Toxicity study and blood pressure–lowering efficacy of whey protein concentrate hydrolysate in rat models, plus peptide characterization

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ABSTRACT

We evaluated the acute (single-dose) and subacute (repeated-dose) oral toxicity of alcalase-hydrolyzed whey protein concentrate. Our acute study revealed no death or treatment-related complications, and the median lethal dose of whey protein concentrate hydrolysate was >2,500 mg/kg. In the subacute study, when the hydrolysate was fed at 3 different concentrations (200, 400, and 800 mg/kg), no groups showed toxicity changes compared with controls. Then, whey protein concentrate hydrolysate was orally administered to spontaneously hypertensive rats. Results revealed significant reductions in blood pressure in a dose-dependent manner, and dosing at 400 mg/kg led to significant blood pressure reduction (−47.8 mm Hg) compared with controls (blood pressure maintained) and the findings of previous work (−21 mm Hg). Eight peptides—RHPEYAVSVLLR, GGAPPAGRL, GPPLPRL, ELKPTPEGDL, VLSELPEP, DAQSAPLRVY, RD-MPIQAF, and LEQVLPRD—were sequentially identified and characterized. Of the peptides, VLSELPEP and LEQVLPRD showed the most prominent in vitro angiotensin-converting enzyme inhibition with half-maximal inhibitory concentrations of 0.049 and 0.043 mM, respectively. These findings establish strong evidence for the in vitro and in vivo potential of whey protein concentrate hydrolysate to act as a safe, natural functional food ingredient that exerts antihypertensive activity.

Key words: angiotensin-converting enzyme (ACE) inhibition, bioactive peptide, hydrolysis, rat study, whey protein concentrate

INTRODUCTION

Hypertension is a sustained increase in blood pressure: systolic blood pressure (BP) >140 mm Hg and diastolic BP >90 mm Hg (American Diabetes Association, 2018). Hypertension is a key risk factor in cardiovascular disease, including coronary heart disease, peripheral arterial disease, and stroke. In 2016, the estimated global incidence of hypertension was 17.9 million, and hypertension accounted for 31% of all deaths worldwide. By 2025, hypertension is projected to affect 1.5 billion people around the world (Kearney et al., 2005). This global increment can be attributed to an aging population that is inadequately equipped for the prevention, diagnosis, and control of the disorder (Olsen et al., 2016). A rise in blood pressure is mainly due to the dual action of angiotensin-I converting enzyme (ACE), which hydrolyzes an inactive decapptide (angiotensin-I) into a potent vasoconstrictor octapeptide (angiotensin-II) and inactivates the vasodilating nonapeptide bradykinin (Guo et al., 2009). By inhibiting the activity of ACE, the formation of angiotensin-II and the breakdown of bradykinin are halted, keeping BP lower and preventing hypertension (van der Ven et al., 2002).

Current hypertension treatment includes synthetic drugs such as angiotensin receptor blockers, direct-acting vasodilators, ACE inhibitors, diuretics, calcium channel blockers, and anti-adrenergics. However, these drugs come with side effects such as fatigue, skin rash, headache, constipation, excessive urination, abnormal heart rate, and dizziness. These limitations have encouraged researchers to seek alternatives that are safe and natural, to replace synthetic agents. Positive results have been garnered from the discovery of bioactive peptides derived from various food proteins. One good source of bioactive peptides is whey protein, which demonstrates considerable nutritional and health benefits for humans. Whey peptides have shown anti-hypertensive (Guo et al., 2009; Tavares et al., 2011;
Morais et al., 2015), immunomodulatory (Wu et al., 2018), antithrombotic (Silveira et al., 2013), antimicrobial (Boyaci et al., 2016), opiate, and antioxidant properties (Madureira et al., 2010).

In our study, whey protein concentrate (WPC), a manufactured dairy product with a protein content of 76.4%, was chosen to generate bioactive WPC hydrolysate by means of enzymatic hydrolysis using alcalase, followed by in vivo studies exploring the acute toxicity (half-maximal lethal dose; LD$_{50}$), subacute toxicity, and BP-lowering efficacy of WPC hydrolysate, along with characterizing 8 peptide sequences identified from WPC.

**MATERIALS AND METHODS**

Materials

We procured WPC from Purefit Company (Selangor, Malaysia). We procured ACE from rabbit lung and its substrate N-hippuryl-His-Leu hydrate powder, from Sigma Chemical Co. (St Louis, MO). We purchased alcalase (EC 3.4.21.62) from Novozymes ( Bagsvaerd, Denmark). Acetonitrile, trifluoroacetic acid and methanol were of HPLC grade. All other chemicals were of analytical grade and were obtained from Fisher Scientific (Waltham, MA) or Merck and Co. (Kenilworth, NJ).

**Enzymatic Hydrolysis of WPC**

Alcalase enzyme was mixed with 2.0 g of WPC at 1:100 ratio (wt/wt) in 100 mL of borate buffer solution (pH 8), and then incubated in a water bath shaker at an agitation rate of 150 rpm and a temperature of 55°C to allow optimum hydrolysis as determined previously (data not shown). At the end of hydrolysis, the mixture was placed in boiling water for 10 min to inactivate the enzyme. After centrifugation at 10,000 × g for 15 min at room temperature, the supernatant was collected and stored at −80°C before freeze-drying. The freeze-dried hydrolysates were used for the acute and subacute toxicity studies, the efficacy studies and the peptide sequencing protocols described below.

**Toxicity Study of WPC Hydrolysate Using Wistar–Kyoto Rats**

**Acute Toxicity (LD$_{50}$).** We determined the acute toxicity of WPC hydrolysate using the limit test described by the Organisation for Economic Co-operation and Development (OECD) Test Guideline 425, with slight modifications (OECD, 2011). We performed a limit test because WPC hydrolysate was likely to be nontoxic; it has GRAS (generally recognized as safe) status from the United States Food and Drug Administration (Boland, 2011) under regulation 21CFR184.1979c. Dosing was 2,500 mg/kg. Before dosing, 5 female Wistar–Kyoto rats with a BW of 150.92 ± 2.12 g (mean ± standard deviation) were fasted overnight from food, but water was freely available. Female rats were selected, because they are more sensitive to LD$_{50}$ than male rats (Lipnick et al., 1995). The sample was administered to 1 rat at a time using oral gavage. Each rat was observed carefully for signs of lethality, if any, within the next 48 h. When the rat survived, the next rat was dosed. All 5 rats were tested.

**Subacute Toxicity (28 d of Repeated-Dose Feeding).** Twenty-eight days of repeated dose feeding was performed according to OECD Test Guideline 407 (OECD, 2008) using Wistar–Kyoto rats aged 8 wk. The animals were randomly divided into 4 groups consisting of 6 males and 6 females each. A control group received distilled water and the treatment groups received WPC hydrolysate at low (200 mg/kg), intermediate (400 mg/kg), and high (800 mg/kg) concentrations. The hydrolysate was dissolved in 1 mL of distilled water and administered using oral gavage on a daily basis for 28 d. We recorded the BW gain and food and water intake of each rat on a weekly basis throughout the dosing period to observe for early signs of toxicity (data not shown).

At the end of the experiment, all rats were anesthetized with an intraperitoneal injection of ketamine at 80 mg/kg of BW and xylazine at 8 mg/kg of BW. A volume of 4.0 mL of blood was withdrawn from the heart via cardiac puncture and immediately transferred to red Vacutainer tubes (Becton, Dickinson and Company, Franklin Lakes, NJ) that contained coagulant. After standing at room temperature to allow clotting for 30 min, the blood was centrifuged at 1,500 × g at 4°C for 10 min. The serum was collected as supernatant and kept at 4°C until clinical chemistry analysis. The main organs, including the kidney, liver, lungs, and reproductive organs (testes for male rats and ovaries for female rats) were harvested, removed of fats, washed using saline buffer, and kept overnight in 10% formalin buffer. The formalin was replaced the next day to facilitate blood drainage from organs. For microscopic slides, formalin-soaked organs were cut down the middle into thin pieces in a cross-sectional manner, dehydrated using alcohol, embedded in paraffin, sliced into thicknesses of 4 to 5 μm, and stained with hematoxylin–eosin dye. All organs from the treatment groups were compared with those of the control group to look for toxicity signs such as cytoplasmic vacuolation, pyknosis/cell necrosis, fading of nucleus, hemorrhage, and swollen cells (Cummings et al., 2012) using a microscope (Nikon Eclipse 50i; Nikon Corp., Tokyo, Japan).
Efficacy Study on WPC Hydrolysate to Reduce Systolic BP in Spontaneously Hypertensive Rats

We evaluated the BP-lowering efficacy of WPC hydrolysate according to the method described by Fuglsang et al. (2003) and Zou et al. (2014), with some modifications. Male spontaneously hypertensive rats, aged 12 wk and with a mean BW of 238.26 ± 9.17 g, were divided into 5 groups of 6 animals each: a negative control group (distilled water), a positive control group (captopril 50 mg/kg), and 3 treatment groups (alcalase-generated WPC hydrolysate at 200, 400 and 800 mg/kg BW). The WPC hydrolysate was dissolved in distilled water, thoroughly mixed, and administered to the rats via oral gavage at a standardized volume of 1 mL/rat. Systolic BP was measured 6 times: a baseline before oral administration (0 h), and at 2, 4, 6, 8, and 24 h after feeding using the CODA tail-cuff method, a noninvasive BP system that allows the identification of pulses at specific rat tail position (Kent Scientific, Torrington, CT). Rats were familiarized with the CODA machine before the experiment. The rats were allowed to move freely into a specially designed restrainer, where their tails were left exposed for cuffing with an occlusive ring. The occlusion cuff was pushed to the base of the tail, and the volume-pressure recording cuff was placed 2 mm away from the occlusion cuff. Before BP measurements, the rats were kept on a warming plate (37°C) for 20 min to ensure sufficient blood flow inside the tail and increase measurement accuracy. A total of 25 readings were taken per measurement cycle because of large BP variation in each rat. Three readings were then selected to calculate the average BP. All spontaneously hypertensive rats were handled gently and humanely to reduce variations in results due to stress.

Animal Care

Both Wistar–Kyoto rats and spontaneously hypertensive rats were purchased from the Animal Experimental Unit, Faculty of Medicine, University of Malaya, Kuala Lumpur, Malaysia, and were acclimatized to laboratory conditions for 1 wk before dosing. All rats were housed in plastic cages, in pairs of the same sex, at a temperature of 22 to 24°C, and with a 12-h light–dark cycle. A standard laboratory diet and tap water were available ad libitum throughout the experiment, except during the acute toxicity study, when fasting was required. All procedures involved in the animal studies were approved by the Institutional Animal Care and Use Committee, Universiti Putra Malaysia (UPM/IA-CUC/AUP No. R078/2015).

Peptide Sequencing Assay

Before sequencing, peptides were separated using reversed-phase HPLC and an isolectric focusing point technique to obtain fractions with less complexity. The fractions demonstrating significantly higher ACE inhibition were then selected for sequencing. During reversed-phase HPLC, the hydrolysate was mixed with deionized water (100 mg/mL), filtered, and loaded onto a C-18 semi-preparative column (9.4 × 250 mm, 5 μm; Agilent Technologies, Santa Clara, CA). Chromatography was performed using 0–10 min 100% mobile phase A (0.1% trifluoroacetic acid in deionized water), followed by gradient elution of mobile phase B (0.1% trifluoroacetic acid in acetonitrile) between 10 and 55 min to achieve 58.8% acetonitrile and holding at 58.8% acetonitrile for 55–67.5 min. Elution was monitored at 205 nm. During isolectric focusing separation, the sample was diluted with deionized water to reach a final volume of 1.8 mL. A gel strip with a pH gradient between 3 and 8 was rehydrated using deionized water. Then, 150 μL of the diluted sample was dispensed into individual wells, and a cover seal was placed on top to prevent sample evaporation. A high voltage (500–4,000 V) was applied to both ends of the gel strip to allow peptides to migrate and reside at their respective pI. Upon completion, samples were retrieved from the liquid phase in each well with different pH.

Peptide sequencing was done using tandem mass spectrometry (MS/MS) as previously described (Zarei et al., 2015). Briefly, potent peptide fractions were freeze-dried and reconstituted with 10 μL of mobile phase A (0.1% formic acid in deionized water), and then spun intermittently at 9,750 × g for 15 min to ensure any insoluble material was pelleted at the bottom of the tube. The supernatant was analyzed using a chip-based nanospray liquid chromatography unit (1200 HPLC-Chip/MS Interface; Agilent Technologies) coupled with an accurate mass quadrupole–time of flight mass spectrometer (Q-TOF; 6520, Agilent Technologies). One microliter of sample was injected onto a HPLC analytical chip (C-18; 160-nL enrichment column; 75 μm × 150 mm) and separated with a gradient elution of mobile phase B (0.1% formic acid in 90% acetonitrile) from 3 to 50% over 30 min. Peptide molecules were then passed through an electrospray ionization source, ionized in positive mode, and fed into a Q-TOF mass spectrometer. Peptides were monitored at a user-defined mass:charge ratio between 110 and 3,000 m/z (MS scan) and were automatically selected for fragmentation into product ions (MS/MS scan, user-defined m/z between 50 and 3,000) by collision-induced dissociation, based on a precursor absolute threshold.
of 200 counts, precursor charge state selection, and preference at Z = 1, 2, 3, and > 3, excluding m/z 922.009798 (Z = 1) and 121.050873 (Z = 1), because both were reference ions. Raw sample data obtained from Q-TOF were searched against the animal species subdirectory of the Swiss-Prot protein database (http://www.uniprot.org/) using the PEAKS studio 6.0 search engine (Bioinformatics Solution Inc., Waterloo, ON, Canada). Database searching was performed using the following parameters: minimum false discovery rate, −10lgP confidence score greater than 20 (a cut-off value to select peptides of high confidence), and “no enzyme” selection, because the alcalase enzyme was not specified. De novo sequencing was conducted according to the following cutoffs: average confidence score higher than 80% for at least 4 consecutive amino acids, and a fragment ion mass tolerance of 0.1 Da. Residues of high confidence, labeled in red and blue, were selected for synthesis.

**Half-Maximal Inhibitory Concentration for ACE-Inhibitory Peptides**

Eight potentially potent peptide samples were chemically synthesized by a commercial peptide supplier (GenScript, Piscataway, NJ) at a final purity of greater than 90%. We determined half-maximal inhibitory concentration (IC$_{50}$) for these peptides using a combined method from 2 previous studies (Cushman and Cheung, 1971, Ferreira et al., 2007), with certain modifications. Peptide stock solutions were diluted to different concentration ranges of 0.015 to 40 mM. The ACE was prepared as stock solution using potassium phosphate buffer (0.01 M, pH 7.0) and stored at −80°C until use. On the experiment day, the ACE was further diluted to 100 mU/mL using sodium borate buffer (pH 8.3). A substrate solution containing 5 mM HHL and 0.3 M NaCl in 0.1 M sodium borate buffer (pH 8.3) was prepared. The assay was initiated by mixing 10 μL of peptide with 10 μL of ACE for 10 min to allow inhibition, followed by the addition of 50 μL of substrate solution. The mixture was incubated for 60 min at 37°C. The enzymatic reaction was terminated by introducing 75 μL of 1 M HCl. After the reaction was halted, 150 μL of pyridine and 75 μL of benzene sulfonyl chloride were added. The solution was vortexed for 1 min and cooled on ice. After cooling, 200 μL solution was transferred to the 96-well plate for absorbance measurements at 410 nm. The experiment was carried out in triplicate. The ACE inhibition was calculated as follows,

\[
ACE\ inhibition\ (\%) = \frac{C - S}{C - B} \times 100,
\]

where C represents the control absorbance (ACE + substrate), S signifies the sample absorbance (peptide + ACE + substrate), and B denotes the blank absorbance (substrate only). The IC$_{50}$ values were then calculated by fitting the data into nonlinear regression from a plot of ACE inhibition (%) versus peptide concentration (mM) using GraphPad Prism 6 (GraphPad Software Inc., San Diego, CA).

**Statistical Analysis**

Using Minitab version 16 (Minitab Inc., State College, PA), statistical analysis and comparison between groups were obtained via one-way ANOVA. Tukey's test was used to identify differences between treatments at the 5% significance level (P ≤ 0.05). Data are presented as mean ± standard deviation from at least 3 determinations.

**RESULTS AND DISCUSSION**

**Acute and Subacute Toxicity of WPC Hydrolysate**

Acute and subacute toxicity provides crucial information that dictates the safety of materials intended for animal or human use. We evaluated the acute oral toxicity of WPC hydrolysate using the limit test procedure, and the results showed that after dosing 5 female Wistar–Kyoto rats with 2,500 mg hydrolysate/kg BW, no animals showed any sign of toxicity and survived the 48-h study period. The LD$_{50}$ for WPC hydrolysate was greater than 2,500 mg/kg, indicating that the sample was relatively nontoxic (i.e., showing no toxicity when dosed below 2,500 mg/kg; OECD, 2011). In the 28-d repeated-dose subacute study, male and female Wistar–Kyoto rats fed at WPC hydrolysate concentrations of 200, 400, and 800 mg/kg of BW demonstrated no toxicity signs, further supporting the safety of WPC hydrolysate for human consumption. Body weight gain, food and water intake, relative organ weight, clinical chemistry parameters, and histopathology assessments during the subacute study are reported below.

**BW, Food Intake, Water Intake, and Relative Organ Weights.** Continuous BW loss, reduced appetite, and fluid intake are signs of toxic effects. In the current study, male and female Wistar–Kyoto rats in all treated groups recorded BW increments during the 28-d dosing period. For male rats, the group receiving intermediate WPC hydrolysate concentration (400 mg/kg) showed significantly lower BW gain from wk 0 to 2 than the control group, but from wk 2 to 4, the same group of rats showed a BW gain of no significant difference from the controls. The BW gained for female
rats in all experimental groups were not significantly different from controls in all weeks. In terms of food and water intake, male rats generally showed higher consumption than female rats due to their larger body size, which required more nutrients to support movement and basal metabolic activity. Food intake in male rats was significantly lower during wk 1 and 4 in the 200 and 400 mg/kg groups; food intake in female rats was lower during wk 1 and 2 in the 400 and 800 mg/kg groups. Male rats showed lower water intake during wk 4 in the 200 and 400 mg/kg groups, and female rats showed significantly higher water consumption during wk 0 and 2 in the 200 mg/kg group, all compared with control (data not shown). All rats were active and healthy and showed no toxic signs, changes in locomotor activity, or behavior during the 28-d study period. The irregular fluctuations in BW gain and food and water intake was likely due to random factors and was irrelevant to the experiment.

We harvested the organs (lung, liver, kidney, testis, and ovary) of the Wistar–Kyoto rats and reported organ weights relative to BW. The ratio of organs to BW was important for explaining differences between the treatment groups and controls, because normalization of organ weight and BW removes any variation because of different BW in individual rats. This approach is supported by many recent works, which reported relative organ weights when evaluating toxicity from lavender essential oil (Mekonnen et al., 2019), lipophilized phenolic compound (Martínez et al., 2018), and herbal plants (Porwal et al., 2017). Figure 1 depicts the relative organ weights for male and female rats. In both sexes, the liver was the heaviest organ, because the liver carries out the principal detoxification processes in the body. The absolute organ weight for male rats was generally higher than that for females (data not shown), because of the larger body size of male rats. Statistically, both male and female rats revealed no significant differences from controls for all organs. We detected no abnormal organ enlargement or shrinkage in the treatment groups fed WPC hydrolysate.

Clinical Chemistry Analysis. As part of the toxicity study, blood was withdrawn to measure biochemical markers that dictate fluctuations in metabolic and physiological function in multiple organs, because blood is the major circulatory medium responsible for transporting nutrients, waste and toxic substances (if any) to excretion sites. Any adverse effects from testing could be easily detected from blood.

We measured levels of alanine transaminase, aspartate transaminase, alkaline phosphatase, total protein, albumin, total bilirubin, creatinine, urea, cholesterol, triglycerides, and electrolytes ($Na^+$, $Cl^-$, $K^+$, $Ca^{2+}$). Different markers are indicative of damage of different organs. Alanine transaminase, aspartate transaminase and alkaline phosphatase are primarily intracellular liver enzymes. When the liver is injured, these enzymes leak into the bloodstream, leading to elevated levels in the blood, as detected from clinical chemistry analysis. Total bilirubin, cholesterol, and triglycerides are also liver markers: high bilirubin (the breakdown product from old red blood cells) is indicative of impaired liver function, and high cholesterol and triglycerides are indicative of fatty liver. Creatinine and urea are markers of kidney function. Creatinine and urea, both waste products from cell metabolism, are excreted via kidney. High levels of creatinine and urea in the blood show retention of waste products in the body and reflect impaired kidney function. Electrolytes are also directly related to renal function, because their concentration is highly regulated by the kidney via osmotic pressure and water–ion ratio. Total protein and albumin are indicators of liver and kidney function. Lowered protein and albumin levels show liver and kidney malfunction, because they reflect the liver’s inability to synthesize protein and the kidney’s failure to retain synthesized protein, causing protein excretion from the body as urine.

Table 1 depicts the clinical chemistry parameters for male and female rats fed WPC hydrolysate at different concentrations. For male rats, all treatment groups showed no significant difference ($P > 0.05$) from controls in the measured parameters, including electrolytes, enzyme markers, and plasma compounds (total protein, albumin, bilirubin, creatinine, urea, cholesterol, and triglycerides). Similarly, we detected no significant difference in female rats, except in potassium and cholesterol. Potassium levels in the 800 mg/kg group and cholesterol levels in the 200 mg/kg group were significantly lower than controls, at 4.26 ± 0.25 mmol/L and 2.23 ± 0.05 mmol/L, respectively. According to the clinical chemistry database for laboratory rats (Boehm et al., 2007; Giknis and Clifford, 2008), the reference levels for potassium and cholesterol are 3.3 to 4.9 mmol/L and 0.7 to 2.5 mmol/L, respectively. Although the experimental data deviated from the controls, they were still within the normal range and thus were of no toxicological significance. It is common to observe large variations in the clinical chemistry of rats, because individual rats have different physical activity levels, emotional behaviors, stress levels, and diet patterns, which cause biochemical values to fluctuate.

Based on our findings for BW, food and water intake, relative organ weights and clinical chemistry analysis, WPC hydrolysate produced no signs of toxicity in male and female Wistar–Kyoto rats. It is apparent that WPC hydrolysate is nontoxic, a finding that was further confirmed by our histopathological observations, below.
Exposure of animals to a test substance for 28 d is usually sufficient to detect over 90% of adverse effects (Greaves, 2012). These effects are conveniently confirmed by histopathological evaluation, and the findings provide crucial safety information. Histopathological studies are performed to evaluate the effect of a test substance on vital organs, at the cellular level. Along with regular observations and clinical chemistry analysis, histopathological studies provide essential information about the overall toxicity of a compound.

In the current study, we harvested 5 organs: lung, liver, kidney, testis, and ovary. We monitored the lung for toxic effects affecting the respiratory system. We monitored the liver and kidney because they represent the major filtration and detoxification organs in the body and are exposed the highest amount of toxic substances from oral ingestion. We evaluated the testis and ovary for signs of toxicity in the reproductive system. Our results showed that all organs were normal and displayed no abnormalities compared with controls (Figures 2 and 3). For lung, both control and
treated samples demonstrated healthy alveolar tissue that was thinly layered to facilitate gas exchange and was infiltrated with lymphocytes to fight off airborne microbes. For liver, both control and treated samples showed healthy lobular structure with the central vein surrounded by hepatocytes (liver single cell unit) that radiated outwards. Kupffer cells were identified as large, distinct nuclei located at the periphery of sinusoid. For kidney, we observed the renal corpuscle (tuft ball-shaped unit) with complete, intact Bowman’s capsule and glomerulus, along with convoluted tubules (rod or spherical in shape). We found no vacuolation of the glomerulus, expansion of Bowman’s space, or formation of cysts, indicating normal kidney histology. For testis, we observed a complete structure of seminiferous tubules with intact content in both control and treated rats. We found no disruption of sperm cells or irregular shape of membrane propria (membranous layer surrounding individual tubule). Finally, for ovary, we observed medulla with thick-walled blood vessels. No ovaries showed toxic signs such as lymphocyte infiltration, oocyte necrosis, or unusual follicle atresia (breakdown of ovarian follicle). The findings from microscopic evaluation supported the nontoxic nature of WPC hydrolysate in Wistar–Kyoto rats after 28 d of administration, supporting its potential as a functional ingredient.

### Efficacy of WPC Hydrolysate in Reducing BP in Spontaneously Hypertensive Rats

We tested WPC hydrolysate in spontaneously hypertensive rats at 3 different concentrations (200, 400, and 800 mg/kg) to measure in vivo BP–lowering efficacy. Figure 4 shows the systolic BP in spontaneously hypertensive rats measured at 0, 2, 4, 6, 8, and 24 h after oral feeding. The aim of monitoring BP up to 24 h is to identify the extent of the BP–lowering effect before values return to baseline. At 24 h, BP readings at all dosages returned to normal, indicating the good degradative and non-cumulative character of WPC hydrolysate after oral ingestion. The negative control group, fed distilled water, showed no significant changes in BP over 24 h, and the positive control group, fed captopril (a well-known antihypertensive drug), showed significant BP–lowering effect after administration. All treatment samples showed significant BP reduction compared with baseline (213.5 mm Hg); we recorded

<table>
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<tr>
<th>Variable</th>
<th>Control</th>
<th>200 mg/kg</th>
<th>400 mg/kg</th>
<th>800 mg/kg</th>
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<tr>
<td>Sodium (mmol/L)</td>
<td>141.33 ± 2.52</td>
<td>143.33 ± 1.53</td>
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<td>143.00 ± 1.00</td>
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<td>Potassium (mmol/L)</td>
<td>5.70 ± 0.35</td>
<td>5.46 ± 0.30</td>
<td>5.20 ± 1.21</td>
<td>5.63 ± 0.50</td>
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<tr>
<td>Chloride (mmol/L)</td>
<td>102.00 ± 1.73</td>
<td>102.33 ± 0.58</td>
<td>103.33 ± 2.08</td>
<td>102.00 ± 0.00</td>
</tr>
<tr>
<td>Calcium (mmol/L)</td>
<td>2.46 ± 0.11</td>
<td>2.30 ± 0.34</td>
<td>2.30 ± 0.10</td>
<td>2.43 ± 0.15</td>
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<td>Alanine aminotransferase (U/L)</td>
<td>42.33 ± 5.13</td>
<td>54.00 ± 12.49</td>
<td>47.66 ± 2.51</td>
<td>46.33 ± 0.57</td>
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<tr>
<td>Aspartate aminotransferase (U/L)</td>
<td>185.00 ± 9.46</td>
<td>183.67 ± 2.08</td>
<td>185.00 ± 4.85</td>
<td>181.33 ± 1.53</td>
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<td>Alkaline phosphatase (U/L)</td>
<td>130.67 ± 13.32</td>
<td>146.33 ± 19.55</td>
<td>127.67 ± 8.96</td>
<td>130.76 ± 9.45</td>
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<td>Total protein (g/L)</td>
<td>63.56 ± 4.55</td>
<td>58.16 ± 12.45</td>
<td>61.46 ± 3.27</td>
<td>64.53 ± 3.64</td>
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<td>Albumin (g/L)</td>
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<tr>
<td>Total bilirubin (µmol/L)</td>
<td>5.66 ± 0.55</td>
<td>4.76 ± 0.92</td>
<td>5.10 ± 0.52</td>
<td>5.23 ± 1.07</td>
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<tr>
<td>Creatinine (mmol/L)</td>
<td>51.33 ± 8.50</td>
<td>42.33 ± 8.32</td>
<td>44.66 ± 4.72</td>
<td>47.33 ± 1.15</td>
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<tr>
<td>Urea (mmol/L)</td>
<td>6.43 ± 0.68</td>
<td>4.83 ± 1.33</td>
<td>7.43 ± 1.70</td>
<td>6.26 ± 0.20</td>
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<tr>
<td>Cholesterol (mmol/L)</td>
<td>2.03 ± 0.05</td>
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<td>2.33 ± 0.65</td>
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<tr>
<td>Triglycerides (U/L)</td>
<td>0.39 ± 0.01</td>
<td>0.53 ± 0.10</td>
<td>0.58 ± 0.16</td>
<td>0.94 ± 0.19</td>
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</table>

**Table 1. Serum biochemical values for male and female Wistar–Kyoto rats fed with whey protein concentrate hydrolysate (at 0, 200, 400, or 800 mg/kg) and killed after 28 d.**

1Values are mean ± SD from 6 rats per group (n = 6).

*Significant difference from control (P < 0.05).
maximum reductions of 29.9 mm Hg for the 200 mg/kg group, 47.8 mm Hg for the 400 mg/kg group, and 57.2 mmHg for the 800 mg/kg group. These BP reductions were dose-dependent: a higher WPC hydrolysate concentration contributed to a stronger BP-lowering effect.

The BP-lowering efficacy of dairy products, particularly from milk proteins, has been reported in the literature. For instance, bovine casein hydrolysate has been shown to reduce BP in spontaneously hypertensive rat up to 42 mm Hg when dosed at 100 mg/kg BW (Yamada et al., 2013), and fermented milk led to a BP reduction of 20 to 34 mm Hg when dosed at 1.5 mL per rat (Muguerza et al., 2006). A study similar to the current study showed that bovine WPC hydrolysate, when produced using a crude enzyme extract from artichoke, led to a maximum BP reduction of 21 mm Hg in spontaneously hypertensive rats when fed at 400 mg/kg (Tavares et al., 2012). At the same concentration, alcalase-hydrolyzed WPC showed a BP reduction of 47.8 mm Hg in the current study. This indicated a significant improvement in BP-lowering efficacy when WPC was treated with alcalase compared with arti-

Figure 2. Light photomicrographs of (a) lung, (c) liver, and (e) kidney cortex for control, and (b) lung, (d) liver, and (f) kidney cortex (20× magnification; hematoxylin and eosin stain) for Wistar–Kyoto rats administered 800 mg/kg whey protein concentrate hydrolysate. Scale bars represent 100 μm in each panel.
choke-derived enzymes. In another similar study, Wang et al. (2012) showed that alcalase-hydrolyzed WPC led to a maximum BP reduction of 37 mm Hg when dosed at 240 mg/kg, also showing improved antihypertensive activity compared with artichoke-derived enzymes. Enzyme selection plays a critical role in the release of antihypertensive peptides. For WPC, alcalase is a better choice for improving the release of these peptides.

Characterization of ACE-Inhibitory Peptides Identified from WPC

Physicochemical Properties. After we demonstrated the nontoxicity and BP-lowering efficacy of WPC hydrolysate in rat models, we subjected it to a sequencing protocol to look for potentially potent ACE-inhibitory peptides. We identified a total of 8 peptides from WPC: RHPEYAVSVLLR, GGAP-PAGIRI, GPPLPRL, ELKPTPEGDL, VLSELPEP, DAQ8APLRVY, RDMPIQAF, and LEQVLPRD. The physicochemical characteristics of these peptides are shown in Table 2. All peptides appeared as crystal white powder with good water solubility, consisted of 7 to 12 amino acids (oligopeptides), had an isoelectric point (pI) between 3.61 and 10.80, and a molecular weight of 748.92 to 1,439.67 Da. The net charge for peptides, measured at pH 7, correlated well with their respective pI: neutral peptides showed a pI close to pH 7, peptides with a negative charge showed pI values in the acidic region, and peptides with a positive charge showed pI values in the basic region. Instability index, which refers to the stability upon gastrointestinal digestion, is calculated using a mathematical formula that sums the instability weight value for dipeptide pairs in the sequence. Stability depends on the strength of the bond between amino acids (i.e., the smallest unit that contributes to a peptide bond). A peptide is considered stable if its instability index is less than 40 (Gasteiger et al., 2005). The current study found 3 stable peptides and 5 unstable peptides in WPC. However, instability does not dictate a peptide’s final ACE-inhibitory functionality upon reaching target site. According to Fujita and Yoshikawa (1999), peptides that are susceptible to gastrointestinal digestion may have improved or reduced ACE inhibition, depending on the nature of the new compound formed after digestion.

IC\textsubscript{50} Values. In terms of ACE inhibition, IC\textsubscript{50} denotes the peptide concentration required to exert half-

Figure 3. Light photomicrographs of male testis for (a) control and (b) treatment, viewed at 20× magnification (hematoxylin and eosin stain), and female ovary for (c) control and (d) treatment, viewed at 4× magnification (hematoxylin and eosin stain). Treatment groups were administered 800 mg/kg whey protein concentrate hydrolysate. Scale bars represent 100 μm in each panel.
maximal inhibitory activity against ACE. A high IC₅₀ implies a peptide with low inhibitory power, and low IC₅₀ implies a low inhibitory power. The IC₅₀ for 8 peptides are reported in Table 3. Two peptides—peptides 5 (VLSELPEP, IC₅₀ = 0.049 mM) and 8 (LEQVLPRD, IC₅₀ = 0.043 mM)—showed the most prominent ACE-inhibitory activity of the 8 WPC peptides. Factors affecting the peptides’ potency included positioning of
the amino acid within the sequence (C- or N-terminal), the types of amino acid present (hydrophobic vs. hydrophilic, aliphatic vs. aromatic) and peptide chain length. The presence of hydrophobic aliphatic amino acids such as leucine, isoleucine, and valine at the N-terminal, and aromatic amino acids such as proline, tyrosine, and tryptophan at C-terminal, would contribute to ACE-inhibitory activity. Also, positively charged lysine and arginine at the C-terminal would improve inhibitory power (FitzGerald et al., 2004; Vermeirssen et al., 2004). The peptides VLSELPEP and LEQVLPRD, showing the lowest IC50 (i.e., the strongest potency), contain valine and leucine at the N-terminal and proline and arginine at the C-terminal, which fit well with the structural requirements for ACE inhibition. Ibrahim et al. (2017) and Tavares et al. (2011), who worked on ACE-inhibitory peptides from goat and bovine whey protein, respectively, discovered several structural similarities with peptides obtained from the current work. For instance, QSLVYPFTGP reported by Ibrahim et al. (2017) demonstrated high amounts of hydrophobic amino acid and considerable amounts of proline residues, and KGYGGVSLEPW, DKVGINYW, and DAQSAPLRVY reported by Tavares et al. (2011) demonstrated aromatic character due to the presence of proline, tyrosine, and tryptophan at the C-terminal.

### CONCLUSIONS

The current study evaluated the acute and subacute toxicity of WPC hydrolysate produced enzymatically using alcalase. Our acute study showed that the median lethal dose (LD50) for WPC hydrolysate was more than 2,500 mg/kg. Daily observation, organ weight, clinical chemistry, and histopathological evaluations from our subacute study showed no significant difference between rats treated with WPC hydrolysate and controls. We concluded that alcalase-hydrolyzed WPC is non-toxic. Following this, we assessed the BP-lowering efficacy of WPC hydrolysate in spontaneously hypertensive rats and the results showed a promising in vivo efficacy that was a significant improvement compared to previous work, supporting the potential of alcalase-generated WPC hydrolysate as a safe, natural antihypertensive ingredient for incorporation into functional foods. We sequentially identified 8 peptides as marker peptides that contributed to ACE inhibition, of which 2 (VLSELPEP and LEQVLPRD) demonstrated IC50 < 0.05 mM and could be further evaluated.

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