Indirect AMP-Activated Protein Kinase Activators Prevent Incision-Induced Hyperalgesia and Block Hyperalgesic Priming, Whereas Positive Allosteric Modulators Block Only Priming in Mice*

Kufreobong E. Inyang, Michael D. Burton, Thomas Szabo-Pardi, Emma Wentworth, Timothy A. McDougal, Eric D. Ramirez, Grishma Pradhan, Gregory Dussor, and Theodore J. Price

School of Behavioral and Brain Sciences, University of Texas at Dallas, Richardson, Texas

Received April 2, 2019; accepted July 16, 2019

ABSTRACT

AMP-activated protein kinase (AMPK) is a multifunctional kinase that negatively regulates the mechanistic target of rapamycin (mTOR) and mitogen-activated protein kinase (MAPK) signaling, two signaling pathways linked to pain promotion after injury, such as surgical incision. AMPK can be activated directly using positive allosteric modulators, as well as indirectly through the upregulation of upstream kinases, such as liver kinase B1 (LKB1), which is a mechanism of action of metformin. Metformin’s antihyperalgesic effects occur only in male mice, raising questions about how metformin regulates pain sensitivity. We used metformin and other structurally distinct AMPK activators narciclasine (NCLS), ZLN-024, and MK8722, to treat incision-induced mechanical hypersensitivity and hyperalgesic priming in male and female mice. Metformin was the only AMPK activator to have sex-specific effects. We also found that indirect AMPK activators metformin and NCLS were able to reduce mechanical hypersensitivity and block hyperalgesic priming, whereas direct AMPK activators ZLN-024 and MK8722 only blocked priming. Direct and indirect AMPK activators stimulated AMPK in dorsal root ganglion (DRG) neuron cultures to a similar degree; however, incision decreased phosphorylated AMPK (p-AMPK) in DRG. Because AMPK phosphorylation is required for kinase activity, we interpret our findings as evidence that indirect AMPK activators are more effective for treating pain hypersensitivity after incision because they can drive increased p-AMPK through upstream kinases like LKB1. These findings have important implications for the development of AMPK-targeting therapeutics for pain treatment.

SIGNIFICANCE STATEMENT

Nonopioid treatments for postsurgical pain are needed. Our work focused on whether direct or indirect AMP-activated protein kinase (AMPK) activators would show greater efficacy for inhibiting incisional pain, and we also tested for potential sex differences. We conclude that indirect AMPK activators are likely to be more effective as potential therapeutics for postsurgical pain because they inhibit acute pain caused by incision and prevent the long-term neuronal plasticity that is involved in persistent postsurgical pain. Our work points to the natural product narciclasine, an indirect AMPK activator, as an excellent starting point for development of therapeutics.

Abbreviations: ACC, acetyl-CoA carboxylase; AMPK, AMP-activated protein kinase; DRG, dorsal root ganglion; LKB1, liver kinase B1; MAPK, mitogen-activated protein kinase; mTOR, mechanistic target of rapamycin; NCLS, narciclasine; PGE₂, prostaglandin E₂; p, phosphorylated; SNI, spared nerve injury.
AMPK Modulation of Incisional Pain

Materials and Methods

**Laboratory Animals.** Animal procedures were approved by The University of Texas at Dallas Institutional Animal Care and Use Committee and were in accordance with National Institutes of Health Guidelines. All the experiments were performed on male or female Institute of Cancer Research (ICR) outbred mice obtained from Envigo (Cambridgeshire, UK) at 4 weeks of age or bred at the University of Texas at Dallas. Mice were housed in the University of Texas at Dallas Animal Care Facility for at least 1 week before the start of behavior testing and surgery. Animals had ad libitum access to food and water and were on a 12-hour noninverted light/dark cycle. Experimenters were blinded to treatment groups in behavioral experiments. Mice were randomized to treatment groups using a random number generator and in such a manner that multiple treatment groups were always found within any individual cage of animals. Male and female mice were housed separately in groups of four per cage.

**Behavioral Testing.** The plantar incision model was used to induce postsurgical pain in mice as described previously (Banik et al., 2006). Mechanical sensitivity was assessed using stimulation of the hindpaw of the mouse with calibrated von Frey filaments from Stoelting Co. (Wood Dale, IL). We used 0.6-, 1.0-, and 1.4-g filaments. The number of responses for each filament was recorded. After baseline testing, mice were treated systemically with an AMPK activator for seven consecutive days. Immediately after day 3 of treatment, plantar incision surgery was performed by making a 5-mm longitudinal incision with a number 11 scalpel blade in the skin of the left hindpaw and the underlying muscle tissue 2 mm below the heel. The wound was closed using a 5-mm suture, followed by a 200-μl subcutaneous injection of gentamicin (5 mg/ml; Sigma-Aldrich, St. Louis, MO). For the sham surgery, mice were placed under isoflurane for 5 minutes, followed by subcutaneous gentamicin. Mice were tested for mechanical hypersensitivity periodically until response frequency returned to baseline levels. After the return to baseline, hyperalgesic priming was tested by giving each animal an intraplantar injection of prostaglandin E2 (PGE2) (100 ng/25 μl). Response frequency after PGE2 was tested 3 and 24 hours postinjection. The different AMPK activators used in this experiment were metformin i.p. (200 mg/kg; LKT Laboratories, St. Paul, MN), ZLN-024 hydrochloride i.p. (30 mg/kg; Terris Pharmaceuticals, UK)/Zhang et al., 2013), NCLS p.o. (5 mg/kg; Santa Cruz Biotechnology, Inc., Dallas, TX) (Julien et al., 2017), and MK-8722 p.o. (a gift from Merck & Co., Kenilworth, NJ) 30 mg/kg (Feng et al., 2017). Metformin and ZLN were dissolved in 0.9% saline, and NCLS was made in (2-hydroxypropyl)-β-cyclodextrin (Sigma Aldrich). MK-8722 was administered in 0.25% methyl cellulose, 5% Tween-80, and 0.02% sodium dodecyl sulfate (Feng et al., 2017). Rotorod testing was done using a series 8 rotorod device from IITC Life Science, Inc. (Woodland Hills, CA). Mice were placed on the rotorod with the setting of four rotations per minute to start with an increase to 40 rotations per minute over the course of 108 seconds. The latency to fall was measured using the device. Testing was done before drug treatment and again on the last day of drug treatment.

**Metformin Pharmacokinetics.** This study was done at Sai Life Science Limited at Hinjewadi, India. Eighteen ICR mice weighing between 20 and 25 g were used, nine males and nine females. Blood samples of approximately 60 μl were collected under light isoflurane anesthesia from sets of three male and female mice at 0.5, 1, and 4 hours after dosing. Plasma was harvested by centrifugation of blood and stored at −70°C until analysis. After blood collection, brain samples were isolated at each time point from the same animals. Brains were dipped three times in ice-cold phosphate-buffered saline, blotted dry, and weighed. Brain samples were homogenized using ice-cold phosphate-buffered saline with twice the volume of brain weight and then stored at −70°C until analysis. Plasma and brain samples were quantified by fit-for-purpose liquid chromatography-tandem...
mass spectrometry methods. Data are shown as nanograms per milliliter in plasma and nanograms per gram of brain weight.

**Neuron Culture.** DRGs were extracted aseptically from 4-week-old male ICR mice in Hanks’ buffered salt solution (HBSS; Invitrogen, Waltham, MA) on ice. The DRGs were dissociated enzymatically at 37°C, first with collagenase A (1 mg/ml, Roche Diagnostics, Basel, Switzerland) for 25 minutes and then collagenase D (1 mg/ml; Roche) that included papain (30 μg/ml; Roche) for 20 minutes. Afterward, a trypsin inhibitor (1 mg/ml; Roche) that contained bovine serum albumin (bovine serum albumin, 1 mg/ml; Fisher) was applied, and the ganglia were mixed to allow further dissociation with a polished Pasteur pipette. The tissue was then filtered through 70-μm nylon cell strainer (Falcon; Corning, NY) and resuspended in Dulbecco’s modified Eagle’s medium F-12 GlutaMax media (Invitrogen) that contained 10% fetal bovine serum (fetal bovine serum; Hyclone Laboratories, Inc., South Logan, UT) and 1× penicillin streptomycin (Pen-Strep; Invitrogen). The media also contained serve growth factor (10 ng/ml; Millipore, Billerica, MA) and 5-fluoro-2'-deoxyuridine + uridine (3.0 μg/ml + 7.0 μg/ml; Sigma) to reduce proliferation of gli and fibroblasts. Neurons were cultured for 7 days on 12-mm glass coverslips (no. 1 thickness; Chemglass Life Sciences, Inc., Vineland, NJ) in a 24-well tissue culture plate (Falcon) coated with poly-D-lysine (Sigma) at 37°C with 95% air and 5% CO₂. On the day of the experiment, drugs were diluted into Dulbecco’s modified Eagle’s medium F-12 plus GlutaMax media and added directly onto the neurons at concentrations indicated in the results for 1 hour. All DRGs from one mouse were used to generate approximately four coverslips of primary cells. Coverslips from multiple independent animals were used in each experiment.

**Immunocytochemistry and Digital Image Analysis.** After AMPK activator treatment, the cells were washed with phosphate-buffered saline and fixed with 10% formalin in phosphate-buffered saline for 30 minutes. Cells were blocked with 10% normal goat serum.
and labeled with anti-peripherin, mouse monoclonal (1:500; Sigma) and phospho-acetyl-CoA carboxylase (ACC) (Ser 79) (p-ACC, 1:1000; catalog no. 3661; Cell Signaling Technologies) overnight at 4°C. Next, cells were washed and incubated with fluorescein-conjugated secondary antibodies (Alexa Fluor, anti-rabbit 488 and anti-mouse 568; Invitrogen) and counterstained with a DAPI stain, 4',6-diamidino-2-phenylindole (DAPI) (Invitrogen) and mounted with Prolong Gold (Invitrogen). Images were taken on an Olympus Fluoview FV1200 laser scanning confocal microscope and analyzed using the colocalization tool within Olympus FV software. The intensity of each channel was adjusted so that only areas that contained a strong signal of 488 and 405 nm filaments were visible. This adjusted imaged contained distinct puncta that could then be counted and analyzed using GraphPad Prism (GraphPad Software, San Diego, CA). Results were reported as the average percent area of p-ACC signal in neurons labeled with peripherin for all representative pictures.

**Western Blotting.** Male mice were anesthetized with isoflurane and spinal cords and DRGs innervating the hindpaw (L3–L5) were removed. Fresh tissues were placed in ice cold lysis buffer (50 mM, Tris pH 7.4, 150 mM NaCl, 1 mM EDTA pH 8.0, and 1% Triton X-100) containing protease and phosphatase inhibitors cocktails (Sigma-Aldrich) and homogenized using beaded homogenization tubes (Bertin Corp; Rockville, MD.). Samples were centrifuged at 14,000 rpm for 15 minutes at 4°C, and the supernatant containing protein extracts was collected. Protein concentrations were assessed using the Pierce BCA protein assay kit (ThermoFisher Scientific) as directed. A total of 10 µg of protein was mixed with Laemmli sample buffer (Bio-Rad, Hercules, CA) and 2-mercaptoethanol and was heated at 95°C for 5 minutes. Samples were loaded into each well of a 10% SDS-PAGE gel along with 15 µl of Precision plus protein kaleidoscope prestained protein standards (Bio-Rad). Proteins were transferred to a 0.45 PVDF membrane (Millipore) at 100 V for 1 hour. Membranes were blocked using 5% nonfat dry milk in 1× Tris-buffered saline-Tween for 2 hours before primary antibody incubation. Primary antibodies used for this experiment were pAMPKα (Thr 172) (1:1000, catalog no. 2535; Cell Signaling Technology, Danvers, MA) and total AMPKα (1:1000, catalog no. 2532; Cell Signaling). Bands were visualized using a Bio-Rad ChemiDoc Touch. Analysis was performed using Image Laboratory version 6.0; Bio-Rad, Hercules, CA.

**Statistics.** Data are shown as mean ± S.E.M., and the numbers of animals or samples used in each analysis are given in the figure legends. GraphPad Prism 7 was used to analyze data and to do curve fitting, and other statistical tests are given in the figure legends. Two-way analysis of variance (ANOVA)s were used to analyze von Frey data. Post-hoc tests used were the Bonferroni’s multiple comparisons test. Significance level was set at α < 0.05. Details on test statistics are given in Supplemental Table 1.

**Results**

**Metformin Decreases Incision-Induced Mechanical Hypersensitivity and Blocked Hyperalgesic Priming in Male but Not in Female Mice.** To assess the effects of metformin on incision-evoked mechanical hypersensitivity in male mice, we first obtained baseline responses for von Frey filament strengths of 0.6, 1.0, and 1.4 g. After baseline, mice were allocated into metformin (200 mg/kg) and vehicle (0.9% saline) interperitoneal treatment groups for 7 days. On day 3 of treatment, plantar incision surgery was performed on all the mice. These mice were tested periodically over the next 2 weeks until they returned to baseline. Once the mice returned to baseline-level sensitivity, animals received 100 ng PGE2 into the affected hindpaw to assess hyperalgesic priming. These mice were tested 3 and 24 hours postinjection. In male mice, metformin reduced acute mechanical hypersensitivity after plantar incision. Metformin also prevented the development of hyperalgesic priming (Fig. 1, A–C). These findings are consistent with our previously published data using the up-down von Frey testing method (Burton et al., 2017). A separate group of sham surgery male mice were allocated into metformin and vehicle treatment groups and baselined for von Frey and rotarod testing to assess possible effects on motor behavior. These mice were treated for 7 days and given a sham surgery on day 3. The mice were tested periodically for von Frey responses, and no differences were observed based on treatment (Supplemental Fig. 1, A–C). After the last day of drug treatment, these mice were tested on the rotarod again to assess the effects of metformin on motor behavior. Metformin had no effect on locomotive behavior in male mice (Supplemental Fig. 1D).

**Fig. 2.** Male and female mice recover from plantar incision surgery at the same rate. (A–C) Plantar incision surgery in male and female mice caused an increase in mechanical hypersensitivity that resolved at the same rate with no significant differences in mechanical hypersensitivity. n = 21 males and n = 18 females.
To test whether similar effects would be observed in female mice, we obtained baseline mechanical sensitivity measurements and randomized them into metformin and vehicle treatment groups under conditions similar to those of males. On day 3 of treatment, plantar incision surgery was performed on these mice. Mechanical hypersensitivity was tested 1, 2, 4, 7, 10, and 14 days postsurgery. Once the mice returned to baseline level sensitivity, animals received 100 ng of PGE$_2$ into the affected hindpaw. Unlike with male mice, metformin had no effect on the acute hypersensitivity caused by incision. Metformin also failed to block hyperalgesic priming in female mice (Fig. 1, D–F). These findings are consistent with our observations in female SNI mice; metformin relieved mechanical hypersensitivity in only male mice. Although a pronounced difference in mechanical hypersensitivity was seen between male and female metformin-treated mice after plantar incision, we did not observe any sex difference in the duration or magnitude of mechanical hypersensitivity in vehicle-treated mice (Fig. 2).

We next sought to determine whether this sex difference with metformin could be explained by pharmacokinetics. Nine male and nine female ICR mice were given a single i.p. dose of 200 mg/kg metformin. A third of the mice were euthanized 30 minutes after injection, another third 1 hour

Fig. 3. NCLS treatment attenuates surgery-induced mechanical hypersensitivity and blocks hyperalgesic priming in male and female mice: (A–C). NCLS treatment starting immediately after baseline measurements dose dependently decreased mechanical hypersensitivity after plantar incision surgery and prevented plantar incision-induced hyperalgesic priming precipitated by PGE$_2$ injection at all doses. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$; n = 4 for the NCLS groups, n = 3 for the vehicle group. (D–F) NCLS treatment also decreased mechanical hypersensitivity after plantar incision surgery and prevented surgery-induced hyperalgesic priming precipitated by PGE$_2$ injection. *$P < 0.05$; **$P < 0.01$; ***$P < 0.001$; n = 6 for the NCLS group, n = 6 for the vehicle group.
after injection, and the remaining 4 hours after injection; brain and plasma were taken at each time point. Metformin was measured in the brain and plasma of each animal and compared for each timepoint and across sexes. Surprisingly, we found that plasma levels of metformin were higher in female mice at 0.5 and 1 hour after intraperitoneal injection (Supplemental Fig. 3A). Brain levels were higher in female mice at all time points (Supplemental Fig. 3B). Brain-to-blood ratios did not differ between sexes (Supplemental Fig. 3C; Supplemental Table 2). Differences in pharmacokinetics for plasma or blood levels of metformin do not explain the sex differences in efficacy seen in the incisional model.

**NCLS Decreased Incision-Induced Mechanical Hypersensitivity and Blocked Priming in Male and Female Mice.** We then tested whether a structurally distinct AMPK activator would show similar sex-specific effects. We obtained baseline mechanical sensitivity measurements from male mice and then sorted them into NCLS (1, 3, or 6 mg/kg) and vehicle ((2-hydroxypropyl)-β-cyclodextrin) groups for oral administration. These mice were treated orally for 7 consecutive days, and plantar incision surgery was performed on day 3 of treatment. These mice were tested for von Frey responses under the same time course as in the metformin experiments. Once the mice returned to baseline sensitivity,

![Fig. 4. Narcielasine induces AMPK activity in male DRG neurons in vitro. Male neuron cultures were treated with vehicle (A), 100 nM (B), or 1 μM NCLS (C) for 1 hour. Representative immunohistochemistry images of the DRG neurons at 40× magnification. Quantification of images shown in (D). One micromolar NCLS increased p-ACC intensity in neuron cultures. Only neurons that were positive for peripherin staining were analyzed. Maximum florescence refers to the maximum florescence intensity per neuron analyzed. **P < 0.01. n = 39 images analyzed per group.](image-url)
animals received PGE₂ into the hindpaw and were tested 3 and 24 hours after injection. In male mice, NCLS reduced the mechanical hypersensitivity after plantar incision in a dose-dependent fashion. All NCLS doses completely blocked development of hyperalgesic priming (Fig. 2, A–C). Because the 3-mg/kg dose showed full efficacy in male mice, we chose to continue our experiments in female mice using only this dose. The experiment in male mice was repeated in its entirety in a group of female mice over the same time course. Once the females returned to baseline, they also received PGE₂ into the affected paw and were tested 3 and 24 hours after surgery. Like the experiment with males, NCLS prevented acute mechanical hypersensitivity in female mice and blocked development of hyperalgesic priming (Fig. 2, D–F). Therefore, indirect AMPK activators can be effective in reducing incision-evoked mechanical hypersensitivity and hyperalgesic priming in both male and female mice. A separate group of sham surgery male mice were allocated into NCLS and vehicle treatment groups and baselined for von Frey and rotarod testing to assess possible effects on motor behavior. These mice were also treated for 7 days but given a sham surgery treatment on day 3. The mice were tested periodically for von Frey responses, and no differences were observed based on treatment (Supplemental Fig. 4, A–C). NCLS also had no effect on rotarod behaviors (Supplemental Fig. 4D).

Whereas NCLS has been shown to activate AMPK in other cell types, it has never been tested for AMPK activity in DRG neurons. To assess this directly, DRGs were cultured from naïve male mice and treated with vehicle, 100 nM, 1 μM NCLS.

Fig. 5. Narciclasine increases AMPK signaling in male and female DRGs. A single dose of NCLS (3 mg/kg) caused a significant increase in AMPK signaling in DRGs for male (A) and female (C) mice 1 hour after injection but did not affect AMPK signaling in liver for males (B) or females (D). NCLS treatment increased the ratio of p-AMPK to total AMPK in male (E) and female (G) DRGs but not in male (F) or female (H) livers. *P < 0.05; Student’s t test; n = 4 per condition in male mice; n = 4 for female NCLS treatment, N = 3 for female vehicle.
for 1 hour. We observed a significant increase in p-ACC intensity with NCLS treatment at 1 μM, demonstrating that NCLS induces AMPK activation in DRG neurons (Fig. 3, A–D). This concentration is higher than previous demonstrations of AMPK activation with 20 nM NCLS in skeletal muscle cells, but those experiments were done over 48 hours of treatment and ours was done with a 1-hour treatment. The discrepancy in the concentration of NCLS needed to activate AMPK may be cell type- or time course-dependent. We conclude from this biochemical study that NCLS is a potent activator of AMPK in DRG neurons.

In addition to these in vitro findings, we used Western blotting to confirm that NCLS can activate AMPK in the DRG in males and females in vivo. Mice of both sexes were given a single dose of NCLS (3 mg/kg) and DRGs, and livers were taken 1 hour postinjection and homogenized for Western blotting. Primary antibodies used were p-AMPK and total AMPK to assess the effects of NCLS on AMPK signaling in these tissues. We observed a significant increase in p-AMPK/total AMPK ratio in both the male and female DRGs with NCLS treatment compared with vehicle (Fig. 4). No difference was found in the p-AMPK/total AMPK ratio in male or female livers with NCLS treatment, which is consistent with previously published data with this compound (Julien et al., 2017).

**Direct Allosteric AMPK Activators Reduce Hyperalgesic Priming in Male and Female Mice without Impacting Acute Incision-Induced Mechanical Hyperalgesia.** Metformin and NCLS activate AMPK via upstream signaling mechanisms. ZLN024 and MK8722 are structurally distinct AMPK activators that bind directly to the kinase to allosterically increase kinase activity, albeit via distinct mechanisms. We used these compounds because they have thoroughly described pharmacokinetics. After establishing baseline mechanical sensitivity measurements from male mice, we then assigned them to ZLN-024 (30 mg/kg) and vehicle (0.9% saline) i.p. treatment groups. These mice received intraperitoneal ZLN-024 for 7 consecutive days, and plantar incision surgery was performed on day 3 of treatment. Once the mice returned to baseline sensitivity, animals

![Fig. 6.](image)

**Fig. 6.** ZLN-024 treatment blocks hyperalgesic priming in male and female mice but had no effect on acute incision-induced mechanical hypersensitivity: (A–F). Although ZLN-024 treatment starting immediately after baseline measurements had no effect on initial mechanical hypersensitivity after plantar incision surgery, drug treatment was effective in preventing plantar incision-induced hyperalgesic priming precipitated by PGE2 injection in male (A–C) and female (D–F) mice. *P < 0.05; **P < 0.01; ***P < 0.001; ****P < 0.0001; n = 6 for the ZLN-024 group, n = 6 for the vehicle group.
received PGE$_2$ into the hindpaw and were tested 3 and 24 hours after injection. In male mice, ZLN-024 did not significantly decrease mechanical hypersensitivity after plantar incision surgery, but 7-day treatment did block hyperalgesic priming (Fig. 5, A–C). For the female cohorts, ZLN-024 also had no effect on the initial surgery-induced mechanical hypersensitivity, but it did block hyperalgesic priming after PGE$_2$ just as with the male cohort (Fig. 5, D–F). Using the same paradigm for testing in male mice, MK8722 did not significantly decrease mechanical hypersensitivity after plantar incision surgery, except a small effect at day 7 after incision; however, 7-day treatment did block hyperalgesic priming (Fig. 6, A–C). In the female cohorts, MK8722 again had no effect on the initial incision-induced mechanical hypersensitivity but also blocked hyperalgesic priming after PGE$_2$, just as with the male cohort (Fig. 6, D–F). These experiments demonstrate that direct activators of AMPK have a strong effect on hyperalgesic priming but do not influence pain hypersensitivity caused by acute incision. A separate group of male mice were allocated into MK8722 and vehicle treatment groups and baselined for von Frey and the rotarod. These mice were treated for 7 days and given a sham surgery on day 3. The mice were tested by the von Frey test throughout this time; after treatment, they were tested on the rotarod again to assess the effects of MK8722 on locomotive behavior. As with metformin and NCLS on sham mice, MK8722 had no effect on von Frey or locomotive behavior in male mice (Supplemental Fig. 5, A–D).

We have previously given two local injections of AMPK activators (one at the time of incision and another 24 hours later) to demonstrate that local activation of AMPK can...
attenuate incision-evoked mechanical hypersensitivity (Tillu et al., 2012; Burton et al., 2017). We assessed whether MK8722 (10-μg dose per injection) would affect incision-evoked pain. In both male (Fig. 7, A–C) and female (Fig. 7, D–F) mice, MK8722 had no effect on acute mechanical hypersensitivity, but local treatment with this direct AMPK activator did attenuate hyperalgesic priming (Fig. 7). This effect is consistent with the systemic dosing experiment with MK8722 but is contrasts with our previous findings with local injection of indirect AMPK activators (e.g., resveratrol) that blocked acute mechanical hypersensitivity and hyperalgesic priming (Tillu et al., 2012; Burton et al., 2017).

Like NCLS, MK8722 activates AMPK in many cell types but has not previously been tested on DRG neurons. DRGs were cultured from naïve male mice and treated with vehicle, 100 nM, or 10 μM MK8722 for 1 hour. We observed a significant increase in p-ACC intensity with MK8722 treatment at 100 nM and 10 μM, demonstrating that MK8722 induces AMPK activation neurons (Fig. 8, A–D). Quantification was done by measuring the maximum florescence (Fig. 8D). Based on this result, we conducted a full-concentration response curve for MK8722 and obtained an approximate EC50 of 900 nM (95% confidence interval, 253 nM–2.62 μM) for AMPK activation in DRG neurons (Fig. 8E). This EC50 for MK8722 is similar to observations in rat primary hepatocytes for AMPK activation (Myers et al., 2017).

**Plantar Incision Decreases pAMPK in the DRG Ipsilateral to Incision Injury.** Male mice went through the same behavioral battery as already described, and then plantar incision surgery was performed. Two hours postsurgery, lumbar DRGs were taken on the ipsilateral and contralateral sides from these mice and homogenized for Western blotting. We used p-AMPK and total-AMPK as the primary antibodies to assess the effects of injury on AMPK signaling. We saw

![Figure 8](https://example.com/figure8.png)

**Fig. 8.** Local MK8722 treatment attenuates hyperalgesic priming in male and female mice but had no effect on acute incision-induced mechanical hypersensitivity: Local injection of MK8722 at the time of incision and again 24 hours later (red arrows) had no effect on initial mechanical hypersensitivity after plantar incision surgery in male or female mice; however, drug treatment was effective in prevention of plantar incision-induced hyperalgesic priming precipitated by PGE2 injection in male (A–C) and female (D–F) mice. *P < 0.05; **P < 0.01; ***P < 0.001; n = 4 for the MK8722 group, n = 4 for the vehicle group in both male and female mice.
a significant decrease in p-AMPK/total-AMPK ratio in ipsilateral DRGs compared with contralateral DRGs (Figs. 9 and 10). This finding is consistent with previously published data demonstrating that pain-inducing stimuli cause a decrease in AMPK signaling in DRG (Atef et al., 2019). Given that many direct AMPK activators require AMPK phosphorylation for their pharmacologic activity, this potentially explains why those compounds are less efficacious for acute incisional pain.

Discussion

One of the key findings of this study is that whereas metformin’s sex differential effect on neuropathic pain (Inyang et al., 2019) can also be observed in the incisional pain model, other AMPK activators did not have a sex-specific effect on pain in our experiments. NCLS, ZLN-024, and MK 8722 all had the same level of efficacy on hyperalgesic priming in male and female mice. Metformin had a robust initial antihyperalgesic effect and blocked hyperalgesic priming in male mice but showed no effect in female mice. These sex differences are not readily explained by the pharmacokinetics of metformin. In fact, female mice had higher plasma and brain levels of metformin than did male mice. Although we do not have an explanation for the sex differences in metformin’s efficacy in incisional or neuropathic pain models in mice, our work does illustrate that other AMPK activators can be effective in the incisional model in both sexes. A possible mechanism that can be explored in future work is sexual...
AMPK phosphorylation is needed for AMPK activation with existing data on MK8722 has not directly assessed whether phosphorylate AMPK. A limitation of this idea is that the other hand, indirect AMPK activators increase AMPK phosphorylation is increased via some other mechanism. On the enhanced AMPK activation unless upstream AMPK phosphorylation is decreased by a pre-rely on phosphorylated AMPK to be able to increase AMPK AMPK. This decrease is because these drugs are thought to potentially decrease the ability of allosteric modulators to activate levels of phosphorylated AMPK would theoretically substantially decrease the activity and is controlled by sex hormones (Cai et al., 2019).

Another key finding of this study is our demonstration that indirect AMPK activators are more efficacious in the acute phase of mechanical hypersensitivity after incisional injury. Whereas the positive allosteric modulators ZLN-024 and MK 8722 showed a robust ability to block hyperalgesic priming, these drugs did little to attenuate the initial mechanical hypersensitivity caused by incision pain. In contrast, pre-treatment with NCLS (in both males and females) and metformin (in males) caused a large initial anti-hyperalgesic effect and more rapid resolution of surgery-induced hypersensitivity in addition to blocking priming. Despite this behavioral difference, we did not see a difference in vitro in the ability of these compounds to activate AMPK. Both NCLS and MK 8722 induced significant AMPK activation in DRG neurons culture with an EC₅₀ for MK 8722 that was consistent with values previously reported in the literature (Myers et al., 2017). It is unlikely that these differences occurred from pharmacokinetic or pharmacodynamic issues in the DRG because we used these compounds at in vivo doses that have been thoroughly vetted previously for AMPK activation. In line with this, we show here that NCLS activates AMPK in the DRG, but not liver, of male and female mice.

How, then, can the difference in acute effects of these different modes of AMPK activation be explained? One potential explanation for the behavioral difference we saw between the indirect AMPK activators and positive allosteric modulators is the decrease in pAMPK in the DRG caused by incision injury, which has been shown previously in diabetic neuropathic pain (Atef et al., 2019). A decrease in DRG neuron levels of phosphorylated AMPK would theoretically substantially decrease the ability of allosteric modulators to activate AMPK. This decrease is because these drugs are thought to rely on phosphorylated AMPK to be able to increase AMPK activity. If AMPK phosphorylation is decreased by a preceding injury, these drugs would achieve a smaller amount of enhanced AMPK activation unless upstream AMPK phosphorylation is increased via some other mechanism. On the other hand, indirect AMPK activators increase AMPK phosphorylation because they act via upstream kinases that then phosphorylate AMPK. A limitation of this idea is that the existing data on MK8722 has not directly assessed whether AMPK phosphorylation is needed for AMPK activation with this compound, but it is clearly a direct activator of the enzyme (Myers et al., 2017).

As previously mentioned, metformin activates AMPK indirectly through the upstream kinase LKB1 (Shaw et al., 2005). NCLS is thought to increase AMPK phosphorylation via a cAMP- and ADP/ATP ratio-dependent mechanism, which may act similarly to resveratrol (Julien et al., 2017). Importantly, resveratrol also inhibits acute incision-evoked pain and blocks the development of hyperalgesic priming (Burton et al., 2017). Therefore, from the perspective of therapeutic development of AMPK activators for the treatment of postsurgical pain, our findings support a focus on the potent indirect AMPK activators. Given that NCLS is far more potent than metformin, this molecule may represent an excellent starting point for further refinement in this space.

Another question arising from our work is why direct allosteric modulators of AMPK are effective in blocking the development of hyperalgesic priming but not acute incision-evoked hypersensitivity. Previous studies have made it clear that hyperalgesic priming can be completely reversed without having any effect at all on the acute pain phase (Asiedu et al., 2011; Price and Inyang, 2015) as we have shown here with ZLN024 and MK8722. Although we have not investigated this finding directly, we favor the hypothesis that these compounds block mTOR and MAPK-dependent protein synthesis sufficiently to attenuate gene expression of proteins that are needed for the development of hyperalgesic priming. This could include the synthesis of proteins, like CREB, that act as retrograde signaling factors from the site of injury to then regulate transcriptional programs that are needed for the transition to a chronic pain state (Melemedjian et al., 2014). Our findings suggest that inhibition of translation of these proteins requires less AMPK activation than does the translation of proteins required for the acute sensitization of nociceptors. This hypothesis can be explored in future studies.

Several shortcomings of our study should be addressed in the future. First, although we have done numerous pharmacologic manipulations in this study, we have not used genetic approaches to determine directly whether AMPK is responsible for the effects we have observed. Second, NCLS's pharmacology is potentially complex and may involve signaling pathways that are independent of AMPK. We cannot rule out this possibility based on our current experiments. Another caveat is that we have relied on evoked testing for pain assessment and have not used other methods, such as paw guarding or mouse grimace scale. Our future work will focus on non-evoked pain measures. Finally, whether the
hyperalgesic priming paradigm truly models development of chronic postsurgical pain is controversial; however, we are not aware of a widely accepted model of chronic postsurgical pain.

In conclusion, direct and indirect AMPK activators block the transition of acute pain to chronic pain that is potentially modeled preclinically by the hyperalgesic priming paradigm, and they can do this in both male and female mice; however, indirect AMPK activators have an acute antihyperalgesic effect that is not seen with direct positive allosteric modulators. Based on these findings, we conclude that indirect AMPK activators, like NCLS, should be further pursued as a treatment option for postsurgical pain that could achieve analgesia and a blockade of development of chronic pain in both sexes. More work is needed to determine the selectivity of NCLS at AMPK; but, given the favorable pharmacologic profile that has already been described for this compound, we propose that it has excellent potential as a starting point for further development.

Acknowledgments
We thank members of the Price, Duasor, and Burton laboratories for input throughout this project.

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2014) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.

References

Inyang KE, Burton MD, Price TJ (2019) Local translation and retrograde axonal transport of CREB and Price TJ (2014) Local translation and retrograde axonal transport of CREB

Prepared for publication: Inyang, Burton, Price.

We thank members of the Price, Dussor, and Burton laboratories for input throughout this project.

Address correspondence to:
Theodore Price, Jr.
Department of Anesthesiology
University of Texas Southwestern Medical Center
4323 Harry Hines Boulevard
Dallas, Texas 75390
E-mail: Theodore.price@utdallas.edu

Authorship Contributions

Participated in research design: Inyang, Burton, Duasor, Price.
Conducted experiments: Inyang, Szoabo-Pardi, Wentworth, McDoig, Ramirez, T. Prehand.

Performed analysis: Inyang, Burton, Price.

Wrote or contributed to the writing of the manuscript: Inyang, Duasor, Price.