent administration of NTG to mice resulted in acute mechanical hyperalgesia with each exposure as well as a progressive and sustained basal hyperalgesia. This chronic basal hyperalgesia occurred in a dose-dependent fashion and persisted for days after cessation of NTG administration. NTG-evoked hyperalgesia was exacerbated by the phosphodiesterase 5 inhibitor sildenafil, also a human migraine trigger, consistent with nitric oxide as a primary mediator of this hyperalgesia. The acute but not the chronic basal hyperalgesia was significantly reduced by the acute migraine preventive therapy topiramate. Chronic NTG-induced hyperalgesia is a mouse model that may be useful for the study of mechanisms underlying progression of migraine from an episodic to a chronic disorder, and for the identification and characterization of novel acute and preventive migraine therapies.

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1. Introduction

Migraine is one of the most common disorders affecting the general population, resulting in a staggering amount of episodic disability and lost productivity worldwide. For a significant percentage of patients, it results in chronic disability [4,28,29,32]. Despite the extraordinarily high prevalence of migraine, our understanding of its pathophysiology is incomplete. Moreover, although there has been significant progress in the acute treatment of migraine attacks, the ability to treat frequent and chronically disabling migraine remains severely limited. There continue to be millions of individuals for whom currently available migraine therapies are either ineffective or poorly tolerated [4,29].

A significant obstacle to the identification of new migraine therapies has been the lack of predictive animal models. There are a number of promising models for acute migraine attack [2,3,12,25], but it has been particularly difficult to study the progression of migraine from an episodic to a chronic disorder. One approach to modeling acute migraine is the quantification of increased sensory sensitivity in response to known migraine triggers. Nitroglycerin (NTG) reliably triggers headache in normal subjects, and it triggers migraine without aura in migraine-susceptible patients [1,9,17,24]. NTG-evoked migraine is a commonly used experimental paradigm in humans (for review, see [15,24,23]). NTG-evoked hyperalgesia in rodents has been developed as a model for sensory hypersensitivity associated with migraine [3,20]. Acute NTG was previously shown to produce thermal and mechanical allodynia in mice that was reversed by the antimigraine therapy sumatriptan [3]. In addition, in a transgenic mouse model of familial migraine, animals expressing a human migraine gene showed an even greater sensitivity to NTG-evoked hyperalgesia [6]. Further, NTG has also been shown to produce light-aversive behavior [20] and increased meningeal blood flow in mice [13,20]. Taken together, these results indicate that the effects of NTG may effectively model migraine-like symptoms in rodents. Here we have extended the NTG-evoked hypersensitivity assays to model the progression of migraine from an acute to a chronic state.
2. Materials and methods

2.1. Animals

Subjects were male and female C57BL/6j mice, weighing 20 to 30 g. Animals were housed in a 12-h light–dark cycle, and food was available ad libitum. All experiments were approved by the University of California Los Angeles Office of Animal Research and the Animal Care Committee at the University of Illinois at Chicago, in accordance with AAALAC guidelines. These experiments adhered to the guidelines of the Committee for Research and Ethical Issues of IASP [33].

2.2. Drug administration

NTG was prepared from a stock solution of 5.0 mg/mL NTG in 30% alcohol, 30% propylene glycol, and water (American Regent). NTG was freshly diluted in 0.9% saline to a dose of 10 mg/kg. The vehicle control used in these experiments was 0.9% saline. We found that there was no significant difference in mechanical thresholds between those observed when 0.9% saline was used vs those observed with 6% propylene glycol, 6% alcohol, and 0.9% saline. All injections were administered as a 10 mL/kg volume. Unless otherwise noted, animals were tested for baseline responses immediately before intraperitoneal (i.p.) injection with NTG. Animals were injected with subsequent drugs (i.p. unless otherwise noted) at 1 h 15 min after NTG injection, then were tested for mechanical or thermal sensitivity 45 min later (2 h after NTG). For chronic experiments, testing occurred every second day over 9 days (5 test days total). For the topiramate experiment, mice were injected with topiramate or vehicle every day for 11 days. On days 3, 5, 7, 9, and 11 of this treatment, basal mechanical sensitivity was determined, and mice received an injection of NTG or vehicle followed by an injection of topiramate/vehicle, and postdrug responses were determined 2 h later. For experiments testing localized intrathecal injections of sumatriptan into the central nervous system, drug (0.06 μg) or 0.9% saline was injected in a final volume of 5.0 μL [22]. Intrathecal injections were performed with a 30-gauge, ½-inch needle at the L4–5 lumbar interspace on lightly restrained, unanesthetized mice.

2.3. CFA-induced inflammatory pain

Inflammatory pain was induced by injecting Complete Freund’s Adjuvant (CFA; 1 mg Mycobacterium tuberculosis [H37Ra, ATCC 25177]/mL of emulsion in 85% paraffin oil and 15% mannide mannoylate; Sigma) into the paw. Before the injection of CFA, baseline mechanical responses were determined. Inflammation was induced by injecting 15 μL of CFA into the plantar surface of the paw, and animals were subsequently tested 72 h later.

2.4. Sensory sensitivity testing

To determine mechanical sensitivity, the threshold for responses to punctate mechanical stimuli (mechanical hyperalgesia) was tested according to the up-and-down method [8]. In brief, the plantar surface of the animal hind paw was stimulated with a series of 8 von Frey filaments (bending force ranging from 0.01 to 2 g). A response was defined as a licking, lifting or shaking of the paw upon stimulation. The first filament tested was 0.4 g. In the absence of a response a heavier filament (up) was tried, and in the presence of a response a lighter filament (down) was tested. This pattern was followed for a maximum of 4 filaments after the first response.

2.5. Statistical analysis

Data are expressed as mean ± SEM. All statistical analyses were performed by Sigmathat software. For all acute pain experiments, 1-way analyses of variance (ANOVA) and for chronic pain experiments, 2-way repeated measures ANOVA was performed. Unless otherwise noted, all experiments were further analyzed by Holm-Sidak post hoc analysis. A significance level of P < .05 was used.

3. Results

3.1. Repeated administration of systemic NTG produces acute and chronic hyperalgesia

Acute administration of NTG has been shown previously to evoke severe mechanical and thermal hyperalgesia [3]. To model the progression to chronic migraine, we administered varying doses of NTG every second day for 9 days, resulting in a total of 5 NTG injection/test days. Mechanical thresholds were tested before and 2 h after NTG administration on each test day. Repetitive intermittent NTG administration over 9 days produced a significant time and dose–dependent chronic basal mechanical hyperalgesia as assessed by testing before each administration of NTG (Fig. 1A). In addition, NTG evoked significant acute mechanical hyperalgesia in a dose–dependent manner on each test day (Fig. 1B). The greatest decrease in basal and posttreatment responses were observed with 10 mg/kg, the dose of NTG previously characterized in an acute study [3]. We chose to further characterize this dose (Fig. 2). Progressively increasing basal hyperalgesia was observed with repeated administration of 10 mg/kg NTG (Fig. 2A). After the final treatment day (day 9), mechanical responses in mice were assessed daily to determine the recovery time for this basal hypersensitivity. Sensory responses returned to the level of naïve mechanical thresholds (day 1) by day 7 after NTG (Fig. 2A). Female mice showed greater basal hyperalgesia in response to chronic NTG, and unlike males, basal sensitization was observed after a single NTG injection (Fig. 2B). In addition, significant acute/post-treatment hyperalgesia was again observed with each administration (Fig. 2C). To determine whether the associative learning of repeated testing contributed to the progression of basal hyperalgesia, we examined basal mechanical thresholds in mice that had received identical intermittent NTG treatment over 9 days, but were only tested on the first day and on day 9 (novice). These novice mice did not show significantly different hyperalgesia on the final day of NTG (novice, 0.15 ± 0.06 vs repeatedly tested, 0.05 ± 0.03).

3.2. The phosphodiesterase 5 inhibitor sildenafil increases acute and chronic mechanical hyperalgesia

To investigate the role of cGMP in NTG-induced basal and evoked hyperalgesia we examined the effects of the phosphodiesterase 5 inhibitor sildenafil, which increases levels of cGMP and augments the effects of nitric oxide as an activator of guanylate cyclase. We treated mice every second day for 9 days (5 test days) with a low dose of NTG (1 mg/kg i.p.), and sildenafil (3 mg/kg i.p., Fig. 3). On each day of testing, chronic treatment with this low dose of NTG alone did not result in chronic basal hyperalgesia (Fig. 3A). However, the combination of this low dose NTG with sildenafil produced significant basal hypersensitivity. Interestingly, by the final day of testing, sildenafil alone had also produced a significant decrease in basal mechanical threshold (Fig. 3A). In addition, the combination of these 2 compounds produced a significantly greater acute hyperalgesia (Fig. 3B), than either compound alone. These results indicate changes in cGMP levels play a role in the hyperalgesia induced by chronic intermittent NTG treatment.
Fig. 1. Chronic NTG evokes and sustains mechanical hyperalgesia in a dose-dependent manner. C57Bl/6J mice were treated every second day with varying doses of NTG (0–10 mg/kg, i.p.) for 9 days. (A) Basal mechanical responses, as assessed before vehicle or NTG administration, significantly decreased in the 3 and 10 mg/kg NTG group during the treatment period; n = 8/group, P < .001 effect of dose and time 2-way repeated measures (RM) ANOVA; the 3 and 10 mg/kg doses were significantly different from vehicle and 1 mg/kg (P < .001). (B) Increasing doses of NTG produced increasing levels of mechanical hyperalgesia, in mice tested 2 h after NTG or vehicle administration; n = 8/group, 2-way RM ANOVA with Holm-Sidak multiple comparisons. Each dose was significantly different from vehicle (P < .001), and each dose was also significantly different from each other (P < .01). Chronic NTG in rodents produces a dose-dependent persistent hypersensitivity.

Fig. 2. Chronic NTG evokes severe and sustained mechanical hyperalgesia. C57Bl/6J mice were treated every second day with vehicle (VEH) or NTG (10 mg/kg, i.p.) for 9 days. (A) Basal mechanical responses, as assessed before vehicle or NTG administration, significantly decreased in the NTG group during the treatment period, and took 7 days to recover after the final NTG injection; n = 7–8/group. (B) Females developed basal mechanical hyperalgesia more quickly than males after chronic NTG treatment; n = 12/group, **P < .01 2-way repeated measures (RM) ANOVA. (C) NTG consistently produced severe mechanical hyperalgesia, in mice tested 2 h after NTG or vehicle administration; n = 7–8/group, ***P < .001 2-way RM ANOVA. Chronic NTG in rodents appears to model the persistent hypersensitivity observed in migraine patients.

Fig. 3. Sildenafil, a phosphodiesterase 5 inhibitor, potentiates hyperalgesia induced by a low dose of NTG. C57Bl/6J mice were treated every second day with vehicle (VEH) or a low dose of NTG (1 mg/kg, i.p.) for 9 days. (A) Basal mechanical responses were assessed before NTG administration, n = 8/group, P < .05 effect of drug and P < .001 effect of time and interaction, 2-way repeated measures (RM) ANOVA; NTG-SIL mice were significantly different from all other groups (P < .01) (B) Mice were injected i.p. with either vehicle (VEH) or 3 mg/kg sildenafil (SIL) 1 h 15 min after NTG/VEH administration, and tested 45 min later (2 h after NTG); n = 8/group, P < .001 effect of drug, 2-way RM ANOVA; NTG-SIL mice were significantly different from all other groups (P < .001). Animals treated with the combination of NTG and sildenafil showed significantly reduced basal and posttreatment responses.
3.3. Sumatriptan alleviates acute NTG-evoked, but not chronic, basal hyperalgesia

We also investigated the effects of the migraine-selective acute therapy sumatriptan in the chronic NTG model. Consistent with previous studies [3], we found that sumatriptan (600 µg/kg, i.p.) reversed the acute hyperalgesia evoked by NTG injection. Systemic Sumatriptan continued to effectively ameliorate acute NTG-evoked hyperalgesia with each injection over 9 days. It did not, however, alter the chronic basal hyperalgesia that occurred over time with chronic intermittent NTG exposure (Fig. 4A, B). These results indicate that while sumatriptan can reverse the acute effects of NTG, it does not reduce the progression of hyperalgesia that occurs with repeated exposure. To address the possibility that sumatriptan non-specifically inhibited nociception, we tested this same dose of sumatriptan within the CFA model of peripheral inflammatory pain. Sumatriptan (600 µg/kg i.p.) was ineffective at reversing CFA-induced mechanical hyperalgesia (Fig. 4C). To investigate the possibility that sumatriptan was acting to inhibit NTG-evoked mechanical hyperalgesia at the spinal level, we injected sumatriptan 0.06 µg (a dose previously shown to have antihyperalgesic effects in certain peripheral and visceral pain models [22]) or vehicle intrathecally into the spinal L4–5 region. In contrast to systemic intraperitoneal administration, intrathecal injection of sumatriptan did not result in any inhibition of NTG-evoked hyperalgesia (Fig. 4D).

3.4. Topiramate inhibited NTG-induced chronic basal hyperalgesia

We then examined the effect of topiramate, a migraine preventive therapy, on acute and chronic basal hyperalgesia evoked by NTG. Mice were treated once daily with topiramate (30 mg/kg, i.p.) for 11 days. On days 3, 5, 7, 9, and 11 basal and acute NTG-evoked mechanical responses were determined. Daily treatment with topiramate significantly inhibited the development of chronic basal hyperalgesia induced by chronic intermittent NTG treatment (Fig. 5A), and it also reduced acute NTG-evoked hyperalgesia (Fig. 5B).

4. Discussion

Our results indicate that chronic intermittent treatment with NTG produces an acute and chronic hypersensitivity which represents a translationally significant model of chronic migraine. Repeated injection of NTG evoked mechanical hyperalgesia, which was blocked by the antimigraine medication sumatriptan. We also found that chronic intermittent treatment with NTG produced a severe and long-lasting basal hyperalgesia, which was slightly more pronounced in female mice. In addition, this basal hypersensitivity was alleviated by treatment with the migraine prophylactic, topiramate. Furthermore, the effects of concomitant treatment with another human migraine trigger, sildenafil, confirmed that this basal hypersensitivity was mediated by increased levels of cGMP.

Several lines of evidence indicate that NTG treatment in rodents could be a predictive model of migraine in humans. NTG is a reliable trigger of migraine in susceptible patients [9, 17, 24]. Interestingly, migraine attacks do not occur immediately after NTG administration with the vasodilatory effects of the drug, but rather after 2 to 6 h [9, 17, 24], consistent with a delayed response to nitric oxide.
oxide not mediated by vasodilation [27]. In mice, the timing of NTG-induced hyperalgesia also shows a delayed response, similar to the time course of headache observed in humans [3]. Systemic administration of NTG has been reported to cause cellular activation in nociceptive pathways, including trigeminal nociceptive pathways [3,26,30,31], providing additional support for relevance to migraine. In addition, NTG has been shown previously to cause light allodynia [20], and to increase meningeal blood flow [13,20] in mice, two hallmarks of migraine. Furthermore, the amelioration of NTG-evoked hyperalgesia by the prototypic antimigraine medication sumatriptan also validates this model for the study of migraine mechanisms. Additionally, acute NTG-induced hyperalgesia has recently been shown to be increased in mice expressing a gene associated with familial migraine [6].

Migraine typically starts as an episodic disorder, but commonly progresses to a frequent or even daily condition [5]. The pathophysiological mechanisms underlying this change are poorly understood. We have extended the previously reported studies on acute NTG-evoked hyperalgesia to show that repetitive intermittent administration of NTG produces a progressive and sustained basal hyperalgesia that persists for days after the last exposure. These results are consistent with clinical observations of patients with chronic migraine in whom allodynia may occur both between and during migraine attacks. In addition, women are far more likely to suffer from chronic migraine than men [32], and our results suggest that female mice may be more sensitive to chronic NTG. Previous studies have also shown that there are estrogen-related discrepancies in the brain regions activated with NTG administration in rats [14], indicating that the chronic NTG model could potentially be a tool to specifically study sex differences in migraine.

The enhancement of the effects of NTG by sildenafil, also an established human migraine trigger [23,18], provides evidence that nitric oxide–mediated increases in cGMP play a primary role in the hyperalgesia evoked by NTG. We found that by the final test day, sildenafil also independently produced basal hyperalgesia with repeated intermittent dosing, suggesting that potentiation of the endogenous activity of guanylate cyclase may be a mechanism for producing chronic hyperalgesia. Drugs that target different aspects of the nitric oxide–cGMP pathway are currently being developed for the treatment of headache [23], and our preclinical data further support the role of this signaling cascade in the development of chronic migraine.

We found that sumatriptan inhibits acute hyperalgesia in response to NTG, similar to the results of previous rodent [3] and human [16] studies. We also found that sumatriptan continued to have this effect with each NTG exposure over time, but that it did not inhibit the resulting progressive chronic basal hyperalgesia. This result is consistent with the observation that while sumatriptan can be consistently effective as an acute migraine therapy, its use does not prevent progression of migraine over time [4]. Since this model tests hyperalgesia in the hind paw, it is possible that the effects of sumatriptan could be mediated by spinal mechanisms. However, we found that intrathecal administration of sumatriptan did not inhibit NTG-evoked mechanical hyperalgesia, in contrast to systemically administered sumatriptan, indicating that within this model sumatriptan is not acting solely through the lumbar spinal cord. In addition, groups studying the dural inflammation model of migraine have also found that inflammatory mediators applied to the dura [12] or dural TRPA1 activation [11] produced mechanical hyperalgesia in both facial and hind paw regions, likely through brainstem relays. Cutaneous allodynia after NTG likely reflects the development of central sensitization which is observed during primary headache, which may be mediated through sensitization of neurons within the thalamus [7].

Topiramate inhibited acute NTG-evoked hyperalgesia, as well as chronic basal hyperalgesia, consistent with its actions as a prophylactic migraine therapy. Topiramate has multiple mechanisms of actions that could be involved in its therapeutic efficacy for migraine, including inhibition of voltage gated sodium channels, enhancement of GABAergic signaling, inhibition of glutamatergic signaling, and inhibition of carbonic anhydrase [21]. It has been proven to be an effective preventive therapy even in the setting of chronic migraine [10]. Moreover, on the basis of limited studies, topiramate has also been suggested to inhibit the progression of migraine to a chronic disorder in patients [19].

In this study we used a dose of NTG (10 mg/kg), which is substantially higher than the equivalent dose used to provoke migraine in humans [24]. It is therefore possible that this dose of NTG in mice could have additional effects apart from those associated with migraine induction in humans. The significant inhibition of NTG–induced hyperalgesia by systemic sumatriptan and topiramate, however, indicates that even if this dose of NTG represents a “supraphysiological” stimulus, it can nonetheless be effectively reversed by standard doses of a proven migraine-specific therapy. Similarly, the ability of topiramate to inhibit the chronic basal hyperalgesia induced by chronic NTG indicates that this model may also be used to test potential migraine prophylactics.

NTG evoked acute and chronic basal hyperalgesia is a promising mouse model for the investigation of migraine mechanisms and therapies. The enhancement of acute NTG-evoked hyperalgesia by a human migraine-associated gene expressed in mice, and the...
inhibition of this hyperalgesia by both acute and preventive migraine therapies provide strong support for the translational value of this model. Ongoing studies of the effects of other human migraine genes, as well as other established acute and preventive therapies, will help to further validate the model as a useful tool for enhancing our ability to understand and treat this frequently disabling disorder.

Conflict of interest statement
The authors report no conflict of interest.

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