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The lethal and sublethal impacts of two tire rubber-derived chemicals on Brook trout (Salvelinus fontinalis) fry and fingerlings

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Key Words: 6PPD-quinone, tire leachate, Brook trout, HMMM, gill morphology

Synopsis: Environmentally relevant 6PPD-quinone exposures cause higher mortality rates in Brook trout Fry inversity of California Riverside, Riverside, CA, USA

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 The lethal and sublethal impacts of two tire rubber-derived chemicals on Brook trout (Salvelinus fontinalis) fry and fingerlings

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fry than fingerlings, altered blood analytes and gill morphology are the likely mechanism of action.

1. Abstract

 Recent toxicity studies of stormwater runoff implicated N-(1,3-dimethylbutyl)-N'-phenyl-p-8 phenylenediamine-quinone (6PPD-quinone) as the contaminant responsible for the mass mortality of coho salmon (*Oncorhynchus kisutch*). In the wake of this discovery, 6PPD-quinone has been measured in waterways around urban centers, along with other tire wear leachates 11 like hexamethoxymethylmelamine (HMMM). The limited data available for 6PPD-quinone have 12 shown toxicity can vary depending on the species. In this study we compared the acute toxicity of 6PPD-quinone and HMMM to Brook trout (*Salvelinus fontinalis*) fry and fingerlings. Our 14 results show that fry are ~3 times more sensitive to 6PPD-quinone than fingerlings. Exposure to HMMM ≤ 6.6 mg/L had no impact on fry survival. These results highlight the importance of 16 conducting toxicity tests on multiple life stages of fish species, and that relying on fingerling life 17 stages for species-based risk assessment may underestimate the impacts of exposure. 6PPD- quinone also had many sublethal effects on Brook trout fingerlings, such as increased interlamellar cell mass (ILCM) size, hematocrit, blood glucose, total CO2, and decreased blood sodium and chloride concentrations. Linear relationships between ILCM size and select blood 21 parameters support the conclusion that 6PPD-quinone toxicity is an outcome of osmorespiratory challenges imposed by gill impairment. udies of stormwater runoff implicated N-(1,3-dimethylbuty
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2. Introduction

 quinone remains unclear. Studies on stormwater runoff as a whole have found that exposure increases hematocrit, decreases plasma sodium and chloride concentrations, decreases blood 61 pH, and disrupts the blood-brain barrier in coho salmon (Blair et al., 2021; McIntyre et al., 62 2021) (Chow et al., 2019). Rainbow trout and Brook trout exposed to 6PPD-quinone also had increased hematocrit levels as well as increases in blood glucose (Brinkmann et al., 2022).

64 6PPD-quinone also decreases swim behavior and increases oxygen consumption in zebrafish, an

 insensitive species (Varshney et al., 2022). *In vitro* studies with Rainbow trout gill cell lines suggest that disruption of mitochondrial respiration through electron transport chain 67 uncoupling may also play a role in the selective toxicity of 6PPD-quinone (Mahoney et al., 68 2022). Changes in the blood, gill, and brain may all be drivers of the effects observed and may play a role in determining species-specific sensitivity to 6PPD-quinone exposure, but more research is needed.

 While previous work has indicated that Brook trout are sensitive to 6PPD-quinone, additional work with this important species is warranted to independently validate those results, to expand on the understanding of potential mechanism of action, and to explore differences in sensitivity across life stages. Most toxicity tests performed on species sensitive to 6PPD- quinone exposure have been conducted using fingerling life stages of fishes, and no data are available on the comparative sensitivity of early life stages (e.g., fry). To fill this data gap, the 77 purpose of this study was to determine if the sensitivity of Brook trout fry is similar to fingerlings, and to assess the impacts of tire wear contaminant exposure on blood chemistry and gill histology to determine a potential mechanism of action for toxicity. Both 6PPD-quinone 80 and HMMM exposures were conducted, as 6PPD-quinone exposure at low concentrations have 81 been shown to cause mortality to Brook trout fingerlings(Brinkmann et al., 2022), and HMMM 82 has been measured in storm water runoff events near urban centers alongside 6PPD-quinone, with limited published toxicity data available. The data generated in the study will be used to 84 better understand the potential risk of these contaminants in Brook trout habitat in close 85 proximity to roadways and urban centers. d.

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⁸⁶ **3. Materials Methods**

⁸⁷ *Organisms*

88 Brook trout (also called Brook char) fry and fingerlings were obtained from a commercial 89 supplier (Brittany Hill Farms, Seeleys Cove, NB) in July 2022. Fish were acclimated at the ⁹⁰ Huntsman Marine Science Centre (St. Andrews, NB) in dechlorinated municipal freshwater for 91 2-3 weeks prior to the launch of any toxicity tests. During the acclimation and holding period, 92 fish were fed a commercial fish food diet and were visually screened daily for disease and 93 deformities. Fry were fed EWOS® #1 crumble (Cargill Incorporated, Surrey, BC, Canada) and 94 were transitioned to 1.0 mm Nutra Spirit produced by Skretting (Vancouver, BC, Canada). The 95 fingerlings were fed a mixture of EWOS 3.0mm Transfer and Skretting's 3.0mm Nutra Olympic 96 feed. Fish were held on a 15:9 light: dark cycle for the duration of the acclimation period. All 97 holding and care for the organisms prior to the trial and exposures were conducted according 98 to the Department of Fisheries and Oceans Animal Care Committee protocol AUP 22-26. The ⁹⁹ negative control validity criteria for survival in all of the exposures was >80%, and this threshold ¹⁰⁰ was met and exceeded for all the experiments conducted in this study as there were no control ¹⁰¹ mortalities in any of the trials. of the launch of any toxicity tests. During the acclimation and
mmercial fish food diet and were visually screened daily for
were fed EWOS® #1 crumble (Cargill Incorporated, Surrey, B
to 1.0 mm Nutra Spirit produced by Skr

¹⁰² *Test materials and exposure methods*

¹⁰³ HMMM was acquired from a commercial supplier (95% purity; Alfa Chemistry, Ronkonkoma, 104 NY, USA). The 6PPD-quinone used in this study was synthesized following the methods ¹⁰⁵ described in Monaghan et al(Monaghan et al., 2021). Stock solutions of both HMMM and 6PPD-106 quinone were prepared using dimethyl sulfoxide (DMSO) and stored frozen at -20°C until use.

151 second trial were collected from all 3 replicates of the highest exposure concentration at both 152 TO and T24 to determine the inter-replicate variability in our trial. Dissolved oxygen, pH, alkalinity, hardness, temperature, and ammonia concentrations were measured at T0 in one barrel, and at T24 in a replicate of each treatment. Fish were assessed for morbidity and mortality using the same criteria described in the fry trials at 1, 3, and 24hrs of exposure in both trials.

3.2.2.1. Blood metrics and gill histology

 The blood and gills were sampled from a subset of individuals in the fingerling exposures. Any surviving fish at the end of the 24hr exposure were euthanized using an overdose of TMS 160 (tricaine methanesulfonate, Syndel, Nanaimo, BC; 400 mg/L), in a staggered manner to ensure an equal time between euthanizing and blood collection. Arterial blood was then collected using a 20-gauge needle inserted into the caudal artery and collected directly into a lithium- heparinized vacutainer. After collection, blood was gently swirled to minimize clotting in the vacutainer. Following blood collection, 100-200μL of blood was loaded into an i-STAT blood 165 analyzer CHEM8+ cartridge (Abbott Point of Care Inc., Union City, CA, USA) using a 1mL syringe. 166 The CHEM8+ cartridge was used to measure sodium, chloride, total $CO₂$, Anion gap, ionized 167 calcium, glucose, urea nitrogen, creatinine, hematocrit, and hemoglobin concentrations in freshly collected blood samples (detection limits are available in **Supplemental Table 1**). The cartridges and analyzer were used according to the manufacturer's specifications, and cartridges have previously been used with fish blood samples to measure ion 171 dysregulation(Chow et al., 2019). Blood glucose was also measured in a larger subset of individuals using the Contour Next Glucose Monitoring System (Bayer, Mishawak, IN, USA), and I. Blood metrics and gill histology
Is were sampled from a subset of individuals in the fingerlin
he end of the 24hr exposure were euthanized using an over
ssulfonate, Syndel, Nanaimo, BC; 400 mg/L), in a staggered
tween e

Analytical chemistry

 Water chemistry samples were collected in 1L polyethylene terephthalate (PET) bottles, frozen immediately after collection, and stored at -20°C. Details on the calibration methods, running

(1) $LC_{50}(t) = LC_{50,} \approx [1-\exp{(-\varepsilon t)}]^{-1}$

214 The non-linear regression model in the package calculates the incipient value (i.e., LC50, ∞)

where t describes the increasing exposure duration in units of hours, and ε describes the rate at

which the organism accumulates damage/repairs in units of hr⁻¹. The same equation was

- 217 applied to the LT50 data to determine the incipient time to lethality (LT50∞), which represents
- the minimum amount of time needed to observe 50% mortality, independent of concentration.

4. Results

Exposure Characterization

4.1.1. Water quality

 Dissolved oxygen remained between 87 – 101 % air saturation in all the treatment groups 223 tested. There were also no treatment or trial specific differences in pH $(6.78 - 8.12)$; which varied between trial and treatments likely due to ammonia), water hardness (12 – 15 mg/L), 225 and alkalinity (16 – 24 mg/L). The water temperature in the fry trial vessels (11.1 – 12.0 °C) was slightly cooler than our fingerling trials (12.0 – 14.8°C) both pre- and post-exposure. Ammonia levels varied considerably between treatment groups and trials (0 – 0.53 mg/L), however all ammonia measurements were below ambient water quality criteria(United States Environmental Protection Agency, 2013). Expanding the characterization

ater quality

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reads no treatment or trial specific differences in pH (6.78

rial and treatments likely due to ammonia), water ha

4.1.2. Water chemistry

 Due to the limit of quantification of the analytical technique used in this study, and the response of the Brook trout, only the highest exposure concentrations were quantified. The

HMMM exposure concentrations remained consistent (~7% decrease) in the test vessel without

fish over the course of the 24hr exposure, however, in the surrogate vessel containing fish

 Figure 1: Concentration response relationship for fry (blue circles) and fingerling (orange triangles) Brook trout following 3 (A) and 24-hrs (B) of exposure to 6PPD-quinone. LC50 values are shown as the vertical lines and presented with the standard errors beside the curves.

4.2.2. Incipient Lethal Level

280 The incipient LC50 (LC50∞) represents the concentration at which the LC50 reaches an

asymptote such that there is no additional change in the value for additional exposure duration

 Figure 2: LC50 over time (A) and LT50 (B) for the fry (blue circles, dashed line) and fingerling (orange triangles, solid line) exposed to 6PPD-quinone. The incipient lethal values (and standard error) and the corresponding epsilon (ε) values for each value and life stage are given in the insert.

Fingerling Blood Chemistry

- 316 that survived the duration of the 24hr test, which may suggest that the sublethal effects on
- 317 blood parameters observed here could be more severe in the mortalities that occurred due to
- exposure.

 Figure 3: Impact of 6PPD-quinone exposure on the hematocrit (A), glucose (B), total CO² (C), chloride (D), and sodium (E) levels in Brook trout fingerling blood after 24hrs. Responses were normalized to control, and the data presented are pooled from both fingerling trials as there were no trial-specific differences in response. The solid horizontal line represents the mean of the control response, and the

- *dashed horizontal lines represent the range in control values ± 2 standard deviations. The Pearson*
- *correlation coefficient (R) and p values for the linear relationships presented are included on each of the*
- *corresponding panels. Shading represents the 95% confidence band around each relationship.*

Histology

- There was a significant increase in the mean relative interlamellar cell mass (ILCM) size in the
- 330 0.5 μ g/L exposed fingerlings (21.1 ± 9.1% SD, n = 10) compared to controls (13.9 ± 4.5%, n = 10)
-

 $(p = 0.045; \text{ Figure 4}).$

- *Figure 4: Lamellar length (longer line) and interlamellar cell mass (ILCM) height (shorter line) in*
- *representative micrographs from representative control (A) and 0.5 µg/L 6PPD-quinone exposed (B) fish.*
- *ILCM sizes are 11 and 39% of lamellar length for upper and lower micrographs, respectively; scale bar =*
- *100 μm.*

337 The ILCM values from the individual fish were not significantly correlated with fish size (total length and body mass) or hematocrit, however, they were significantly correlated with numerous blood plasma endpoints (**[Figure](#page-23-0)**). Increases in blood glucose (**Figure 5A**) and total CO₂ (Figure 5B) correlated with increases in relative ILCM height (R = 0.64, p = 0.018, and R = 0.7 and p = 0.011, respectively). Increases in relative ILCM height also correlated with decreases in blood sodium (**Figure 5C)** and chloride (**Figure 5D**) at 24hrs of exposure (R = -0.67, p = 0.013, and R = -0.58 and p = 0.037, respectively).

 Figure 5: Correlation of ILCM with significant blood chemistry parameters, glucose (A), total CO² (B), sodium (C), and chloride concentrations (D). The open circles are control fish, and the filled circles are fish that were exposed to 0.5 µg/L 6PPD-quinone for 24-hours. The Pearson correlation coefficient (R) and p values for the linear relationships presented are included on each of the corresponding panels. Shading represents the 95% confidence band around each relationship.

5. Discussion

 Since the discovery of 6PPD-quinone as the contaminant predominantly responsible for urban runoff mortality syndrome there has been a push to test the sensitivity of other fishes to tire wear particles, leachate, and compounds (e.g., HMMM). HMMM exposure caused no effects to fry Brook trout at concentrations as high as 6.6 mg/L, suggesting that acute toxicity may be attributed to only specific tire leachate contaminants. Brook trout fingerlings were considered one of the most sensitive species/life stage exposed to 6PPD-quinone to date (Brinkmann et al., 2022), and our fry and fingerling Brook trout data support these results. Despite the differences between our study and Brinkmann et al.(Brinkmann et al., 2022) in exposure set up (208-L steel drums lined with BPA-free low density polyethylene bags vs. 150-L fiberglass tank), analytical chemistry methods (LC-MS/MS vs. UHPLC-MS/MS), fish stock (Seeleys Cove, NB vs. Coleman, AB), and sample size (three replicates with n = 10 vs. two replicates with n = 4), our results are 362 in very good agreement (24-hr LC50s of 0.5 and 0.59 µg/L in this study and Brinkmann et al.(Brinkmann et al., 2022) respectively) which supports the use of both exposure methods in generating comparable toxicity data. Our study expanded on the results presented in Brinkmann et al.(Brinkmann et al., 2022) by including Brook trout fry, and our study indicates 366 that data collected from fingerling sized fish alone may underestimate the species sensitivity. In future studies, additional testing on earlier life stages may be required to ensure the data generated is protective of not only the fingerling life stage, but also more sensitive life stages like the fry . r specific tire leachate contaminants. Brook trout fingerling:
ensitive species/life stage exposed to 6PPD-quinone to dat
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 (0.09 µg/L) of measured values. Peak exposure concentrations of 6PPD-quinone have been associated with storm water runoff(Challis et al., 2021; Johannessen et al., 2022b), which suggests exposures are more likely to be transient and linked with brief, pulsed, dynamic run-416 off events. The metrics determined through the application of these models to our LC and LT50 values provide critical inputs needed to understand the impact of exposure duration on observed effects with this novel contaminant.

419 Blood chemistry parameters were only measured in fish which survived the 24hr exposure. ⁴²⁰ Dysregulation of ion transport was evident in 6PPD-quinone fish in a concentration dependent 421 manner. This same effect has been observed in coho salmon exposures to stormwater runoff 422 (Chow et al., 2019; McIntyre et al., 2018). The dose-dependent decrease in concentration of the ⁴²³ dominant blood ions (sodium and chloride) in our 6PPD-quinone-exposed fish indicates a 424 progressive loss of ability to maintain stable plasma ion levels, i.e., osmoregulatory distress, and 425 is consistent with results observed in coho exposed to tire leachate (with up to 2.4 µg/L of ⁴²⁶ 6PPD-quinone) in McIntyre et al. 2021(McIntyre et al., 2021). A concomitant increase in blood 427 glucose demonstrates that the fish were trying to deal with this by mobilizing energy reserves 428 to increase aerobic metabolism. Freshwater fish are hyperosmotic and therefore constantly ⁴²⁹ losing ions by passive diffusion across any permeable surfaces in contact with the water. ⁴³⁰ Maintaining a stable osmotic gradient and ion profile requires them to actively osmoregulate, ⁴³¹ accounting for a significant proportion of their resting metabolism, a situation that becomes ⁴³² less tenable when stressed and can ultimately lead to metabolic exhaustion and death. The 433 increase in hematocrit and blood glucose due to 6PPD-quinone exposure observed in this study ⁴³⁴ has been previously demonstrated with Rainbow and Brook trout(Brinkmann et al., 2022), and with this novel contaminant.
Darameters were only measured in fish which survived the
ion transport was evident in 6PPD-quinone fish in a concen
ne effect has been observed in coho salmon exposures to st
9; McIntyre et al.

 The increased size of the interlamellar cell mass (ILCM) in the surviving 6PPD-quinone-exposed 440 fish provides further evidence of osmoregulatory distress. This remodeling of the gills slows 441 passive diffusion of ions between the blood perfusing the lamellae and the water passing between them (Wood and Eom, 2021), and here may represent a compensatory response to limit ion loss in these surviving fish. Gill lamellae need a very large and permeable surface area to support respiratory gas exchange, but this then makes them a primary site for the loss of 445 ions by diffusion and uptake of water by osmosis in freshwater fish. Healthy fish counter this 446 through active osmoregulation. When this poses a serious metabolic challenge, they can 447 decrease the amount of permeable surface area in contact with water by increasing the size of the protective ILCM. However, this also reduces the functional surface area for gas exchange to support aerobic respiration, an outcome commonly referred to as the 'osmorespiratory compromise' (Wood and Eom, 2021). The fish are now faced with increased oxygen demand (i.e., stress response) but a reduced ability to extract oxygen from the water. Our finding that 452 ILCM size increased in direct proportion to the changes in blood sodium, chloride, glucose, and CO₂ levels, taken together with observations of gasping behaviors, gill flaring, and erratic swimming before the onset of mortality within 3-6 hours of exposure, provides compelling evidence for an inability to meet the aerobic demands while attempting (and failing) to osmoregulate adequately when exposed to 6PPD-quinone. The results of this study highlight her evidence of osmoregulatory distress. This remodeling o
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nese surviving fish. Gill lamellae need a very la

- the importance of toxicity testing with early life stages of sensitive species and potential
- mechanism of action for toxicity for this novel contaminant.

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Highlights

- Brook trout fry and fingerlings were exposed to 2 different tire-wear contaminants.
- No effects were observed in HMMM fry exposures.
- Fry were 2-3x more sensitive to 6PPDq than the fingerling life stage.
- 6PPDq exposure resulted in concentration dependent changes in blood chemistry.
- 6PPDq changed the gill morphology, resulting in osmorespiratory compromise.

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Declaration of interests

 \boxtimes The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

 \Box The authors declare the following financial interests/personal relationships which may be considered as potential competing interests:

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