L-cysteine modulates visceral nociception mediated by the CaV2.3 R-type calcium channels

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Abstract

CaV2.3 channels are subthreshold voltage-gated calcium channels that play crucial roles in neurotransmitter release and regulation of membrane excitability, yet modulation of these channels with endogenous molecules and their role in pain processing is not well studied. Here, we hypothesized that an endogenous amino acid l-cysteine could be a modulator of these channels and may affect pain processing in mice. To test this hypothesis, we employed conventional patch-clamp technique in the whole-cell configuration using recombinant CaV2.3 subunit stably expressed in human embryonic kidney (HEK-293) cells. We found in our in vitro experiments that l-cysteine facilitated gating and increased the amplitudes of recombinant CaV2.3 currents likely by chelating trace metals that tonically inhibit the channel. In addition, we took advantage of mouse genetics in vivo using the acetic acid visceral pain model that was performed on wildtype and homozygous Cacna1e knockout male littermates. In ensuing in vivo experiments, we found that l-cysteine administered both subcutaneously and intraperitoneally evoked more prominent pain responses in the wildtype mice, while the effect was completely abolished in knockout mice. Conversely, intrathecal administration of l-cysteine lowered visceral pain response in the wildtype mice, and again the effect was completely abolished in the knockout mice. Our study strongly suggests that l-cysteine-mediated modulation of CaV2.3 channels plays an important role in visceral pain processing. Furthermore, our data are consistent with the contrasting roles of CaV2.3 channels in mediating visceral nociception in the peripheral and central pain pathways.

Keywords R-type channels · Hyperalgesia · Inflammatory pain · Acetic acid · Visceral nociception

Introduction

Voltage-gated calcium channels (VGCCs) play essential roles in the nervous system including gene expression (L-type), neurotransmitter release (N-, P/Q-, and R-type), and membrane depolarization (T-type). The family of VGCCs consists of the pore-forming α1 subunit that provides the binding sites for agonists and antagonists. The CaV2.3 (former α1E-subunit) or R-type channel is a subtype of the VGCC family that was discovered and cloned relatively late, partly due to its resistance to the conventional calcium channel antagonists [16, 22]. Subsequent studies have shown that neuronal CaV2.3 channels in the periphery, spinothalamic, and thalamocortical tracts contribute to gene translation and cell membrane excitability [18]. Immunohistochemistry experiments have shown the abundance of CaV2.3 protein in the dorsal root ganglia (DRG) and reticular nucleus of thalamus (RTN), as well as neocortex, suggesting their possible role in peripheral and central nociception and underlying sensory processing [5, 18, 23].

The role of endogenous redox and chelating agents, including amino acids and peptides in regulating pain processing, has been a sought-after subject in pain research. Electrophysiological studies have reported enhanced firing frequency in neurons in the presence of endogenous reducing amino acid l-cysteine [7]. L-cysteine could be found locally in the interstitial spaces in relatively high concentrations under conditions of metabolic stress or tissue injury [11]. Although previous studies have suggested a few specific pathways for the action of these substances,
the thorough mechanism remains elusive and sometimes controversial. For example, l-cysteine in the extracellular environment may oxidize cysteine groups on the ion channel membrane thereby manipulating channels' natural dynamics [4, 21]. On the other hand, there are arguments on the chelating properties of l-cysteine rather than oxidizing, where chelating agents regulate trace metals in the environment thus manipulating function of an ion channel [8].

Our group previously reported that CaV2.3 channel function in sensory neurons can be inhibited by trace metals and facilitated with certain endogenous chelating agents such as glutamate [19]. However, functional significance of such modulation has not been previously studied. Hence, we hypothesized that modulation of the CaV2.3 channels via endogenous redox/chelating agents could affect pain processing in mice. Herein, using human recombinant CaV2.3 subunit stably expressed in human embryonic kidney cells (HEK-293) and mouse genetics, we have studied the role of CaV2.3 channels and an endogenous amino acid such as l-cysteine on sensitization of visceral pain responses.

Materials and methods

Human embryonic kidney 293 cells

Human variant of CaV2.3 was stably transfected in HEK-293 cells that were grown in DMEM (Sigma-Aldrich D5796-6X500ML) supplemented with 10% fetal calf serum (Sigma-Aldrich F0679-500ML), 100 U/ml penicillin (Sigma-Aldrich P4333-100ML), 0.1 mg/ml streptomycin (Sigma-Aldrich P4333-100ML), and 0.5 mg/ml gentamycin (Sigma-Aldrich A1720-1G), and 0.2 mg/ml hygromycin-B to stabilize channel expression (Sigma-Aldrich H3274-250MG) [18].

Electrophysiology

Electrophysiological experiments were performed using the conventional patch-clamp technique in the whole-cell configuration at room temperature (22 °C) using Cornerstone PC-ONE amplifier and Digidata 1322A with pClamp 10.7 software (Molecular Devices). Data were filtered at 2 kHz and digitized at 5 kHz. Cell capacitance, membrane resistance, access resistance, and holding potential were measured using the built-in Membrane Test function. The average access resistance was around 4 MΩ. Currents were recorded in the following external solutions (in mM, all purchased from Sigma-Aldrich): 160 TEA-Cl, 10 BaCl2, 0.1 EGTA, and 10 HEPES, pH adjusted to 7.4 with TEA-OH. Pipettes for recordings (around 4 MΩ resistance) were filled with internal solution of the following composition (in mM): 100 Cs-methane sulfonate, 14 phosphocreatine, 10 HEPES, 9 EGTA, 5 Mg-ATP, 0.3 Tris-GTP, pH adjusted to 7.3 with Cs-OH. The recording chamber was constantly perfused with the extracellular solution at 1 ml/min.

Animals

Experimental procedures with animals were performed according to a protocol approved by the Institutional Animal Care and Use Committee of the University of Colorado Anschutz Medical Campus, Aurora, CO, USA. Treatments of animals adhered to guidelines set forth in the NIH Guide for the Care and Use of Laboratory Animals. All efforts were made to minimize animal suffering and to use only the necessary number of animals to produce reliable scientific data. Wild-type and homozygous Caena1e mutant male C57BL/6 J littermates were obtained from the Mutant Mouse Resource and Research Centers (MMRRC) and used for pain behavior studies as outlined elsewhere [15]. All animals were maintained on a 14/-10-h light/dark cycle with food and water ad libitum. Behavioral assessments for in vivo experiments were done in a blinded fashion.

Acetic acid visceral pain model

Visceral pain was induced by injection of 300 µl acetic acid (in 0.6% saline) intraperitoneally (i.p.). L-cysteine was administered either systemically at 10 mg/kg subcutaneously (s.c.), or 36 µg/300 µl intraperitoneally (i.p.), or intrathecally. For intrathecal (i.t.) administrations, animals were anesthetized with 2% isoflurane, and 1.2 µg/10 µl l-cysteine solution in saline was injected between L4 and L5 vertebrae about 20 min prior to i.p. acetic acid injections. The correct i.t. injection site was confirmed by observing subjects’ tail flick. Writhing reflex of subjects were recorded and scored for 30 min post acetic acid injections [2].

Analysis

Data were analyzed using Clampfit 10.7 (Molecular Devices), MS Excel 2007 (Microsoft), and GraphPad Prism 9. Smooth curves in the figures represent fits to the average data, whereas the mean values in the table were calculated from fits to individual cells. Data are reported as the mean±SEM. We used either one-way ANOVA, to compare multiple mean values, or a paired Student’s t-test when evaluating the effect of acute application of agents. The voltage dependence of activation and inactivation was determined by fitting the peak current–voltage (I–V) data from each cell with a Boltzmann-Goldman–Hodgkin–Katz (GHK) equation

\[ G(V) = G_{\text{max}} \frac{1}{1 + \exp[-(V - V_{\text{gp}})/k]} \]

and

\[ I(V) = \frac{I_{\text{max}}}{1 + \exp \left[ -\frac{V - V_{\text{gp}}}{k} \right]} \].
Results

L-cysteine augments calcium currents in Ca_{2.3}-transfected HEK-293 cells at physiologically relevant concentrations

HEK-293 cells were perfused with external solution alone (baseline) and 50 µM l-cysteine in external solution (perfusion) consecutively to study the effect of l-cysteine on the gating of Ca_{2.3} channels (Fig. 1). Current densities were obtained by normalizing the peak current amplitude of each cell to its capacitance and averaging across all examined cells and plotted against test potentials accordingly. Figure 1a depicts inward traces for the baseline and perfusion, where red traces are recorded at –10 mV test potential in either experiment. We noted that 50 µM l-cysteine robustly increased barium current from the baseline about twofold (peaking at -10 mV) with only partial recovery in most of the tested cells. Peak current densities at –10 mV test potential are shown in Fig. 1b. Figure 1c shows the I–V curve for the baseline (black circles) and perfusion (red squares) where 50 µM l-cysteine has significantly shifted the peak to hyperpolarized potentials by about

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5 mV. To accurately determine the R-type calcium channels gating properties in the presence of 50 µM l-cysteine, activation and inactivation protocols were applied (Fig. 1d, e). While no significant difference was observed in the inactivation kinetics between two groups, l-cysteine shifted $V_{50}$ of activation to hyperpolarized potentials by about 10 mV. Analyzing 10–90% rise time in I–V curves in Fig. 1f revealed that l-cysteine significantly speeds up $\text{Ca}_{\text{V}2.3}$ currents compared to the baseline. We fitted and analyzed activation and inactivation curves (Fig. 1g, h respectively) for each cell and averaged among the test potentials to obtain fast and slow time constants. Overall, Fig. 1 proved the significant effect of 50 µM l-cysteine on facilitation of channel gating and augmenting R-type calcium current densities. In order to investigate the interaction of l-cysteine with $\text{Ca}_{\text{V}2.3}$ channels under higher pathologically relevant concentrations (e.g., inflammation), we repeated our experiment using 500 µM l-cysteine.

**L-cysteine augments R-type calcium currents at high concentrations**

While 50 µM l-cysteine proved to be an effective endogenous R-type calcium current modulator at low concentrations, the question about the role of pathologically relevant higher doses of l-cysteine that may exist under pain or inflammation remains unclear. Figure 2 summarizes patch-clamp electrophysiological recordings in $\text{Ca}_{\text{V}2.3}$-transfected HEK-293 cells perfused with 500 µM l-cysteine. Figure 2a depicts barium current traces for baseline and perfusion. Red traces are recorded at −10 mV test potential where perfusion has increased the inward current compared to the baseline. Similar to lower concentration, we found that at 500 µM l-cysteine significantly increased current from the baseline about twofold (Fig. 2a, b) peaking at −10 mV. In Fig. 2c, we show that 500 µM l-cysteine was applied and washed in the course of 16 min to find out that it reversibly increased R-current amplitude with a relatively fast on and off kinetics. Figure 2d shows the I–V curve for the baseline (black circles) and perfusion (red squares) where 500 µM l-cysteine has significantly shifted the peak to hyperpolarized potentials by about 10 mV. The $\text{Ca}_{\text{V}2.3}$ channel gating in the presence of 500 µM l-cysteine was determined by running activation (Fig. 2e) and inactivation (Fig. 2f) protocols. Similar to 50 µM concentration, 500 µM l-cysteine perfusion left inactivation kinetics of $\text{Ca}_{\text{V}2.3}$ channels practically intact; however, $V_{50}$ of activation was shifted to hyperpolarized potentials by over −13 mV. Figure 2g and 2h investigate the possible role of 500 µM l-cysteine in recovery from inactivation and deactivation time constants of $\text{Ca}_{\text{V}2.3}$ channels; however, no significant effect was observed.

**L-cystine blocks $\text{Ca}_{\text{V}2.3}$ currents**

In order to validate if l-cysteine manipulates currents arising from $\text{Ca}_{\text{V}2.3}$ channels by oxidation mechanisms, we perfused $\text{Ca}_{\text{V}2.3}$-transfected HEK-293 cells with its oxidized analog l-cystine at 500 µM (Fig. 3). L-cystine consists of two cysteine molecules bridged by an oxidized thiol group. Figure 3a depicts current traces for the baseline and perfusion. Red traces are recorded at −10 mV test potential. Interestingly, we found that 500 µM l-cystine did not increase $\text{Ca}_{\text{V}2.3}$ currents, and instead it partially blocked the currents. L-cystine inhibited peak amplitudes of $\text{Ca}_{\text{V}2.3}$ currents by 50% in an apparently irreversible manner, suggesting a possible chemical nature of its mechanism (Fig. 3a, b). Figure 3c shows I–V curve for the baseline (black circles) and perfusion (red squares); while 500 µM l-cystine helps currents reach apex at lower membrane potentials, it also partially blocks the currents. Similar to l-cysteine, activation curve of the $\text{Ca}_{\text{V}2.3}$ channels reveals that l-cystine shifted $V_{50}$ of activation by −10 mV (Fig. 3d). Although 10–90% rise time was trending lower after perfusing l-cystine, it was not significantly different (Fig. 3e). We also fitted and analyzed activation (Fig. 3f) and inactivation curves (Fig. 3g) for each cell and averaged among the test potentials to obtain fast and slow time constants; no significant difference was observed.

**Chelating agents increase $\text{Ca}_{\text{V}2.3}$ currents**

We previously demonstrated that chelating mechanisms play a role in enhancing T-type calcium currents in sensory neurons [8]. In order to examine whether chelating mechanisms may be contributing to current manipulation arising from recombinant $\text{Ca}_{\text{V}2.3}$ channels, we perfused cells with a metal-chelating agent EDTA (Fig. 4). As expected, perfusing cells with EDTA at either 50 or 100 µM concentrations significantly increased currents in HEK-293 cells (Fig. 4a, b); however replacing EDTA in the bath with 50 µM l-cysteine did not increase the current any further (Fig. 4c, d). The data presented in Fig. 4 suggest that introducing a chelating agent to $\text{Ca}_{\text{V}2.3}$ channels augments their currents similar to l-cysteine. Furthermore, we conclude that l-cysteine manipulates $\text{Ca}_{\text{V}2.3}$ currents most likely through chelating constitutively bound trace metals rather than oxidizing mechanisms (Fig. 4e, f).

**In vivo pain studies show that $\text{Ca}_{\text{V}2.3}$-deficient mice show hyperalgesia and different responses to injections of l-cysteine**

In order to investigate a possible role of $\text{Ca}_{\text{V}2.3}$ channels in visceral pain processing, we compared pain behavior of wildtype (WT) and $\text{Ca}_{\text{V}2.3}$ null (2.3 KO) littermates.
We used pain model consisting of i.p. injections of 300 µl 0.6% acetic acid (A.A.) in saline and recorded the writhing reflexes of the subjects for 30 min [3]. Figure 5a shows the time course of pain behavior in male WT (black circles) and 2.3 KO mice (red squares). The writhing responses were greater in mice lacking Cav2.3 channels at each time point up to fourfold (Fig. 5b), showing a significant role of these channels in this pain model for the first time. Since our patch-clamp data showed that l-cysteine has strong interactions with Cav2.3 calcium channels (see Figs. 1, 2 and 3), we proposed that it may be able to manipulate the pain behavior of WT subjects differently than in KO mice. Figure 6a compares the effect of systemic subcutaneous (s.c.) administration of l-cysteine (see the “Materials and Methods” section).
and methods” section) in WT and 2.3 KO mice. While the hyperalgesic effect in WT subjects was significant, it was abolished in the 2.3 KO mice. In Fig. 6b, the experiment was repeated by administering a very small dose of l-cysteine peripherally using i.p. injections together with injections of A.A. Similar to s.c. route, we found that l-cysteine administered i.p. increased pain reflex in WT littermates but not in 2.3 KO mice. Figure 6a and 6b together suggest that not only Cav2.3 channels are involved in nociception, but also augmenting R-type calcium currents systemically or peripherally will increase pain sensitization. Next, we injected under anesthesia WT and 2.3 KO subjects intrathecally (i.t.) in the lower lumbar spinal cord with l-cysteine prior to the conducting
Fig. 4 The effect of metal-chelating agent EDTA on recombinant CaV2.3 currents. 

a At 50 µM, EDTA significantly increased inward currents in 3 cells (paired two-tailed t-test, \( P=0.02 \)), replacing EDTA with 50 µM l-cysteine in the bath did not cause an additional increase in currents in the same cells. 

b At 100 µM, EDTA significantly increased currents in 6 cells (paired two-tailed t-test, \( P=0.003 \)), replacing EDTA with 50 µM l-cysteine in the bath did not cause a further increase of currents in the same cells. 

c, d Traces of inward CaV2.3 currents arising at −10 mV test potential for 50 µM (c) and 100 µM (d) EDTA followed by 50 µM l-cysteine solution perfusion. 

e, f Perfusing 3 cells with 50 µM EDTA (e) and 6 cells with 100 µM EDTA (f) significantly increased the inward current and diminished any further l-cysteine effects. All currents are normalized to the maximal current density achieved with perfusion of EDTA.

Fig. 5 Role of CaV2.3 channels in inflammatory visceral nociception. 

a Time course of pain reflex counts in acetic acid pain model in WT and CaV2.3 KO male mice with 5-min intervals (two-way ANOVA \( P=0.005 \) with Sidak multiple comparisons). 

b Total cumulative writhing reflex counts of 7 male WT and 2.3 KO mice (two-tailed t-test, \( P=0.004 \)).
visceral pain testing (Fig. 6c). Interestingly, in contrast to systemic and peripheral injections, we found that central administration of l-cysteine using i.t. route reduced the pain response in the WT subjects while the effect was abolished in Ca,2.3-deficient mice. Hence, our data demonstrate opposing roles of Ca,2.3 channels in the periphery and spinal cord in visceral pain transmission demonstrated through injections of l-cysteine. Interestingly, we noted that baseline writhing responses upon i.t. injections of l-cysteine were decreased in both WT and mutant cohorts when compared to baseline responses in our experiments with i.p. and s.c. injections. This may be due to use of relatively brief isoflurane anesthesia that is necessary for i.t. injections.
Discussion

Herein, we first demonstrated that endogenous amino acid l-cysteine facilitates activation of Ca_{v}2.3 channels and consequently enhances its current density likely by a metal-chelating mechanism. Further, we have demonstrated the opposing effects of intrathecally and systemically administered l-cysteine modulation of Ca_{v}2.3 channels in A.A.-induced abdominal inflammatory pain model. We found that l-cysteine administrated i.t. partially blocks pain transmission in the spinal cord of WT littermates, while its effect was abolished in Ca_{v}2.3 KO mice. Meanwhile, s.c. and i.p. administration of l-cysteine induced pain sensitization in WT subjects, but not in Ca_{v}2.3 KO littermates.

**Ca_{v}2.3 currents are augmented with l-cysteine**

Previous studies have investigated the interaction between l-cysteine and Ca_{v}3.2 isoform of T-type calcium channels in the dorsal root ganglion (DRG) neurons [7–9]. It was shown that similar to this study, l-cysteine acts via chelating trace metals and increases T-current activation kinetics, and consequently increases subthreshold excitability. Knowing that R-type and T-type calcium channels have dynamic properties and structural similarities, we investigated the interactions between R-type calcium channels and endogenous l-cysteine. We hypothesized that l-cysteine may have the similar effect on these channels. Thus, using Ca_{v}2.3-transfected HEK-293 cells, we recorded the inward currents using whole-cell patch-clamp recording. As expected, the results for both 50 and 500 µM L-cysteine showed increased current densities and facilitation of Ca_{v}2.3 channel gating as demonstrated by a leftward shift in current activation curves.

**L-cystine partially blocks Ca_{v}2.3 currents**

Looking at the amino acid construct of the outer membrane segment of the Ca_{v}2.3 channel, there are a few sites that resemble the cysteine residue of T-type calcium channels [10, 19]. Thus, we designed experiments to introduce cystine directly to the recording bath and study its effect on recombinant Ca_{v}2.3 currents. In contrast to l-cysteine, we found that l-cystine partially blocked Ca_{v}2.3 currents. Looking at the opposite results and what we observed with l-cysteine, we concluded that the mechanism of l-cysteine action is not oxidizing the thiol group on cysteine residues of Ca_{v}2.3 channels.

**L-cysteine acts on Ca_{v}2.3 channels likely through chelating the trace metals**

Trace metals are highly regulated in the nervous system and serve as neuronal excitability co-factors [4, 6]. The abundance of certain trace metals (Zn^{2+} and Cu^{2+}) has been reported in neurological disorders namely epilepsy, stroke, Alzheimer’s disease, and Parkinson’s [1, 4]. In contrast to mineral divalent ions, trace metals have extremely low concentrations in the nervous system (70–150 µM for Zn^{2+} and Cu^{2+}) where a family of intracellular proteins, called metalloproteins, regulate them [6, 21]. While the vast body of trace metals are bound to these proteins, the free ion (chelatable) concentrations can reach up to 30 µM for Zn^{2+} and 1.7 µM for Cu^{2+}. These “chelatable” free trace metal bodies can be found in the synaptic clefts in the cortex, limbic system, and basal ganglia [1]. In order to investigate the role of l-cysteine in chelating trace metals in the recording environment, we perfused cells with a known chelating agent EDTA and then replaced the bath solution with 50 µM l-cysteine. While chelating agent EDTA increased the Ca_{v}2.3 current upon perfusion, subsequent l-cysteine perfusion effect was effectively occluded. Each experiment resulted in a finite current increase; however, once l-cysteine was introduced to the bath, a significant current increase was observed. Taken together, these results shed light on the action mechanism of l-cysteine and strongly suggest the metal-chelating properties of l-cysteine are the cause of current manipulation in Ca_{v}2.3 channels.

**L-cysteine sensitizes visceral pain in WT mice**

Although functional role of Ca_{v}2.3 channels has been investigated in the previous studies [12, 13, 17], their thorough physiological properties remain elusive. The Ca_{v}2.3 calcium channels are widely distributed throughout the thalamus (GABAergic reticular nucleus), cortex, and the spinal cord [22]. The distribution of Ca_{v}2.3 channels in the main pain processing pathway, from dorsal horn spinal cord to the cortex, made us curious on their role in an abdominal inflammatory pain model. Somewhat surprisingly, our results show that Ca_{v}2.3 KO mice exhibited more prominent visceral pain responses when compared to the WT littermates. The lower pain threshold in the global 2.3 KO subjects could be the result of removal of Ca_{v}2.3 channels in the brain, spinal cord, and/or the periphery of null mice. Moreover, Ca_{v}2.3 channels could serve as a novel analgesic target, thus we were interested to see if regulating the systemic level of l-cysteine would have any major effects on blocking pain in transgenic mice. We observed that systemic application of l-cysteine 20 min prior to the pain behavior assay or peripheral applications at the same time with A.A. increases pain responses in WT subjects significantly; however, the effect is largely diminished in KO subjects. These results together demonstrate the specificity of l-cysteine action to Ca_{v}2.3 channels. However, dissecting their pharmacological target (central vs. periphery) requires further experiments. Our results are also not sufficient to exclude the possibility of...
the effect of l-cysteine on signaling pathways on other voltage-gated calcium channels, and Caᵥ3.2 isoform of T-type channels in particular [7–9, 14, 20]. Since both Caᵥ2.3 and Caᵥ3.2 channels are expressed in peripheral and central pain pathways, we conclude that the effects of L-cysteine on pain processing in vivo are likely mediated by both of these channels in concert. In addition, our results strongly suggest that Caᵥ2.3 channels are not saturated with endogenous levels of chelators of trace metals in vivo, since we were still able to modulate nociception with exogenous application of L-cysteine in WT mice.

L-cysteine partially blocks pain transmission in the spinal cord

In order to distinguish the role of Caᵥ2.3 channels in the spinal cord, we injected WT and KO littermates with l-cysteine i.t. 20 min prior to the pain behavior task. We found that WT subjects injected with l-cysteine i.t. had less prominent visceral pain reflexes compared to vehicle (saline). In contrast, the effect of i.t. injections of l-cysteine in the KO subjects was subliminal and there was no significant difference between drug and the vehicle, which again demonstrates the specificity of l-cysteine action on Caᵥ2.3 channels in the spinal cord. Although these results prove that spinally expressed Caᵥ2.3 channels are manipulated upon exposure to l-cysteine, further experiments are required to clarify the plausible mechanisms of interactions of l-cysteine with Caᵥ2.3 channels in the dorsal horn of the spinal cord, a main pain processing region.

Conclusion

Herein, we have demonstrated the crucial role of Caᵥ2.3 channels in visceral pain transduction, and shown the acting mechanism of an endogenous chelating agent, l-cysteine, through chelating trace metals in the spinal cord to block pain transmission. In contrast, systemic and peripheral administration of l-cysteine augmented abdominal writing responses suggesting different roles for peripheral and central Caᵥ2.3 channels in visceral pain processing.

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Declarations

Conflict of interest The authors declare no competing interests.

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