Presence of the carcinogen ptaquiloside in fern-based food products and traditional medicine: Four cases of human exposure

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ABSTRACT

Ptaquiloside (PTA) is a natural carcinogen present in many ferns. Brackens (Pteridium sp.) contain PTA and are classified by WHO/IARC as ‘... possibly carcinogenic to humans’, however, these ferns are used in food, traditional medicine and as food supplements around the world. This study aimed to outline the presence of PTA in different human exposure routes by using and validating an LC-MS based protocol to test the contents of PTA in commercial products, the degradation product Pterosin B (PtB) and wild specimens from Europe, Asia and North America. The Limit of Detection of the protocol was 0.024 μg g⁻¹ for PTA and 0.028 μg g⁻¹ for PtB. PTA and PtB were present in most wild specimens (PTA: BD - 6300 ± 520 μg g⁻¹; PtB: BD - 449 ± 1 μg g⁻¹) while commercial products made from fronds, as well as fronds prepared as traditional Chinese medicine, were in the range 44 ± 3 to 666 ± 33 μg g⁻¹ for PTA and BD to 1653 ± 184 μg g⁻¹ for PtB. This study did not find PTA/PtB in rhizomes and products made thereof nor in homoeopathic products based on bracken. Boiling or drying bracken showed to reduce PTA some degree but cannot remove it completely. Interestingly, crosiers with no PTA/PtB were found in the USA, indicating a potential for commercial production of PTA-free fronds.

1. Introduction

Ptaquiloside (PTA) is the most prominent illudane-type glycoside (Table 1), which are found in a variety of ferns, but the toxicity and health effects have been most intensively studied in bracken ferns, a group of large perennial ferns with a global distribution. Farm animals tend to browse only on bracken ferns in times of food scarcity, as the group of large perennial ferns with a global distribution. Farm animals first syndromes affect cattle and buffaloes in paddocks or open grazing areas with bracken, while the latter is commonly encountered with free-ranging sheep in bracken infested land. The diseases are generally reported in countries like Brazil, Venezuela, Australia, New Zealand, Italy, Spain, Portugal, England and Scotland (GBIF.org., 2021; Marrs and Watt, 2006; O’Connor et al., 2019).

Bracken is suspected for causing human cancers and has therefore been classified as ‘possibly carcinogenic to humans’ by the WHO/IARC. Humans are exposed to the compounds when consuming bracken based traditional medicine (e.g. traditional Chinese medicine) or via ingesting bracken contaminated milk, meat, spores or drinking water, which is common in Asia (e.g. China, Korea and Japan) and North and South America (e.g. Brazil, USA and Canada) (Alonso-Amelot et al., 1993; Fletcher et al., 2011; GLIFWC, 2014; Kristanc and Kreft, 2016; Marliere et al., 2002; O’Driscoll et al., 2016; Skrbic, 2020; Virgilio, 2015). Due to the adverse health effects of bracken, the genus is placed on many national lists of plants not to be consumed by humans, and in Denmark, all parts of Bracken are classified as not acceptable for use as dietary supplements, in herbal teas or food (Gry et al., 1998). Bracken and PTA have also been placed on the WHO/IARC urgency list of potentially carcinogenic compounds/products to be re-evaluated (WHO/IARC, 2021).

Ptaquiloside is present in all parts of bracken ferns; fronds (leaves), rhizomes (below-ground stems), spores and roots, but the specific amount relies on the annual growth cycle and growing conditions. The fronds are known to have the highest concentration in spring, with contents reaching from 1000 to 50,000 μg g⁻¹ (dry-weight), while the content in rhizomes is negligible most of the year. The content may however increase in autumn due to nutrient translocation from fronds to storage rhizomes (bracken is a deciduous perennial fern). Bracken stands with no PTA have been found in New Zealand. There is a substantial variation in the distribution between illudane glycosides within, as well as between species of bracken. The content of PTA in spores and roots found in this is considered low compared to what is found in the fronds (Kisielius et al., 2019; Rasmussen et al., 2008, 2013, 2015).

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Ptaquiloside is relatively stable in bracken processed as comestibles despite undergoing acid, as well as alkaline hydrolysis, to form pterosin B (PtB; Table 1). The rate of hydrolysis is pH-dependent, leaving a window of stability for PTA in the range pH4.5 to 6.5. Degradation of PTA is strongly temperature-dependent and boiling for a long time will cause degradation of PTA (Ayala-Luis et al., 2006). Glycosidase enzymes can also cause PTA degradation. PtB becomes the end-product of the above-mentioned reactions, but other intermediates exist: Ptaquilosin and the so-called Bracken Dienone (O’Connor et al., 2019). PtB is present at similar levels as PTA in bracken fronds and can be used as an indicator of earlier presence of PTA in the compound (Rasmussen and Pedersen, 2017). Ptaquilosin and Bracken Dienone are believed to be the ultimate carcinogens (alkylating metabolites) and are highly unstable under acid and neutral pH. They are only found in neutral to alkaline matrices such as milk (Aranha et al., 2019).

Most studies addressing PTA in wild food products have been carried out in Japan or South America, but there is a lack of knowledge concerning the content of PTA in commercial products such as dried bracken and traditional medicines. The purpose of this study was to outline the presence of PTA in different human exposure routes by testing the hypothesis: Food products, natural remedies and traditional medicine made with bracken fern contain the carcinogen PTA. An LC-ESI-MS based dilute-and-shoot method has been validated to explore four exposure scenarios: Case 1) Using fresh bracken crosiers for food (fresh or after blanching); Case 2) Preparing traditional Chinese medicinal products using fresh bracken; Case 3) Collecting wild bracken in North America; Case 4) Exposure to PTA from commercial food products and European natural remedies obtained from online shops. The study is based on PTA/PtB standards/samples were kept at -20°C, as well as the accuracy of the aqueous extraction method, have been demonstrated elsewhere (Kisielius et al., 2020; Rasmussen and Pedersen, 2017).

The precision of the analytical method was reported as the RSD of 7–11 repetitive peak area measurements for 3 analytical standards: Ptaquiloside (PubChem CID: 53297436; Table 1); Pterosin B (PubChem CID: 115049; Table 1). The precision of retention time and peak asymmetry were recorded as part of the peak area measurements. The linear range was limited to Common Bracken (Pteridium aquilinum (L.) Kuhn) and PTA, as this is the dominant illudane-type glycoside in this bracken species (Kisielius et al., 2019). PtB was included as it is formed from PTA under natural conditions as well as due to sample handling.

2. Materials and methods

2.1. Chemicals and essential laboratory consumables

HPLC grade acetonitrile (Hiperpur HPLC hypergradient grade; Panreac, Barcelona, Spain); Sodium hydroxide, sodium acetate, trifluoroacetic acid (TFA) and hydrochloric acid (pro analysis; Sigma-Aldrich, United Kingdom). Deionised water was used for preparing reagents and plant extracts (except otherwise stated in the protocol below). LCMS-grade water was used for preparing eluents (ELGA Purelab® Classic, ELGA VEOLIA, Kruger Aquacare, Glostrup, Denmark). PTA was obtained from the University of Copenhagen (purity: 103 ± 10%; Professor Hans Christian Bruun Hansen, Denmark). Analytical standards of PtB were made from pure PTA (Rasmussen and Pedersen, 2017). PTA/PtB standards/samples were kept at -18 °C when not in use. 0.45 μm Q-Max Syringe Filter (cellulose acetate, sterile, Fresenette Aps, Knebel, Denmark). Polyamide 6 resin (analytical grade; Fluka, Steinheim, Switzerland). Whatman filter vials (0.20 μm filter vial; Whatman Mini-UniPrep PTFE filter vial, GE Healthcare UK Limited, United Kingdom).

2.2. Chemical compounds studied in this article

Ptaquiloside (PubChem CID: 53297436; Table 1); Pterosin B (PubChem CID: 115049; Table 1).

2.3. Method development and validation

The analytical method was based on Rai et al. (2017) and further optimised for use in food science using loganine as an internal standard. The method was validated according to the ICH guidelines (ICH, 2005) for PTA and PtB in terms of linearity, limit of detection (LOD), limit of quantification (LOQ), precision (analytical standards; samples) and matrix effects for different sample types (accuracy; PTA only). The ruggedness/storage stability of PTA/PtB standards and samples (-20 °C), as well as the accuracy of the aqueous extraction method, have been demonstrated elsewhere (Kisielius et al., 2020; Rasmussen and Pedersen, 2017).

The precision of the analytical method was reported as the RSD of 7–11 repetitive peak area measurements for 3 analytical standards: cPTA = 4.4/48/88 μg L⁻¹; cPtB = 4.4/44/88 μg L⁻¹) and a single bracken extract (cPTA = 76 μg L⁻¹; cPtB = 68 μg L⁻¹). In addition, the precision of loganine, used as an internal standard, was also measured (cLOG = 100 μg L⁻¹), and the precision of retention time and peak asymmetry were recorded as part of the peak area measurements. The linear range was explored in the expected relevant concentration range for PTA and PtB. Linearity was determined using 8 calibration levels in triplicate for PTA/PtB and singular for LOG (cPTA = 4.4–8.8 · 107.6–26.4 · 35.2–44–66–88 μg L⁻¹; cPtB = 4.4–8.8 · 17.6–26.4 · 35.2–44–66–88 μg L⁻¹; cLOG = 10–50· 100—150–200–200 μg L⁻¹); LOD/LOQ were calculated as 3/10 times the RSD of the resulting concentration from 7 repetitive injections (4.4 μg L⁻¹; concentration calculated from the response factor of the external/internal calibration curves). Analytical accuracy and check for matrix effects for PTA were determined using a representative sample of bracken fern extract spiked with PTA at 4 levels, corresponding to 0, 33, 66 and 100% of the PTA concentration (0–60 μg L⁻¹). The same setup was used to spike extracts of homoeopathic ampoules, globuli and rhizome flour. The extraction protocol was optimised for extracting PTA and PtB with respect to: Extraction solution (water; 50% methanol); pH of extraction solution (pH5.5; natural pH of extract); extraction time (20/40/60 min); sample-solute ratio (40 mg–40 mL; 100 mg to 40 mL).

Table 1

Properties of ptaquiloside and pterosin B.

| Compound:          | Ptaquiloside | Pterosin B |
|--------------------|--------------|------------|
| Molecular mass (avg., Da): | 398.447      | 218.292    |
| $\text{Log}_{10}K_{\text{ow}}$: | -0.95        | 3.19       |
| Water solubility (25 °C, mg/L): | 20,400          | 213        |
| Bioaccumulation estimate ($\text{Log}_{10}$BCF): | 0.50            | 0.27        |
| Bioactivity (Yahara et al., 2016; O’Connor et al., 2019): | Acute toxic; genotoxic. | Sik3 pathway inhibitor. |

* Estimate from US Environmental Protection Agency’s EPISuite™ (EPA, 2018).
2.4. Analytical set-up and identification protocol, LC-ESI-MS

The PTA and PtB were made on an Agilent 1260 Infinity HPLC System in combination with an Agilent 6130 Single Quadrupole (LC-MS). PTA and PtB were separated at 1.0 mL min \(^{-1}\). The ranges were chosen to represent expected sample solute ratios (40/100 mg to 40 mL), using 2 extract dilution factors which was then compared to MS spectra of pure samples (Figs. S1–2).

2.5. Supporting methods

pH was measured in plant extracts, soil suspensions and water using an ordinary pH combination electrode (section 2.8).

2.6. Case 1: Using fresh bracken crosiers for food - fresh or after blanching

Crosiers (46 unfolded young fronds; 15–40 cm; \(P. \text{aquilium}\)) were harvested in May 2017 in Ravnholt Forest (North of Copenhagen, Denmark). The identity of the ferns has previously been confirmed as Common Bracken in the mature state according to Frederiksen et al. (2012) and GBIF.org (2021). The fronds were kept cool (by a cooling box) and transported to the laboratory to be processed by blanching within 5 h (short time blanching or steaming are used worldwide with the crosiers). In addition, the Japanese method of adding wood ash/baking powder to the process water to increase alkalinity and its effect was also explored. The fronds were cut into approximately 5 cm long pieces and thoroughly mixed before randomly split into 5 sub-batches (each of approximately 50 g). Two batches were kept in reserve while the remaining material was processed as follows: 1) Reference; 2) Blanched for 2 min in approximately 0.5 L boiling 0.005 M CaCl\(_2\); 3) Blanched for 2 min by boiling 0.005 M CaCl\(_2\) plus commercial baking powder. The blanched material was cooled to room temperature immediately after cooking by soaking in cold tap water, pH was measured twice in all batches. PTA and PtB were extracted using a modified method by Caceres-Pena et al. (2013): Approximately 20 g fresh crosier was extracted with 100 mL 90 °C deionised water for 5 min in a kitchen blender, following centrifugation of 3 times 10 mL aliquots for 10 min at 9000 rpm and PTA/PtB measured in the supernatant (triplicate; final validated method). The samples were prepared within 15 min of cooking and kept at –18 °C until analysis.

2.7. Case 2: Preparing traditional Chinese medicinal products using fresh bracken

In March 2016, a sampling trip to the Shatin District of Hong Kong was conducted to collect crosiers and young unfurling ferns in nature. The identity of the specimens was confirmed to be Common Bracken (Frederiksen et al., 2012). The ferns were airdried for five days immediately after collection at approximately 25 °C in a well-ventilated room, mimicking a traditional way of drying ferns in private homes. The dried ferns were brought back to the University College Copenhagen and processed, from which the content of PTA/PtB was determined in triplicate using the final validated method. Samples were kept at –18 °C until analysis.

2.8. Case 3: Collecting wild bracken in North America

Eight sites used for collecting crosiers for consumption were located and bracken identified by staff from Great Lakes Indian Fish & Wildlife Commission (USA). Each site was divided into 1–3 homogeneous sub-sites (A–C) based on sun exposure (sunny; partial shade; full shade) and the soil texture, which were estimated based on field determination (Soil Survey Staff, 2014). A total of 7–10 young, unfolding crosiers were collected from each sub-site (May 2017), cut into 2–5 cm long pieces and dried for 24–32 h at 35–38 °C in a vegetable drier (9-tray Excalibur Food Dehydrator; Sacramento, California). The samples were subsequently ground in a Magic Bullet grinder (NutriBullet, LLC, USA). In addition, soil samples were collected from each sub-site and dried at air temperature before pH was measured in a 1:2.5 w/v 0.01 M CaCl\(_2\)-suspension. The dried ferns were shipped to the University College Copenhagen, and the content of PTA/PtB was determined using the final method. Samples were kept at –18 °C until analysis. The pH of the soil samples was determined by Great Lakes Indian Fish & Wildlife Commission (USA; single determination; Soil Survey Staff (2014)).

2.9. Case 4: Exposure to PTA from commercial food products and European natural remedies obtained from online shops

A list of Asian supermarkets and grocery shops in the Copenhagen region of Denmark, in addition to the major cities of Denmark, was outlined (autumn 2015/spring 2016), all of whom were visited or contacted by phone or email to inquire about bracken-based food products. From here, only noodles made of bracken rhizomes were obtained as it was the only product available. A variety of bracken-based food products were obtained from shops in the USA, Japan and China through international online shops (dried crosiers, rhizome flour and warabi sprinkle; 10 products in total). The products were produced in either Korea by Sam Heung Trade, Doo Me San, Assi and Choripdong, in Japan by Ohsawa, Itoku Food or Uji Maccha or in China from Zishu.The content of PTA/PtB in the products was determined using the final method in triplicate. Samples were kept at –18 °C until analysis.

Internet searches were used to identity European natural remedies containing bracken, and two providers were identified: WALA Heilmittel GmbH (Germany) and Remedia Homeopathy (Austria). From both providers, a variety of homeopathic solutes and pills (globules) were obtained using Remedia Homeopathy’s online shop (www.remedia-homeopathy.com) and the web-pharmacy, Apo-Rot (www.apo-rot.dk), and a total of 13 different products were purchased. As low concentrations of PTA and PtB were expected, solutes were analysed in triplicate according to the protocol using minimal dilution, and maximum sample injections were completed for qualitative screening (one sample out of each set of triplicates). Globules were dissolved in deionised water according to the protocol and analysed in triplicate according. Samples were kept at –18 °C until analysis.

2.10. In silico estimates and statistical analysis

The octanol-water partition coefficient, water solubility and bio-accumulation estimates were estimated from US Environmental Protection Agency’s EPISuite™ (EPA, 2013) and retrieved using ChemSpider (Royal Society of Chemistry (RSC, 2018)). Linear regression and descriptive statistics were performed in Microsoft® Excel® for Office 365 MSO (incl. data analysis package), while one-way analysis of variance and Tukey post hoc test were performed in Pisces Conservation Ltd® QED vs. 1.5.5.503.
A PTA-transformation ratio was calculated for all products to indicate how pre-processing and cooking affected the PTA-related toxicity:

$$\text{PTA}_{\text{trans}} = \frac{c_{\text{PTA}}/2}{(c_{\text{PTA}}/2) + c_{\text{PtB}}} \; \text{μg g}^{-1}$$  \hspace{1cm} (1)$$

The concentration of PTA was divided by two, as this is the approximate ratio of the molar weight between PTA and PtB. A PTA_{\text{proc}} value close to 1.0 indicated the limited transformation of PTA into PtB (change of molar balance) without taking intermediates or other transformation products into account. The value must be interpreted along with the processing effect regarding the total content of PTA/PtB, as complete destruction of the compounds may take place. It must be noticed that PtB may be present in unprocessed samples as well.

3. Results

3.1. Validation of analytical method incl. sample extraction protocol

The performances of the developed analytical methods are presented in Tables 2 and S1. The full chromatographic resolution was obtained using 53% MeOH and 47% 0.5 mM sodium acetate eluent at 1 mL min^{-1}, from which no interfering compounds were observed for any of the analytes in the chromatograms (Figs. S1–2). The performance of the external standard method for quantification of PTA and PtB was found to be equivalent to the internal standard method using loganine (LOG) as an internal standard.

PTA and PtB were linear with up to 1500 and 1250 μg L\(^{-1}\) respectively regarding the external and internal standard. Moreover, LOG proved linear in the range test (up to 200 μg L\(^{-1}\)) with linearity (r) ranging from 0.9991 to 0.9997. The calibration range used for quantification of PTA and PtB was set to 4.4 \(\pm 0.02\) for both compounds). During simple processing (blanching for 2 min), the PTA content was affected by the pH, which at pH 6.5 showed an approximately 17% reduction in PTA and an approximately 65% reduction at pH 7.9. Both treatments displayed a transformation ratio (PTA_{\text{trans}}) of 0.92, compared to 0.98 in the control. Hence, blanching increases the transformation of PTA to PtB and, at the same time, reduces the total content of PTA. However, the increase of PtB does not count for the decrease in PTA on a molar basis, indicating the formation of other metabolites or reaction products, like ptaquilosin, DNA-adducts or chloro-pterosin not quantified in the project.

The extraction of PTA and PtB was optimised and validated with respect to extraction solution composition, pH and extraction time (Tables S3–7), and the maximum yield of PTA and PtB was achieved after 40 min with no apparent effect using a buffer of pH 5.5 in aqueous extractions. A test using a buffered methanolic extraction solvent (the eluent: 53% MeOH, 47% 0.5 mM sodium acetate) revealed a negative effect on the PTA recovery compared to aqueous extraction. In conclusion, optimal PTA and PtB recovery is obtained using aqueous extraction for 40 min. This procedure was used for testing the sample-solvent ratio and the effect of the dilution factor on the yield (Tables S6–7). The results demonstrate that the dilution factor is important, possibly due to matrix interference in the MS detector, as approximately 25–30% lower concentration of PTA and PtB was measured using a dilution factor of 2.3 compared to 25. In contrast, no significant effects were found going from 40 mg to 40 mL to 100 mg to 40 mL.

The final extraction protocol was as follows: 40–100 mg sample was extracted using 40 mL deionised water for 40 min in a centrifuge tube, and the extraction was diluted 25 times resulting in a final composition of the analytical solution of 100 μL L\(^{-1}\). LOG, 0.1 M ammonium acetate and 48% MeOH. The samples were prepared and filtered in Whatman Mini-UniPrep PTFE 0.2 μm filter vials and kept at \(-20^\circ\)C until analysis. The LOD of the entire protocol was 0.024 μg \(^{-1}\) PTA and 0.028 μg \(^{-1}\) PtB. The accuracy is approximately 100%, while the precision of the complete extraction method including the instrumental analysis was 7.8–12.7% for PTA and 6.8–9.5% for PtB (depending on the sample-solvent ratio; Tables S2, S6–7).

3.2. Case 1: Using fresh bracken crosiers for food - fresh or after blanching

The fresh, unprocessed crosiers which were used as a control had a content of 6300 ± 520 μg \(^{-1}\) PTA (Table 3). There were significant effects of the treatments on the content of PTA and PtB (p < 0.05 for both compounds). During simple processing (blanching for 2 min), the PTA content was affected by the pH, which at pH 6.5 showed an approximately 17% reduction in PTA and an approximately 65% reduction at pH 7.9. Both treatments displayed a transformation ratio (PTA_{\text{trans}}) of 0.92, compared to 0.98 in the control. Hence, blanching increases the transformation of PTA to PtB and, at the same time, reduces the total content of PTA. However, the increase of PtB does not count for the decrease in PTA on a molar basis, indicating the formation of other metabolites or reaction products, like ptaquilosin, DNA-adducts or chloro-pterosin not quantified in the project.

### Table 2

| Analyte     | Linear range (μg L\(^{-1}\)) | Calibration curve | Linearity (r) | Calibration range (μg L\(^{-1}\)) | LOD (μg L\(^{-1}\)) | LOQ (μg L\(^{-1}\)) |
|-------------|-----------------------------|-------------------|---------------|----------------------------------|---------------------|---------------------|
| **External standard methods** | | | | | | |
| Ptaquiloside | 0–1500 | 646.29 \(\pm\) 213.07 | 0.9999 | 4.4–88.0 | 0.6 | 1.9 |
| Pterosin B | 0–1250 | 788.43 \(\pm\) 228.24 | 0.9997 | 4.4–88.0 | 0.7 | 2.4 |
| Loganine | 0–200 | 916.16 \(\pm\) 3187.6 | 0.9995 | 10.0–200.0 | ND | ND |
| **Internal standard method (LOG 100 μg L\(^{-1}\))** | | | | | | |
| Ptaquiloside | 0–1500 | 0.0074 \(\pm\) 0.0005 | 0.9991 | 4.4–88.0 | 0.6 | 2.0 |
| Pterosin B | 0–1250 | 0.0096 \(\pm\) 0.0001 | 0.9997 | 4.4–88.0 | 1.0 | 3.2 |

- Limit of detection.
- Limit of quantification.
- Correlation coefficient.
- Only tested in relevant concentration range when used as an internal standard.
- Loganine.
Table 3

| Treatment          | pH of water | Ptaquiloside (μg g⁻¹)* | Pterosin B (μg g⁻¹)* | PTAtrans |
|--------------------|-------------|------------------------|----------------------|----------|
| Control (not       | ND          | 6292 ± 521*            | 68±4*               | 0.98     |
| processed)         |             |                        |                      |          |
| Blanched (2 min)   | 6.50 ± 0.08 | 5261 ± 132             | 249±8               | 0.92     |
| Blanched (2 min)   | 7.91 ± 0.09 | 2196 ± 67              | 79±3*               | 0.93     |
| ANOVA P-value      |             | 9.4E-6                 | 0.7E-8              |          |

A to D Mean contents with different letters are statistically different (Tukey post hoc test of each compound, α = 0.05).

Table 4

| Sample type        | Ptaquiloside (μg g⁻¹) | Pterosin B (μg g⁻¹) | PTAtrans |
|--------------------|-----------------------|---------------------|----------|
| Crosiers           | 638 ± 90*             | 775 ± 158*          | 0.29     |
| Unfurling pinnae   | 736 ± 57*             | 312 ± 19*           | 0.55     |
| Mature pinnae      | 294 ± 128*            | BD                   | 1.00     |
| ANOVA P-value      | 7.7E-6                | 1.2E-4              |          |

A to D Mean contents with different letters are statistically different (Tukey post hoc test of each compound, α = 0.05).

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3.3. Case 2: Preparing traditional Chinese medicinal products using fresh bracken

Whole crosiers and pinnae were processed as traditional medicine after harvested in the wild, with a range of PTA from 294 to 638 μg g⁻¹ (Table 4). The mature pinnae displayed the lowest amount of PTA, while crosiers and unfurling had approximately the same amount. No PtB was found in the mature pinnae in contrast to crosiers and unfurling pinnae, which showed rather high contents of PtB (177–775 μg g⁻¹). Significant differences in the content of PTA and PtB was observed for the different growthstages (p < 0.05 for both compounds). The transformation ratio was lowest in the crozier and reached unity in the mature pinnae. The result is somewhat surprising as older plant parts are believed to have higher contents of pterosins, although this could also be a result of the processing. The crosiers and the unfurling pinnae had higher water content compared to the mature pinnae, which indicate that the older plants die more quickly and have lower enzymatic activity than the young plant, resulting in no formation of PtB. Assuming all PTA in the fresh crosier was present in the dried material as PTA or PtB, the original content ranged between 280 and 2030 μg g⁻¹.

3.4. Case 3: Collecting wild bracken in North America

Twenty bracken stands were sampled from eight different locations used for harvesting crosiers in spring (Table 5). The crosiers are usually prepared fresh in the region, but for practical reasons, the PTA and PtB were measured in dried material. The content of PTA ranged from 0 to 273 μg g⁻¹ while PtB ranged from 37 to 449 μg g⁻¹, and the transformation ratio varied 0.06 to 0.48 despite using strict protocols in the laboratory, and it is, therefore, possible that a portion of PtB was produced before the harvest. The results demonstrate that crosiers may include substantial amounts of PTA, though considering that the drying process transforms PTA into PtB, the content of PTA in crosiers might be even larger. Interestingly, a high number of sites did not have a detectable content of PTA nor PtB, and they can, therefore, be used to harvest “safe” bracken, provided no other toxins of the illudane-type are present in the ferns.

The content of PTA/PtB in crosiers with PTA/PtB was not strongly correlated with environmental co-factors such as soil pH, light exposure nor soil type. However, the average pH of stands with PTA/PtB was 5.2 ± 0.5, while the locations with no PTA/PtB had a soil pH of 4.6 ± 0.3, indicating that soil pH may influence the presence of toxins in the crosiers. Looking at the normalised PTA equivalents for the positive sites, a statistically significant difference is found between the sunny and the fully shaded sites, with the partially shaded sites in-between indicating lower contents of PTA in crosiers harvested from sunny locations compared to crosiers emerging in the shade of trees and bushes (Table S8).

3.5. Case 4: Exposure to PTA from commercial food products and European natural remedies obtained from online shops

In total, 22 commercial food products and natural remedies were tested for PTA and PtB (Table 6), though only dried bracken crosiers from Korea were found to contain the compounds. The reminding part of the products (homeopathic solutes, pills, rhizome flour or sprinkle) contained no PTA nor PtB above the Limit of Detection and can be considered safe in relation to PTA toxicity.

The content of PTA in the dried crosiers ranged from 44 to 666 μg g⁻¹ with a low ptaquiloside transformation ratio (0.04–0.19), indicating the general transformation of PTA into PtB. The content of PtB was found to be in the range 473 to 1653 μg g⁻¹. Assuming all PTA present in the crosiers were either present as PTA or transformed into PtB, the PTA content in fresh Korean crosiers ranged from approximately 900 to 3200 μg g⁻¹.

4. Discussion

Humans and bracken ferns have co-existed for millennia, and bracken has played a prominent part in human culture in many parts of the world. A brief contemporary review of bracken ethnobotany in Europe, Asia and North America can be found in Supplementary Information (Section 3). Historical founds date the use of bracken back centuries ago, in which bracken has been found in the stomach of a 5300-year-old glacier mummy, Otzi (Maixner et al., 2018). In Europe, the fern rhizomes have mainly been used as a food supplement during war and famine, in which flour was made from the rhizomes, while fronds have been used as bedding, in religious rituals, as bleach, thatch and a source of potash. Crosiers, however, do not seem to have been used for any purpose at all (Brondgegaard, 1978; Kristanc and Kreft, 2016; Rymer, 1976). Brackens is continually considered an ingredient in traditional medicine and the local diet for people in some parts of the world, particularly in Asia, from where numerous industrial products can be obtained (Table S and Supplementary Information Section 3). Bracken is, therefore, grown industrially all over the world to satisfy market needs in countries like China, Japan and Korea (Liu et al., 2012).
Commercial bracken products as well as freshly harvested crosiers. The alkaline hot-water treatment (\( O_\text{bracken} \) is known to contain natural genotoxic illudane glycosides mirrored in the number of recipes and videos. Obtained via colleagues in the USA. Internet searches revealed a high number of hits on bracken and bracken recipes, including a high number that bracken is collected in nature, that bracken can be obtained from Asia and Europe was an easy feat, while products from Amazon-based in the USA could not be directly shipped and were, therefore, sometimes using alkaline water. This study demonstrates that fresh parts of the world typically include soaking and boiling/steeping, including crosiers, contained PTA as starting point. These findings support results from other studies concerned with bracken, where drying is used as a preparation method for sampling (e.g., Rasmussen et al., 2020; Rasmussen et al., 2015). Industrial drying processes seem to lower the content of PTA as indicated in the transformation rates found in commercial products, as well as in home-dried crosiers using kitchen-scale driers (Tables 4–5). Hence, drying seems to be an effective mean for lowering PTA, provided no equally carcinogenic compounds are formed during the process.

Several products, mainly those based on rhizomes, were found to contain no PTA. The content may vary depending on the season, and high levels have only been observed in autumn. Rhizomes and rhizome-based products could be considered safe concerning PTA toxicity and when prepared from rhizomes collected in winter, which seems to be the practice in China. This also applies to European natural remedies, which was not surprising, as these were all homeopathic products (Table 5).

5. Conclusion

In this study, an LC-ESI-MS dilute-and-shoot method was validated regarding quantification of PTA and PtB in food products, traditional medicine and natural remedies, along with a novel protocol for extracting PTA and PtB. The analytical method proved as successful as other LC-MS/MS methods in relation to PTA/PtB quantification, and the combined method was used to explore human exposure to bracken toxins in four cases:

Case 1. - Using fresh bracken crosiers for food - fresh or after blanching: Fresh bracken crosiers displayed very high contents of PTA. Blanching reduced the PTA significantly, and the effect was enhanced when adding baking powder to increase the pH. The formation of carcinogenic degradation products may, however, take place.

Case 2. - Preparing traditional Chinese medicinal products using fresh bracken: PTA was found in the dried materials and could cause danger to the consumers. The preparations had PTA in range with brackens

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Table 5

| Location | Ptaquiloside (\( \mu g \frac{g^{-1}}{g^{-1}} \)) | Pterosin B (\( \mu g \frac{g^{-1}}{g^{-1}} \)) | \( PTA_{\text{trans}} \) | pH | Light exposure | Soil type |
|----------|---------------------------------------------|------------------------------------------|----------------|-----|----------------|-----------|
| Site 1:  |                                             |                                           |                 |     |                |           |
| 01A      | 128 ± 30                                   | 262 ± 12                                 | 0.20 ± 6.2      | Sunny | Loam          |
| 01B      | 218 ± 35                                   | 449 ± 1                                  | 0.20 ± 5.3      | Partial Shade | Clay        |
| 01C      | 273 ± 72                                   | 405 ± 88                                 | 0.25 ± 5.2      | Full Shade | Not recorded |
| Site 2:  |                                             |                                           |                 |     |                |           |
| 02A      | 24 ± 3                                     | 49 ± 1                                   | 0.20 ± 6.0      | Partial Shade | Sand/Loam/Clay |
| 02B      | 67 ± 2                                     | 57 ± 12                                  | 0.37 ± 5.9      | Full Shade | Loam/Sand |
| 02C      | 43 ± 6                                     | 42 ± 7                                   | 0.34 ± 6.1      | Sunny | Loam/Sand |
| Site 3:  | 03A                                         | BD                                        | –               | 4.4  | Sunny | Sand/Clay |
|          | 03B                                         | BD                                        | –               | 4.2  | Partial Shade | Sand/Clay |
| Site 4:  |                                             |                                           |                 |     |                |           |
| 04A      | BD                                         | BD                                        | –               | 4.0  | Full Shade | Sand/Clay |
| Site 20: | 20A                                         | BD                                        | –               | 4.5  | Partial Shade | Silt/Loam |
| Site 21: | 21A                                         | 35 ± 2                                   | 276 ± 37       | 0.06 ± 3.9 | Full Shade | Silt/Loam |
|          | 21B                                         | 25 ± 0                                   | 161 ± 20       | 0.07 ± 4.3 | Sunny | Silt/Loam |
|          | 21C                                         | 71 ± 0                                   | 241 ± 10       | 0.13 ± 4.2 | Partial Shade | Silt/Loam |
| Site 40: | 40A                                         | 75 ± 0                                   | 62 ± 0         | 0.38 ± 4.8 | Partial Shade | Sand |
|          | 40B                                         | 67 ± 0                                   | 37 ± 0         | 0.48 ± 4.9 | Full Shade | Sand |
|          | 40C                                         | BD                                       | –              | 4.7  | Sunny | Sand |
| Site 41: | 41A                                         | BD                                       | –              | 5.2  | Full Shade | Loam |
|          | 41B                                         | BD                                       | –              | 4.9  | Partial Shade | Sand |

\( ^\text{a} \) As pr. dry weight.
\( ^\text{b} \) Below Limit of Detection.
\( ^\text{c} \) Soil Survey Staff (2014).
collected from other parts of the world. This is the first report on PTA in the Chinese bracken.

Case 3. - Collecting wild bracken in North America: PTA was present in many specimens, thereby posing a danger for people collecting bracken in the investigated region. However, some bracken did not contain PTA indicating a possible option for growing PTA-free safe bracken, provided other illudane glycosides are not present. This is the first extensive report on PTA in the North American bracken.

Table 6

| Product | Ptaquiloside (μg g⁻¹) | Pterosin B (μg g⁻¹) | PTA_{mean} |
|---------|----------------|----------------|------------|

**Homeopathic solutes:**
- WALA. SALIX RHUS COMP. Ampoules N1. Germany.
- WALA. AQUILINUM COMP. Ampoules N1. Germany.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Dilution C4. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Dilution C200. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Dilution LM2. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Dilution LM6. Austria.

**Homeopathic pills:**
- WALA. SALIX RHUS COMP. Globuli relati N1. Germany.
- WALA. AQUILINUM COMP. Globuli relati N1. Germany.
- WALA. CONCHAE COMP. Globuli relati N1. Germany.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Globuli C4. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Globuli 50 MK. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Globuli LM2. Austria.
- REMEDIA HOMEOPATHY. Pteridium aquilinum Globuli LM6. Austria.

**Dried crosiers:**
- Sam Heung Trade. Dried fern brake. Korea. 175 ± 16 1653 ± 184 0.05
- Doo Me San. Dried bracken. Korea. 666 ± 33 1417 ± 125 0.19
- Assi dried fern bracken (pink package). Korea. 74 ± 16 830 ± 217 0.04
- Assi dried fern bracken (green package). Korea. 47 ± 3 473 ± 44 0.04
- Choripdong. Dried bracken. Korea. 44 ± 3 553 ± 47 0.04

**Rhizome flour and noodles:**
- Obasawa Rhizome flour. Japan. BD BD –
- Ioku Food Warabi Mochi Kit (flour). Japan. BD BD –
- Zishu Zhanhuangyuan noodles. China. BD BD –
- Warabi mochi sprinkle:
  - Ioku Food. Warabi Mochi Kit (powder). Japan. BD BD –
  - Uji Maccha. Warabi Mochi (powder). Japan. BD BD –

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