An unbiased and efficient assessment of excitability of sensory neurons for analgesic drug discovery

Zainab A. Mohammed, Katerina Kaloyanova, Mohammed A. Nassar*

Abstract
Alleviating chronic pain is challenging, due to lack of drugs that effectively inhibit nociceptors without off-target effects on motor or central neurones. Dorsal root ganglia (DRG) contain nociceptive and non-nociceptive neurones. Drug screening on cultured DRG neurones, rather than cell lines, allows for the identification of drugs most potent on nociceptors with no effects on non-nociceptors (as a proxy for unwanted side effects on central nervous system and motor neurones). However, screening using DRG neurones is currently a low-throughput process, and there is a need for assays to speed this process for analgesic drug discovery. We previously showed that veratridine elicits distinct response profiles in sensory neurones. Here, we show evidence that a veratridine-based calcium assay allows for an unbiased and efficient assessment of a drug effect on nociceptors (targeted neurones) and non-nociceptors (nontargeted neurones). We confirmed the link between the oscillatory profile and nociceptors, and the slow-decay profile and non-nociceptors using 3 transgenic mouse lines of known pain phenotypes. We used the assay to show that blockers for Na1.7 and Na1.8 channels, which are validated targets for analgesics, affect non-nociceptors at concentrations needed to effectively inhibit nociceptors. However, a combination of low doses of both blockers had an additive effect on nociceptors without a significant effect on non-nociceptors, indicating that the assay can also be used to screen for combinations of existing or novel drugs for the greatest selective inhibition of nociceptors.

Keywords: Calcium imaging, Veratridine, Voltage-gated channel blockers, Nav1.7 Knockout, Nociceptors, Mouse DRG, Capsicin, Drug screening

1. Introduction
Chronic pain has significant negative impacts at personal, social, and economic levels, affecting millions of people worldwide.10,19 Several drug classes are used to treat chronic pain, but their effectiveness is limited by either lack of potency or adverse effects. Therefore, the treatment of severe or chronic pain is a challenging clinical need.

The search for new analgesics focuses on targets in pain neurones in the dorsal root ganglia (DRG). Dorsal root ganglia contain a heterogeneous population of neurones that can be classified according to expression of sensory markers into 11 subtypes, but they can be broadly classified as nociceptors and non-nociceptors (proprioceptors and touch neurones). Drug screens use cell lines expressing the nociceptive target of interest. Although this platform allows for high-throughput screening, it provides no information on drugs’ effects on non-nociceptors. This information would weigh the potency on nociceptors against adverse effect on non-nociceptors before costly and laborious in vivo testing. However, screens on DRG neurones are currently not practical because they are low throughput, and there is not a simple protocol to distinguish between drugs’ effects on nociceptors vs non-nociceptors.

Voltage-gated sodium channels (VGSCs) are critical for the excitability in DRG neurones. We discovered that the VGSC opener, veratridine, produces distinct calcium responses in cultured sensory neurones. We found that most nociceptors show an oscillatory response (OS), whereas most non-nociceptors show a slowly decaying response (SD). Figure 1. Here, we provide evidence that the OS and SD veratridine-response profiles can be used as readouts in an assay to evaluate the effect of drugs on nociceptors and non-nociceptors. We first showed that the OS population is drastically reduced in a mouse model where most nociceptors are ablated. We then determined the pattern of veratridine responses in a Na1.7 knockout mouse, a pain-free model. Subsequently, we used this pattern as a reference criterion against which 2 VGSC blockers, against the Na1.7 and Na1.8 channels, were evaluated. These channels are validated targets for analgesic drug development with several promising compounds identified in recent years, but lack of in vivo potency and selectivity on nociceptors has contributed to failure in clinical trials. The 2 VGSC blockers were unable to match the Na1.7 knockout pattern when applied separately but showed an additive effect when applied together. This suggests that our assay can also be used to screen for combinations of drugs that act on different targets in nociceptors for effective pain relief.

2. Methods

2.1. Dorsal root ganglia culture
CS7BL6 mice were sacrificed according to Schedule 1 of the Animal (Scientific procedure) Act 1986. Dorsal root ganglia from

Sponsors or competing interests that may be relevant to content are disclosed at the end of this article.
Z.A. Mohammed and K. Kaloyanova contributed equally to this work.
Department of Biomedical Science, University of Sheffield, Sheffield, United Kingdom
*Corresponding author. Address: Biomedical Science, Firth Court, Western Bank, University of Sheffield, Alfred Deny Building, C-floor, Room C223, S10 2TN Sheffield, United Kingdom. Tel.: +44-1142222392; fax: +44-1142222787. E-mail address: m.nassar@sheffield.ac.uk (M.A. Nassar).

PAIN 161 (2020) 1100–1108
Copyright © 2020 The Author(s). Published by Wolters Kluwer Health, Inc. on behalf of the International Association for the Study of Pain. This is an open access article distributed under the terms of the Creative Commons Attribution-Non Commercial-No Derivatives License 4.0 (CCBY-NC-ND), where it is permissible to download and share the work provided it is properly cited. The work cannot be changed in any way or used commercially without permission from the journal.
http://dx.doi.org/10.1097/j.pain.0000000000001802
all spinal levels were dissected and collected in phosphate-buffered saline at room temperature. Phosphate-buffered saline was then replaced with 1-mL dissociation solution containing Dulbecco’s Modified Eagle’s Medium/F12 (DMEM/F12) with Glutamax (Gibco, United Kingdom), 1× penicillin/streptomycin mix (Gibco), dispase (1 mg/mL, Sigma, United Kingdom), and collagenase Type XI (0.6 mg/mL, Sigma) and incubated at 37˚C and 5% CO2 for 90 minutes. Dorsal root ganglia were triturated with a P1000 pipette tip 10 times at 60 and 90 minutes. After the second trituration, the cell suspension was carefully layered on top of 15% bovine serum albumin (Melford, United Kingdom), dissolved in DMEM/F12 containing penicillin/streptomycin, and centrifuged at 800g for 10 minutes with the minimum deceleration speed. The cell pellet was then washed in a culture medium composed of DMEM/F12 containing 10% fetal bovine serum and penicillin/streptomycin (all from Gibco). Cells were pelleted again and then resuspended in 60-μL culture medium. A 3-μL “drop” was placed on the centre of a glass coverslip coated with D-polyornithine (20 μg/mL, Sigma). Coverslips were flooded with a 1 mL of culture medium after 15 to 30 minutes. Cells were imaged 24 to 48 hours after plating. Dorsal root ganglia from 1.8-DTA, Nav1.7KO, and Nav1.8KO were placed in ice-cold Hibernate-A medium (Gibco) containing penicillin/streptomycin while being transported from UCL (about 5 hours). Dorsal root ganglia were then dissociated as above.

### 2.2. Calcium imaging

All recordings were performed at room temperature (22-25˚C). Dorsal root ganglia neurons were loaded with 2 μM Fura-2AM (ThermoFisher, United Kingdom) in standard Ringer solution (140 mM NaCl, 4 mM KCl, 2 mM CaCl2, 10 mM HEPES, 5 mM glucose, and pH = 7.4 with NaOH) for 30 minutes at 37˚C. Coverslips were then washed with ringer solution and left for 15 minutes at 37˚C/5% CO2 after which the Ringer was replaced. Coverslips were stored in a dark container at room temperature until used. Cells were perfused at a flow rate of 3 mL/minute with Ringer solution for at least 5 minutes to establish a stable calcium baseline. Ringer with 40 mM KCl was perfused at the end of recordings to identify viable neurons. Cells were imaged with a 40× objective and a Hamamatsu C4742-95 camera. Cells were excited with 350 and 380 nm for ratiometric measurement of intracellular calcium using Caim Dual OptoLED system. Simple PCI 6 software was used for data acquisition, background subtraction, and Fura-2AM ratiometric measurement (F350/380 nm).

All drugs were made in standard Ringer solution from stock solutions of the following concentrations: veratridine (5 mM in ethanol, Abcam ab120279), capsaicin (10 mM in ethanol, Tocris 0462), α, β-methylene ATP (10 mM in water, Sigma M6517), allyl isothiocyanate (AITC; 100 μM, Sigma 377430), 4,9-anhydrotetrodotoxin (300 μM in water, Tocris 6159), PF-04856264 (10 mM in DMSO, Sigma 11916), and A-803467 (10 mM in DMSO, Abcam Ab120282).

### 2.3. Data and statistical analysis

Neurons were included in the analysis if they respond to 40 mM KCn. On rare occasions, neurons responded to veratridine but not KCn (or the KCn response was not clear due to the calcium signal not returning to baseline after the application of last agonist). We defined a response as an increase in (F350/380) ratio of >6 SD above the baseline. Differences in fluorescence (ΔF/F₀) were calculated according to the following formula: F350/380 ratio during agonist application (F) minus the average F350/380 ratio of the 2.5 minutes before agonist application (F₀). Mean values from each experiment (each N is a culture from one mouse) were compared with each other by one-way analysis of variance (ANOVA) with Sidak’s post-test. All statistical analysis and comparisons were performed by GraphPad Prism software (version 7.00 for Windows).

### 3. Results

#### 3.1. The oscillatory response population is highly diminished in the Nav1.8-DTA (nociceptor-ablated) model

To confirm the link between neurons with the OS profile and pain behaviour, we examined DRG from the Na1,8Cre-DTA (1.8-DTA) mouse. The Na1,8 channel is expressed in 80% to 90% of neurons. The 2 minor profiles, ID and RD, are mostly nociceptors but can be excluded from an assay for simplicity as both account for less than 5% to 10% of all neurons.

- **Figure 1.** Assessments of the excitability of nociceptors and non-nociceptors based on veratridine-response profiles. Dorsal root ganglia (DRG) contain a heterogeneous population of sensory neurons. Nociceptors are typically small in size, express the Na1.8 channel, and respond to nociceptive compounds (eg, ATP and capsaicin). Most nociceptors responded to veratridine with an oscillatory (OS, rapid decay (RD), or intermediate decay (ID) profiles. Non-nociceptors are typically large in size, do not express the Na1.8 channel, and do not respond to nociceptive compounds. Most non-nociceptors respond to veratridine with a slow-decay (SD) profile. The percentages of the OS and SD veratridine profiles plus the percentage of veratridine-irresponsive (VTD-) neurons can be used for an efficient assay of the excitability DRG neurons. Changes in the SD population reflect changes in non-nociceptors (blue, typically 15%-20% of all neurons). Changes in the OS population (red, typically 30%-40% of all neurons) reflect changes in nociceptors. Changes in the veratridine-irresponsive population (gray, typically 30%-40% of all neurons) can reflect sensitisation of high-threshold and normally “silent” neurons. The 2 minor profiles, ID and RD, are mostly nociceptors but can be excluded from an assay for simplicity as both account for less than 5% to 10% of all neurons.
nociceptors. In the 1.8-DTA mouse, Na\textsubscript{1.8}-expressing neurons are ablated (Fig. 2A) leading to the loss of most nociceptors and a profound loss of pain. Therefore, we hypothesised that the veratridine-response profile that pertains to nociceptors (i.e., OS) would be reduced in the 1.8-DTA DRG. First, we determined the percentage of remaining nociceptors in 1.8-DTA DRG using the 3 nociceptive agonists (capsaicin, αβ-methyleneadenosine triphosphate, and isothiocyanate) to identify all subtypes of nociceptors as described previously. The percentage of neurons responding to any of the nociceptive agonists was drastically reduced from 75% in controls to just 8% in 1.8-DTA (Fig. 2B). This shows that most neurons in 1.8-DTA DRG are non-nociceptors with very few nociceptors left.

The ablation of most nociceptors caused opposing changes in the percentages of the veratridine OS and SD populations. The OS population decreased from 34% in controls to 7% (Fig. 2C). By contrast, the SD population increased from 17% in controls to 66%. This confirms that the OS population represents most nociceptors, whereas the SD population represents most non-nociceptors. Of note, the total number of veratridine-irresponsive neurons decreased from 39% in control to 23% in 1.8-DTA (Fig. 2C). The veratridine-irresponsive population can be either nociceptors (i.e., respond to the nociceptive agonists but not veratridine, yellow section in Fig. 2D) or non-nociceptors (i.e., respond to neither the nociceptive agonists nor veratridine, orange section in Fig. 2D). The observed decrease in veratridine-irresponsible neurons was in fact due to the loss of almost all veratridine-irresponsible nociceptors, a population that accounted for 26% of all neurons in controls but only 0.8% in 1.8-DTA.

Results from the 1.8-DTA mouse link the reduction of the OS population in vitro to the previously described pain deficits in vivo. Therefore, drugs can be screened based on their ability to reduce the OS population (an indication of potency on nociceptors) without an effect on the SD population (an indication of unwanted effects on non-nociceptors), Figure 1.

3.2. A reference veratridine-response pattern for a potent and safe analgesic

Next, we established the reference criterion by which drug potency and selectivity on nociceptors can be quickly ascertained.

![Figure 2.](image-url)
in our assay. We base this criterion on the change to the percentages of the OS and SD populations in the Na\textsubscript{v1.7} knockout mouse where pain behaviour is lost due to a decrease in the excitability of nociceptors (as opposed to their ablation as in 1.8-DTA). Na\textsubscript{v1.7} is a VGSC that is highly expressed in nociceptors and is a critical determinant of their excitability. Deletion of Na\textsubscript{v1.7} in DRG neurons leads to profound loss of pain without adverse motor or central nervous system (CNS) effects.\textsuperscript{14,17} Therefore, the changes in the OS and SD populations in the Na\textsubscript{v1.7} knockout represents the effects of a potent and safe analgesic. Therefore, drugs can be screened by our assay based on the similarity of their effect on DRG neurons to Na\textsubscript{v1.7} deletion.

To validate the use of the pattern of veratridine responses in Na\textsubscript{v1.7}KO as the criterion for a “good” analgesic, we compared it with the pattern of a would-be “bad” analgesic, a drug affecting non-nociceptors and CNS neurons. For this purpose, we used a Na\textsubscript{v1.6} channel blocker. Although conditional Na\textsubscript{v1.6} knockout suggests it contributes to pain,\textsuperscript{3} and it is not a potential analgesic target because of its role in CNS and motor neurons.\textsuperscript{12,18} In DRG, Na\textsubscript{v1.6} contributes more to sodium current in large (non-nociceptive) than in small (nociceptive) neurons.\textsuperscript{3} Therefore, we hypothesized that a Na\textsubscript{v1.6} blocker will reduce the OS and SD populations, but the effect on the SD population will be greater. Low concentrations of the tetrodotoxin metabolite, 4,9-anhydro-\textsuperscript{6}TTX (4,\textsuperscript{9}TTX), preferentially blocks Na\textsubscript{v1.6} (IC\textsubscript{50} 120 times lower than Na\textsubscript{v1.7}).\textsuperscript{20} As expected, 300 nM 4,\textsuperscript{9}TTX greatly increased the percentage of veratridine-irresponsive neurons from 42% to 68% (Fig. 3B). Unlike Na\textsubscript{v1.7} deletion, 4,\textsuperscript{9}TTX reduced both the SD and OS populations, but the reduction in the SD population (0.48 of control) was greater than the reduction in the OS population (0.62 of control). Therefore, drugs that reduce the SD population may act on non-nociceptive neurons with potential unwanted physiological effects.

To test whether our assay has the potential to detect increases in neuronal excitability (ie, as a readout for changes that lead to hyperexcitability, Fig. 1), we examined veratridine responses in the Na\textsubscript{v1.8} knockout mouse (Na\textsubscript{v1.8}KO). Na\textsubscript{v1.8} is also an important determinant of the excitability of nociceptors. The global Na\textsubscript{v1.8}KO mouse, however, showed a compensatory increase in Na\textsubscript{v1.7} expression.\textsuperscript{5} This is the most likely reason why the Na\textsubscript{v1.8}KO mouse does not show a major loss of pain. Because Na\textsubscript{v1.8} is expressed in nociceptors, then it is logical to assume that the compensatory increase in Na\textsubscript{v1.7} will occur in nociceptors. Therefore, we hypothesised that this compensatory increase in Na\textsubscript{v1.7} will primarily affect the OS population. In

**Figure 3.** Reference veratridine-response patterns for “safe” and “unsafe” analgesic drugs. (A) Deletion of Na\textsubscript{v1.7} causes a major loss of pain without adverse CNS or motor effects. The veratridine-response pattern of the Na\textsubscript{v1.7}KO represents that of a safe and potent analgesic. Na\textsubscript{v1.7} deletion leads to a decrease in responsiveness to veratridine (increase VTD) population due to a decrease in the OS population but not the SD population. VTD− CTR = 51.0 ± 3.51 vs VTD− KO = 80.4 ± 2.8; SDCTR = 9.1 ± 1.9 vs SDKO = 7.5 ± 1; OSCTR = 28.2 ± 2.2 vs OSKO = 6.7 ± 1.3; IDCTR = 7.4 ± 0.4 vs IDKO = 3.2 ± 0.6; RDCTR = 4.0 ± 0.6 vs RDKO = 1.9 ± 0.5%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 1448 neurons from 6 floxed-control mice and 1630 neurons from 6 Na\textsubscript{v1.8}KO. ANOVA, analysis of variance.

(B) 300 nM of the Na\textsubscript{v1.6} blocker 4,\textsuperscript{9}TTX reduces responsiveness to veratridine (increases VTD−) through decreases in both the SD and OS populations. The decrease in SD is greater than that in OS. VTD− CTR = 41.6 ± 3.1 vs VTD− 4,\textsuperscript{9}TTX = 68.0 ± 1.7; SDCTR = 20.68 ± 2.6 vs SD4,\textsuperscript{9}TTX = 10.0 ± 1.5; OSCTR = 28.7 ± 1.8 vs OS4,\textsuperscript{9}TTX = 17.6 ± 2.2; IDCTR = 5.4 ± 0.9 vs ID4,\textsuperscript{9}TTX = 3.1 ± 0.5; RDCTR = 3.5 ± 1.1 vs RD4,\textsuperscript{9}TTX = 1.4 ± 0.3%, One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 635 untreated and 927 treated neurons from 6 C57B6 mice. (C) The compensatory increase in Na\textsubscript{v1.7} channels in nociceptors of the Na\textsubscript{v1.8}KO increases responsiveness to veratridine (decreases VTD−) due to an increase in the OS population. VTD− CTR = 55.4 ± 3 vs VTD− KO = 57.1 ± 3.2; SDCTR = 6.9 ± 1.7 vs SDKO = 9.1 ± 1.1; OSCTR = 26.7 ± 2.0 vs OSKO = 40.3 ± 2.7, IDCTR = 7.0 ± 0.5 vs IDKO = 7.65 ± 1.1; RDCTR = 3.6 ± 0.6 vs RDKO = 5.7 ± 0.5%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 1493 neurons from 5 littermate-control mice and 1028 neurons from 4 Na\textsubscript{v1.8}KO. ANOVA, analysis of variance.
Nav1.7KO, the percentage of veratridine-irresponsive neurons decreased from 55% in littermate controls to 37% in Na\textsubscript{v1.8KO} (Fig. 3B). This increase in veratridine responsiveness came from the increase in the OS population from 27% in controls to 40%. The changes in the other populations were small and highly insignificant. In other words, the compensatory increase in Na\textsubscript{v1.7} expression caused previously silent neurons to respond to veratridine, and that these additional neurons had the OS profile, the profile of nociceptors. These results show that our assay is sensitive to increases in Na\textsubscript{v1.7} and thus can be used to detect pathological changes in Na\textsubscript{v1.7} function (eg, gain of function mutations or painful conditions).

Collectively, our results confirm that most nociceptors respond to veratridine with the OS profile, whereas most non-nociceptors respond to veratridine with the SD profile. Only the OS population is reduced when nociceptors are ablated (1.8-DTA) or lose a critical excitability determinant (Na\textsubscript{v1.7KO}). Importantly, the veratridine-response pattern of the Na\textsubscript{v1.7KO} can be used as the reference for potent and safe analgesic action, a reference pattern that potential analgesics can be evaluated against before-detailed in vitro characterisation or testing.

### 3.3. Evaluating the potency and selectivity of subtype-specific voltage-gated sodium channel blockers using the Na\textsubscript{v1.7KO} as reference

Several blockers of Na\textsubscript{v1.7} and Na\textsubscript{v1.8} channels have been developed and are currently in clinical trials. We used our assay to evaluate a candidate analgesic drug for each. PF-04856264 (PF-048) is from the arylsulfonamide class of selective Na\textsubscript{v1.7} blockers. PF-048 reduced responses to veratridine at concentrations similar to or lower than those used in standard cell line–based FLIIPR assays, and shows that our assay can identify nonpore blockers such as arylsulfonamides. PF-048 at 1 \( \mu M \) increased the percentage of veratridine-irresponsive neurons (Fig. 4A) from 46% to 58% through a decrease in the OS population, which decreased from 26% to 17% (ie, 0.65 of control). PF-048 at 5 \( \mu M \) (Fig. 4B) had a greater effect, increasing veratridine-irresponsive neurons from 44% to 72% through a larger decrease in the OS population from 28% to 12% (0.43 of control). However, 5 \( \mu M \) PF-048 decreased the SD population from 13% to 6%. Because a reduction in the SD population was not observed in Na\textsubscript{v1.7KO}, this suggests that PF-048 will affect non-nociceptors at a dose that does not even reduce the OS population to the Na\textsubscript{v1.7KO} level of 0.25 of control.

The Na\textsubscript{v1.8} blocker A-803467 (A-80) decreases the excitability of DRG neurons. At 0.1 \( \mu M \), it reduced the OS population from 25% to 12% (Fig. 4C) but without an overall increase in veratridine-irresponsive neurons (51% vs 55%). A-80 at 0.1 \( \mu M \) did not silence the OS population but rather prevented its multiplex oscillatory behaviour, converting them to the ID and RD profiles that respond to veratridine with a single peak (Fig. 1). A-80 at 0.3 \( \mu M \) increased veratridine-irresponsive neurons from 41% to 66% (Fig. 4D), through significant decreases in the OS population (from 28% to 12%) and the SD population (from 19% to 12%). These data suggest that A-80 is more potent in reducing the excitability of the OS population than PF-048, but it also affects the SD population at the higher concentration.

### 3.4. A combination of low concentrations of PF-04856264 and A-803467 have an additive effect on nociceptors

We suggest that our assay can be used not only to screen for novel drugs, but also to identify effective combinations of existing ones. As a proof of concept, we used a combination of 1 \( \mu M \) PF-048 and 0.1 \( \mu M \) A-80 (Fig. 5A). When applied separately, neither reduced the SD population significantly, but their reduction of the OS population is considerably less than Na\textsubscript{v1.7} deletion. The combination reduced the OS population from 30% to 11%, greater than either alone. Importantly, the combination still did not significantly reduce the SD population. A direct comparison of the decreases in the OS and SD populations between the VGSC blockers used and Na\textsubscript{v1.7} deletion (Fig. 5B) clearly shows that the combination produced the closest effect on the OS population (reduction to 0.38 of control) to that observed in Na\textsubscript{v1.7KO} (0.25 of control).

### 4. Discussion

Cell lines provide a high-throughput platform for analgesic drug discovery. However, their use leaves a knowledge gap that must be addressed on DRG neurons using lower throughput methods such as calcium imaging and patch clamping. The knowledge gap includes information on the effect of a drug on nociceptors as a whole (rather than just the molecular target of interest) and the concentrations needed for a potent effect on nociceptors with a minimal effect on non-nociceptors (as a proxy for unwanted side effects on CNS and motor neurons). We discovered that nociceptors and non-nociceptors respond to veratridine with distinct response profiles. Here, we show that a veratridine-based assay allows for a simultaneous assessment of drugs’ action on both populations and therefore provides an efficient and more informative assay for screening for analgesics on sensory neurons. First, we discuss the results from our findings and then discuss the key advantages of our assay.

We used the 1.8-DTA mouse that has well characterised pain deficits to demonstrate the link between the loss of the OS population and loss of pain. In 1.8-DTA neurons, we recorded a loss of 89% of functionally defined nociceptors (Fig. 2B), which is in agreement with the expression of Na\textsubscript{v1.8} in 80 to 90% of nociceptors. Our identification of nociceptors is based on responsiveness to one or more of 3 nociceptive agonists. This definition will encompass 6 of the putative 11 molecular subtypes (the NP1-3, PEP1-2, and TH types) of sensory neurons as defined in one single-cell RNA-seq study. By our functional definition, non-nociceptors will equate to the 5 neurofilament-positive subtypes (NF1-5). Ablation of nociceptors leads to the loss of 78% of the OS population, confirming that most nociceptors respond to veratridine with the OS profile. In the Na\textsubscript{v1.7KO} mouse, nociceptors are not ablated, but their excitability is greatly reduced to the extent that the pain loss in the Na\textsubscript{v1.7KO} is very similar to that of the 1.8-DTA. Not surprisingly, deletion of Na\textsubscript{v1.7} “silenced” 75% of the OS population (Fig. 3A), an almost identical percentage to that lost in 1.8-DTA.

Given that Na\textsubscript{v1.7} is expressed in all DRG neurons, why was there no significant reduction in the SD population in Na\textsubscript{v1.7KO}? Veratridine primarily activates TTX-sensitive VGSCs, and indeed, we previously showed that TTX silences all OS and SD neurons. However, although Na\textsubscript{v1.7} is the main TTX-sensitive channel in nociceptors, non-nociceptors rely on other TTX-sensitive subtypes, namely Na\textsubscript{v1.1}, Na\textsubscript{v1.2}, and Na\textsubscript{v1.6}. In the absence of Na\textsubscript{v1.7}, these subtypes allow veratridine to activate non-nociceptors. The predominance of other TTX-sensitive subtypes in non-nociceptors explains why deletion of Na\textsubscript{v1.7} in all DRG neurons has no effect on touch or proprioception. This is also why 4,9TTX, which preferentially blocks Na\textsubscript{v1.6} channels, affected the SD population more than the OS population (Fig. 3B).
Figure 4. Evaluating the effect of PF-04850264 and A-803467 on nociceptors and non-nociceptors. (A) 1 μM PF-048 increases the percentage of veratridine-irresponsive neurons through a reduction of the OS but not the other 3 populations. VTD – CTR = 45.9 ± 2 vs VTD – 1 PF = 68.1 ± 4; SDCTR = 16.9 ± 2 vs SD1PF = 13.6 ± 1.7; OSCTR = 26.2 ± 1.7 vs OS1PF = 16.8 ± 3; IDCTR = 7.5 ± 1.3 vs ID1PF = 7.0 ± 1.4; RDCTR = 4.1 ± 1.6 vs RD1PF = 3.6 ± 0.9%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 813 untreated and 681 treated neurons from 7 C57Bl6 mice. (B) 5 μM PF-048 increases the percentage of veratridine-irresponsive neurons through a reduction of both the OS and SD populations. 5 μM PF-048 is equally potent on the SD and OS profiles reducing both to about 50% of control values. VTD – CTR = 44.1 ± 2.7 vs VTD – 5 PF = 71.8 ± 1.5; SDCTR = 13.2 ± 1.6 vs SD1PF = 6.3 ± 1.0; OSCTR = 28.1 ± 2.4 vs OS1PF = 12.4 ± 1.6; IDCTR = 7.9 ± 1.6 vs ID5PF = 5.9 ± 1.1; RDCTR = 6.2 ± 1.8 vs RD5PF = 3.8 ± 0.7%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 605 untreated and 338 treated neurons from 5 C57Bl6 mice. (C) 100 nM A-80 does not change the percentage of veratridine-responsive neurons but reduces the OS population by about 50%. VTD – CTR = 51 ± 4.9 vs VTD – 100A80 = 54.8 ± 4.9; SDCTR = 12.9 ± 3.2 vs SD100A80 = 14.4 ± 2; OSCTR = 25.1 ± 2.6 vs OS100A80 = 11.61 ± 2.6; IDCTR = 5.2 ± 1.3 vs ID100A80 = 10.7 ± 2.5; RDCTR = 6.6 ± 0.8 vs RD100A80 = 7.3 ± 1.6%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 607 untreated and 594 treated neurons from 6 C57Bl6 mice. (D) 300 nM A-80 increases the percentage of veratridine-irresponsive neurons through a reduction of both the OS and SD populations. The reduction in the OS population is slightly greater than that in the SD profile. VTD – CTR = 40.8 ± 2.3 vs VTD – 300A80 = 66.2 ± 2.4; SDCTR = 19.1 ± 1.7 vs SD300A80 = 11.5 ± 2; OSCTR = 28.0 ± 1.3 vs OS300A80 = 12.2 ± 1.2; IDCTR = 8.5 ± 1.5 vs ID300A80 = 6.8 ± 1.5; RDCTR = 4.8 ± 0.9 vs RD300A80 = 3.3 ± 0.7%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 1197 untreated and 1294 treated neurons from 7 C57Bl6 mice. ANOVA, analysis of variance.

We propose to use veratridine responses of the Na1.7KO as the criterion VGSC blockers need to match for in vivo potency and safety. The pattern of veratridine responses we observed in vitro is due to the deletion of Na1.7 and are unlikely to be influenced by the reported increase in endogenous opioid agonism in the spinal cord. This is because veratridine responses are recorded from disassociated neurons after 1 day in culture (that lacks opioid-releasing spinal cord cells) and under constant Ringer flow. Our assay detected a significant increase in the number of neurons responding to veratridine in Na1.8KO, and these “unsilenced” neurons have the OS profile, demonstrating for the first time the effect of the compensatory increase in Na1.7 expression. All of the above strongly confirms the link between nociceptors’ excitability and the OS population and suggests that a reduction in the OS but not the SD population correlates with a reduction in pain without motor/CNS effects.

Our assay has several advantages over cell line-based assays and patch clamping. First, a cell line-based assay examines the effect of a condition or a drug on one target at a time, whereas our assay evaluates the overall effect on DRG neurons, including non-nociceptors. For example, while A-80 was potent in reducing excitability, we noted clear cytotoxic effects at concentrations >0.5 μM (not shown). The use of veratridine as a single “pan” activator of all sensory neurons has advantages over the commonly used potassium chloride (KCl). Dorsal root ganglion neurons respond to veratridine with distinct profiles that can be used to identify the main populations, whereas KCl produces a similar profile in all neurons. In addition, while 80% of Na1.7KO neurons did not respond to veratridine, they were all KCl-positive (Fig. 3A), strongly suggesting that KCl responsiveness is a poor predictor of the excitability of DRG in in vitro assays.

Second, our assay allows for changes in the population of high-threshold or “silent” sensory neurons that are observed in vivo, to be monitored in vitro. This is possible due to the way veratridine activates sensory neurons. Veratridine acts on open VGSC at the resting membrane potential to prevent their inactivation. Neurons with low expression of threshold VGSC channels (eg, Na1.7 in DRG) or and have a hyperpolarised membrane potential will have few open VGSC channels at resting potentials and will not respond to veratridine. We observed that between 30% and 50% of cultured DRG neurons do not respond to 3 minutes of 30 μM veratridine. Interestingly, these are similar to the percentages (38%-48%)
of high-threshold “silent” DRG neurons that did not respond to 10- to 100-mA current injection in vivo but were unmasked by inflammation. Because two-thirds of veratridine-irresponsible neurons responded to nociceptive markers (yellow segment in Fig. 2D), we speculate that the veratridine-irresponsible population includes the high-threshold silent nociceptors observed in vivo. However, we have no data to support this hypothesis at present.

Third, our assay allows the range of plasma concentrations for a potent action on nociceptors without an effect on non-nociceptors to be determined efficiently. Both PF-048 and A-80 reduced the OS population with A-80 being 10 times more potent than PF-048. However, at the higher concentration, both reduced the SD population, an effect not observed in either NaV1.7 or NaV1.8 KO; both techniques reduced the SD population, an effect not observed in vivo. Notice that the combined action of the lower doses of the PF and A80 produced the closest reduction of the SD population to NaV1.7 deletion. ANOVA, analysis of variance; VGSC, voltage-gated sodium channel.

Figure 5. The additive effects of a combination of PF-04856264 and A-803467 on nociceptors. (A) A combination of 1 μM PF-048 and 100 nM A-80 increases the percentage of veratridine-irresponsible neurons through a decrease of the OS population only. VTD-CTR = 40.8 ± 3.5 vs VTD-A80PF = 71.5 ± 5.7; SDCTR = 19.2 ± 2.5 vs SDA80PF = 12.2 ± 1.5; OSCTR = 30.4 ± 2.4 vs OSA80PF = 11.5 ± 2; IDCTR = 5.7 ± 0.8 vs IDA80PF = 2.4 ± 0.8; RDCTR = 3.7 ± 0.4 vs RDA80PF = 2.1 ± 1.1%. One-way ANOVA with Sidak’s test. Pie charts represent mean values in the histogram. Data from 980 untreated and 1114 treated neurons from 7 C57Bl/6 mice. (B) Comparison of the changes in the OS and SD populations caused by VGSC blockers to those of the NaV1.7KO. The higher doses of PF and A80 and 4,9TTX caused a significant reduction in the SD population. Notice that the combined action of the A80PF drug combination (Fig. 5) and making use of a nonwash protocol. The potential of this assay to screen for drugs for other classes of ion channels (calcium and potassium) needs to be established.

Future work will focus on adapting the assay to high-throughput platforms by automating data analysis (including calculation of response onset, amplitude, and area under the curve) and making use of a nonwash protocol. The potential of this assay to screen for drugs for other classes of ion channels (calcium and potassium) needs to be established.

For drug screening, the assay is most useful when the primary screening criterion is the decrease in the OS population where the screening window is large, a 75% drop in the 1.7KO and 62% in the A80PF drug combination (Fig. 5). Although a decrease in the total number of veratridine-responsive neurons can be used as the screening criterion, the screening window is narrower (A decrease of 38% in the 1.7KO). In all cases, the screening criterion should include a requirement for minimal effect on the SD population.

In summary, we described an assay to assess the excitability of nociceptors and non-nociceptors. The assay can be used in other applications (Fig. 6) in addition to drug development and screening. It can be used for in-depth, long-term characterisation of excitability changes in knockout mice and models of pain pathologies (diabetes, cancer, and aging). It can be used to optimise protocols to differentiate human stem cells into the different classes of sensory neurons. Meanwhile, all this may have the added advantage of reducing the number of animals needed for in vitro and in vivo testing.
Unbiased, medium-/high-throughput, and high content assessment of excitability of DRG neurons

Drugs screening
- Validation of hits from cell lines on DRG neurons

Stem cell models
- Validation of stem cell differentiation protocols

Rodent models
- Long-term characterisation of changes in DRG in diabetes, aging, and cancer

Figure 6. Applications of the veratridine-based calcium assay. The assay is a very efficient method to characterise changes in a heterogeneous population of neurons and therefore has several applications. The assay can be used to identify lead analgesic drugs either by validating hits from cell line–based screens on all types of DRG neurons or identification of hits by a direct screen on DRG neurons. The assay can be used to assess how stem cell–derived neurons compare to primary neurons of the same type more efficiently than by patch clamping. The assay can be used to compare neurons derived from patients’ IPSC with known or unknown genetic mutations. The assay is suited to characterise pathologies that develop over time as in diabetes, aging, or cancer. Finally, the assay can be used to efficiently characterise changes in DRG from the large number of transgenic strains generated by phenotyping consortia. DRG, dorsal root ganglia.

Conflict of interest statement
The authors have no conflicts of interest to declare.

Acknowledgements
The authors are grateful for John N Wood, UCL, for provision of the 1.8-DTA, Na\textsubscript{v}1.8KO, and Na\textsubscript{v}1.7KO transgenic mice. The authors are grateful for Yusef and Younes Nassar for proofreading the manuscript.

Author contributions: M.A. Nassar conceived, designed, and supervised experiments. Z.A. Mohammed and K. Kaloyanova codesigned, carried out, and analysed all experiments. M.A. Nassar contributed to preparations of DRG cultures. M.A. Nassar wrote the manuscript with contributions from Z.A. Mohammed and K. Kaloyanova.

Article history:
Received 4 October 2019
Received in revised form 20 December 2019
Accepted 7 January 2020
Available online 9 January 2020

References
[1] Abrahamsen B, Zhao J, Asante CO, Cendan CM, Marsh S, Martinez-Barbera JP, Nassar MA, Dickenson AH, Wood JN. The cell and molecular basis of mechanical, cold, and inflammatory pain. Science 2008;321: 702–5.
[2] Akopian AN, Sivilotti L, Wood JN. A tetrodotoxin-resistant voltage-gated sodium channel expressed by sensory neurons. Nature 1996;379: 257–62.
[3] Chen L, Huang J, Zhao P, Persson AK, Dib-Hajj FB, Cheng X, Tan A, Waxman SG, Dib-Hajj SD. Conditional knockout of Na\textsubscript{v}1.6 in adult mice ameliorates neuropathic pain. Sci Rep 2018;8:3845.
[4] Deuis JR, Wingerd JS, Winter Z, Durek T, Dekan Z, Sousa SR, Zimmermann K, Hoffmann T, Weidner C, Nassar MA, Alewood PF, Lewis RJ, Vetter I. Analgesic effects of GpTx-1, PF-04856264 and CNV1014802 in a mouse model of Na\textsubscript{v}1.7-mediated pain. Toxins (Basel) 2016;8:E78.
[5] Djouhri L, Fang X, Okuse K, Wood JN, Berry CM, Lawson SN. The TTX-resistant sodium channel Nav1.8 (SNS/PN3): expression and correlation with membrane properties in rat nociceptive primary afferent neurons. J Physiol 2003;550:739–52.
[6] Emery EC, Luiz AP, Wood JN. Nav1.7 and other voltage-gated sodium channels as drug targets for pain relief. Expert Opin Ther Targets 2016;20: 975–83.
[7] Habib AM, Wood JN, Cox JJ. Sodium channels and pain. Handb Exp Pharmacol 2015;227:39–56.
[8] Jarvis MF, Honore P, Shieh CC, Chapman M, Joshi S, Zhang XF, Kort M, Carroll W, Marron B, Atkinson R, Thomas J, Liu D, Krambis M, Liu Y, McGarvaughty S, Chu K, Roesloffs R, Zhong C, Mikusa JP, Hernandez G, Gauvin D, Waide C, Zhu C, Pai M, Scaino M, Shi L, Drizin I, Gregg R, Matulenko M, Hakeem A, Gross M, Johnson M, Marsh K, Wagoner PK, Sullivan JP, Faltynek CR, Krafte DS. A-803467, a potent and selective Nav1.8 sodium channel blocker, attenuates neuropathic and inflammatory pain in the rat. Proc Natl Acad Sci U S A 2007;104:9520–5.
[9] Kingwell K. Nav1.7 withholds its pain potential. Nat Rev Drug Discov 2019;18:321–23.
[10] Leadley RM, Armstrong N, Reid KJ, Allen A, Misso KV, Kleijnen J. Healthy aging in relation to chronic pain and quality of life in Europe. Pain Pract 2014;14:547–58.

[11] McCormack K, Santos S, Chapman ML, Kratke DS, Mannion BE, West CW, Krambis MJ, Antonio BM, Zelimer SG, Printzenhoff D, Padilla KM, Lin Z, Wagoner PK, Swan NA, Stupple PA, de GM, Butt RP, Castle NA. Voltage sensor interaction site for selective small molecule inhibitors of voltage-gated sodium channels. Proc Natl Acad Sci U S A 2013;110: E2724–32.

[12] Meisler MH, Plummer NW, Burgess DL, Buchner DA, Sprunger LR. Allelic mutations of the sodium channel SCN8A reveal multiple cellular and physiological functions. Genetics 2004;122:37–45.

[13] Michaelis M, Habler HJ, Jaenig W. Silent afferents: a separate class of primary afferents? Clin Exp Pharmacol Physiol 1996;23:99–105.

[14] Minett MS, Falk S, Santana-Varela S, Bogdanov YD, Nassar MA, Heegaard AM, Wood JN. Pain without nociceptors? Nav1.7-independent pain mechanisms. Cell Rep 2014;6:301–12.

[15] Minett MS, Pereira V, Sikandar S, Matsuyama A, Loliqner S, Kanellopoulos AH, Mancini F, Iannetti GD, Bogdanov YD, Santana-Varela S, Millet Q, Baskozos G, MacAllister R, Cox JJ, Zhao J, Wood JN. Endogenous opioids contribute to insensitivity to pain in humans and mice lacking sodium channel Nav1.7. Nat Commun 2015;6:8967.

[16] Mohammed ZA, Doran C, Grundy D, Nassar MA. Veratridine produces distinct calcium response profiles in mouse Dorsal Root Ganglia neurons. Sci Rep 2017;7:45221.

[17] Nassar MA, Stirling LC, Forlani G, Baker MD, Matthews EA, Dickenson AH, Wood JN. Nociceptor-specific gene deletion reveals a major role for Nav1.7 (PN1) in acute and inflammatory pain. Proc Natl Acad Sci U S A 2004;101:12706–11.

[18] O’Brien JE, Meisler MH. Sodium channel SCN8A (Nav1.6): properties and de novo mutations in epileptic encephalopathy and intellectual disability. Front Genet 2013;4:213.

[19] O’Connor AB. Neuropathic pain: quality-of-life impact, costs and cost effectiveness of therapy. Pharmacoeconomics 2009;27:95–112.

[20] Rosier C, Lohberger B, Hofer D, Steinecker B, Quasthoff S, Schreibmayer W. The TTX metabolite 4,9-anhydro-TTX is a highly specific blocker of the Nav1.6 voltage-dependent sodium channel. Am J Physiol Cell Physiol 2007;293:C783–9.

[21] Shields SD, Ahn HS, Yang Y, Han C, Seal RP, Wood JN, Waxman SG, Dib-Hajj SD. Nav1.8 expression is not restricted to nociceptors in mouse peripheral nervous system. PAIN 2012;153:2017–30.

[22] Tay B, Stewart TA, Davis FM, Deuis JR, Vetter I. Development of a high-throughput fluorescent no-wash sodium influx assay. PLoS One 2019; 14:e0213751.

[23] Usoskin D, Furlan A, Islam S, Abdo H, Lonnerberg P, Lou D, Hjerrild-Leffler J, Haeggstrom J, Knarchenko O, Kharchenko PV, Linnarsson S, Emfors P. Unbiased classification of sensory neuron types by large-scale single-cell RNA sequencing. Nat Neurosci 2015;18:145–53.

[24] Viventi S, Dottori M. Modelling the dorsal root ganglia using human pluripotent stem cells: a platform to study peripheral neuropathies. Int J Biochem Cell Biol 2018;100:61–8.

[25] Yekkirala AS, Roberson DP, Bean BP, Woolf CJ. Breaking barriers to novel analgesic drug development. Nat Rev Drug Discov 2017;16:545–54.