RESEARCH ARTICLE

A fruit fly model for studying paclitaxel-induced peripheral neuropathy and hyperalgesia [version 2; referees: 1 approved, 1 approved with reservations]

Previously titled: A fruit fly model for studying paclitaxel-induced pain

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Abstract

Background: Paclitaxel-induced peripheral neuropathy is a common and limiting side effect of an approved and effective chemotherapeutic agent. The cause of this nociception is still unknown.

Methods: To uncover the mechanism involved in paclitaxel-induced pain, we developed a Drosophila thermal nociceptive model to show the effects of paclitaxel exposure on third instar larvae.

Results: We found that paclitaxel increases heat nociception in a dose-dependent manner, and at the highest doses also obstructs dendritic repulsion cues.

Conclusions: Our simple system can be applied to identify regulators of chemotherapy-induced pain and may help to eliminate pain-related side-effects of chemotherapy.

Keywords

Drosophila, fruit fly, paclitaxel, nociception, pain, CIPN
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Author roles: Hamoudi Z: Conceptualization, Formal Analysis, Investigation, Methodology, Project Administration, Supervision, Visualization, Writing – Original Draft Preparation, Writing – Review & Editing; Khuong TM: Formal Analysis, Investigation, Writing – Original Draft Preparation; Cole T: Formal Analysis, Methodology; Neely GG: Conceptualization, Funding Acquisition, Methodology, Project Administration, Supervision, Writing – Review & Editing

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Introduction
Chemotherapy-induced peripheral neuropathy (CIPN) is a dose-limiting side effect of many effective cancer treatments (Burton et al., 2007), and can have a lasting impact on the quality of life of cancer survivors (Hausheer et al., 2006 and Shimozuma et al., 2012). A meta-analysis of 31 studies from over 4000 chemotherapy-treated patients revealed that CIPN was prevalent in 68.1% of patients in the first month following chemotherapy, in 60% of patients at 3 months, and in 30% at 6 months or more (Seretny et al., 2014).

Paclitaxel has a potent ability to cause CIPN (Addington & Freimer, 2016; Reyes-Gibby et al., 2009). Derived from the bark of the western yew, Taxus brevifolia, it is an approved and effective treatment against breast, ovarian, lung and Kaposi sarcoma (Chang et al., 1993; Gill et al., 1999; Holmes et al., 1991; McGuire et al., 1989; Wani et al., 1971). Patients treated with paclitaxel experience side effects as early as one to three days following treatment (Lipton et al., 1989; Reyes-Gibby et al., 2009). Common symptoms are hypalgesia, hyperalgesia, allodynia, tingling, numbness, and shooting pain (Boland et al., 2003). Paclitaxel has a direct effect on Schwann cells, promotes axonal degeneration, and can cause mitochondrial damage (André et al., 2000; Cavaletti et al., 1995; Sahenk et al., 1994), however the molecular mechanisms causing pain are still largely unknown.

While much knowledge has been gained about the genetics of pain from vertebrate systems, high-throughput dissection of pain is possible using the fruit fly Drosophila melanogaster (Neely et al., 2010). When challenged with a noxious thermal stimulus, third instar larvae exhibit an aversive escape response that has been utilised to identify conserved genes required for nociception (Babcock et al., 2009; Neely et al., 2010; Tracey et al., 2003). This nociceptive response is a result of activating class IV multidendritic-dendritic arborisation (md-da) sensory neurons at the site of stimulation (Hwang et al., 2007). Previously in Drosophila, paclitaxel has been reported to be toxic in somatic cells, and causes loss of axons in peripheral nerves. (Bhattacharya et al., 2012; Cunha et al., 2001). However, its effects on nociception have not yet been evaluated. Here, we examined the effects of paclitaxel exposure on the fruit fly larval nociception system, and observed a robust and dose-dependent increase in pain perception. This system is amenable to high throughput screening and genetic manipulation (Honjo et al., 2016), and may help define why chemotherapies such as paclitaxel cause pain.

Methods
Drosophila treatment
All flies were reared at 25°C and 65% humidity over a 12-hour light-dark cycle. Six female and two male Canton S Drosophila melanogaster were mated on food medium (5.4% sucrose, 3.6% yeast, 1% agar, 1.2% nipagin, and 0.6% propionic acid) treated with ethanol (vehicle), 0 µM, 0.1 µM, 0.5 µM, 2.5 µM, 5 µM or 10 µM paclitaxel (Taxol®; Catalog No. A4393) purchased from ApexBio (Houston, USA). A stock of 1000 µM paclitaxel in ethanol was prepared and diluted in food medium accordingly to create the different drug concentrated food.

F0 Flies were discarded two days after mating and F1 larvae were left to grow for another three days. On the sixth day, early third instar were collected to assess nociception or dendritic morphology.

Behavioural assay
For the thermal nociceptive assay (Tracey et al., 2003), distilled water was added to experimental vials to soften the food and release the foraging third instar larvae. The softened, liquid food was then passed through mesh to catch the larvae to be transferred to a 100mm petri dish sprayed with distilled water. The larvae were touched laterally on abdominal segments four to six with a heat probe (soldering iron with narrow tip) set to 42°C or 46°C. The rolling response was measured in seconds with a cut-off of 10 seconds. For each drug concentration, five repeats were performed, with 30–40 larvae per repeat.

Live confocal microscopy and image analysis
Third instar larvae (ppk-Gal4,20xUAS-mCD8-GFP) were collected, washed, and placed dorsal side up on a microscope slide, immobilized in 1:5 (v/v) diethyl ether to halocarbon oil and covered with a 22 × 50 mm glass coverslip (Das et al., 2017). A Nikon C2 Confocal microscope was used to image GFP-expressing class IV md-da sensory neurons at abdominal segment 2 (A2), under a 20x magnification. Images of Z-stack sections were captured at 1024 × 1024 pixel resolution and representative images were captured at 2048 × 2048 pixel resolution, both with 2x averaging. Z-stack images were converted to maximum intensity projection using Imagem and automated Sholl analysis was performed on these images. Terminal branches were counted manually. 13 animals were imaged for each treatment. All experiments were conducted in a blinded manner.

Statistical analysis
Data represent mean ± SEM and are compared to vehicle control. Analysis was done using GraphPad Prism 5. Statistical analysis for response time was done using Kruskal-Wallis, followed by Dunn’s pairwise test for multiple comparisons. Statistical analysis for area under the curve mean, terminal branches,
Results

Our goal here was to develop a reproducible paradigm to investigate the effects of paclitaxel on nociception in the fly larvae. Based on previous studies for toxicity (Bhattacharya et al., 2012; Cunha et al., 2001), we selected paclitaxel doses below the lethal limit (Figure 1A), and then tested larval nociception using a heat probe set to a low intensity noxious heat (42°C; Figure 1B), which is mildly noxious to fly larvae (Babcock et al., 2009). Our dose-response study revealed 2.5 µM paclitaxel was sufficient to induce significant hyperalgesia, with a maximal hyperalgesia effect observed at 10 µM (Figure 1C, d = 0.54). Concentrations higher than 10 µM paclitaxel were 100% lethal (not shown). Paclitaxel did not significantly alter heat nociception latency to a 46°C heat stimulus across any of the doses (Figure 1D, d = 0.17). Vehicle (ethanol) control and normal (no ethanol) control showed a response time of 5.71 sec (±0.23 SEM; n=173) and 5.62 sec (±0.20 SEM, n=180, not shown), respectively (42°C; Figure 1E). At low concentrations of 0.1 µM (5.21 sec ± 0.23 SEM; n=150) and 0.5 µM (5.44 sec ± 0.26 SEM; n=131) paclitaxel did not affect response profiles, however, concentrations of 2.5 µM paclitaxel (4.22 sec ± 0.19 SEM; n=180; p<0.001) and higher altered response distribution and significantly enhanced nociceptive latency (42°C; Figure 1E). The fastest latency response was observed at 10 µM paclitaxel (3.84 sec ± 0.24 SEM; n=140; p<0.001) with a 36.6% increase in response time relative to vehicle control (Figure 1C).

To evaluate if paclitaxel exposure caused robust morphological differences in peripheral pain sensing neurons, we fed genetically labelled (ppk-Gal4,20xUAS-mCD8-GFP) larvae paclitaxel and imaged the sensory neuron structure (Figures 2A–B). Treating larvae with 10 µM paclitaxel affected its repulsive cues with like neurons, overlapping and forming a closed circular structure (Figure 2B, orange box) compared to vehicle control (Observed in 5 paclitaxel treated animals compared to 0 control animals, Fisher’s Exact Test p < 0.05). In some paclitaxel treated larvae we observed very short dendritic arbors with lower GFP intensity (Figure 2B’, open arrowhead). This was not observed in vehicle control larvae (Figure 2A’). We next used Sholl analysis to quantify branch distribution with a focus on number of intersections as a function of distance from the cell soma. This revealed increased branching closer to the cell soma in paclitaxel treated larvae compared to control (Figure 2C). Area under the curve (AUC) was also calculated for each animal (Figure 2D). Treatment with paclitaxel significantly increased the area under the curve compared to vehicle control (Figure 2D, p < 0.05). We also determined maximum branch number and its critical radius and found paclitaxel treatment compared to vehicle control did not have a significant effect on maximum branch number (62.62 ± 2.69; n=13 control and 61.28 ± 2.72; n=13 paclitaxel) or critical radius (177.1 ± 6.78; n=13 control and 192.1 ± 7.70; n=13 paclitaxel) (Figures 2E–F). Finally, paclitaxel did not significantly affect terminal branch number compared with vehicle control (Figure 2G).
Figure 1. Paclitaxel induces heat-hyperalgesia in Drosophila larvae. Schematic representation of the A) experimental design and B) thermal nociceptive assay in Drosophila larvae. C–D) Average nociceptive latency (in seconds) in response to a 42°C or 46°C thermal stimulus, respectively. Increased paclitaxel concentration significantly induces heat-hyperalgesia in third instar larvae at 42°C. Note concentrations higher than 10 µM paclitaxel were 100% lethal. E) Percentage response to each time point in seconds to 42°C thermal stimulus. All values represent mean ± SEM. p values were generated using Kruskal-Wallis, followed by Dunn’s pairwise test for multiple comparisons. Significance is relative to vehicle control. Five repeats were performed for each drug concentration with roughly 30 larvae each (n = 130–180 animals).
Figure 2. Paclitaxel obstructs dendritic repulsion cues. Representative images (A–B) and quantification (C–G) of ppk-Gal4,20xUASmCD8-GFP larvae following vehicle control or 10 µM paclitaxel treatment. Images are of class IV md-da neurons at abdominal segment A2, under a 20x magnification. Scale bar represents 100 µm. Paclitaxel treatment obstructs dendritic repulsion cues (B’, shaded arrowhead), compared to vehicle control (A’).

C. Branch distribution using Sholl analysis.

D. Area under the curve.

E–F) Maximum branch numbers and critical radius reported by Sholl analysis.

G) Branch terminal numbers. Values represent mean ± SEM (n = 13 animals). n.s. p > 0.05, t tests and post hoc comparisons: *p < 0.05.
approaches to dissect this mechanism and identify regulators of chemotherapy pain. Together this work can lead to a better understanding of how the pain arises, and potentially avoid these severe side effects while more effectively targeting the underlying disease.

Data availability

Dataset 1: Larval response time in seconds to 42°C heat stimulus. Paclitaxel fed larvae were touched with a 42°C heat probe and their response time was measured in seconds with a cut-off of 10 seconds. Different treatments were tested: food control, ethanol (vehicle) control, 0.1 µM, 0.5 µM, 2.5 µM, 5 µM, and 10 µM paclitaxel. Five repeats were performed (n = 130 - 180). DOI, 10.5256/f1000research.13581.d191023 (Hamoudi et al., 2018a).

Dataset 2: Larval response time in seconds to 46°C heat stimulus. Paclitaxel fed larvae were touched with a 46°C heat probe and their response time was measured in seconds with a cut-off of 10 seconds. Different treatments were tested: food control, ethanol (vehicle) control, 0.1 µM, 0.5 µM, 2.5 µM, 5 µM, and 10 µM paclitaxel. Five repeats were performed (n = 130 - 180). DOI, 10.5256/f1000research.13581.d191022 (Hamoudi et al., 2018a).

Dataset 3: Dendritic morphology of third instar pppk-Gal4,20xUASmCD8-GFP. Confocal images of vehicle control and 10 µM paclitaxel treated larvae. Images represent class IV md-da neurons at abdominal segment A2. Images are at 20x magnification with 2x averaging. Scale bar represents 100 µm. DOI, 10.5256/f1000research.13581.d222127 (Hamoudi et al., 2018c).

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The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.
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Current Referee Status: ✔️ ❓

Version 1

Referee Report 22 March 2018

doi:10.5256/f1000research.14752.r30145

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This study examined the effects of Paclitaxel exposure on Drosophila larval nociception system and the authors propose that their model is suitable for high throughput screening and further mechanistic studies. The study is overall an interesting and clearly written, however, I do have the following concerns:
1. The dose response effect of thermal stimulation was only at 42 degrees. There was no discussion or explanation why this effect was not seen at 46 degrees.
2. The behavior experiment was based on thermal stimulation. I would be interested why mechanical stimulation was not chosen since mechanical sensitivity is common among patients who develop Paclitaxel induced peripheral neuropathy?
3. It is not clear to me what the timeline is between the exposure of the larvae with paclitaxel and performing microscopic studies.
4. There should be at least a short discussion about the result.

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?
Yes

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

If applicable, is the statistical analysis and its interpretation appropriate?
I cannot comment. A qualified statistician is required.

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

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.
I have read this submission. I 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 09 Oct 2018

Greg Neely, University of Sydney, Australia

Thank you for your comments.

Response to comment #1: This effect is not seen at 46°C. At this temperature intensity, larvae respond rapidly (~1.5 seconds) and it is difficult to see even faster responses. To look for hyperalgesia, we instead lowered the heat stimulus intensity to 42°C, which is at the threshold for nociception in this system, and where nociceptive responses take on average ~5 seconds to elicit.

Response to comment #2: The type IV multidendritic nociceptor neurons that transduce heat nociception also transduce mechanical nociception, as these neurons are multimodal. We have tried on numerous occasions to generate reproducible data for mechanical nociception but so far in our hands this assay does not work well enough for us to feel comfortable publishing. Given the multimodal nature of type IV multidendritic nociceptor neurons, we reasoned that thermal hyperalgesia is a good readout for the overall sensitization of these sensory neurons.

Response to comment #3: The animals are born into paclitaxel containing food, and then early third instar are collected at day 6 to assess nociception or dendritic morphology. This information was provided in the methods, however we have further clarified this aspect.

Response to comment #4: We have now written a short discussion, please see discussion section.

Competing Interests: No competing interests were disclosed.

Referee Report 01 March 2018

doi:10.5256/f1000research.14752.r30150

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The authors characterize the thermal nociception in *Drosophila* larvae that have been cultured on a range of paclitaxel concentrations. Using a heat probe to elicit the rolling defense behavior, they find that while paclitaxel has no effect on response times with a 46°C probe, it shortens probe response times when the larvae have been grown on 2.5 µm paclitaxel or above.

The authors and readers might like to consider the following comments on and questions about the 23 Jan 2018 version.
1. The Title describes a model for studying paclitaxel-induced pain, however the assay uses a heat probe to induce pain, and paclitaxel lowers the sensitivity to the probe, thus modeling the hyperalgesia component of paclitaxel CIPN. Would the Title better serve the reader if edited to focus on this side-effect specifically?

2. In Abstract–Results, the authors write: "We found that paclitaxel increases pain perception in a dose-dependent manner, without overt morphological changes." Changing "perception" to "sensitivity" would eliminate the baggage of the former word.

3. In Abstract–Conclusions: "Our simple, high throughput model can be combined with genomics approaches to identify regulators of chemotherapy-induced pain to eliminate its adverse side effects." However, they have not established that this is high-throughput by most common definitions of the term, nor do they show anywhere in the paper that it can be combined with genomics. The Conclusions would be improved if rephrased to better reflect what the data show.

4. "High-throughput" is a phrasal adjective that requires a hyphen.

5. In Introduction it says "This system is amenable to high throughput screening" however, this is not shown in the present manuscript nor is a reference cited in support of the statement.


7. I encourage the authors to use estimation statistics instead of significance testing. This would involve presenting and discussing the effect sizes. For example, in Figure 1c, it looks like 2.5 µm paclitaxel has the effect of reducing response time by ~1.5 s.

8. It would be nice to calculate standardized effect sizes (e.g. Cohen's d) of the paclitaxel effects; this would allow the authors and readers to estimate sample sizes needed for a screen (and thus possible throughput rates).

9. In Results, the authors write "Thus we establish that paclitaxel sensitizes larvae to heat pain via enhancing sensory neuron or higher order nociception, and not via inducing overt morphological changes." Is it true that enhancing sensory neuron or higher-order nociception are the only two alternatives? If not, this sentence should be rephrased.

10. The Conclusions section reads more like an overview of future plans for the assay. Could it be rewritten to more closely address the paper's findings?

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?

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?

Partly

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

Yes

Are the conclusions drawn adequately supported by the results?

No

Competing Interests: No competing interests were disclosed.
I have read this submission. I believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Author Response 09 Oct 2018

Greg Neely, University of Sydney, Australia

Thank you for your comments.

Response to comment #1: That’s a reasonable point. Since first submission, we have new data that shows our model also involves peripheral neuropathy, so taken together, we have updated the title to capture this aspect and address the reviewer’s comment. The new title is “A fruit fly model for studying paclitaxel-induced peripheral neuropathy and hyperalgesia”. We hope this is acceptable.

Response to comment #2: Done.

Response to comment #3: We have revised this section and now state “Our simple system can be applied to identify regulators of chemotherapy-induced pain”.

Response to comment #4: Done.

Response to comment #5: We have now included a reference for this statement.

Response to comment #6: Done.

Response to comment #7: We thank the reviewers for their comment and we have incorporated the estimation statistics into our analysis and added all the data points.

Response to comment #8: We have calculated the effect size, added it to the graphs (1C and 1D), and we have now also mentioned this in the results section. Moreover, we have changed the data representation to show all the data points.

Response to comment #9: Good point, this has been revised as suggested.

Response to comment #10: We have now added a discussion section.

Competing Interests: No competing interests were disclosed.

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