SURVIVAL TIME AND HISTOLOGICAL OUTCOME OF MAJOR ORGANS FOLLOWING HONEY ADMINISTRATION TO ACUTE PARAQUAT-INTOXICATED RATS: A PRELIMINARY STUDY

SUK PENG TANG, HASNAN JAAFAR, SIEW HUA GAN, KUTTULEBBAI N. S. SIRAJUDEEN, SITI AMRAH SULAIMAN

Objective: The objective of this research was to investigate the possible protective effect of Tualang honey (TH) in acute paraquat (PQ) toxicity in rats.

Methods: A total of 48 male Sprague-Dawley rats aged eight weeks old were used. Oral PQ and TH were administered at 225 mg/kg and 0.2 g/kg, respectively. The effects of single and multiple TH treatments on PQ-intoxicated rats were then investigated. Single TH treatment groups received TH at 0.5 (PQ+TH0.5h), 2 (PQ+TH2h) or 6 (PQ+TH6h) hours following PQ administration. Multiple TH treatment groups received TH at 0.5, 2 and 6 h (PQ+THtrp) or further daily treatment for the following six days (PQ+TH7d) after PQ administration (n=6 per group). The survival time of the rats was recorded until day 28 before sacrifice, which was followed by a histological examination.

Results: Treatment with TH did not improve the survival rate of PQ-intoxicated rats. However, the median survival time of rats that received multiple TH treatments was significantly longer compared to that of the PQ+TH6h group. TH treatment was found to improve the histological outcomes of PQ-intoxicated rats, particularly in the lungs.

Conclusion: Our findings suggest the potential role of honey in delaying the toxic effects of PQ.

Keywords: Paraquat, Acute toxicity, Tualang honey

Although paraquat (PQ) is a widely used herbicide, the challenges resulting from PQ exposure are reported all over the world and are mainly caused by suicidal intent, accidental poisoning or occupational exposures [1]. PQ is known to stimulate the production of various reactive oxygen species (ROS) via a single electron redox cycle in vivo [2, 3]. ROS readily attack key cellular structures and molecules, thus causing cellular deleterious effects that form the basis of various disease conditions [4, 5].

To date, there is no specific antidote clinically available for PQ poisoning. The conventional approach in treating PQ poisoning focuses on three main areas, which include the prevention of absorption from the gastrointestinal tract, the enhancement of elimination of PQ from the body and the administration of therapies directed against toxicity. Nevertheless, these treatment methods have been disappointing, with very high mortality rates [1, 6-9]. Many studies have been conducted in an effort to search for an effective antidote for PQ poisoning with several being directed towards the use of antioxidants because PQ induces its toxic effect via oxidative stress-mediated mechanisms [10-14].

Honey, which is one of the oldest foods consumed over the centuries with potential therapeutic properties because of its high antioxidant capacity, has gained substantial research interest in the past decade [15]. Honey contains both aqueous and lipophilic antioxidants, and interaction between these antioxidants suggests honey’s potential as an ideal natural antioxidant that can act at various cellular sites in the case of PQ poisoning [16, 17]. Tualang honey (TH) is a wild honey harvested from Tualang trees found in many Asian rainforests. Various studies have been conducted in an effort to evaluate the possible medicinal uses of TH [18-24]. One of the proposed therapeutic values is attributed partly due to its antioxidant properties. A study by Mohamed et al. [25] showed that TH contains phenolic compounds with strong antioxidant activities. However, to date, the protective effects of honey on PQ toxicity have not been investigated. Therefore, the aim of this study is to investigate the potential beneficial effects of TH in acute PQ poisoning in an animal model.

TH (AgroMas®) used in this study was supplied by the Federal Agricultural Marketing Authority (FAMA), Kedah, Malaysia. The honey samples used in this study were filtered, concentrated to 20% (w/v) water content at 40 °C and sterilized by gamma irradiation at 25 kGy [Sterilgamma (M) Sdn. Bhd., Selangor, Malaysia]. The same batches of honey were used throughout the analysis. The chemical characteristics including the antioxidant properties of the investigated honey have been previously described [25].

Male Sprague-Dawley rats (n=48) aged eight weeks old were purchased from the Animal Research and Service Center, Health Campus, USM, Kubang Kerian, Kelantan, Malaysia. All animals were individually housed in polypropylene cages in a well-ventilated room maintained at a 12 h light/dark cycle and 25±2 °C room temperature. Food pellets and water were given ad libitum unless otherwise stated. All rats were acclimatized to the room condition for at least 1w prior to the experiment. This study was approved by the Animal Ethics Committee, Universiti Sains Malaysia [Approval No.: USM/Animal Ethics Approval/2009/(145)[140]], which is in accordance with the Institutional Guideline for the Care and Use of Animals for Scientific Purposes.

The rats were randomly divided into eight groups of six rats each. Groups A and B were administered 0.5 ml of double deionized water (ddH2O), whereas Groups C, D, E, F, G and H were administered PQ (Sigma-Aldrich, St Louis, MO, USA; 225 mg/kg, p.o.) at the beginning of the study (t=0). The rats were then treated with either ddH2O or TH (0.2 g/kg, p.o.) at the specific time intervals described in table 1. The doses of PQ and TH were determined based on preliminary studies conducted in our laboratory (data not shown).
with ketamine (90 mg/kg) and xylazine (5 mg/kg) at the end of the experimental period. The lungs, kidneys and liver of all rats (moribund and surviving) were collected and fixed in 10% formalin solution. They were then processed for subsequent histopathological analysis by hematoxylin and eosin (H and E) staining.

Kaplan-Meier survival analysis with Log-Rank pairwise comparison test was performed using the IBM Statistical Package for the Social Sciences (SPSS) software version 22.0 (Armonk, NY, USA) to evaluate the median differences between the experimental groups.

This study is the first to report on the effect of honey administration on PQ intoxication in rats. Table 2 shows the effects of administering various honey dosing regimens on the median survival time of PQ-intoxicated rats. Overall, TH treatment did not improve the survival rate of PQ-intoxicated rats, regardless of the time of administration, with mortality rates remaining high (>60%). However, the rats that received triple honey treatments at 0.5, 2 and 6 h (group PQ+THtrp) or continuous daily honey treatment for seven days (group PQ+TH7d) following PQ administration had a significantly longer median survival time compared to animals that received only a single honey treatment at six hours following PQ intoxication (group PQ+TH6h). In comparing the groups that received only single honey treatments (at 30 min, 2 h or 6 h), there was a trend in which the earlier the animals were administered the honey treatment, the longer the observed median survival time.

### Table 1: Rat grouping and type of treatment in all experimental groups

<table>
<thead>
<tr>
<th>Group</th>
<th>Treatment received after PQ administration*</th>
</tr>
</thead>
<tbody>
<tr>
<td>A (N)</td>
<td>ddH2O at 2 h</td>
</tr>
<tr>
<td>B (TH)</td>
<td>TH at 2 h</td>
</tr>
<tr>
<td>C (PQ)</td>
<td>ddH2O at 2 h</td>
</tr>
<tr>
<td>D (PQ+TH0.5h)</td>
<td>TH at 0.5 h</td>
</tr>
<tr>
<td>E (PQ+TH2h)</td>
<td>TH at 2 h</td>
</tr>
<tr>
<td>F (PQ+TH6h)</td>
<td>TH at 6 h</td>
</tr>
<tr>
<td>G (PQ+THtrp)</td>
<td>TH at 0.5, 2, and 6 h</td>
</tr>
<tr>
<td>H (PQ+TH7d)</td>
<td>TH at 0.5, 2, and 6 h and once daily for the subsequent 6 d</td>
</tr>
</tbody>
</table>

*Groups A and B (non-PQ treated) were administered ddH2O (0.5 ml, p. o.), whereas groups C, D, E, F, G and H (PQ-treated) were administered PQ (225 mg/kg, p. o.) at t=0. The dose of TH used was 0.2 g/kg, n=6 per group. [Abbreviations: ddH2O, double distilled water; PQ, paraquat; TH, Tualang honey]

All rats were closely observed for a maximum period of 28 d before being sacrificed. The survival time after PQ administration of each rat was recorded until day 28. All moribund animals or animals showing severe signs of toxicity were sacrificed. All surviving animals were sacrificed by decapitation under deep anaesthesia.

### Table 2: The effects of administering various honey dosing regimens on the median survival time of PQ-intoxicated rats

<table>
<thead>
<tr>
<th>Group</th>
<th>Survival (%)</th>
<th>Median (95% CI) (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PQ+TH0.5h</td>
<td>16.7</td>
<td>3.50 (2.53, 4.47)</td>
</tr>
<tr>
<td>PQ+TH2h</td>
<td>16.7</td>
<td>4.00 (1.84, 6.16)</td>
</tr>
<tr>
<td>PQ+TH6h</td>
<td>16.7</td>
<td>3.10 (1.82, 5.02)</td>
</tr>
<tr>
<td>PQ+THtrp</td>
<td>16.7</td>
<td>2.50 (1.78, 3.22)</td>
</tr>
<tr>
<td>PQ+TH7d</td>
<td>16.7</td>
<td>4.50 (2.10, 6.90)**</td>
</tr>
</tbody>
</table>

*p<0.05; **p<0.01 compared to group PQ+TH6h (Log Rank pairwise comparison)
congested liver sinusoids [fig. 3(A)]. Additionally, there was an increase in the number of binuclear hepatocytes in all PQ-intoxicated groups, but the increment is comparatively smaller in groups that received multiple honey treatments. Similar histological changes were observed in the liver of rats found dead or sacrificed earlier within four to seven days (96-168 h) following PQ intoxication [fig. 3(B)]. In the rats that survived until the end of the experimental period (28 d following PQ-intoxication), no evidence of major histological changes was observed among the experimental groups [fig. 3(C)]. Liver injury may promote regenerative activity, which is relatively rare in healthy adult hepatocytes [28]. The increase in the major dividing cell type during liver regeneration may be a result of the cellular response to PQ-induced hepatotoxicity. PQ-induced karyomegaly and binucleation in the mouse liver was previously shown in caspase-2 knockout mice that had a reduced ability to withstand oxidative stress and were more susceptible to various oxidative stimuli [29].

Altogether, these findings suggest that treatment with TH may reduce the toxic actions of PQ. Further investigation of the oxidative stress parameters such as the enzyme activities of superoxide dismutase, catalase, and glutathione peroxidase, as well as malondialdehyde content, should be conducted to elucidate the exact mechanisms. Nevertheless, the individual rats responded differently to PQ, with the earliest fatalities recorded at 10 h post PQ intoxication, followed by within days and 1 w, with some animals even surviving until the end of the experimental period. The inter-individual differences in responses made comparison between the rats a challenge. Additionally, the histological findings based on early death (< 96 h), delayed death (up to 1 w) and survival for the experimental period represented a small number of rats in each experimental group. Therefore, the relatively smaller sample size in this study may not reflect the actual response to PQ or TH. Moreover, although TH seemed to prolong the rats’ survival time to a certain extent and improved the histological outcomes compared to animals that received PQ only, the mortality rate remained high. Therefore, a new therapeutic approach is still needed in which the use of honey can be incorporated with other therapeutic agents (i.e., honey as an adjunctive treatment).

Fig. 1: Representative photomicrographs of H and E stained lung section from rats sacrificed or found dead (A) within four days (≤ 96 h), (B) within four to seven days (96-168 h) and (C) at the end of the experimental period (day 28) following PQ intoxication (scale bar: 200 µm; original magnification: ×100). The major histopathological changes include distorted alveolar configuration or large distended air spaces (*), vascular congestion and pulmonary edema (alveolar filled with pink/pale-eosinophilic transudate), diffuse alveolar wall thickening (thick arrow) with interstitial inflammation and edema, infiltration of inflammatory cells in the alveolar space and mild congestion (double arrow). [ad, alveolar duct; bv, blood vessel; PQ, paraquat group; TH, Tualang honey (single-dose honey treatment at 0.5 (TH0.5h), 2 (TH2h) or 6 h (TH6h); multiple-dose honey treatment at 0.5, 2 and 6 h (THtrp) or 7 d of consecutive treatment (TH7d)]

Fig. 2: Representative photomicrographs of H and E stained kidney section from rats sacrificed or found dead (A) within four days (≤ 96 h), (B) within four to seven days (96-168 h) and (C) at the end of the experimental period (day 28) following PQ intoxication (scale bar: 50 µm; original magnification: ×400). The major histopathological changes include features of acute tubular necrosis [pyknotic nuclei (P), loss of nuclei (arrow), or vacuolated cytoplasm (V)], tubular protein cast (#) and mild congestion (double arrow) were observed in the PQ-intoxicated rats with or without honey treatments. [G, glomerulus; L-tubular lumen; PQ, paraquat group; TH, Tualang honey (single-dose honey treatment at 0.5 (TH0.5h), 2 (TH2h) or 6 h (TH6h); multiple-dose honey treatment at 0.5, 2 and 6 h (THtrp) or 7 d of consecutive treatment (TH7d)
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CONFLICT OF INTERESTS

The authors declare that there is no conflict of interest regarding the conduct of this study.

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Overall, although treatment with single or multiple doses of TH at various times of administration failed to improve the survival rate in acute PQ-intoxicated rats, it may reduce or delay the toxic effects of PQ as evidenced by the post-mortem histological analyses in the major organs in the honey-treated groups.

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