Probing the peripheral immune response in mouse models of oxaliplatin-induced peripheral neuropathy highlights their limited translatability [version 2; peer review: 1 approved, 1 approved with reservations]

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Abstract

Background: Chemotherapy-induced peripheral neuropathy (CIPN) is a disabling side effect of various chemotherapeutic agents, including oxaliplatin. It is highly prevalent amongst cancer patients, causing sensory abnormalities and pain. Unfortunately, as the underlying mechanisms remain poorly understood, effective therapeutics are lacking. Neuro-immune interactions have been highlighted as potential contributors to the development and maintenance of CIPN, however, whether this is the case in oxaliplatin-induced peripheral neuropathy (OIPN) is yet to be fully established.

Methods: In this study we used flow cytometry to examine the peripheral immune response of male C57BL/6 mice following both single and repeated oxaliplatin administration. In animals exposed to repeated dosing, we also undertook mechanical and thermal behavioural assays to investigate how oxaliplatin alters phenotype, and conducted RT-qPCR experiments on bone marrow derived macrophages in order to further inspect the effects of oxaliplatin on immune cells.

Results: In contrast to other reports, we failed to observe substantial changes in overall leukocyte, lymphocyte or myeloid cell numbers in dorsal root ganglia, sciatic nerves or inguinal lymph nodes. We did however note subtle, tissue-dependant alterations in several myeloid subpopulations following repeated dosing. These included a significant reduction in MHCII antigen presenting cells in the sciatic nerve and an increase in infiltrating cell types into the inguinal lymph nodes. Though repeated oxaliplatin administration had a systemic effect, we were unable to detect a pain-like behavioural phenotype in response to either cold or mechanical stimuli. Consequently, we cannot comment on whether the observed myeloid changes are associated with OIPN.

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Any reports and responses or comments on the article can be found at the end of the article.
Conclusions: Our discussion puts these results into the wider context of the field, advocating for greater transparency in reporting, alignment in experimental design and the introduction of more clinically relevant models. Only through joint concerted effort can we hope to increase our understanding of the underlying mechanisms of CIPN, including any immune contributions.

Keywords
Pain, neuropathy, chemotherapy, oxaliplatin, immune, flow cytometry, behaviour, reproducibility

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Introduction

Chemotherapy induced peripheral neuropathy (CIPN) is an unpleasant and debilitating side-effect of numerous neurotoxic chemotherapeutics, including epothilones, proteasome inhibitors, taxanes, vinca alkaloids and platinum-based agents (Starobova & Vetter, 2017). Characteristically, CIPN presents itself in a ‘glove and stocking’ fashion, initially affecting the extremities before progressing proximally (Wolf et al., 2008). It typically induces a range of sensory abnormalities including paresthesias and dysesthesias, which can be exacerbated by warm or cool temperatures, impaired vibration and even sensory loss (Staff et al., 2017; Zajączkowska et al., 2019). Moreover, various painful sensations have also been attributed, including burning, shooting or electric-shock-like pain (Boland et al., 2010) and increased sensitivity to both mechanical and thermal stimuli (Flatters et al., 2017; Zajączkowska et al., 2019). Importantly, risk of developing CIPN is known to increase with total cumulative dose; thus it is often a dose limiting factor (Burton et al., 2007), ultimately affecting survival.

Prevalence of CIPN is markedly high, acutely affecting 60–70% of patients, with 30% continuing to suffer symptoms 6 months following cessation of chemotherapy (Seretny et al., 2014). One such agent with a particularly high incidence of inducing long-lasting peripheral neuropathy is oxaliplatin, a platinum-based chemotherapeutic commonly used in the treatment of cancers of the digestive tract, including colorectal, oesophageal, stomach, liver and pancreatic (Zajączkowska et al., 2019). Staggeringly, oxaliplatin-induced peripheral neuropathy (OIPN) has been reported to affect almost 80% of patients two years following the end of treatment (Park et al., 2011). In fact, increased incidence of numbness or tingling of the hands and feet has been noted to persist in oxaliplatin-treated patients for 6 years following cessation of chemotherapy (Kidwell et al., 2012), highlighting it as one of the more disabling agents.

Similarly, sensory deficits, primarily in the form of evoked pain-like behaviours, have been observed in various rodent models of CIPN, using a myriad of oxaliplatin-based designs (Currie et al., 2019; Gadgil et al., 2019; Höke & Ray, 2014). Both acute and more chronic OIPN studies report significantly increased sensitivity to both cold (Descoeur et al., 2011; Gauchan et al., 2009b; Joseph & Levine, 2009; Joseph et al., 2008; Ling et al., 2007b; Nassini et al., 2011; Renn et al., 2011) and mechanical (Gauchan et al., 2009a; Nassini et al., 2011; Renn et al., 2011) stimuli within the first week of exposure. Furthermore, this altered nociception often appears to outlast the treatment itself – a phenomenon comparable to clinically observed ‘coasting’, whereby neuropathy worsens or newly develops following the end of treatment (Staff et al., 2017). For example, in rodents, it has been repeatedly shown that both single (Gauchan et al., 2009a; Gauchan et al., 2009b; Ling et al., 2007b; Nassini et al., 2011) and repeated (Ling et al., 2007a; Xiao et al., 2012) administration of oxaliplatin produce behavioural deficits which persist for at least one week following the final dose. Importantly, and in line with clinical observations, similar phenotypes are also displayed in models utilising other, non-platinum based, chemotherapeutics such as paclitaxel and vincristine (Authier et al., 2003; Flatters & Bennett, 2004; Gauchan et al., 2009a; Kiguchi et al., 2008b; Makker et al., 2017; Muthuraman et al., 2008; Old et al., 2014; Shen et al., 2015; Xiao et al., 2007), indicating the presence of common underlying mechanisms.

Despite CIPN being widely reported in both clinical and pre-clinical settings, exactly what these underlying mechanisms are is not fully understood, though development is thought to be multi-factorial (Starobova & Vetter, 2017). Potential contributors, including dysregulation of calcium homeostasis, axon degeneration, mitochondrial dysfunction, oxidative stress and alterations to ion channels and the immune response have all been proposed (Flatters et al., 2017; Starobova & Vetter, 2017). In fact, this latter notion, which suggests a role for neuro-immune interactions, has garnered considerable attention in recent years (Lees et al., 2017), with studies demonstrating altered immune responses to various chemotherapy drugs. Centrally, multiple groups have reported an increase in spinal microglia (Burgos et al., 2012; Kiguchi et al., 2008a; Peters et al., 2007; Ruiz-Medina et al., 2013; Shen et al., 2015), though there has been some debate regarding whether microglia or astrocytes contribute to the pathogenesis of CIPN (Robinson et al., 2014; Zhang et al., 2012). Peripherally, there is strong evidence supporting the involvement of immune cells. For example, repeated administration of vincristine is reported to induce macrophage infiltration into the sciatic nerves and DRG (Kiguchi et al., 2008b; Old et al., 2014). Similarly, following two 18mg/kg i.v. injections of paclitaxel, increased expression of the macrophage marker CD68 has been observed in these peripheral tissues (Peters et al., 2007). In fact, even a single 6mg/kg dose of paclitaxel has been shown to result in increased incidence of leukocytes in the DRG, including macrophage, monocyte, neutrophil and T cell populations (Liu et al., 2014). Furthermore, in the case of both chemotherapy agents, disruption of the inflammatory response is associated with improvement of behavioural deficits, highlighting the importance of such immune cells in the development of painful phenotypes (Liu et al., 2014; Old et al., 2014).

Although oxaliplatin is one of the most utilised drugs in CIPN models (Currie et al., 2019), its effects on the immune response have been relatively poorly investigated. Of the
limited number of studies conducted, many have focused only on tissues of the central nervous system (Cho et al., 2016; Di Cesare Mannelli et al., 2014; Janes et al., 2015), while the handful of peripheral approaches in the literature have largely not employed flow cytometry techniques, relying on immunohistochemical analysis (Di Cesare Mannelli et al., 2013; Li et al., 2016) and whole tissue qPCR (Marmiròli et al., 2017). In potentially the only peripherally focused, flow cytometry-heavy investigation of OIPN, oxaliplatin-induced increases were noted specifically in circulating T cell populations, though such differences were not detected in either the lumbar DRG or sciatic nerves (Makker et al., 2017). However, their flow cytometric investigations did not extend to myeloid cell types in the periphery, such as macrophages, which have been implicated in the development and even maintenance of neuropathic pain in a variety of models (Kiguchi et al., 2019; Ristoiu, 2013), including nerve injury (Liang et al., 2020; Yu et al., 2020).

We are therefore lacking more detailed accounts regarding the specific effects of oxaliplatin on the peripheral immune response, and in particular, on innate cell types. Furthermore, the fact reports to date have been somewhat conflicting, makes interpretation of the current OIPN literature ambiguous. For example, lumbar DRG expression of the macrophage/microglial marker ionized calcium binding adaptor molecule 1 (IBA1) has been shown to both increase (Li et al., 2016) and remain unaltered (Makker et al., 2017) following repeated oxaliplatin administration. Meanwhile, another study completely failed to detect macrophages in either the L4–5 DRG or sciatic nerves following 3 weeks of oxaliplatin treatment (Di Cesare Mannelli et al., 2013).

The aim of this study is therefore to utilise immunological approaches in order to clarify whether there is peripheral dysregulation of the innate and/or adaptive immune systems in response to both single and repeated oxaliplatin administration, and if so, speculate how this may contribute to the generation and maintenance of painful peripheral neuropathies.

**Methods**

**Ethical considerations**

All experiments described were carried out in accordance with the United Kingdom Home Office Legislation (Animals (Scientific Procedures) Act, 1986, 2021) and were approved by the Home Office to be carried out at King’s College London under project license number P57A189DF. This study is reported in line with the Animal Research: Reporting of In Vivo Experiments (ARRIVE) guidelines (Percie du Sert et al., 2020) and a completed Full ARRIVE 2.0 Checklist has been provided (Hore et al., 2021a).

All efforts were made to ameliorate harm to animals used in this study. This was achieved in a number of ways, including:

- Provision of extra food and diet supplements to mice who were undergoing repeated dosing and were at risk of losing weight.
- Regularly weighing mice and keeping a close eye on their health to ensure that any animals displaying signs of illness could be assisted rapidly and not left to suffer.
- Ensuring new needles were used for each animal during dosing and rotating injection side in animals who underwent repeated dosing.
- Regular handling of animals prior to and during behavioural testing to familiarise them with being handled and minimise any distress.

**Animals**

Male C57BL/6JOlA-Hsd mice were purchased from Envigo at 8–13 weeks of age and acclimatised to the animal unit for one week prior to any procedures. Mice that were purchased at 13 weeks were singly housed within one week of arrival due to fighting between cage mates. For all experiments, mice were housed in standard individually ventilated cages (Tecniplast) in groups of five maximum at a 12h light–dark cycle, with *ad lib* access to food and water.

**Chemotherapy induced peripheral neuropathy (CIPN) models**

We conducted two separate studies, designed to measure the effect of oxaliplatin versus vehicle treatment. In all instances, mice were primarily housed in groups of 5 and within each cage 2–3 mice were appointed to either the oxaliplatin or vehicle treatment group, to allow for cage-mate controls. We did not employ any computer-based randomisation method to assign mice to a particular group. Rather, animals were arbitrarily chosen by the experimenter delivering the treatment. In the case of the 13-week old mice which were subjected to repeated administration, animals were assigned to treatment group prior to being singly housed due to fighting. Mice were subjected to one of the following dosing regimens and in both instances, were returned to their home cage immediately after injection.

**Single administration (n=6 mice):** The oxaliplatin solution was made up on the day of injection by diluting 1.2mg of oxaliplatin powder (Sigma, #O9512) in 2ml of 5% glucose to give a final concentration of 0.6mg/ml. A single 6mg/kg dose of oxaliplatin, or an equal volume of 5% glucose, was administered intraperitoneally (i.p.) and mice were weighed prior to, and for 2 days following injection.

**Repeated administration (n=10 mice):** The oxaliplatin solution was made up at the beginning of each dosing week by diluting the stock oxaliplatin (5mg/ml) (Guy’s Cancer Centre, Accord) 1:10 with 5% glucose. Mice were administered a single 3mg/kg dose of oxaliplatin or an equal volume of 5% glucose, daily for 5 days – this constituted 1 cycle of injections. This was repeated for 4 cycles, with mice given a dosing break of 1 week after the first 2 cycles. Note that on the final week of dosing mice were only injected 4 times (Monday-Thursday), as 2 mice...
Mice were acclimatised to the testing arena – a 20cm diameter incremental hot/cold plate surrounded by a transparent acrylic cylinder (Ugo Basile) – for five minutes on one day prior to testing. On the 7 testing days, animals were acclimatised to a switched-off plate for 2 minutes before being transferred to an identical cold plate set at 10°C. Mice were observed for a response (jumping, hind paw shaking or hind paw licking) and their latency to respond was recorded. If a mouse made a jump response it was immediately removed from the arena and returned to its home cage. A maximum latency of 90 seconds was set to prevent damage to the plantar skin. To ensure that the correct latencies were noted, each time point was recorded, and the videos were re-scored. If the live and re-scored latencies differed, the re-scored time was taken. Any mice that failed to respond were awarded the maximum latency of 90 seconds.

**Tissue processing**

In accordance with the approved methods of euthanasia set out in the licence we were working under, either 4 days (single dose model) or 38–39 days (repeated dosing model) following their first injection, mice were deeply anaesthetised via overdosage of pentobarbital (Euthatal; Merial, Lot# P02601A) administered i.p.. Once unresponsive, animals were perfused with 10ml of 1x PBS to avoid blood contamination. Following sacrifice, a laminectomy was performed in order to expose the lumbar spinal cord. To ensure the correct DRG were taken, the sciatic nerves were exposed and followed up towards the spinal cord to identify and dissect out L3-L5 DRG into F12 (Gibco, # 21765-029). The sciatic nerves themselves were then dissected out into a petri dish containing F12 and cut to 0.5cm. Lastly, the right and left inguinal lymph nodes were exposed and dissected out into F12, trying to separate them as much as possible from the surrounding fat. Tissues were kept in F12 on ice until all animals in a given batch were processed, they were then dissociated largely in accordance with previously described methods (Liang et al., 2020). Briefly, tissues were transferred into 50µl of digestion mix and incubated at 37°C, shaking at 220RPM for 45 minutes (see Table 1 and Table 2 for digestion mixes used for each tissue type). To achieve optimal digestion, nerves and inguinal lymph nodes were chopped into small pieces with spring scissors (50 and 30 chops, respectively) prior to incubation. Following digestion, samples were centrifuged, supernatants removed, and the remaining pellets were resuspended in 100µl of FACS buffer (see Table 3 for composition). Samples underwent dissociation via repeated up-down pipetting using a P200 (30x for DRG and inguinal lymph nodes; 50x for sciatic nerves) and

<table>
<thead>
<tr>
<th>Reagent (supplier, cat #)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>F12 (Gibco, 21765-029)</td>
<td>-</td>
</tr>
<tr>
<td>Dispase II (Sigma Aldrich, 04942078001)</td>
<td>3mg/ml</td>
</tr>
<tr>
<td>Collagenase type IA (Sigma Aldrich, C9891)</td>
<td>12.5mg/ml</td>
</tr>
<tr>
<td>DNase I (Sigma Aldrich, 10104159001)</td>
<td>10mg/ml</td>
</tr>
</tbody>
</table>

**Table 1. Digestion mix used in processing of L3-5 dorsal root ganglia (DRG) samples.**
were then filtered through the 35µm cap of a BD Falcon 12 × 75 mm tube with cell strainer cap (BD Biosciences, # 352235) into a 96 well v-bottom plate (Thermo Scientific, # 612V96). Finally, the plate was centrifuged, supernatants discarded and the remaining pellets underwent antibody staining as described in the following.

**Flow cytometry**

Staining of samples for flow cytometry was conducted as described previously (Liang *et al.*, 2020). Briefly, to distinguish live cells, samples were incubated in a fixable yellow viability dye (Invitrogen, # L34959) for 30 minutes, followed by 30 minutes incubation in a mix of directly conjugated antibodies and Fc block (see Table 4 for antibody panel and concentrations). Following centrifugation, the staining mix was removed and remaining pellets were incubated for 5 minutes in 4% paraformaldehyde (PFA) for fixation. Once fixed, samples were centrifuged, PFA removed and pellets resuspended in 200µl of FACS buffer. Flow cytometry was conducted on a BD Fortessa at the NIHR BRC flow core facility at King’s College London, with compensation controls employed as described previously. All analysis was carried out using FlowJo version 10.6.0 software (see Extended Figure 1 for gating strategies employed (Hore *et al.*, 2021a)).

**Harvesting and culture of bone marrow derived macrophages (BMDMs)**

Harvesting and culture of BMDMs was conducted as described previously (Liang *et al.*, 2020). Briefly, femur and tibia bones from both hind limbs were collected and the bone marrow was flushed out with cold PBS. Following centrifugation and filtration, the cell suspension was plated onto 15cm

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**Table 2. Digestion mix used in processing of sciatic nerve and inguinal lymph node samples.**

<table>
<thead>
<tr>
<th>Reagent (supplier, cat #)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>F12 (Gibco, 21765-029)</td>
<td>-</td>
</tr>
<tr>
<td>Collagenase type IA (Sigma Aldrich, C9891)</td>
<td>6.25mg/ml</td>
</tr>
<tr>
<td>Pronase (Millipore, 53702)</td>
<td>0.2%</td>
</tr>
<tr>
<td>Hyaluronidase (ABNOVA, P52330)</td>
<td>0.4%</td>
</tr>
</tbody>
</table>

**Table 3. Fluorescence-activated cell sorting (FACS) buffer composition.**

<table>
<thead>
<tr>
<th>Reagent (supplier, cat #)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBSS (Gibco, 14175095S)</td>
<td>-</td>
</tr>
<tr>
<td>BSA (Sigma-Aldrich, A3983)</td>
<td>0.4%</td>
</tr>
<tr>
<td>HEPES (Gibco, 15630080)</td>
<td>15mM</td>
</tr>
<tr>
<td>EDTA (Invitrogen, 15575038)</td>
<td>2mM</td>
</tr>
</tbody>
</table>

**Table 4. Previously optimised (Liang *et al.*, 2020) antibody panel used for all flow cytometry experiments.**

<table>
<thead>
<tr>
<th>Laser</th>
<th>Colour</th>
<th>Epitope</th>
<th>Cell Type</th>
<th>Final Dilution</th>
<th>Mono/ polyclonal</th>
<th>Species</th>
<th>Cat #</th>
</tr>
</thead>
<tbody>
<tr>
<td>UV</td>
<td>BUV395</td>
<td>Ly6G</td>
<td>Neutrophils</td>
<td>1:300</td>
<td>Monoclonal</td>
<td>Rat anti-mouse</td>
<td>BD Bioscience, #563978</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Violet</td>
<td>AmCyan</td>
<td>Live/ Dead</td>
<td>-</td>
<td>1:1000</td>
<td>N/A</td>
<td>N/A</td>
<td>Invitrogen, #L34959</td>
</tr>
<tr>
<td></td>
<td>BV650</td>
<td>Ly6C</td>
<td>Monocytes</td>
<td>1:1500</td>
<td>Monoclonal</td>
<td>Rat anti-mouse</td>
<td>BioLegend, #128049</td>
</tr>
<tr>
<td>Blue</td>
<td>FITC</td>
<td>CD45</td>
<td>Leukocytes</td>
<td>1:1200</td>
<td>Monoclonal</td>
<td>Rat anti-mouse</td>
<td>BioLegend, #103108</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Myeloid lineage</td>
<td>1:1200</td>
<td>Monoclonal</td>
<td>Rat anti-mouse, human</td>
<td>BioLegend, #101215</td>
</tr>
<tr>
<td>PE</td>
<td>β -TCR</td>
<td>T cells (αβ chain)</td>
<td>1:300</td>
<td>Monoclonal</td>
<td>Armenian hamster anti-mouse</td>
<td>BioLegend, #109207</td>
<td></td>
</tr>
<tr>
<td>Red</td>
<td>APC-Cy7</td>
<td>MHCII</td>
<td>Activated macrophages &amp; dendritic cells</td>
<td>1:1200</td>
<td>Monoclonal</td>
<td>Rat anti-mouse</td>
<td>BioLegend, #107628</td>
</tr>
<tr>
<td></td>
<td>APC</td>
<td>δ-TCR</td>
<td>T cells (γδ chain)</td>
<td>1:300</td>
<td>Monoclonal</td>
<td>Armenian hamster anti-mouse</td>
<td>BioLegend, #118116</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Fc block</td>
<td>1:20</td>
<td>Monoclonal</td>
<td>Rat anti-mouse</td>
<td>BioLegend, #101302</td>
</tr>
</tbody>
</table>
Petri dishes and incubated for 5–7 days to allow for differentiation into mature naïve macrophages. After 5–7 days of incubation, cells were gently dislodged with a cell scraper (Greiner, #541–070) and incubated for a further 24hrs in DMEM (Gibco, #32430–027) + MCSF (PeproTech, #315–02). The following day, cells were incubated for 4hrs in 10ng/ml of TNFα (BioLegend, #575202), while unstimulated cells were incubated in plain DMEM as a control. For RNA extraction, we used an RNeasy Microkit 50 (Qiagen, #74004) following manufacturer’s instructions. mRNA quantity was evaluated using a Qubit 3 Fluorometer (Invitrogen).

Reverse transcription quantitative real-time polymerase chain reaction (RT-qPCR) of BMDMs
1ng of the extracted RNA was used to synthesize cDNA using the Smart-Seq2 protocol, 16 cycles of amplification (Picelli et al., 2014). Where appropriate, the resultant cDNA was diluted down to 1ng/µl with double distilled H₂O (ddH₂O). 1ng of cDNA was used in standard SYBR Green RT-qPCR reactions, whereby 1µl of cDNA was added to a mix comprised of 5µl LightCycler® 480 SYBR Green I Master (Roche, #42352720), 1µl of relevant primer mix (10µM) and 3µl of ddH₂O. Samples were run on a LightCycler® 480 Instrument II (Roche, #05015243001) to probe for genes of interest (see Table 5 for primer sequences). All primers were tested for their efficiency and specificity prior to use. The housekeeping gene GAPDH was used to calculate ΔΔCt values. All reactions were run in duplicate with water used as a negative control.

Outcome measures
In this study, the following outcome measures were assessed, and comparisons were made between oxaliplatin and vehicle treated mice.

Behavioural assays: Cold plate = latency to jump or latency of hind paw response, namely shaking or licking (seconds), von Frey = 50% hind paw withdrawal threshold (grams).

Flow cytometry experiments: Total number of live single cells per population of interest.

BMDM experiments: Total macrophage cell number, based on haemocytometer counts under a light microscope. Expression of genes associated with DNA damage, apoptosis and cellular stress.

Table 5. List of primers used to probe for genes associated with DNA damage, apoptosis and cellular stress.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Function</th>
<th>Primer sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>GAPDH forward</td>
<td>Housekeeping</td>
<td>GGCCTTCGGTGTTCCCTAC</td>
</tr>
<tr>
<td>GAPDH Reverse</td>
<td></td>
<td>TGTCACTCATCTTTGGAGGGTT</td>
</tr>
<tr>
<td>TRP53 forward</td>
<td>Cell cycle regulator, apoptosis inducible</td>
<td>GTCAACGACATGACGGAGG</td>
</tr>
<tr>
<td>TRP53 reverse</td>
<td></td>
<td>TCTTCCAGATCTGGGATAC</td>
</tr>
<tr>
<td>GADD45A forward</td>
<td>DNA damage/stress</td>
<td>CGGAAGGAGTGGACGATG</td>
</tr>
<tr>
<td>GADD45A Reverse</td>
<td></td>
<td>TTATGGGGGTCTACG</td>
</tr>
<tr>
<td>PUMA forward</td>
<td>Pro-apoptotic gene</td>
<td>GCCGGGAGCACAGAAGA</td>
</tr>
<tr>
<td>PUMA reverse</td>
<td></td>
<td>AGTCCCCATGAAGATATTGCAGATGAC</td>
</tr>
<tr>
<td>FOS forward</td>
<td>Cell stress transcription factor</td>
<td>CGGTGTACACGCGGACTA</td>
</tr>
<tr>
<td>FOS reverse</td>
<td></td>
<td>TTGGCATGACAGGACAGA</td>
</tr>
<tr>
<td>SESN2 forward</td>
<td>Cell growth and survival regulator</td>
<td>TCCGGATGCCATTCGAGAT</td>
</tr>
<tr>
<td>SESN2 reverse</td>
<td></td>
<td>TCCGGGTGTAAGACCATCAC</td>
</tr>
<tr>
<td>DRAM1 forward</td>
<td>DNA damage/autophagy</td>
<td>TCATCTCTACGTGGTGC</td>
</tr>
<tr>
<td>DRAM1 reverse</td>
<td></td>
<td>CTGCACCAGAAATGCAGAG</td>
</tr>
<tr>
<td>MDM2 forward</td>
<td>P53 regulator</td>
<td>TGGATGTCAGCTACG</td>
</tr>
<tr>
<td>MDM2 reverse</td>
<td></td>
<td>TCCAGAGGCTTTAACACCTCAG</td>
</tr>
<tr>
<td>PTEN forward</td>
<td>Cell growth regulator</td>
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</tr>
<tr>
<td>PTEN reverse</td>
<td></td>
<td>CGGTGTCAATAGTCTCAG</td>
</tr>
<tr>
<td>F4/80 forward</td>
<td>Myeloid cells/macrophages</td>
<td>TGACCTCACCAGTGCTCTA</td>
</tr>
<tr>
<td>F4/80 reverse</td>
<td></td>
<td>TTCCCCAGAATCCAGTCTTTCC</td>
</tr>
</tbody>
</table>
cellular stress, based on ΔΔCt values calculated from data generated in RT-qPCR experiments.

*Systemic effect of oxaliplatin*: Body weight (grams)

**Statistical analysis**

For analysis of the flow cytometry data, comparisons between the oxaliplatin and vehicle groups were carried out using either an unpaired t-test or Mann-Whitney test, depending on normality (Shapiro-Wilk test). Data from two animals were excluded, or partially excluded from analysis - in accordance with the ARRIVE 2.0 reporting guidelines (Percie du Sert et al., 2020), details of these exclusions can be found in Extended Table 1 (Hore et al., 2021a). For behavioural tests and weights, repeated measures two-way ANOVAs were conducted, followed by Sidak’s multiple comparisons tests to assess differences between the two groups at a given timepoint and between timepoints within each group. For BMDM experiments, paired and multiple t-tests were used. In all cases significance was set at \( p \leq 0.05 \). All statistics were performed using GraphPad Prism version 9.0.0 software.

Power functions were calculated for non-parametric and parametric two-tailed t-tests, and for repeated measures ANOVAs (between factors) for a range of different effect and sample sizes, to help assess the likely sensitivity of our flow cytometry experiments and behavioural tests (Extended Figure 4 and Figure 5 (Hore et al., 2021a)). For this we used G*Power version 3.1.9.7 software with the following parameters – t tests (Means: Difference between two independent means), y axis: power (1-\( \beta \) err prob), as a function of: Effect size \( d \) (from 0 through to 2), \( \alpha \) err prob: 0.05, for a range of sample sizes from \( n=6 \) to \( n=16 \) or \( n=6 \) to \( n=18 \). F tests (ANOVA: Repeated measures, between factors), y axis: power (1-\( \beta \) err prob), as a function of: Effect size \( f \) (from 0 through to 1), \( \alpha \) err prob: 0.05, for a range of sample sizes from \( n=6 \) to \( n=16 \).

Effect sizes mentioned in the discussion and Extended Table 2 (Hore et al., 2021a) were obtained by calculating Cohen’s \( d \) using the equation \((\text{Mean}_1 - \text{Mean}_2)/\sqrt{(SD_1^2 + SD_2^2)/2}\).

**Open-access software alternatives**

**GraphPad Prism**: R is a language and environment for statistical computing which can be used to conduct all statistical analysis carried out in this study and create accompanying graphs.

**FlowJo**: All flow cytometry analysis can be conducted using Flowing Software 2.5.1, which is available to download for free.

**Results**

A single dose of 6mg/kg oxaliplatin resulted in short-term weight loss which resolved within 2-days following injection

In order to gain an indication of whether oxaliplatin was having a negative effect on the health of these mice - suggesting effective administration - the body weights of all animals were monitored prior to injection i.e. day 0, and for 2 days following injection i.e. day 1 and 2 (Figure 1 (Hore et al., 2021a)). The oxaliplatin group lost a significant amount of weight within 1 day of injection (day 0 - day 1: \( p=0.0170 \)), however, mice had, on average, returned to their pre-injection weights by the second day following dosing. On the other hand, their vehicle treated counterparts steadily gained weight following injection (day 0 – day 2: \( p=0.0129 \)). These data provide some indirect evidence for oxaliplatin having been successfully administered. However, they also suggest that a single dose of oxaliplatin negatively affects the health of a mouse for only a short period of time after administration. No behavioural assays were conducted for this acute administration model, and flow cytometry was performed 4-days following injection i.e. once animals had returned to their pre-injection weights.

Acute oxaliplatin administration did not alter total leukocyte number in lumbar DRG, sciatic nerves or inguinal lymph nodes

Based on reports from the literature that even acute oxaliplatin-based CIPN models induce prolonged behavioural deficits (Descoeur et al., 2011; Gauchan et al., 2009a; Gauchan et al., 2009b; Joseph et al., 2008; Ling et al., 2007b; Nassini et al., 2011), we hypothesised that if neuro-immune interactions are implicated in this phenomenon, any associated changes in peripheral immune profile should be observed in such a model. All samples were processed in a single batch, 4 days following a single i.p. injection of oxaliplatin (6mg/kg) or an equal volume of vehicle, and run together on a flow cytometer, using a

![Days post-injection](image-url)
Repeated oxaliplatin administration resulted in failure to gain weight

As we largely failed to detect alterations in the peripheral immune response at any level following acute administration of oxaliplatin, we proceeded to investigate if we could detect changes using a paradigm with more continuous drug delivery. We used 4 dosing cycles over a period of a month (19 injections in total). Such a model better reflects the clinical situation, where patients on average undergo 4 to 8 cycles of treatment over a period of 3–6 months (Cancer Research UK, 2018). We monitored the weight of all mice throughout treatment and observed a reduction in the weights of oxaliplatin treated mice within 24 hours of the first injection, a trend which persisted until the end of the experiment. Although the oxaliplatin group was not significantly lighter comparing the start and end of the experiment (p=0.5122), they were significantly lighter than their vehicle counterparts by day 9, i.e. the 7th treatment day (p=0.0087) and remained so until the end of the experiment (p=0.0047). No recovery of weight loss was observed in the treatment break week (days 15 and 18). Meanwhile, vehicle treated mice gradually gained weight over the testing period, though were not significantly heavier by the end of the experiment (p=0.1093) (Figure 3).

Although we did not perform tissue specific assays to assess platinum concentrations, as other studies have done (Canta et al., 2011; Marmiroli et al., 2017), these results strongly indicate that oxaliplatin was having a systemic effect and therefore is likely to have reached the tissues of interest in this study i.e. sciatic nerves, lumbar DRG and inguinal lymph nodes.

Repeated oxaliplatin administration did not alter total leukocyte number in lumbar DRG, sciatic nerves or inguinal lymph nodes

For this experiment, mice were processed in two batches (n=3/day) 38–39 days following their first injection of oxaliplatin (3mg/kg), or equal volume of vehicle, and all samples were run together on a flow cytometer (see Table 4 and Extended Figure 1 for panel and gating strategies employed). As was the case with the acute model, our data failed to show any differences in total leukocyte number (CD45+ cells) between oxaliplatin and vehicle groups (Figure 4 (Hore et al., 2021b)). Average cell numbers were comparable to those obtained from mice who had only received a single dose of oxaliplatin (Figure 2).

We observed substantial inter-animal variability and therefore considered whether we might be able to detect a relationship between cell numbers and processing batch or drug treatment group. We were particularly interested in the latter question, given that chronic OIPN is evident in only about 70% of patients (Molassiotis et al., 2019; Park et al., 2013). However, we did not detect any convincing correlations with either of these variables. Simple examination of the data (Figure 4) already indicates that there is no clear group difference in variability, and while some batch-associated variability was present, this was restricted to the sciatic nerve. To be sure, we also correlated total leukocyte number with behavioural data from each animal to see if there was a relationship between an animal’s sensitivity to mechanical and cold stimuli and its immune profile, but we were unable to detect any such trend (Extended Table 3).

Repeated oxaliplatin administration caused subtle changes to a subset of myeloid lineage subpopulations

Despite observing no major immune disruption, our data did show subtle alterations in a subset of the myeloid lineage cell...
types we investigated, though effects were tissue dependant (Figure 5). Within the sciatic nerves and DRG, we noted a reduction in the overall number of CD11b+ myeloid cells in oxaliplatin treated mice. In the sciatic nerve specifically, a

**Figure 3.** Repeated oxaliplatin treatment significantly impacted weight gain. Oxaliplatin treated mice started to lose weight within 1 day of dosing and became significantly lighter than the vehicle group by day 9. This trend continued throughout dosing until 6 days following the last injection i.e. ‘Day 39’, where oxaliplatin mice on average weighed 4.3 grams less than their vehicle treated counterparts. Note that days 15 and 18 were time points during the non-dosing week. Mice used for flow cytometry batches 1 and 2 were collected on days 38 and 39, respectively. Data displayed as mean ± SD, (days 1-30: n=10, days 31-32: n=9-10, day 33: n=8-9, day 39: n=5-6), RM Two-way ANOVA with Sidak’s multiple comparisons test revealed a significant main effect of group (F (1, 18) = 35.11, p<0.0001) and interaction with time (F (20, 346) = 22.66, p=0.0001). ** p<0.01, *** p<0.001, **** p<0.0001.

**Figure 4.** Total leukocyte numbers did not differ between vehicle and oxaliplatin treated mice in any tissue type following repeated administration of oxaliplatin. Analysis of total number of CD45+ live single cells from flow cytometry of lumbar 3–5 dorsal root ganglia, sciatic nerves and inguinal lymph nodes. Points of the same shape within each group were processed in the same batch. Data displayed as individual animals ±SEM, (n=6). Unpaired t-test or Mann-Whitney U test, depending on whether the data were normally distributed (Shapiro-Wilk test).
Figure 5. Following repeated oxaliplatin treatment total myeloid cell numbers did not differ between groups in any tissue type, however some myeloid subpopulations were dysregulated in a tissue dependant manner. Flow cytometry revealed no differences in total number of myeloid cells (CD45+/CD11b+) or infiltrating monocytes (CD45+/CD11b+/Ly6G-/MHCII-, Ly6C+) between groups for any tissue type. However, repeated oxaliplatin treatment appeared to reduce the number of MHCII antigen-presenting cells (CD45+/CD11b+/Ly6G-/MHCII+, Ly6C-) in the sciatic nerves and resident macrophages (CD45+/CD11b+/Ly6G-/MHCII-, Ly6C-) in the inguinal lymph nodes. Furthermore, in the inguinal lymph nodes both the double positive population (CD45+/CD11b+/Ly6G-/MHCII+, Ly6C+), which are likely infiltrating monocytes differentiating into resident populations, and the neutrophil population (CD45+/CD11b+/-Ly6G+) were significantly upregulated in the oxaliplatin group. Analysis of flow cytometry results from (A) Lumbar 3-5 dorsal root ganglia; (B) Sciatic nerves; (C) Inguinal lymph nodes. Data displayed as individual animals ±SEM (n=6). Unpaired t-test or Mann-Whitney U test, depending on whether the data were normally distributed (Shapiro-Wilk test), * p<0.05, ** p<0.01.

decrease in all myeloid cell subpopulations was noted, with exception of the MHCII-/Ly6C- double negative population, in which the opposite was observed (Figure 5B). However, none of these findings reached statistical significance, with the exception of the MHCII+ population in nerve (p = 0.0260). Trends for all myeloid subpopulations were comparable between the sciatic nerve and DRG, though were consistently more exaggerated in the sciatic nerve.

Meanwhile, an inverse picture was observed for the inguinal lymph nodes (Figure 5C). In this tissue type, an increase in total myeloid cells was observed for the oxaliplatin group, albeit statistically non-significant. Similarly, cell number directionality for individual subpopulations was in direct contrast to the DRG and sciatic nerves, with increases observed in oxaliplatin treated mice for the Ly6C+, MHCII+, MHCII+/Ly6C+ and Ly6G+ subpopulations, though this only reached significance for the latter two (p=0.0041 and p=0.0152, respectively). Meanwhile, a significant decrease was noted in MHCII-/Ly6C-cell numbers (p=0.0022).

Finally, no differences between oxaliplatin and vehicle groups were observed for lymphoid lineage cell types in any tissue (Extended Figure 2 (Hore et al., 2021a)).

Comparing single versus repeated administration of oxaliplatin, we observed clear similarities in results between the two, with trends in both myeloid and lymphoid subpopulations consistent between paradigms in the sciatic nerve and DRG (Figure 5, Extended Figures 2 and 3 (Hore et al., 2021a)). These similarities, however, were not evident in the lymph nodes (Figure 5, Extended Figures 2 and 3 (Hore et al., 2021a)). For instance, acute oxaliplatin administration significantly downregulated the number of MHCII+ cells in oxaliplatin treated mice (p=0.0230, Extended Figure 3C (Hore et al., 2021a)), while there was no significant change and, if
anything, an increase in this cell-type in lymph nodes after repeated administration (Figure 5C).

Repeated oxaliplatin administration did not result in altered sensitivity to either mechanical or cold stimuli

The fact we were unable to detect stark differences in the peripheral immune response was mirrored in the results of behavioural assays conducted on mice which underwent repeated oxaliplatin administration. Mice were tested for responses to both cold and mechanical stimuli at multiple timepoints from 1–36 days following their first injection. Results from the 10°C cold plate assay, assessed by latency to respond (hind paw shake, hind paw lick or jump), indicated that oxaliplatin treated mice did not display altered sensitivity to cold stimuli (Figure 6A). Although the oxaliplatin group consistently showed reduced latencies throughout the testing paradigm, at no point did this differ significantly to the vehicle group. Furthermore, both groups displayed similar trajectories over time, with an initial slight decrease in response latency which had resolved by the end of the experiment (day 36). Similarly, using the up-down von Frey method (Figure 6B), we were unable to detect differences in sensitivity to mechanical stimuli between oxaliplatin and vehicle treated mice. Both groups displayed a slight reduction in 50% withdrawal threshold over the first week of testing, which levelled out at around 0.4g from 11 days following first injection until the end of testing. For both behavioural tests, this absence of an oxaliplatin-effect was observed regardless of whether behaviour was conducted on dosing days, during the week-long dosing break (days 15 and 17) or after dosing had ceased (day 36).

Bone marrow derived macrophages (BMDMs) harvested from mice receiving repeated oxaliplatin administration did not differ from controls

In addition to behavioural and flow cytometry experiments, we also conducted in vitro work on mouse derived BMDMs, in order to see whether cell number and phenotype were altered by repeated oxaliplatin treatment. BMDMs from batch 1 mice (n=3) were harvested 38 days following their first injection, meanwhile, batch 2 mice (n=2) were processed 2 weeks later, thus harvesting of BMDMs from these animals took place 52 days after first injection. Prior to stimulation, cells were resuspended in PBS and counted on a haemocytometer under a light microscope. When all samples were taken together (n=5), results indicated that overall macrophage numbers did not differ significantly between the 2 treatment groups (Figure 7A). In order to assess whether oxaliplatin treatment alters the phenotype of macrophages in response to a pro-inflammatory stimulus, BMDMs from each mouse were stimulated with TNFα and qRT-PCR was used to probe for a series of marker genes associated with DNA damage, apoptosis and cellular stress. Our data showed no differences between oxaliplatin and vehicle groups, regardless of whether BMDMs had been stimulated with TNFα or not (Figure 7B). Altogether, our in vitro work failed to detect an effect of oxaliplatin on the phenotype of BMDMs. This is line with our flow cytometry findings, where we also failed to detect a change in total CD11b+ cell counts in various tissues.

Discussion

We set out to test whether alterations in neuro-immune interactions could be observed during the development or maintenance of OIPN in mice. However, despite observing clear signs that oxaliplatin was having a systemic effect, we were unable to detect any substantial oxaliplatin-induced alterations in either pain-like behaviours or the peripheral immune response. Neither acute nor repeated administration of this chemotherapeutic agent majorly influenced the immune profile in neuropathic pain-relevant peripheral tissues including sciatic nerves and lumbar DRG, or in the peripheral inguinal lymph nodes. In keeping with the lack of immune changes, though in contrast to much of the literature, we also failed to observe oxaliplatin-induced pain-like phenotypes in mice receiving multiple doses, namely increased sensitivity to cold and mechanical stimuli. There are many possible reasons for our failure to detect an effect – including limitations in our own experimental design, but also the suitability and robustness of current animal models of CIPN (Currie et al., 2019). In the following, we will discuss these issues and highlight possibilities for future investigations.

No behavioural phenotype observed in mice undergoing repeated oxaliplatin treatment

We failed to detect a behavioural phenotype in mice undergoing repeated cycles of oxaliplatin treatment over a 5-week period, with sensitivity to both mechanical and cold stimuli comparable between drug and vehicle groups. What potential limitations might account for this absence of any effect? Our choice of behavioural assays was based on a multitude of studies which have observed phenotypes consistent with CIPN in rats or mice using very similar von Frey and cold plate paradigms to the ones employed here (Currie et al., 2019). Furthermore, our data were collected in a blinded fashion by a seasoned experimenter with more than 4 years of experience working with mice. We therefore consider it unlikely that we failed to select appropriate tests or that we carried them out incorrectly.

There are of course subtle differences in how von Frey and cold plate tests are carried out at different laboratories. For example, in the cold plate assay we used “latency to respond” as a read-out, whereas previous studies have relied on “threshold to respond” (Descoeur et al., 2011; Renn et al., 2011) or the number of responses during a set time period (Ta et al., 2009). It is therefore possible that very small, specific effects might have been missed in this current study, but still unlikely that a major pain phenotype would have gone unnoticed.

Another limitation is that we cannot be absolutely sure that oxaliplatin reached and damaged sensory neurons. We have circumstantial evidence, since the drug clearly had a significant systemic effect, impeding weight gain. However, we did not employ tissue specific assays. It would therefore have been beneficial to measure platinum concentration in nerve or DRG via atomic absorption spectrometry (Canta et al., 2011;
Figure 6. Behavioural assays revealed no difference in sensitivity to either cold or mechanical stimuli between vehicle and oxaliplatin treated mice at any time point 1–36 days following first injection. (A) Although oxaliplatin treated mice had a consistently lower threshold to a 10°C cold stimuli, this did not differ significantly from the vehicle group at any time point. (B) Mechanical threshold, as assessed using the von Frey test, decreased slightly over time, but did not differ between groups at any time point. Data displayed as mean ±SEM, (24hrs-23days: n=10, 30-36days: n=8-9). RM two-way ANOVA reveals no significant main effect of group (cold plate: (F (1, 18) = 1.401, p=0.2519), von Frey: (F (1, 18) = 0.1151, p=0.7384)) nor an interaction with time (cold plate: (F (6, 102) = 0.2796, p=0.9454), von Frey: (F (9, 156) = 1.330, p=0.0543)).

Beyond our own study limitations, there are also general issues with how data from animal models of CIPN are being generated and reported. For starters, how exactly OIPN is being induced varies widely from publication to publication. When designing this study, we searched the relevant literature to no avail in hopes of finding a ‘gold standard’ model. Studies typically administer doses of anything between 2–6mg/kg, though a single dose of just 0.5mg/kg has also reportedly proven effective (Joseph et al., 2008). There is also great variation in injection regimens and routes of administration, e.g. intraperitoneal (i.p.) versus intravenous (i.v.). Though increased sensitivity to both cold and mechanical stimuli are consistently reported, seemingly irrespective of model (Extended Table 4), it is important to note that studies investigating the effects of different doses of a given chemotherapeutic agent, including oxaliplatin, have observed that higher doses induce stronger pain-like responses to cold stimuli (Descour et al., 2011; Joseph et al., 2008). Such findings highlight the difficulty in comparing results between laboratories and the need for adoption of standard dosing schedules (Flatters et al., 2017) or at least better within-study comparison of current dosing schedules. The latter could help researchers adapt their dosing regimens to specific questions they might have. For more extensive comparison of the numerous rodent OIPN models used in the current literature, please refer to Extended Table 4 and reviews by Hopkins et al. (2016) and Authier et al. (2009).

In addition to differences in model induction, one also needs to consider differences in mouse genetic background, since they have been shown to vary in their susceptibility to oxaliplatin as detectable by behavioural, morphological and neurophysiological assessments (Marmiroli et al., 2017). In fact, of particular interest to this study, the findings of Marmiroli and colleagues suggest that C57BL/6 mice are one of the least susceptible strains, displaying comparatively minimal signs of OIPN, including no cold hyperalgesia.
Figure 7. Repeated oxaliplatin administration does not alter number or phenotype of bone marrow derived macrophages (BMDMs). (A) No significant difference in the overall number of cells between oxaliplatin and vehicle groups (p=0.6419). Note that these counts were taken prior to the stimulation step, where a subset of cells were incubated with TNFa. Paired t-test. (B) No significant differences were observed between oxaliplatin and vehicle treated groups for any of the genes investigated. This was true for both unstimulated and stimulated BMDMs (all genes p=>0.12 and p=>0.16, respectively). Multiple t-tests. Data displayed as individual animals ±SEM, (n=5).

Finally, there are issues with how behavioural tests are being conducted. There are clear inter-study inconsistencies, e.g. with the same group reporting use of a 10°C stimulus to assess cold allodynia in two publications (Ling et al., 2007b; Ling et al., 2007a), but thermal hyperalgesia in another (Descoeur et al., 2011). Furthermore, temperatures used to assess cold hyperalgesia vary widely - even without performing an in-depth search of the literature, we observed a range from -4.2°C (Ta et al., 2009) to 10°C (Descoeur et al., 2011). An additional concern is that investigators continue to use relatively low n numbers in behavioural tests despite high inter- and even intra-animal variability anticipated in most assays. For example, of 1051 mechanical monofilament experiments and 190 cold plate experiments conducted in the CIPN literature, the average sample size was n=8 per group (personal communication with (Currie et al., 2019)). However, with such a sample size it is unlikely that one would be able to detect small effect sizes. For example, using a repeated measures ANOVA with four time points, the minimum effect size that you would be well powered to detect is f=0.6. Similarly, using n=8 in an unpaired t-test, the minimum effect size one would expect to detect is d=1.5. Instead, sample sizes of more than double this (i.e. n=16) are required for well-powered detection of effect sizes d=1 or larger. (Extended Figure 4 (Hore et al., 2021)). These estimates are based on general parameters and will of course vary depending on the specifics of a particular experiment (such as
the precise number of repeated measures). Nevertheless, it is likely that many studies in the field, including our own, will not be designed to detect small differences and will be vulnerable to both false positive and false negative results. In hindsight, we should have refrained from basing our sample sizes on previously published work and instead determined our n-numbers based on the smallest effect size of interest (Albers & Lakens, 2018).

The issue of blinding, or lack thereof, is also an important consideration. Although the hope is that every group conducts behavioural tests with certain practices in place, realistically this cannot be monitored. Strikingly, two recent systematic reviews examining 337 (Currie et al., 2019) and 650 (Gadgil et al., 2019) CIPN publications revealed that, respectively, only 51.3% and 44% declared that experimenters were blind to treatment group, suggesting that roughly half of all reports in the field run the risk of significant experimenter bias. Lastly, it has been demonstrated that data which confirm the null hypothesis fail to be published in the CIPN field, with many more studies than expected reporting behavioural alterations (Currie et al., 2019).

Indeed, given the statistical framework we all use, even if all studies were powered to detect relevant effect sizes 95% of the time, we would still expect 5% of them to return false negative results. Our study might have been one such instance – and for future meta-analyses, it is crucial to publish and capture each of them.

**Translational potential of current animal models of CIPN**

Beyond issues surrounding the execution of animal work in this field, there have been more fundamental concerns as to the suitability of using chemotherapeutics in animals as models for human CIPN. Notably, due to the fact robustness of oxaliplatin-based models is poor across strains and unclear across sex, a recent systematic review did not judge any current OIPN models to be appropriate, preferring instead those involving administration of paclitaxel or cisplatin via a clinically relevant route (Gadgil et al., 2019).

It is true that numerous discrepancies exist between human and animal phenotypes (Currie et al., 2019). For example, a key characteristic of CIPN in patients is sensory loss (Gupta & Bhaskar, 2015; Rice et al., 2018), while animal models almost exclusively report increased sensitivity, particularly to mechanical stimuli (Hopkins et al., 2016; Lees et al., 2017). Moreover, a recent clinical study of OIPN patients reported that the peak prevalence in mechanical deficits was only reached at 6 months following the end of chemotherapy treatment (Molassiotis et al., 2019). In contrast, animal studies tend to observe increased sensitivity to mechanical stimuli within one week of oxaliplatin administration (Gauchan et al., 2009a; Joseph & Levine, 2009; Nassini et al., 2011; Renn et al., 2011). Finally, acute CIPN reportedly affects 68% of patients on average (Seretny et al., 2014), while animal models using a range of chemotherapeutic agents consistently report behavioural phenotypes in 90–100% of subjects (Gadgil et al., 2019). Taken together, it therefore appears that type, incidence and onset of symptoms are inconsistent between human CIPN patients and animals in CIPN models. Some of these discrepancies could again be exacerbated by reporting bias, e.g. with studies omitting ‘non-responders’ from their reports without any accompanying explanation or erroneously expecting and therefore detecting a mechanical phenotype due to lack of blinding.

There have been efforts to improve on translatability. For example, tumour bearing animals have been used to investigate the effects of chemotherapy in other contexts (Hong et al., 2018; Lin et al., 2012). However, to the best of our knowledge, only one such study has attempted to directly study CIPN (Boyle et al., 2001), despite appropriate models existing for over 20 years (Boyle et al., 1999). And of course, attempts have been made to better mirror the chronic nature of chemotherapy treatment (Di Cesare Mannelli et al., 2013; Marmiroli et al., 2017) which typically lasts between 3–6 months and involves 4–8 cycles of treatment (Cancer Research UK, 2018). Accordingly, with 4 cycles over 5 weeks (5% of the lifespan of a mouse), we have tried to adopt such an improved approach here.

Finally, although repeated dosing mimics the clinical situation more accurately, it is important to acknowledge that animal welfare also poses an important consideration. The published literature would lead us to believe that similar phenotypes are produced in both mice and rats, following just a single dose of oxaliplatin (Extended Table 4). Thus, if there is truly no difference in outcomes, we would of course advocate preferential adoption of acute models, which eliminate unnecessary stress and discomfort associated with repeated injections. However, a clear absence of reports stating that a pain-like phenotype was not observed makes it difficult to gauge the proportional “success” of each type of dosing regimen. Moreover, it is unclear what “success” would even constitute, given that sensory loss – while commonplace in the clinic – is not acknowledged in the animal literature. As such, it is currently difficult to determine what the best model for studying OIPN would be, not least since the most appropriate dosing regimen might differ depending on the particular question under investigation.

**Oxaliplatin only induced minor changes in immune cell numbers in sensory nerves and lymph nodes**

With the aforementioned issues in mind, perhaps our flow cytometry experiments were never fit to answer the question of whether neuro-immune interactions are associated with the development and maintenance of OIPN. Still, we can at least conclude that even repeated oxaliplatin treatment did not have a striking effect on leukocyte numbers, with total myeloid and lymphoid cell counts comparable between oxaliplatin and vehicle groups in all tissue types investigated. However, it should be noted that power functions indicate our experiments would likely have only been sensitive enough to detect effect sizes of roughly $d=1.8$ or larger. In other words, we should have had an 80% chance of detecting differences in datasets where 96.4% of the oxaliplatin-treated samples had cell counts higher than the mean of all vehicle samples (Extended Figure 5 (Hore et al., 2021a)). Though in actuality we were able to detect effect sizes as low as $d=1.25$.
(Extended Table 2 (Hore et al., 2021a)), it is likely that we would have missed many other, smaller effects, given the constraints of our current study design.

Despite these limitations, our results are largely in accordance with previous work which found no major alterations in peripheral immune cell composition following repeated oxaliplatin administration. Immunohistochemical analysis revealed that expression of the macrophage/microglial marker IBA1 and the pan-T cell marker CD3, was comparable in the lumbar DRG and sciatic nerves of drug and saline treated mice (Makker et al., 2017). Prior flow cytometry data on total leukocyte numbers is lacking. While some studies have employed this technique to study immune cells after oxaliplatin administration, they focused on specific immune subpopulations and failed to report on total leukocyte numbers, despite inclusion of an appropriate antibody to do so i.e. CD45 (Makker et al., 2017; Stojanovska et al., 2018).

While total myeloid numbers were unchanged, we did observe some subtle effects in specific myeloid sub-populations, though this was tissue dependent. In DRG and sciatic nerve, we noted a reduction in MHCII expressing cells in oxaliplatin treated animals. Although this trend was consistent between models, it only reached significance in the sciatic nerves of mice undergoing repeated administration. In support of this finding, RT-qPCR analysis of whole DRG has revealed downregulation of the MHCII genes H2-Ab1 and H2-Eb1 following 4 weeks of i.v. oxaliplatin administration, an outcome which may suggest oxaliplatin has the ability to selectively damage MHCII+ cells (Marmiroli et al., 2017). However, the consequences of this reduction in the number of MHCII antigen-presenting cells are unclear. They are known to be essential for the initial activation of CD4+ T cells (Holling et al., 2004), but we failed to detect any striking changes in lymphocyte numbers in any tissue type investigated (Extended Figure 2 (Hore et al., 2021a)). A prior publication described increased CD4+ and CD8+ T cells in the blood after oxaliplatin treatment (Makker et al., 2017), but like us, failed to detect changes in these populations, or in overall T cell number (CD3+ cells), in the DRG or sciatic nerves. Such results indicate that even if oxaliplatin has the capacity to induce a heightened adaptive immune response, the downstream effects of this are either not large enough to be detected by current study designs or do not manifest in the peripheral nervous tissues relevant for OIPN.

In inguinal lymph nodes, we observed a different picture; following repeated oxaliplatin treatment an increase in almost all myeloid subpopulations was evident, with this reaching significance for the Ly6G+ and MHCII+/Ly6C+ populations. Meanwhile we observed a significant downregulation in MHCII+/Ly6C- cells in the oxaliplatin group. Thus, oxaliplatin appeared to cause a reduction in resident myeloid cells but an increase in infiltrating cell types such as monocytes and neutrophils. In contrast, previous work has reported a significant reduction in macrophages and dendritic cells in the mesenteric lymph nodes of oxaliplatin treated mice (Stojanovska et al., 2018), suggesting that, once more, the effects of oxaliplatin may be tissue-specific. However, like us, Stojanovska and colleagues did not find any alterations in T cell number. Such results indicate that a myeloid cell shift in the lymph nodes, in either direction, does not appear to have a large effect on adaptive immune cell numbers.

Unlike in the sciatic nerve and DRG, observed trends in total cell number for each population were not consistent between single and repeated administration models in the lymph nodes. Specifically, in the acute model the only difference between groups was noted in the MHCII+ population, where we observed a significant reduction in oxaliplatin-treated mice. These inconsistencies may suggest that, at least within the lymphatic system, immune cell composition is differentially affected by dose, though we would need to repeat these experiments to be certain that they are robust.

As a final limitation of our flow cytometry data, we would like to note that in keeping with the bulk of pre-clinical work, we used inferential statistics on small sample sizes (n = 6) and did not adjust for multiple comparisons when comparing different cell populations. While this is convention, in actuality it provides a false veneer of certainty where there is none. A small individual study such as ours does not capture enough information to conclude much about the probability of the data reflecting the underlying distribution. Only through reporting of all future studies of this nature (including those that support the null hypothesis) can we begin to build up a sample size large enough to definitively shape our view of how oxaliplatin affects immune cells “in general”.

Conclusions

Based on results from this study, we cannot reliably comment on whether neuro-immune interactions are involved in OIPN as we detected no behavioural phenotype and thus no evidence for peripheral neuropathy, even when oxaliplatin was administered in repeated cycles over long periods of time. At least in mice, therefore, we have found this model to be somewhat less robust than other peripheral pain models we have employed in the past, such as partial sciatic nerve ligation (PSNL) (Denk et al., 2016; Saunders et al., 2018) and intra-plantar administration of complete Freund’s adjuvant (Lopes et al., 2017; Saunders et al., 2018). While the absence of any distinct oxaliplatin-induced effects makes our data hard to interpret, we nevertheless decided to publish them here to help fight the publication bias widely evident in the current CIPN literature.

While our behavioural data were inconclusive, our flow cytometry experiments were somewhat easier to interpret. Specifically, in our experiments, oxaliplatin did not appear to have striking effects on peripheral myeloid and lymphoid cell types in lumbar DRG, sciatic nerves or associated lymph nodes. There were only minor changes in myeloid sub-populations,
some of which were consistent between our single and repeated administration experiments and with prior literature.

We did not examine the effects on microglia, the resident macrophages of the central nervous system, which play a prominent role in many chronic pain conditions (Suter, 2016). As it stands, the literature on the effects of oxaliplatin on microglia is conflicting, with a number of immunohistochemical studies reporting increased expression of IBA1 in the dorsal horn of the spinal cord (Cho et al., 2016; Di Cesare Mannelli et al., 2013; Di Cesare Mannelli et al., 2014), while others report no difference (Janes et al., 2015; Makker et al., 2017; Zheng et al., 2011).

Finally, our discussions highlighted general limitations with animal models of CIPN. In that context, we think it would be beneficial to streamline, at least to some extent, protocols between laboratories, increase reporting of methodological details, and make efforts to more closely mimic the types and timing of symptoms experienced by CIPN patients. Furthermore, where feasible, more focus should be put on conducting experiments using CIPN patient derived samples, like blood and surgically resected or post-mortem tissues. Findings from these studies could then be used to test the translational potential of various findings made in animals, moving us one step closer towards understanding the mechanisms underlying CIPN and aiding development of more effective therapeutics.

Data availability
Underlying data
Open Science Framework: Probing the peripheral immune response in mouse models of oxaliplatin-induced peripheral neuropathy highlights their limited translatability. https://doi.org/10.17605/OSF.IO/K2SHA (Hore et al., 2021a).

This project contains the following underlying data:

- Figure 4 and Figure 5 and Extended data Figure 2: 8 FCS files of fluorescent minus one ‘FMO’ controls used for gating purposes in flow cytometry experiments (FCS)
- Figure 4 and Figure 5 and Extended data Figure 2: Total numbers gating – repeated dosing (WSP). This is a FlowJo workspace where all samples were gated and total cell numbers were generated.


This project contains the following underlying data:

- Figure 2, Extended data Figures 2 and 3: 36 FCS files generated in flow cytometry experiments on nerve, DRG and lymph node tissues (FCS). Note that vehicle animals are labelled as ‘PBS’.
- Figure 2, Extended data Figures 2 and 3: 8 FCS files of fluorescent minus one ‘FMO’ controls used for gating purposes in flow cytometry experiments (FCS)
- Figure 2, Extended data Figures 2 and 3: Total numbers gating – single dose (WSP). This is a FlowJo workspace where all samples were gated and total cell numbers were generated.

Extended data
Open Science Framework: Probing the peripheral immune response in mouse models of oxaliplatin-induced peripheral neuropathy highlights their limited translatability. https://doi.org/10.17605/OSF.IO/K2SHA (Hore et al., 2021a).

This project contains the following extended data:

- **Extended Figure 1 (PDF).** Representative gating strategies employed for flow cytometry experiments. (A) DRG (sample displayed: vehicle, chronic model, second processing day), (B) Sciatic nerves (sample displayed: vehicle, chronic model, second processing day), (C) Inguinal lymph nodes (sample displayed: oxaliplatin, chronic model, second processing day). For all tissue types, gating was as follows: CD45+ vs FSC-A for CD45+ events; SSC-A vs FSC-A for CD45+ cells i.e. leukocytes; FSC-W vs FSC-A for single cells; Live/Dead- vs FSC-A for live cells; FSC-A vs CD11b for myeloid cells (CD11b+) OR Ly6G vs CD11b for neutrophils (CD11b+, Ly6G+) and lymphoid cells (CD11b-, Ly6G-). From the myeloid cell population with neutrophils excluded (CD11b+, Ly6G-): MCHII versus Ly6C for MCHII antigen presenting cells (MCHII+, Ly6C-) resident macrophages (MCHII-, Ly6C-), infiltrating monocytes (MCHII, Ly6C+) and a double positive population which we propose to be infiltrating monocytes differentiating into resident populations (MCHII+, Ly6C-). From the lymphoid population (CD11b-, Ly6G-): βTCR vs γ8TCR
for αβ+ T cells (βγTCR+, γδTCR-) and γδ+ T cells (γδTCR+, βγTCR-).

- **Extended Figure 2 (PDF).** Total numbers of lymphoid and lymphoid subpopulation cells did not differ between vehicle and oxaliplatin treated mice in any tissue type either after acute or repeated oxaliplatin administration. Analysis of total number of lymphoid (CD45+/CD11b-/Ly6G-), αβ+ T cells (CD45+/CD11b-/Ly6G-βγTCR+, γδTCR-) and γδ+ T Cells (CD45+/CD11b-/Ly6G-βγTCR-, γδTCR+) from flow cytometry of (A) Lumbar 3–5 dorsal root ganglia: (i) acute administration, (ii) repeated administration. (B) Sciatic nerves: (i) acute administration, (ii) repeated administration. (C) Inguinal lymph nodes: (i) acute administration, (ii) repeated administration. Data displayed as individual animals ±SEM, (n=5–6). Unpaired t-test or Mann-Whitney U test, depending on whether data was normally distributed (Shapiro-Wilk test).

- **Extended Figure 3 (PDF).** Total myeloid cell numbers did not differ between vehicle and oxaliplatin treated mice in any tissue type 4 days following a single dose of oxaliplatin (6mg/kg). Flow cytometry revealed no significant difference in total number of myeloid cells (CD45+/CD11b+) for any tissue type. Similarly, no significant dysregulation was observed in any myeloid lineage subpopulation investigated, with the exception of MHCII antigen presenting cells (CD45+/CD11b+/Ly6G/MHCII+, Ly6C-), which were downregulated the inguinal lymph nodes of oxaliplatin treated mice. Analysis of flow cytometry results from (A) Lumbar 3–5 dorsal root ganglia. (B) Sciatic nerves. (C) Inguinal lymph nodes. Data displayed as individual animals ±SEM, (n=5–6). Unpaired t-test or Mann-Whitney U test, depending on whether data was normally distributed (Shapiro-Wilk test), * p<0.05, ** p<0.01.

- **Extended Figure 4 (PDF).** Plotted power functions give an idea of the smallest effect sizes we would have been likely to detect in our flow cytometry experiments with n = 6 (80% chance of detection for any x-axis values at or to the right of the dotted black vertical line). These power calculations are for a parametric unpaired t-test (A) and a non-parametric Mann-Whitney test (B), to mirror those used in our analyses.

- **Extended Table 1 (DOC).** List of animals excluded from experimental analysis.

- **Extended Table 2 (DOC).** Effect sizes of all comparisons which resulted in statistically significant differences. Selected non-significant examples from the myeloid cell population (CD45+/CD11b+) are also given for comparative purposes. Extended Figure 5 provides the minimum effect sizes for which these statistical tests would have been well-powered (i.e. 80%), given our total sample size of 12.

- **Extended Table 3 (XLSX).** Data for correlation of total leukocyte number (CD45+ live single cells) in each tissue type investigated with behavioural data, to determine if there is a relationship between sensitivity to mechanical and cold stimuli and immune profile.

- **Extended Table 4 (DOC).** A comparison of all the oxaliplatin models used in publications referenced throughout this paper and the resulting behavioural phenotypes reported.

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

**Reporting guidelines**

Open Science Framework: The ARRIVE guidelines 2.0: author checklist for ‘Probing the peripheral immune response in mouse models of oxaliplatin-induced peripheral neuropathy highlights their limited translatability’. https://doi.org/10.17605/OSF.IO/F9CKW (Hore et al., 2021a)

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

**Acknowledgments**

We would like to thank Professor Andrew Rice for his expertise and advice. We also wish to thank our placement BSc students Covadonga Mont Fernandez and Raquel Llama Hernandez for aiding with the BMDM experiments conducted during this study and Dr Naomi Shinotsuka for her help harvesting and culturing BMDMs.


Open Peer Review

Current Peer Review Status: ✓  ?

Version 2

Reviewer Report 15 June 2021

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I think the manuscript has greatly improved and now is ready for indexing.

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Neurobiology of pain

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Version 1

Reviewer Report 29 April 2021

https://doi.org/10.21956/wellcomeopenres.18340.r43304

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Hore and colleagues describe their investigation of oxaliplatin-induced peripheral neuropathy following single or repeated oxaliplatin administration. In the repeated dosing model they assessed for mechanical and thermal pain-related behaviours. They also investigated whether there were changes to peripheral immune cells in these models. This study did not identify changes in leukocyte, lymphocyte or myeloid cell numbers in DRG, sciatic nerves or inguinal lymph nodes. Small changes in myeloid cells were identified. Importantly, the authors highlight that their results are not consistent with what has been reported in the field.

We applaud you, this is an excellent example of transparent reporting, and it is encouraging to read such study, especially as the findings do not agree with the published literature. This is a really important step to tackle publication bias (as the authors note). Further, you have shared the raw data from your study (via Open Science Framework).

You consider in detail the impact of study design limitations (e.g. small sample size) and report your results with this in mind. This is an important study for the field highlighting the need to appropriately power experiments due to inter and intra animal variability of behavioural assays. You provide excellent detail on the measures used to ameliorate harm to animals used in the study and you give details of attrition (not common across the CIPN literature).

Some thoughts for you to consider:

1. Where you state the study is reported in line with the ARRIVE guidelines, we would encourage you to be specific that the citation is to their checklist.

2. Randomisation - you use the term “arbitrarily assigned”? It is unclear whether this is true randomisation?

3. You do not adjust your p-value for multiple testing, we would encourage this (e.g. figure 5). Although given that outcomes were not sufficiently powered or no primary outcome is stated, inferential statistics are probably not appropriate.

4. Related to the above point, you based your sample size on previous studies. If previous studies were underpowered (which is likely) this leads to a persistent problem that is likely a domain-wide issue. However, you do note the limitation of using a small sample size in your discussion.

5. Where live and re-scored latencies differed, the re-scored time was taken but is there no possibility this value was erroneous?

6. For the power functions, it is unclear which data were used. You describe “published behavioural tests”, where do these come from? These findings are first described in the discussion, they should probably be included in the results sections.

7. You have essentially used “Weight” as an outcome, we suggest including this in the outcome measures list.

8. Where you state “data not shown”, is it possible to add this to the OSF project for
completeness?

9. We appreciate that “negative data” are how such findings are often described but we think this is unnecessarily pejorative. Given the working hypothesis (no formal hypothesis is stated), you have accepted the null hypothesis. We also assert that the limitations in experimental design you describe could lead to either accepting or rejecting the null hypothesis, it is just that in cases like this we try to find reasons for the findings observed. We would encourage you to re-frame this as all studies, irrespective of their findings, should be taking a similar approach as you have done here.

10. You suggest that we should having a standard dosing schedule for oxaliplatin. We can imagine that different dosing schedules may be required in certain circumstances and it may be more useful would be to better understanding of the effect of different dosing schedules?

**Is the work clearly and accurately presented and does it cite the current literature?**
Yes

**Is the study design appropriate and is the work technically sound?**
Partly

**Are sufficient details of methods and analysis provided to allow replication by others?**
Yes

**If applicable, is the statistical analysis and its interpretation appropriate?**
Partly

**Are all the source data underlying the results available to ensure full reproducibility?**
Yes

**Are the conclusions drawn adequately supported by the results?**
Yes

**Competing Interests:** We were contacted by Franziska Denk about the methods used in Animal models of chemotherapy-induced peripheral neuropathy: A machine-assisted systematic review and meta-analysis (plos.org) because the authors wanted to include discussion of them but did not want to misrepresent them. We were however still able to provide an impartial review of the study.

**Reviewer Expertise:** experimental validity, preclinical bio-curation, meta-research

We confirm that we have read this submission and believe that we have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however we have significant reservations, as outlined above.
Thank you for the supportive feedback regarding our manuscript. We greatly appreciate you taking the time to bring to our attention a number of important points and hope that we have addressed your concerns with our revised manuscript or with our responses below.

1. Where you state the study is reported in line with the ARRIVE guidelines, we would encourage you to be specific that the citation is to their checklist. Apologies for this important oversight. In the ‘Ethical Considerations’ section we have now cited the Percie du Sert et al. (2020) paper these guidelines derive from. Additionally, the link “Hore et al 2021a” provided in our MS takes you to OSF where we have uploaded a PDF of the Full ARRIVE 2.0 Checklist where we note the section of our manuscript in which each recommendation is addressed (Files > Archive of OSF storage > ARRIVE guidelines 2.0 - Author Checklist).

2. Randomisation - you use the term “arbitrarily assigned”? It is unclear whether this is true randomisation? As we did not employ computer-based randomisation, we cannot term our approach true randomisation and have thus deemed it “arbitrary assignment”. We have now explained more thoroughly how animals were assigned in the methods section to clarify what we mean by the term “arbitrary”.

3. You do not adjust your p-value for multiple testing, we would encourage this (e.g. figure 5). Although given that outcomes were not sufficiently powered or no primary outcome is stated, inferential statistics are probably not appropriate. We completely agree. We followed the convention of most pre-clinical work by using inferential statistics on samples sizes that are arguably too small to be meaningful (given the expected effect sizes) and using unadjusted p-values when adjusted ones would have been more appropriate. To highlight this potential issue, we have now added a paragraph to our discussion – just before the "Conclusion" section.

4. Related to the above point, you based your sample size on previous studies. If previous studies were underpowered (which is likely) this leads to a persistent problem that is likely a domain-wide issue. However, you do note the limitation of using a small sample size in your discussion. This is very true and indeed, in hindsight, we should not have based sample sizes on previous publications, but instead on smallest effect size of interest, as highlighted in Albers and Lakens (2018). We have now added this to our discussion in the section on behavioural phenotypes.

5. Where live and re-scored latencies differed, the re-scored time was taken but is there no possibility this value was erroneous? This is a possibility; however, we would argue that scoring from the video will always provide a more accurate value than live scoring. During re-scoring, videos could be paused and re-watched to ensure a true response was recorded. Indeed, in many cases, where responses were subtle, videos were re-watched multiple times to ensure accuracy of the
latency recorded. Additionally, in some instances a live score was not recorded as it was difficult to observe the animal without being too intrusive or no clear responses were made during the 90 seconds. In these instances, scoring solely from the video recording was the only option. Therefore, in order to be consistent, we decided it would be most appropriate to take the re-scored values for all instances.

6. For the power functions, it is unclear which data were used. You describe “published behavioural tests”, where do these come from? These findings are first described in the discussion, they should probably be included in the results sections.

Our power functions are simple sensitivity analyses. We are not using any particular data/studies, instead we are determining the mere mathematics of which effect size a given statistical test is powered to detect given different samples sizes, ranging from what is a below average n number for a behavioural study in the field (e.g. n = 6) to what is an above average number for a behavioural study (e.g. n = 16). We know from your work that n = 8 is average in the field.

Given the theoretical nature of these sensitivity analyses, we therefore did not really consider them results as such. We have now tried to clarify this in our “statistical analysis” section by separating the paragraph on actual observed effect sizes based on our data from that on the theoretical sensitivity analyses. We also dropped the word “published” – confusing in this context.

7. You have essentially used “Weight” as an outcome, we suggest including this in the outcome measures list.

Apologies for this oversight, we have now added this to our outcome measures list.

8. Where you state “data not shown”, is it possible to add this to the OSF project for completeness?

Of course - we have now uploaded Extended Table 3 to OSF, where data comparing overall leukocyte numbers to behavioural phenotype for each animal is provided.

9. We appreciate that “negative data” are how such findings are often described but we think this is unnecessarily pejorative. Given the working hypothesis (no formal hypothesis is stated), you have accepted the null hypothesis. We also assert that the limitations in experimental design you describe could lead to either accepting or rejecting the null hypothesis, it is just that in cases like this we try to find reasons for the findings observed. We would encourage you to re-frame this as all studies, irrespective of their findings, should be taking a similar approach as you have done here.

All uses of the term “negative data” have now been altered to avoid using what we agree to be a pejorative term.

10. You suggest that we should having a standard dosing schedule for oxaliplatin. We can imagine that different dosing schedules may be required in certain circumstances and it may be more useful would be to better understanding of the effect of different dosing schedules?

This is a very good point, thank you for bringing it to our attention. Though we do comment on the fact that different dosing regimens all appear to result in similar behavioural
phenotypes (please see our newly added Extended Table 4), we also note that higher doses reportedly induce stronger pain-like responses. Thus, it is clear that more investigation to compare different dosing schedules is required in order for researchers to be able to decide upon the most appropriate regimen for their particular study. Accordingly, we have now worked this into our discussion under the section “No behavioural phenotype observed in mice undergoing repeated oxaliplatin treatment”.

**Competing Interests:** No competing interests were disclosed.

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Reviewer Report 12 April 2021

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Margarita Calvo

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The article "Probing the peripheral immune response in mouse models of oxaliplatin-induced peripheral neuropathy highlights their limited translatability" is a very clear report of a negative study on Oxaliplatin induced neuropathy in a mouse model. I agree with the authors that publication of data like this is very important for the field as it help to improve models of neuropathic pain and because it reduces publication bias.

The methods and the analysis are very detailed, allowing for reproducibility of the data, which is very welcome and not very common.

My only suggestion would be to include in the discussion data regarding OIPN in rats, and how this compares to mouse models. It is important to try to understand which model of OIPN is best fitted to reproduce the clinical problem, and therefore, a comparative with data published in other rodents might be very useful. I would also suggest to make a detailed comparison of the protocols used to administer oxaliplatin in the different models, as this varies across laboratories and might be key determinants of development of neuropathy.

I agree that nothing can be said in terms of immune system recruitment in OIPN with this model, as it does not reproduce the neuropathy seen in patients.

Maybe it is worth in this article to give suggestions to other researchers on what are the best models for studying OIPN in the opinion of the authors: species to use, protocol and dose of oxaliplatin administration, and outcomes to measure. This may help other researchers to use adequate models, and to avoid models which have proven not to mimic the clinical condition, in
order to understand the mechanism behind OIPN.

Is the work clearly and accurately presented and does it cite the current literature?
Partly

Is the study design appropriate and is the work technically sound?
Yes

Are sufficient details of methods and analysis provided to allow replication by others?
Yes

If applicable, is the statistical analysis and its interpretation appropriate?
Yes

Are all the source data underlying the results available to ensure full reproducibility?
Yes

Are the conclusions drawn adequately supported by the results?
Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Neurobiology of pain

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 02 Jun 2021

Zoe Lee Hore, King’s College London, London, UK

Thank you for the positive feedback and for bringing to our attention some additions that could be made to improve the utility of our manuscript to the scientific community working in this area.

To address the concerns you have raised, we have made the following changes to the manuscript:

Firstly, we have now added Extended Table 4 to our Extended Data on OSF, which compares all of the models used in papers we have referenced in our manuscript. This table highlights differences in administration route, dose and the resulting phenotypes observed in both single and repeated dose models in mice and rats. In addition, we have referenced two review papers which have compared a number of oxaliplatin-based models used across the two species.

In relation to these comparisons, we have now noted in our discussion that models used
between rodent species all appear to result in similar phenotypes, whether the drug is administered acutely or in repeated cycles. We also make the point that while no models in the current OIPN literature appear to mimic the clinical problem exactly, repeated dosing at least mimics clinical administration schedules. However, as it stands, the literature is clearly “missing” data on when model induction failed. It is therefore difficult to gauge the proportional “success” of each type of dosing regimen. As such, we unfortunately do not feel as though we can currently make a very informed choice on the most appropriate model for studying OIPN. We state this clearly in the text.

**Competing Interests:** No competing interests were disclosed.