**Abstract**

This research investigated the reno-protective effect of *Thunbergia laurifolia* Linn. (TL) in a lead-induced toxicity test through the modulation of cell signaling pathways. The study carried out to evaluate the effect of TL leaf extracts in Swiss Albino mice exposed to lead acetate (PbAc). Prior to in vivo study, a probable kidney-protective effect of the plant leaf extract was presumed through an activity-specific (PASS) molecular docking analysis. In animal model study, albino mice were divided in seven groups and co-treated with PbAc and TL (100, 200 mg/kgBW) or vitamin E (100 mg/kgBW) for 38 days, whereas the untreated control, TL control, and vehicle control groups received sodium acetate, PbAc, sodium acetate plus mineral oil, respectively. At the end of treatment, blood and kidney tissue were collected for investigating Pb concentration, estimating biochemical profile, evaluating oxidative stress and inflammatory parameters. The histopathological change of kidney along with apoptosis was assessed from kidney sections using H & E staining and TUNEL assay. Pb-exposed mice were found to be increased concentration of Pb in the blood and kidney sample, which further led to increased MDA levels in the plasma, blood, and tissue. Followed by kidney damage, increased expression of TNF-α, iNOS, and COX-2 in kidney tissues were noticed, which were related to elevated TNF-α in the systemic circulation of Pb-treated mice. Co-treatment with TL or vitamin E significantly reduced altered structure and apoptosis of kidney tissues. Downregulation of inflammatory markers especially TNF-α, iNOS, and COX-2 with simultaneous improvement of renal function through reduced plasma BUN and creatinine levels demonstrate that TL act as a potential dietary supplement to detoxify Pb in kidney showing an antioxidant and anti-inflammatory effect.

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**1. Introduction**

Lead (Pb) poisoning is an occupational health hazard especially in the third world and industrializing countries (Azeh Engwa et al., 2019). Widespread usage of Pb in different industries including automobiles, paints, ceramics, plastics, and similar industries makes its abundance in the environment to cause environmental pollution (Centers for Disease control and Prevention, 2020). It can exist for a long time in gasoline, industrial processes such as Pb smelting coal combustion, Pb-based paints, Pb containing pipes, battery recycling, grids, and bearings due to its non-degradability. Intoxication with Pb followed by deposition in the body is related to different diseases such as neurotoxicity or neurodegenerative diseases, immunological disorders, gastrointestinal diseases, hematological diseases, and reproductive complexity, etc.
of cytosolic Ca\textsuperscript{2+} also evidenced with Pb-induced kidney toxicity (Rana et al., 2010, 2018). Moreover, the hindrance enhanced lipid peroxidation, reduced antioxidant activity, and kidney dysfunction (Rana et al., 2020, 2018). Therefore, the development of therapeutics from alternative sources could meet the growing demand to overcome the Pb-associated health complexity.

Thunbergia laurifolia Linn. (TL, Acanthaceae) is a trumpet vine, widely distributed in Southeast Asian countries and native to India (Kanchanapoom et al., 2002). Antidote properties of TL have made it well known in Thailand, where dried leaves, dried roots, fresh leaves, and bark are used as traditional medicine (Rocejanasaroj et al., 2014). An earlier study reported that TL is active against pesticides, arsenic, and strychnine toxicity (P. Tejasen and Thongthapp, 1986). Moreover, a study from our lab unveiled that TL is effective against the cognitive dysfunction of mice with Pb exposed (Phyu and Tangpong, 2013). Tangpong and Satarug 2010 reported that TL act as a chelating agent against neuronal cell death and memory loss in Pb exposed mice (Tangpong and Satarug, 2010). According to the accumulated evidences, TL also possesses a wide range of pharmacological properties, for example, anti-mutagenic, anti-inflammatory and antipyretic properties along with antioxidant, anti-inflammatory, antimicrobial, neuroprotective, anti-proliferative, hepatoprotective, and antidiabetic activity (Chan et al., 2011; Tangpong and Satarug, 2010; Wonchalee et al., 2012). Furthermore, it is also used in the treatment of drug addiction (Chan et al., 2011).

Currently, there is no existing study and insight mechanisms on the TL effect against Pb induced kidney toxicity in the mice model. Thus, this study was designed to investigate the role of TL against lead acetate (PbAc) induced renal toxicity and its underlying mechanism.

2. Materials and methods

2.1. In silico study

2.1.1. Prediction of activity spectra (PASS) analysis

To predict the probable kidney protective effect of TL, a well-known computer-aided tool called “prediction of activity-specific for substances or PASS” program was used. PASS can predict a widespread spectrum of biological activity based on the structural formula of the organic substances. The 2D structure (SDF file) of identified phytochemical constituents of TL leaves (Junsi and Siripongvitukorn, 2016) such as α-Spinosterol (PubChem CID 5281331), β-Sitosterol (PubChem CID 222284), Caffeic acid (PubChem CID 689043), Protocatechuic acid (PubChem CID 72), Gallic acid (PubChem CID 370), Apigenin (PubChem CID: 5280443), Giraffone acid (PubChem CID: 159930), Lutein (PubChem CID: 5281243), 3′-O-β-D-glucopyranosyl-stilberricoside (PubChem CID: 6325264), Stigmasterol (PubChem CID: 5280794), Benzyl β-glucopyranoside (PubChem CID: 11076492), 6-C-glucopyranosyl apigenin (PubChem CID: 442658), 6,8-di-β-D-glucopyranosyl apigenin (PubChem CID: 3084407), 3-Hexenyl-beta-D-glucopyranoside (PubChem CID: 5318046), Hexanol beta-D-glucopyranoside (PubChem CID: 6428285) were uploaded in PASS online (way2drug).

The output of probable activity was presented as Pa (active) and Pi (inactive). The output of probable activity was presented as Pa (active) and Pi (inactive) (Filimonov et al., 2014). Here, Pa > Pi was considered to be an active compound and Pa greater than 0.3 were believed to be experimentally effective.

2.1.2. Molecular docking study

To predict the insight of the protective mechanism, binding efficiency of phytoconstituent to IP3R (PDB:3UJO) that contributes to [Ca\textsuperscript{2+}] dysregulation (Wang et al., 2015), to activate Nrf-2 following release from KEAP1 (PDB:5CGG) during oxidative stress were studied. Besides, the interaction among phytoconstituents and inflammatory mediators including TNF-α (PDB: 2AZ5), INOS (PDB: 3E67), COX-2 (PDB: 5KIR), and proteins of MAPK pathways P38 Kinase (PDB: 2YIS), JNK (PDB: 3OY1), and NF-kB activity regulator, IkB kinase-β (PDB: 4KIK) were screened. Published crystal structures of aforementioned protein as PDB format were imported into Mastro software (Scodinger v11). To process the proteins for docking study, proteins preparation wizards were imported to receptor grid generation by applying OPLS5 force field and 0.30 Å was set for the maximum heavy atom root-mean-square-deviation (RMSD).

To process the ligand, 2D structures of water soluble phytoconstituents, such as Caffeic acid (PubChem CID 689043), Protocatechuic acid (PubChem CID 72), Gallic acid (PubChem CID 370), were converted to 3D structure using the ligprep wizards (Scodinger v11) with a default setting of OPLSS force field (pH 7.0 ± 2.0, allow 32 stereoisomers per ligand). After that, prepared proteins were processed to receptor grid generation by applying OPLSS force field, and the ligands were allowed to dock flexibly using the ligand docking module. The resultant scores were extracted as an excel file, and the lowest docking score was considered as the best docking.

The drug likeness properties of docked phytoconstituents were analyzed by the QikProp module of Schrodinger. Briefly, the 3D structures processed during the ligand preparation were input into QikProp wizard and then analyzed with the default setting. A compound was considered to possess characteristics resembles the available drug, if it met any three parameters out of four. The parameters were molecular weight < 500, hydrogen bond donor < 5, hydrogen bond acceptor < 10, and predicted octanol/water partition coefficient (QLogP) value (-)2.0 – 6.5.

2.2. In vivo study

2.2.1. Preparation of TL leaves extract

TL leaves (10 kg) were collected from Nakhon Si Thammarat, Thailand, during the month between April to May, and was identified by an expert botanist. A voucher specimen of the sample has been preserved in the University Herbarium with the accession number BKF 186138. The leaves were oven-dried at 60 °C for 4 h. Dried leaves were pulverized (Philip, Germany), soaked into water and autoclaved at 121 °C for 15 min. Afterward, the suspension was kept in dark at 4 °C for 2 days, filtered using Whatman No. 1, and then centrifuged at 4000 rpm for 15 min. The supernatant was collected and was kept at −40 °C for 12 h before lyophilization (EYELA, Tokyo, Japan). The yield value of TL extract was 9.65% in powder form and stored at −30 °C for further use.

2.2.2. Animals and experimental design

Forty eight-week-old ICR male mice (30–33 g) were purchased from National Laboratory Animal Center, Salaya district, Nakhon Pathom, Thailand. These mice were housed at polypropylene cages in the animal center of Walailak University, Thailand. The mice were maintained under controlled environmental conditions (22 ± 2 °C, 12:12 h light:dark cycle) and provided with food and water ad libitum.
were allowed to access standard food and water ad libitum for 2 weeks followed by standard laboratory conditions such as temperature (23 ± 2 °C), humidity (50 ± 5%), and light/dark cycle 12 h. To evaluate the effect of TL against Pb-induced nephropathy, mice were treated according to Phyu and Tangpong 2013, where Pb significantly impaired cognitive behavior and oral administration of TL extracts markedly improved cognitive function (Phyu and Tangpong, 2013). In this study, the experiments were randomly divided into seven independent groups (n = 6) and were continued for 38 days. The treatments strategies in different groups were as follows:

**Group 1:** Untreated Control, (received only 1% CH3COONa in drinking water).

**Group 2:** PbAc (1% in DW) (received only 1% PbAc in drinking water).

**Group 3:** TL 200 mg/kgBW (received only 200 mg/kgBW of TL water extract; P.O.).

**Group 4:** Pb + TL 100 mg/kgBW (received 1% PbAc in drinking water + 100 mg/kgBW of TL water extract; P.O.).

**Group 5:** Pb + TL 200 mg/kgBW (received 1% PbAc in drinking water + 200 mg/kgBW of TL water extract; P.O.).

**Group 6:** Pb + Vit-E 100 mg/kgBW (received 1% PbAc in drinking water + 200 mg/kgBW of Vit-E in mineral oil, P.O.).

**Group 7:** Vehicle Control (received mineral oil, P.O.).

Here, P.O (Per os) dictates the oral administration by gavage.

### 2.2.3. Sample collection

The mice were anesthetized using Nembutal sodium (65 mg/kg BW) and sacrificed via left ventricle puncture. Blood was obtained and then collected in a 2.5 mL K3EDTA containing tube. Afterwards, a portion of the right kidney was removed and homogenized (Sonics VCX70, USA) in a mixture of cold PBS and cocktail of protease inhibitors (leupeptin, pepstatin, and aprotinin) prior to centrifugation at 15,000 × g for 15 min. The supernatant was stored at −80 °C for further analysis. The leftover portion of the right kidney was minced and homogenized in a cold lysis buffer containing Radio Immunoprecipitation Assay (RIPA) buffer, PMSF (Sigma), protease inhibitor cocktail (Roche), and PhosSTOP (Roche). Then, the homogenate was centrifuged at 15,000 × g for 15 min at 4 °C. Protein concentration was determined by the Bradford reagent (Bio-Rad, USA).

### 2.2.4. Relative kidney weight

Bodyweight of mice was recorded before sacrifice. The left kidney weight of mice was recorded during organ collection. The relative kidney weight was calculated using the following formula (Almeer et al., 2019):

\[
\text{Relative kidney weight} = \frac{\text{Left Kidney Weight}}{\text{Body weight}} \times 100
\]

### 2.2.5. Determination of Pb concentration in blood and tissue samples

The Pb content of blood and tissue samples was determined following the protocol of the previously published article (Rana et al., 2020). Briefly, the samples were diluted with a mixture of 0.2% Triton-X 100, (NH4)2HPO4, and water. A graphite furnace assembled with atomic absorption spectrometry (AAS) (Zeeman Atomic Absorption Spectrophotometer Z-5000, Hitachi, Japan) was used to determine the Pb concentration. The detection wavelength was set at 283.3 nm and the Pb concentration was calculated from the standard curve of pure Pb. The Pb content of blood and tissue samples was expressed as µg/dL and µg/g protein, respectively.

### 2.2.6. Determination of oxidative stress-related parameters

Oxidative stress is a common mechanism of cellular injury via lipid peroxidation. Thiobarbituric acid reactive substances (TBARS), a secondary byproduct of oxidative stress manifest as an indicator of LPO. For TBARS analysis, a mixture of trichloroacetic acid (TCA) was mixed and centrifuged at 3000 rpm for 15 min. Then, the resultant supernatant was mixed with 15% thiobarbituric acid (TBA) following the boiling for 30 min. After centrifugation at 3000 rpm for 15 min, the absorbance of supernatant was measured at 595 nm. The level of TBARS was calculated against a standard curve of Malondialdehyde (1, 1, 3, 3-tetraethoxyxopropane) (Rana et al., 2020).

The level of superoxide dismutase (SOD) and catalase (CAT) activity was assessed according to the protocol of Rana, Tangpong, and Rahman, 2020 (Rana et al., 2020). In case of SOD, 1 ml of Tris-EDTA buffer (pH 8.2) and pyrogallol solution (0.2 mM) were mixed. Thereafter, the sample (50 µL) was added to the mixture and absorbance was recorded at 420 nm for each 30 sec of 3 min duration using UV–vis spectrophotometer (Jasco V– 630, Japan). The data were expressed as U/g of protein. For CAT activity, 2 ml of PBS was mixed with 10 µL of tissue homogenate. After that, 1 ml of 30 mM H2O2 solution was added to the mixture. The decomposition rate of H2O2 was recorded against PBS (blank) at 240 nm for each 30 sec of 3 min. The CAT activity was expressed as katal (k)/mg of protein.

### 2.2.7. Determination of kidney function parameters

To assess the kidney function, the blood urea nitrogen (BUN) and creatinine were determined using a bioassay system reagent kit (Hayward, CA, USA) in an automatic chemistry analyzer (KONE Lab20, Tokyo, JP). Besides, urine samples were collected for analyzing the creatinine and protein levels. Protein level was measured using UV–vis spectrophotometer (Jasco V– 630, Japan).

### 2.2.8. Determination of inflammatory markers

#### 2.2.8.1. Tumor necrosis factor-alpha (TNF-α) by ELISA.

The level of plasma TNF-α was investigated using an enzyme-linked immunosorbent assay (ELISA) kit of R&D System, Minneapolis, MN, USA. The experiment was conducted following the supplied protocol by the manufacturer.

### 2.2.8.2. Determination of TNF-α, COX-2, and iNOS expression by western blot

The protein expression was determined following the previously published report (Rana et al., 2020). In brief, a 40 µg of protein was loaded into each well of SDS polyacrylamide gel and run for electrophoresis (Bio-Rad Laboratories, Inc., California, USA) at 100 V. Afterward, the separated protein was transferred to a nitrocellulose membrane (Roche Diagnostics Corporation, Indianapolis, IN, USA) at 300 mA, 4 °C for 1.5 h. Then, the membrane was blocked with 5% skim milk in TBST (20 mM Tris–HCl, pH 7.4, 150 mM NaCl, 0.02% Tween 20). The membrane was then incubated overnight at 4 °C with primary antibody: TNF-α, COX-2, and iNOS (Cell Signaling Technology, Massachusetts, USA) followed by the washing with TBST 3X for 5 min. Following incubation with secondary antibody, the membrane was washed 3X for 5 min. After that, the intensity of protein band was determined by enhanced chemiluminescence detection kit (Bio-Rad, USA). The relative expression of protein was expressed as fold change compared to β-actin as control (Cell Signaling Technology, Massachusetts, USA).

### 2.2.9. Assessment of kidney histology and apoptosis

For analyzing kidney histology, tissue was fixed in 4% neutral-buffered formalin, dehydrated in a series of graded alcohol and then embedded in paraffin. The tissue was sectioned (4 µm) into a glass slide using Leica CM1950 Cryostat (Leica Microsystem,
Germany). The slides were hydrated and stained with hematoxylin and eosin (H & E) followed by air-drying. The photograph was investigated and captured using a light microscope (Olympus EX51, Olympus Corporation, Tokyo, Japan).

To investigate the extent of apoptosis, the frozen kidney tissue (-30 °C) was sectioned (7 μm) using a microtome (Leica Microsystems CM1950, Wetzlar, Germany) and fixed on glass slides using frozen mounting media (FSC 22 Mounting Media, Leica Biosystems, Harbourfront Center, Singapore). An in situ apoptosis detection kit (Abcam, Cambridge, UK) was used to evaluate the TUNEL positive cells in kidney section. The slides were investigated under a light microscope (Olympus EX51, Olympus Corporation, Tokyo, Japan), and the result was expressed as the number of TUNEL positive cells per 100 fields.

2.3. Statistical analysis

All data were expressed as mean ± standard error mean (SEM). Significant differences variables among groups were measured by one-way analysis of variance (ANOVA) followed by Newman–Keuls post hoc test. Here, a data are presented as mean + SEM (n = 6). The multiple comparisons were analyzed by one-way analysis of variance followed by Newman–Keuls post hoc test. Here, a P < 0.05 vs untreated control, b P < 0.05 vs PbAc treated group.

3. Results

3.1. PASS prediction and molecular docking analysis

PASS analysis identified a total 15 phytoconstituents from TL leaves (See the supplementary Table S1). The result showed that individually these compounds were antitoxic/antidote, anti-oxidative stress element stimulant (HMOX1 expression enhancer; NF-E2-related factor 2 stimulants), calcium regulator, kidney function stimulant. Moreover, these phytoconstituents also possess antioxidant and anti-inflammatory activity (See the supplementary Table S2).

Based on the PASS analysis and solubility in water, polyphenol compounds (e.g. caffeic acid, protocatechuic, and gallic acid) were selected for molecular docking which had better interaction with the assigned protein of interest (See the supplementary Figure S1). The docking values of phytoconstituents interaction with proteins were summarized in Table 4. For instance, protocatechuic acid which docked with IP3R possesses the highest score 5.93 and 5.9, and the interacting residues are ARG 511, TYR 567, LYS569, ARG 269, ARG 265. Gallic acid with −5.9 docking score interacted with KEAP1 at ARG 483, SER 508. The order of compounds that showed the highest binding affinity to TNF-α, iNOS, COX-2, p38k, JNK, and NF-κB were protocatechuic acid (-5.66), protocatechuic (-6.258), gallic acid (-7.81), caffeic acid (-6.70), gallic acid (-6.62), and caffeic acid (-7.03), respectively.

The analyzed data of drug likeliness properties revealed that all of the phytochemicals that interacted with the assigned protein aforementioned earlier were resemble to be drug candidates (Table 5).

3.2. TL attenuated aberrant relative kidney weight

To validate the approach of treatment, the mice body weight, kidney weight, and body Pb content were investigated (Table 1). Study showed that no significant difference was observed between the mice body weight of Pb treated group and untreated control. In contrast, both kidney weight and % of relative kidney weight parameters were significantly different (1.38-fold) between untreated control and Pb treated group, while co-treatment of TL attenuated the parameters dose-dependently (1.04 and 0.92 for 100 and 200 mg/kgBW) (Table 1). On the other hand, the increment of blood and kidney Pb burden were closely associated with the elevated relative kidney weight in Pb treatment. Co-treatment with TL reduced the bodily Pb deposition in dose–response and significant manner except for blood Pb content (Table 1). The significant blood Pb content indicated that Pb deposition was crucial to elevate the % of relative kidney weight and uncovered as a probable pathological index in Pb-induced kidney toxicity.

3.3. TL mitigated the Pb-induced oxidative stress

To investigate the plausible mechanism of pathogenesis, the oxidative stress status was evaluated (Fig. 1). Treatment with Pb downregulated the activity of superoxide dismutase (SOD) and catalase (CAT) in kidney tissue as shown 1.36 and 3.36-fold, accordingly (Fig. 1A-B). Co-treatment of TL 200 mg/kgBW with PbAc significantly (P < 0.05) augmented the activity of SOD and CAT, however, vitamin E co-treatment failed to induce the SOD, and CAT in significant a passion. Furthermore, TL alone also enhanced CAT but not SOD activity (Fig. 1A-B). Besides, the plasma, RBC, and tissue TBARS levels were increased by 2.17, 2.0, and 1.32-fold, accordingly, when mice were treated with Pb (Fig. 1C-E). Coherent to the antioxidant activity, co-treatment of TL at both doses (100 and 200 mg/kgBW) remarkably (P < 0.05) suppressed the plasma, RBC, and tissue lipid peroxidation in response to oxidative stress (Fig. 1C-E). These findings dictated that TL co-treatment is effective in revising the reduced antioxidant enzymes (SOD and CAT activity) and increased lipid peroxidation.

3.4. TL modulated inflammation by suppressing the inflammatory mediators

To investigate the association of inflammation, plasma TNF-α level and tissue protein expression of TNF-α, COX-2, and iNOS were assessed (Fig. 2). The PbAc treated group exhibited (P < 0.05) almost 1.69-fold increased plasma TNF-α compared to untreated control group (Fig. 2A). Similarly, PbAc also upregulated the inflammatory tissue proteins expression such as TNF-α, COX-2, and iNOS were increased to 1.63, 3.17, and 1.57-fold, respectively.

Table 1
Effect of TL extract on mice body weight, kidney weight, and Pb concentrations of PbAc treated mice.

| Groups            | Mice body weight | Kidney weight | % of relative kidney weight | Blood Pb (μg/dL) | Kidney Pb (μg/g protein) |
|-------------------|------------------|---------------|----------------------------|------------------|-------------------------|
| Untreated Control | 39.96 ± 0.80     | 0.34 ± 0.02   | 0.81 ± 0.03                | 1.83 ± 0.26      | 1.43 ± 0.08              |
| PbAc (15 in DW)   | 39.96 ± 1.24     | 0.45 ± 0.02   | a 1.12 ± 0.04              | 65.35 ± 0.83     | 52.70 ± 0.27             |
| TL 200 mg/kgBW    | 44.39 ± 0.77     | 0.40 ± 0.01   | b 0.90 ± 0.03              | 1.69 ± 0.48      | 1.37 ± 0.06              |
| Pb + TL 100 mg/kgBW| 35.78 ± 1.68    | 0.37 ± 0.02   | ab 1.04 ± 0.04             | 63.12 ± 1.59     | 45.45 ± 2.14             |
| Pb + TL 200 mg/kgBW| 39.20 ± 0.80    | 0.36 ± 0.01   | b 0.92 ± 0.04              | 62.96 ± 1.25     | 42.50 ± 0.97             |
| Pb + Vit-E 100 mg/kgBW | 36.68 ± 1.04 | 0.37 ± 0.02   | a 1.01 ± 0.06              | 64.89 ± 2.43     | 52.90 ± 2.13             |
| Vehicle Control   | 42.60 ± 0.80     | 0.33 ± 0.01   | 0.77 ± 0.04                | 1.89 ± 0.56      | 1.45 ± 0.09              |

Data are presented as mean ± SEM (n = 6). The multiple comparisons were analyzed by one-way analysis of variance followed by Newman–Keuls post hoc test. Here, a P < 0.05 vs untreated control, b P < 0.05 vs PbAc treated group.
As expected, co-treatment with TL at doses of 100, 200 mg/kgBW or vitamin E showed a significant reduction of circulating TNF-level compared to PbAc treated group (P < 0.05) (Fig. 2A). In addition, the expression of TNF-α, COX-2, and iNOS in the kidney were also downregulated in a dose-dependent manner when co-treated with TL or vitamin E (Fig. 2B-D). Thus, these findings illustrated that TL or vitamin E treatment can significantly attenuate the inflammation through suppressing the inflammatory mediators of PbAc induced kidney.

3.5. TL improved kidney function attenuating bodily biochemicals

To evaluate the renal function, the BUN and creatinine levels were evaluated in plasma sample, while the protein and creatinine levels, protein/creatinine ratio were estimated from urine sample (Table 2). Results showed that a significant (P < 0.05) increment of plasma BUN (1.37-fold) and creatinine (1.65-fold) levels were recorded in the mice treated with PbAc. Co-treatment of TL at both doses with PbAc declined the plasma BUN and creatinine levels significantly (P < 0.05), which were similar to that of vitamin E (1.93 and 1.4-fold, respectively).

Urine analysis study indicated that co-treatment of TL or vitamin E with PbAc significantly (P < 0.05) reduced the biochemical profile such as protein, creatinine levels, and protein/creatinine ratio than that of PbAc treated group (Table 2). These outcomes reflected the probability of TL’s effects against Pb induced kidney dysfunction.

3.6. Renoprotective and anti-apoptotic effect of TL against Pb

To investigate the magnitude of kidney damage, kidney tissues were stained with H&E and TUNEL assay (Figs. 3 and 4). The figure of control group represents the normal structure of glomerulus, proximal and distal in comparison to PbAc treated kidney, which shows the profound (+ + +) glomerular swelling, degenerative tubules, and interstitial inflammation (Fig. 3 and Table 3). These architectural changes of kidney tissue were returned to normal when co-treated with TL or vitamin E as shown by the grading score 0 (Fig. 3 and Table 3).

Successively, findings of the TUNEL assay also corroborated the outcome of H&E staining. Likewise, the degeneration of nephron architecture, an outnumber (5.75-fold) of cell death were reported in PbAC treated group compared to untreated control (P < 0.05) (Fig. 4). Co-treatment of TL at different doses declined the dead cell number dose-dependently. Interestingly, the outcome was promising for TL 200 mg/kgBW (2.97-fold vs PbAc), which was better than...
vitamin E (1.79-fold vs PbAc). In addition, there was no toxic effect was observed by TL 200 mg/kgBW or vitamin E vehicle, respectively (Fig. 4). Thus, TL has the ability to restore the architectural alterations of kidney by Pb.

4. Discussion

This study exerted the ability of PbAc to induce kidney toxicity by initiating the oxidative stress, inflammation, and cell death, which is resulted in kidney dysfunction and aberrant architecture of nephron. TL was effective in reversing all alterations that caused by PbAc. The possible underpinning protective mechanism could be due to its capability to activate Nrf-2, decrease cytosolic Ca²⁺, downregulate the inflammatory mediators and pathways (MAPK and NF-κB).

Lead intoxication is a global concern and is closely associated with industrialization. Therefore, its remedies such as chelating therapy are requisite based on the magnitude of toxicity (Rana et al., 2020) Persistent usage of antidote alone (CaNa₂EDTA, Dimercaprol, Dimercaprol) or in combination (CaNa₂EDTA with Dimercaprol) can develop adverse effects such as encephalopathy,
Fig. 3. Histopathological alterations in the kidney of experimental mice. Kidney tissues were stained with hematoxylin and eosin staining. Histological alteration of kidney was mainly observed in the cortex region, both in proximal (PT) and distal tubules (DT). Untreated Control, TL 200 mg/kgBW, and Vehicle Control groups showed normal renal architectures. In contrast, only PbAc exposed group (PbAc 1% in DW) showed the presence of tubular necrosis such as cloudy swelling of PT, shrinkage of both DT and PT. However, co-treatment with TL (100 and 200 mg/kgBW) and vitamin E (100 mg/kgBW) significantly restored the renal alterations compared to only PbAc exposed group (PbAc 1% in DW).

Fig. 4. Evaluation of the kidney apoptosis of PbAc treated mice kidney by terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) assay. (A)-Quantification of TUNEL positive cells in the kidney tissue section. The values are presented as mean ± SEM (n = 6). The multiple comparisons were analyzed by one-way analysis of variance followed by Newman–Keuls post hoc test. Whereas, a P < 0.05 vs untreated control, b P < 0.05 vs PbAc treated group. (B)- TUNEL positive cells were indicated by the red arrow in kidney tissue section (original magnification x400).
kidney damage, cardiac disorder (tachycardia), hepatic enzyme upregulation, hematological disorder, essential metal ion (Zn) depletion, and vomiting (Schroder et al., 2015). Therefore, researchers are looking for alternative sources of therapeutics like herbal medicine to mitigate the clinical conditions based on their ethnobotanical study, which was also the aim of our study. The research on PbAc toxicity has been under investigation since the long, but the toxicodynamics of Pb on the kidney is yet to be developed. In general, oxidative stress is termed as the commonly employed mechanism in Pb-induced nephrotoxicity; an outcome of excessive ROS (Kelainy et al., 2019) exclusively superoxide anion (Gargouri et al., 2013) production. In this context, we postulated that attenuation of oxidative stress and its downstream player by TL could be the plausible protective role against PbAc induced kidney toxicity. Our current study demonstrated that co-administration of TL with PbAc reduced the oxidative stress with enhanced SOD and CAT activity as well as reduced Pb accumulation, which were in agreement with a previous study (Hasanein and Riahi, 2018). Hence, PASS analysis of TL phytoconstituents (Supplementary file 1) and the previous study from our lab revealed that TL is a potent antioxidant and could possess the chelating effect as a form of antioxidant activity (Rana and Tangpong, 2017). The phenolic compound present in the TL leaves exclusively caffeic acid, gallic acid and protocatechuic acid are previously reported to protect kidney against oxidative stress enhancing the endogenous antioxidant (Kohansal et al., 2019; Rehman and Suitana, 2011; Yamabe et al., 2015). However, there is an ambiguity to explain the failure of TL to reduce blood Pb deposition.

Shreds of evidence suggested that Pb intoxication alters the histological integrity of the kidney as well as kidney functions. According to our current study, TL restored the PbAc induced all histological alterations such as hydropic degeneration changes in tubules, glomerular damage and inflammatory cellular infiltration. These results were in agreement with the previous studies (Mao et al., 2016; Zhang et al., 2019). Consistently, tissue apoptosis analysis strongly corroborates with the histopathological findings and kidney dysfunctions. For example, higher number of cell death (TUNEL positive) was evidenced in PbAc treated mice, while TL effectively protected the cell from death. This result could be due to the counteracting effect of TL against ROS mediated activation of caspase-3 and caspase-9 (He et al., 2016; Monime et al., 2016; Siddarth et al., 2018), exclusively by protocatechuic acid, caffeic acid and gallic acid (Mard et al., 2015; Trumbeckaite et al., 2017; Yamabe et al., 2015). To activate caspase cascades, DNA fragmentation is a prerequisite (Mcllwain et al., 2013), which was actively achieved by PbAc in the kidney (Kelainy et al., 2019). Our previous report uncovered that TL can protect DNA from PbAc induced toxicity (Rana and Tangpong, 2017). However, the current study did not exclude any other means of programmed cell death, e.g pyroptosis.

Table 3

| Groups                  | Hydropic degeneration changes in tubules | Glomerular damage | Inflammatory cellular infiltration |
|-------------------------|-----------------------------------------|-------------------|-----------------------------------|
| Untreated Control       | 0                                       | 0                 | 0                                 |
| PbAc (15 in DW)         | + ++                                    | ++                | ++                                |
| TL 200 mg/kgBW          | 0                                       | 0                 | 0                                 |
| Pb + TL 100 mg/kgBW     | +                                       | +                 | +                                 |
| Pb + TL 200 mg/kgBW     | 0                                       | 0                 | 0                                 |
| Pb + Vic-E 100 mg/kgBW  | 0                                       | 0                 | 0                                 |
| Vehicle Control         | 0                                       | 0                 | 0                                 |

Histological evaluations of kidney tissues (n = 6). Here, +++ represents profound damage or diffuse damage to more than 50% of the tissue, ++ indicates moderate damage or damage scattered between 10 and 50% of tissue, + is considered as mild damage or random in<10% of the tissue, and 0 = no damage.

The interplay between inflammation, ROS, and Pb toxicity is well documented (Wang et al., 2016). Pb mediated ROS abundance can function as signaling messengers to NF-κB (Gloire et al., 2006). Activated NF-κB is then translocated to the nucleus, where it regulates the transcription of inflammatory genes such as TNF-α, COX-2, and iNOS ((Bhaskar et al., 2011; Gloire et al., 2006; Nair et al., 2006), which is coherent with our current findings. Moreover, ERK/JNK/p38 pathway is also involved in Pb-induced ROS mediated inflammation and tubular damage (Wang et al., 2016). According to Meissner, Viehmann, and Kurts, 2019, Danger-associated molecular pattern molecules (DAMPs) generated from damaged kidney could also induce the inflammation (Meissner et al., 2019). In our current study, TL attenuated kidney inflammation through the downregulation of inflammatory mediators possibly by interfering with NF-κB, MAPK pathway. Accumulated study reported that caffeic acid, gallic acid and protocatechuic acid are able to limit the inflammation by reducing cytokines like IL-1β, TNF-α, monocyte chemoattractant protein 1 (MCP-1) and myeloperoxidase (MPO) activity (Chao et al., 2010; Owumi et al., 2020; Semaining et al., 2015).

According to previous studies, TL leaf also contains water soluble phenolic compounds along with other water insoluble constituents (Chan et al., 2011; Oonsivilai et al., 2008; Ratchadaporn, 2006). To explore the possible pathway of the protective effect of TL, PASS analysis of the available phytoconstituents of TL was employed. Based on PASS analysis and our recent review, we assumed that the activity of TL phytoconstituents against PbAc, might be due to the capability to induce Nrf-2 and to limit Ca2+ ion in the cytosol (activation of IP3R), or to modulate the MAPK, NF-κB pathways. In the meantime, Trumbeckaite and colleagues reported that ester of caffeic acid and phenethyl alcohol is capable to induce mitochondrial calcium uptake and protects kidney cells from ischemia-induced necrosis (Trumbeckaite et al., 2017). Besides, protocatechuic acid, caffeic acid and gallic acid also possess the activity to inhibit the activation of NF-κB and MAPK pathways (Ahad et al., 2015; Huang et al., 2018; Ma et al., 2018; Takakura et al., 2018). In agreement with our previous study, the current docking study suggested that among phytochemicals polyphenols were the most active constituents to interact with a high affinity toward IP3R, Keap-1, JNK, p38, iNOS, TNF-α, COX-2, which support our hypothesis (Rana et al., 2020). Concomitant to current findings, previous studies also demonstrated that phenolic content is highly effective against Pb induced toxicity via intervening NF-κB and MAPK activation (Gargouri et al., 2013; Wang et al., 2016).

In a nutshell, the above findings suggested that PbAc caused kidney damage and impaired function through oxidative stress, inflammation, and apoptosis. TL ameliorated all the aberrant indices via inducing Nrf-2 and suppressing IP3R, MAPK, NF-κB pathways. Therefore, TL might be applicable to treat Lead-mediated kidney complications, though extensive studies and clinical trials are needed.
Table 4
Phyto-constituents of TL extract attenuates oxidative stress and inflammation via interacting with IP3R, KEAP-1, TNF-α, iNOS, COX-2, p38 kinase, JNK, and NF-kB.

| Compound names | IP3R (PDB: 3UJ0) | KEAP-1 (PDB: 5CGJ) | TNF-α (PDB: 2AZ5) | iNOS (PDB: 3E67) | COX-2 (PDB: 5KIR) |
|----------------|------------------|--------------------|--------------------|------------------|------------------|
|                | Docking score   | Glide energy       | Interacting Residues | Docking score   | Glide energy       | Interacting Residues | Docking score   | Glide energy       | Interacting Residues | Docking score   | Glide energy       | Interacting Residues |
| Caffeic acid   | –4.16           | –22.10             | ARG 269 ARG 511 ARG 504 LYS 569 | –5.05           | –25.40             | SER 555 GLN 530 | –5.52           | –26.02             | LEU B:120 TYR B:119 SER B:160 | –4.977           | –26.18             | GLU 371          |
| Protocatechuic acid | –5.93          | –27.62             | ARG 511 TYR 567 LYS 569 ARG 269 ARG 265 | –5.75           | –21.68             | ARG 483 SER 508 | –5.66           | –21.20             | TYR B:151 SER B:60 | –6.258           | –25.08             | TRP 366 | SER 436 | –7.26 | –27.65 | NONE |
| Gallic acid    | –5.68           | –26.42             | ARG 265 ARG 269 ARG 511 TYR 567 LYS 569 | –5.9            | –23.72             | ARG 483 SER 508 | –5.57           | –22.47             | TYR B:119 SER B:60 | –4.97            | –22.73             | GLU 371          | –7.81            | –32.34             | MET 522 |

| Compound names | P38 KINSAE (PDB: 2Y1S) | JNK (PDB: 3OY1) | NFκB (PDB: 4KIK) |
|----------------|------------------------|-----------------|------------------|
|                | Docking score   | Glide energy       | Interacting Residues | Docking score   | Glide energy       | Interacting Residues | Docking score   | Glide energy       | Interacting Residues |
| Caffeic acid   | –6.70           | –28.78             | ALA 111 GLY 110 MET 109 HID 107 LYS 53 | –5.91           | –29.33             | MET 149 LYS 93 | –7.03           | –30.33             | CYS 99 GLU 97 ASP 166 LYS 44 |
| Protocatechuic acid | –6.41          | –25.62             | GLY 110 MET 109 HID 107 | –5.87           | –24.20             | LEU 144 LYS 93 | –6.10           | –24.03             | ASP 103 CYS 99 |
| Gallic acid    | –6.47           | –26.78             | GLY 110 MET 109 HID 107 | –6.62           | –29.66             | ILE 70 MET 149 | –5.99           | –24.78             | ASP 103 CYS 99 |
Table 5
Drug likeliness properties of TL extract.

| Compound name          | PubChem CID | Drug-likeliness properties | MW (g/mol) | HB donor | HB acceptor | QPlogPo/w |
|------------------------|-------------|----------------------------|------------|----------|-------------|-----------|
| Caffeic acid           | 689,043     |                            | 180.16     | 3        | 3.5         | 0.557     |
| Protocatechuic acid    | 72          |                            | 154.122    | 3        | 3.5         | 0.032     |
| Gallic acid            | 370         |                            | 170.121    | 4        | 4.25        | -0.568    |

QikProp module of Maestro 11 (Schrödinger) was used to project the drug-likeliness properties of these seven compounds. Here, MW represents Molecular weight; HB donor represents donation of hydrogen bond by solute; HB acceptor, number of hydrogen bonds received by the solute; QPlogPo/w, Predicted octanol/water partition coefficient. Compounds that meet at least three properties such as MW < 500, HB donor < 5, HB acceptor < 10, QPlogPo/w (−) 2.0−6.5, were considered to possess drug-like properties.

Ethical Approval

Approval for the use of animals was given by the Animal Care and Use Committee (ACUC) of Walailak University, Thailand. Animals were handled as stipulated by the guidelines for the use of animals set by the ACUC of the university, with the Animal Ethics Approval Certificate number: 002/2013.

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CRediT authorship contribution statement

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Declaration of Competing Interest

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.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.sjbs.2020.08.016.

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