Detection of Adulteration in *Hydrastis canadensis* (Goldenseal) Dietary Supplements via Untargeted Mass Spectrometry-Based Metabolomics

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Abstract:

Current estimates report that approximately 25% of U.S. adults use dietary supplements for medicinal purposes. Yet, regulation and transparency within the dietary supplement industry remains a challenge, and economic incentives encourage adulteration or augmentation of botanical dietary supplement products. Undisclosed changes to the dietary supplement composition could impact safety and efficacy; thus, there is a continued need to monitor possible botanical adulteration or mis-identification. Goldenseal, *Hydrastis canadensis* L. (Ranunculaceae), is a well-known botanical used to combat bacterial infections and digestive problems and is widely available as a dietary supplement. The goal of this study was to evaluate potential adulteration in commercial botanical products using untargeted metabolomics, with *H. canadensis* supplements serving as a test case. An untargeted ultraperformance liquid chromatography-mass spectrometry (LC-MS) metabolomics analysis was performed on 35 *H. canadensis* commercial products. Visual inspection of the chemometric data via principal component analysis (PCA) revealed several products that were distinct from the main groupings of samples, and subsequent evaluation of contributing metabolites led to their confirmation of the outliers as originating from a non-goldenseal species or a mixture of plant materials. The obtained results demonstrate the potential for untargeted metabolomics to discriminate between multiple unknown products and predict possible adulteration.

Keywords: Metabolomics | Goldenseal | Adulteration | Dietary supplements | Mass spectrometry

Article:

***Note: Full text of article below***
Detection of adulteration in *Hydrastis canadensis* (goldenseal) dietary supplements via untargeted mass spectrometry-based metabolomics

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**ABSTRACT**

Current estimates report that approximately 25% of U.S. adults use dietary supplements for medicinal purposes. Yet, regulation and transparency within the dietary supplement industry remains a challenge, and economic incentives encourage adulteration or augmentation of botanical dietary supplement products. Undisclosed changes to the dietary supplement composition could impact safety and efficacy; thus, there is a continued need to monitor possible botanical adulteration or mis-identification. Goldenseal, *Hydrastis canadensis* L. (Ranunculaceae), is a well-known botanical used to combat bacterial infections and digestive problems and is widely available as a dietary supplement. The goal of this study was to evaluate potential adulteration in commercial botanical products using untargeted metabolomics, with *H. canadensis* supplements serving as a test case. An untargeted ultraperformance liquid chromatography-mass spectrometry (LC-MS) metabolomics analysis was performed on 35 *H. canadensis* commercial products. Visual inspection of the chemometric data via principal component analysis (PCA) revealed several products that were distinct from the main groupings of samples, and subsequent evaluation of contributing metabolites led to their confirmation as originating from a non-goldenseal species or a mixture of plant materials. The obtained results demonstrate the potential for untargeted metabolomics to discriminate between multiple unknown products and predict possible adulteration.

**1. Introduction**

Dietary supplements have become a focal point in personal medicinal care, with natural products becoming increasingly prevalent within the industry. Approximately 25% of Americans take a dietary supplement as part of their everyday health regimen [Asher et al., 2017; Newman and Cragg, 2016; Smith et al., 2016]. In the United States, the prevailing regulatory structure at the Food and Drug Administration (FDA) views herbal supplements as food rather than pharmaceuticals [Dietary Supplement Health and Education Act, 1994]. As such, the evaluation and reporting of adverse events remains with the manufacturer; the FDA does not generally perform pre-market testing on dietary supplements [Dietary Supplement Health and Education Act, 1994]. However, the FDA can act to remove any adulterated supplements from the market if proof of adulteration has been established [Dietary Supplement Health and Education Act, 1994]. As an example of some of the regulatory challenges surrounding dietary supplements, four stimulants - two 1,3-dimethylamylamine (DMAA) analogs and two banned stimulants (1,3-dimethylamylamine and 1,3-dimethylbutylamine) - were found in a study analyzing weight loss supplements [Choen et al., 2017]. The FDA had banned the 1,3-DMAA stimulant and removed any supplements containing the compound in August 2016 due to increased incidence of correlated emergency room visitations and because the conditions for it to be legally marketed had not been met [Choen et al., 2017]. However, analysis performed on other weight loss products still on the market after 2016 revealed analogs of 1,3-DMAA in five out of six commercial products tested [Choen et al., 2017]. This is just one case that illustrates how dietary supplements can be adulterated with potentially harmful compounds.

Goldenseal, *Hydrastis canadensis* L. (Ranunculaceae), is among the top 40 herbal supplements sold in the United States [Smith et al., 2016]. It is used to treat gastrointestinal disturbances, eye infections, and inflammation [Cicero and Baggioni, 2016; Le et al., 2013; Leyte-Lugo et al., 2017]. Root extracts of this botanical have demonstrated antimicrobial and antibacterial [Cicero and Baggioni, 2016; Le et al., 2013; Leyte-Lugo et al., 2017], as well as cytotoxic properties in vivo [Karmakar and Biswas, 2010; Le et al., 2013]. The root contains several benzylisoquinoline alkaloids, including berberine, hydrastine, and canadine [Cicero and Baggioni, 2016; Karmakar and Biswas, 2010; Le et al., 2013; Le et al., 2014; Leyte-Lugo et al., 2017]. Berberine is the...
most abundant alkaloid in H. canadensis roots and the antimicrobial
activity of H. canadensis has generally been attributed to this compound
[Brown and Roman, 2008; Junio et al., 2011]. Several flavonoids have
also shown to contribute to the antimicrobial activity of goldenseal,
working synergistically with the alkaloid berberine [Britton et al.,
2018; Junio et al., 2011].

Goldenseal dietary supplements are often harvested from wild pop-
ulations and those are available in limited quantities; a cultivated
plant takes years to fully mature and is expensive to farm [McGraw
et al., 2003; Tims, 2016]. Thus, there is an economic incentive to
adulterate goldenseal dietary supplements, and adulteration has be-
come a pertinent issue with the dietary supplement industry. Adul-
teration can involve spiking plant material with synthetic compounds,
using a different species in the same genus, or substituting a completely
different species in place of the stated one [Tims, 2016]. Several plants
that have been used to adulterate goldenseal supplements are barberry,
Berberis vulgaris L. (Berberidaceae), Chinese goldthread, Coptis chinensis
Franch. (Ranunculaceae), and Oregon grape, Mahonia aquifolium
(Pushr) Nutt. (Berberidaceae). These plants also produce berberine in
high quantities, yet possess distinctly different metabolic profiles from
that of goldenseal [McGraw et al., 2003; Tims, 2016; Weber et al.,
2003]. Material from berberine-producing plants has been known to
have been incorporated into dietary supplement capsules in lieu of
goldenseal [Pengelly et al., 2012; Tims, 2016; Weber et al., 2003].
Goldenseal, specifically the alkaloid hydrastine, has been shown to
inhibit two major metabolic enzymes, CYP2D6 and CYP3A4, which
metabolize approximately half of the drugs currently on the market
[Gupta et al., 2015]. Goldenseal dietary supplements could thus affect
the absorption, distribution, metabolism, and excretion of certain drugs
taken concomitantly, and the presence of an unknown species could
precipitate additional drug interactions that would not otherwise be
expected [Gupta et al., 2015; Gurlay et al., 2008]. The adulterated
supplement could also demonstrate different biological activity, or have
other side effects [Ciceri and Baggioni, 2016].

There have been several published methodologies used to detect
adulteration of goldenseal specifically. One method used HPLC-UV to
analyze and quantify analytes in the plant material, finding non-gold-
enseal constituents palmatine, coptisine, and jatrorrhizine [Tims,
2016]. The other method utilized GC-MS to quantify metabolites in
different commercial root products [Weber et al., 2003] using
various extraction solvents (hexane, chloroform, methanol, ethanol,
and water). Palmatine, jatrorrhizine, and coptisine, all compounds
found in plant species that also contain berberine, were found in one
out of three of the commercial samples tested [McGraw et al., 2003].
Utilizing FT-IR data in combination with different chemometric tech-
niques an in silico limit of detection was established at 5% adulteration,
depending on the adulterant species [Liu et al., 2018]. Comparing bot-
nanical material can also be achieved via genomic methods, e.g., DNA
barcoding. Barcoding has been shown to be an effective tool in au-
thentication of botanical and dietary supplements that are comprised of
fresh, dried, or powdered material, where intact DNA sequences are still
present [Coutinho Moraes et al., 2015; Little, 2014; Raja et al., 2017].
However, the DNA barcoding approach is more difficult when applied
to botanical extracts, as the manufacturing process often leads to re-
moval or degradation of DNA or contamination with rice filler. DNA
barcoding is not feasible for processed botanical products where the
DNA is either not present or potentially highly degraded, or where there
are two or more species present [Coutinho Moraes et al., 2015; New
York State Office of the Attorney General, 2015; Parveen et al., 2016].

Metabolomics approaches are applied to characterize multiple small
molecule metabolites in a biological sample set simultaneously, typi-
cally involving spectroscopic or spectrometric analyses, with nuclear
magnetic resonance (NMR) spectroscopy or mass spectrometry (MS)
being the most common analytical inputs [Sun et al., 2018]. Untargeted
metabolomics can be employed to analyze datasets when little is known
about the composition of the sample set and when the variance between
samples could be attributed to several sources [Kellogg et al., 2016; Tao
et al., 2018; Cappello et al., 2018]. Metabolomics can be used to dis-
tinguish one group of samples from another based on unique chemical
profiles, and has been applied to a wide scope of biological and chem-
ical applications, including identification of toxicological or disease
biomarkers [Sun et al., 2018], natural product drug discovery [Kellogg
et al., 2016], identification of secondary metabolites in Gram negative
bacteria [Depke et al., 2017] and characterization of botanicals (black
tea, green tea, ginseng, coffee) [Guo et al., 2018; Kellogg et al., 2017;
Lu et al., 2013; Souard et al., 2018; Zhang et al., 2018].

In this study, liquid chromatography-mass spectrometry (LC-MS)
data were utilized to perform untargeted metabolomics analysis on a
range of commercial goldenseal products, comprised of either the aerial
portion (leaves, stem), root, or rhizome. A variety of plant material
references (both goldenseal and other berberine-producing species)
were utilized to identify possible adulterated products. Though several
targeted techniques have been used to find adulterants in dietary sup-
plements [Simmler et al., 2017; Steuer et al., 2017], there has not yet
been a study performed to detect adulteration in a sample set of com-
mercial products, ostensibly derived from the same botanical, with an
untargeted approach. Untargeted metabolomics approaches have been
employed to discriminate between different botanical species as well as
variations in the geographic origin of materials [Kang et al., 2008;
Mncwangi et al., 2014], and studies adulterating pure botanical mate-
rial have shown discriminating patterns that can be discerned using
untargeted techniques [Dowlatabadi et al., 2017; Geng et al., 2015;
Geng et al., 2017]. However, little attention has been paid to com-
mercial botanical products, where neither the geographical prove-
nance, cultivation conditions, nor the harvesting and production spe-
cifications are known. This unknown information introduces a degree
of variability in product composition and could complicate efforts to
discern patterns and identify outliers, while an untargeted approach to
commercial products facilitates analysis without any a priori hy-
potheses on the nature of possible adulterants. All samples employed in
this study were commercial products marketed as Hydrastis canadensis
dietary supplements, and the only information regarding their com-
position was the label provided on the package by the manufacturer.
Additionally, studying commercial supplements highlights the direct
connection between possible adulteration and naïve consumption by
the consumer. The goal of these studies was to employ untargeted
metabolomics analyses to distinguish potential adulteration in a batch
of 35 samples of commercial H. canadensis products simultaneously.

2. Materials and methods

2.1. General methods

All solvents and chemicals used were of reagent or spectroscopic
grade, as required, and obtained from Thermofisher Scientific
(Waltham, MA, USA). Berberine and hydrastine standards were pur-
chased from Sigma-Aldrich (St. Louis, MO, USA) and were found to
have a purity of 99% and 98% respectively. A canadine reference was
isolated and purified from H. canadensis as described previously [Leyte-
Lugo et al., 2017] and demonstrated purity of 79%. Purity was de-
termined via LC-UV.

2.2. Sample selection and reference materials

Commercial goldenseal products were selected based on their po-
pularity in online consumer sales reports [Amazon.com, 2017]. The 35
products included 19 capsules, six tinctures, eight powdered bulk ma-
terials, and two bagged teas (Table 1). Each sample was randomly
coded with an internal reference number (beginning with the letters
GS) to maintain manufacturer anonymity (Supplemental, Table S1).

Reference materials were obtained from commercial suppliers as
well as harvested by the investigators. Hydrastis canadensis leaf (GS-35)
Table 1

Botanical and physical characteristics of the 35 commercial goldenseal products. Preparation refers to the post-harvest treatment a sample received (drying, extraction followed by drying, or freeze-drying); formulation represents how the final material was packaged for the consumer. Botanical source relates to the physiological portion of the plant which was harvested and incorporated into the final product. Botanical source as reported by manufacturer.

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Products (%)</th>
<th>Formulation</th>
<th>Products (%)</th>
<th>Botanical Source*</th>
<th>Products (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dried material</td>
<td>24 (69)</td>
<td>Capsule</td>
<td>19 (54)</td>
<td>Root</td>
<td>25 (72)</td>
</tr>
<tr>
<td>Botanical extract</td>
<td>9 (26)</td>
<td>Tincture</td>
<td>6 (17)</td>
<td>Rhizome</td>
<td>4 (11)</td>
</tr>
<tr>
<td>Freeze-dried material</td>
<td>2 (5)</td>
<td>Powder/loose material</td>
<td>8 (24)</td>
<td>Herb/leaf</td>
<td>4 (11)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Tea</td>
<td>2 (5)</td>
<td>Aerial parts</td>
<td>1 (3)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Not specified</td>
<td>1 (3)</td>
</tr>
</tbody>
</table>

and root (GS-36) samples were purchased from ChromaDex (Irvine, CA, USA). In addition, H. canadensis material was collected in August 2016 from William Burch in Hendersonville, North Carolina (NC, N 35°24.277′, W 082°20.993′, 702.4 m elevation), and a voucher specimen was deposited with the Herbarium of the University of North Carolina at Chapel Hill (accession: NCU583414) and authenticated by Dr. Alan S. Weakly. Leaf (GS-37) and root (GS-38) samples were dried in air at room temperature for several weeks prior to extraction.

Reference material from common goldenseal adulterants was also purchased from ChromaDex. These samples included Capsis chinensis rhizome (GS-39) and root (GS-40) samples, Mahonia aquifolium leaf (GS-41) and root (GS-42) references, and Berberis vulgaris root (GS-43) samples. All reference materials were obtained as dried powders, and performed in triplicate. Samples were shaken for 24 h then 20 mL of methanol were added. Extractions were performed in triplicate. Samples were weighed into scintillation vials (200 mg of material per sample) and 20 mL of methanol were added. Extractions were performed in triplicate. Samples were shaken for 24 h then filtered with 13 mm Puradisc Whatman (GE Healthcare, Chicago, IL, USA) syringe filters. Drying of extracts was accomplished under N₂ gas, and they were stored at room temperature prior to analysis.

2.3. Sample extraction

Samples were weighed into scintillation vials (200 mg of material per sample) and 20 mL of methanol were added. Extractions were performed in triplicate. Samples were shaken for 24 h then filtered with 13 mm Puradisc Whatman (GE Healthcare, Chicago, IL, USA) syringe filters. Drying of extracts was accomplished under N₂ gas, and they were stored at room temperature prior to analysis.

2.4. Mass spectrometry analysis

Liquid chromatography tandem to mass spectrometry (LC-MS) data were acquired utilizing a Q Exactive Plus quadrupole-Orbitrap mass spectrometer (ThermoFisher Scientific) with an electrospray ionization (ESI) source coupled to an Acquity UPLC system (Waters, Milford, MA, USA). Samples were resuspended in CH₃OH to a concentration of 1 mg/mL (expressed as mass of extract per volume solvent). Injections of 3 μL were performed on an Acquity UPLC BEH C₁₈ column (1.7 μm, 2.1 × 50 mm, Waters) with a flow rate of 0.3 mL/min using a binary solvent gradient of H₂O (0.1% formic acid added) and CH₃CN (0.1% formic acid added): initial isocratic composition of 95:5 (H₂O:CH₃CN) for 1.0 min, increasing linearly to 0:100 over 7 min, followed by an isocratic hold at 0:100 for 1 min, gradient returned to starting conditions of 95:5 and held isocratic again for 2 min. The positive/negative switching ionization mode of the mass spectrometer was utilized over a full scan of m/z 150–2000 with the following settings: capillary voltage, 5 V; capillary temperature, 300 °C; tube lens offset, 35 V; spray voltage, 3.80 kV; shear gas flow and auxiliary gas flow, 35 and 20 units, respectively. Extracted ion chromatographs were obtained from the XCalibur software (ThermoFisher Scientific).

2.5. Metabolomic analysis

The LC-MS data were analyzed, aligned, and filtered using MZmine 2.28 software (http://mzmine.github.io/) with a slightly modified version of a previously reported method [Kellogg et al., 2017]. The following parameters were used for peak detection: noise level (absolute value), 1 × 10⁵ counts; minimum peak duration 0.5 min; tolerance for m/z intensity variation, 20%. Peak list filtering and retention time alignment algorithms were performed to refine peak detection. The join algorithm was used to integrate all the chromatograms into a single data matrix using the following parameters: the balance between m/z and retention time was set at 10.0 each, m/z tolerance was set at 0.001, and retention time tolerance was defined as 0.5 min. The peak areas for individual ions detected in triplicate extractions were exported from the data matrix for further analysis. Samples that did not contain a particular marker ion were coded with a peak area of 0 for that variable to maintain a consistent number of variables throughout the dataset. Chemometric analysis was completed using Sirius version 10.0 (Pattern Recognition Systems AS, Bergen, Norway). Transformation from heterogeneous to homoscedastic noise was carried out by a fourth root transformation of spectral variables. Principal component analysis (PCA) was used for untargeted metabolomics profiling of the goldenseal samples with Sirius software. The 95% confidence interval was calculated using Hotelling’s T² in R with the R package ‘car’ [Fox and Weisberg, 2011]. Heatmap construction was performed on the log-transformed, mean-centered peak area data for each relevant metabolite, using the R package ‘gplots’ [Warnes et al., 2016].

2.6. Compound identification

Variables, unique m/z value and retention time (m/z-RT) pairs, present in the loadings plot were used to confirm and explain the variance in the corresponding scores plot. These ions were identified by using exact mass (< 5 ppm) and retention time. The compounds (berberine [1], canadine [2], hydrastine [3], cotisine [4], palmatine [5], jatrorrhizine [6], and dihydrocoptisine [7]) are all known and well documented (Fig. 3). The m/z-RT pairs were compared and confirmed with literature values.

3. Results and discussion

3.1. Identification of goldenseal outliers by untargeted mass spectrometry metabolomics

Untargeted metabolomic analysis of the goldenseal samples using LC-MS yielded 5423 marker ions (unique retention time—m/z ion pairings) for 117 objects (35 commercial goldenseal samples and four goldenseal reference materials extracted in triplicate), which were statistically modeled using PCA. The extraction replicates of each goldenseal product were overlaid on the PCA plot (Supplemental Figure S5), indicating repeatability of the extraction technique and subsequent LC-MS analysis. The three extractions were averaged for subsequent PCA analysis. The 4-component PCA model accounted for 68.0% of the variance in the sample set.

Inspection of the data based upon the botanical source of each sample (e.g., leaf, aerial portion, root, rhizome, whole plant) indicated two distinct sample clusters located in different regions of the two-dimensional space prescribed by principal component 1 (PC1, 25.0% variability explained) and principal component 2 (PC2, 17.3% variability explained) (Fig. 1). The authenticated goldenseal reference samples (commercial and vouchered leaf and rhizome material) also clustered with their commercial counterparts (aerial (green) and root/
3.2. Tentative identification of unique compounds in outlier samples based upon PCA loadings plot

The separation observed in the PCA scores plot of PC1 versus PC2 (Fig. 1) can be explained through metabolites highlighted in the corresponding PCA loading plots (Fig. 2). The loadings plot graphically estimates the degree to which each variable (RT-m/z pair) contributes to the separation of three samples in the PCA scores plot; the greater the magnitude of a variable's loading score, the more it contributes to that principal component. Graphing a corresponding PCA loadings plot highlights marker ions that are associated with the observed clustering [Lever et al., 2017].

The PCA loadings plot highlighted hydrastine (m/z 384.1459 [M + H]+) and canadine (m/z 340.1537 [M + H]+) as having significant contributions towards the first principal component (PC1) due to their large magnitude in the x-direction. The loadings plot also included the 13C isotope peaks for each of the metabolites, lending additional confidence to the identification and significance of these compounds. A dimer of hydrastine was also present. Hydrastine and canadine are two of the main alkaloids present in goldenseal, root and leaf, and were found to be missing in the outlier samples [Le et al., 2013; Le et al., 2014]. The large cluster of goldenseal supplements, both of leaf and root/rhizome, were separated from the outliers along the positive x-axis due to the presence of these major alkaloids. These data suggest that the outliers could possess a possible mixture of plant material and/or a lack of *Hydrastis canadensis*.

A series of outliers were observed to lie along the y-axis direction, contributing to the observed variance along the second principal component (PC2). PC2 was also responsible for discriminating outlier samples from goldenseal supplements (Fig. 1). The loadings plot revealed several metabolites that were present in higher concentrations in the adulterated samples, and thus were dominant peaks in the positive direction. These included coptisine (m/z 321.0954 [M + H]+), palmitine (m/z 352.1535 [M + H]+), their 13C isotopes (m/z 322.1072 [M + H]+ and 353.1571 [M + H]+), respectively, and dihydrocoptisine (m/z 323.1121 [M + H]+) (Fig. 3). All of these compounds have been previously shown to be present in other berberine containing species, specifically *Coptis chinensis*, *Berberis vulgaris*, and *Mahonia aquifolium* [Ivanovska and Philipov, 1996; Pengelly et al., 2012; Račková et al., 2004; Weber et al., 2003; Yang et al., 2017]. Their presence in the outliers (GS-07, GS-20, and GS-33) and absence in other goldenseal supplements and reference material was supported by the heat map (Fig. 5) and the corresponding stacked mass spectrometry chromatograms (Fig. 6).

3.3. Adulteration analysis with reference materials

Reference materials for non-*H. canadensis* species were extracted and incorporated into the metabolomics analysis. These included *Coptis chinensis* rhizome (GS-39) and root (GS-40), *Mahonia aquifolium* leaf (GS-41) and root (GS-42), and *Berberis vulgaris* root (GS-43). The resulting dataset contained 5573 marker ions for 135 objects (i.e., 35 goldenseal products, nine reference materials, and a process blank, all prepared by extraction in triplicate). After analysis of the reproducibility of the extraction method the average response of the triplicate extractions was taken to yield a final 5423 × 45 dataset (RSD < 5%).

PCA analysis yielded a 4-component model explaining 62.6% of the variance contained within the dataset. Examining the position of the non-goldenseal reference material within the PCA scores plot, the standards for *M. aquifolium* herb and root (GS-41 and GS-42, respectively) clustered closely to sample GS-07. Sample GS-33 was located in close proximity to the *C. chinensis* reference materials (GS-39 and GS-40) and all three were shifted from the *H. canadensis* clusters, suggesting GS-33 was a dietary supplement formulated from *C. chinensis* instead of the labeled *H. canadensis*. Additionally, GS-07 was found in a similar region to the *B. vulgaris* (GS-43) reference material, implying there was a mix of plant material present. After examining the position of the samples in the PCA scores plot, the corresponding loadings plot and mass spectrometry chromatograms provided additional substantive evidence as to the variables (ions) responsible for the observed variation.

Liquid chromatography-mass spectrometry (LC-MS) chromatograms provided corroborating evidence as to the nature of the three adulterated samples (Fig. 6). The base peak chromatogram for the *H. canadensis* reference material (GS-36 and GS-38) contained three main peaks...
in the positive ionization mode: berberine, m/z 336.1229 \([\text{M}]^+\), hydrastine, m/z 384.1440 \([\text{M+H}]^+\), and canadine, m/z 340.1545 \([\text{M+H}]^+\), all of which are characteristic marker compounds for goldenseal [29–30]. Berberine was consistently present across all root/rhizome samples regardless of putative botanical origin (Fig. 5); however, hydrastine and canadine levels were found to vary considerably. In the base peak chromatogram for GS-07, GS-20 and GS-33 (Fig. 6A, B, and 6C, respectively), the peaks for canadine and hydrastine were not present. Three additional peaks – tentatively identified as palmatine (m/z 352.1543 \([\text{M+H}]^+\)), coptisine (m/z 320.0917 \([\text{M+H}]^+\)), and dihydrocoptisine (m/z 322.1074 \([\text{M+H}]^+\)) based on accurate mass determination – were present in GS-07 (Fig. 6A), which coincided with ions having the same m/z (< 5.0 ppm) and retention time as the corresponding ions in reference material from B. vulgaris (GS-43) [Ivanovska and Philipov, 1996; Weber et al., 2003]. The second outlier supplement, GS-20, had a large berberine peak present but was lacking
hydrastine and canadine peaks (Fig. 6B). The high berberine content was believed to be responsible for the clustering of GS-20 with GS-41 and GS-42 (M. aquifolium leaf and root, respectively). As shown in the extracted ion chromatograms, M. aquifolium root also contains jatrorrhizine (m/z 338.1392 [M]+) and palmatine (m/z 352.1543 [M+H]+) both of which were found in GS-20 (Fig. 6C) [Račková et al., 2004; Weber et al., 2003]. The final outlier sample, GS-33, also displayed a spectral profile distinct from that of goldenseal [Weber et al., 2003; Yang et al., 2017] (Fig. 6C). Canadine and hydrastine were not present, but palmatine, coptisine, and dihydrocoptisine were also detected in the sample. This was consistent with C. chinensis, commonly known as Chinese goldthread. The overlap observed in these chromatograms with non-goldenseal alkaloids, and the absence of two principal goldenseal marker compounds (hydrastine and canadine) (see Table 2), supported the hypothesis from the untargeted metabolomic analysis that samples GS-07, GS-20, and GS-33 were adulterated.

4. Conclusion

Currently “there is a need to develop or extend existing analytical approaches to identify unexpected adulterants”, specifically in the dietary supplement industry [Pawar et al., 2017]. Untargeted metabolomics analysis of commercial goldenseal dietary supplements efficiently identified three samples as potentially being adulterated, without prior knowledge of composition or suspicion of adulteration. The altered status of samples GS-07, GS-20, and GS-33 was substantiated by incorporating standard reference materials into the metabolomics data analysis. This second, more detailed study revealed that these outliers possessed few goldenseal metabolites, instead containing other, non-goldenseal components. GS-07 appeared to contain B. vulgaris botanical material based upon the metabolite correlation found in the PCA scores plot (Fig. 4) and mass spectrometry chromatograms of relevant reference materials (Fig. 6A). GS-20, clustered closely to GS-42 (M. aquifolium root) and GS-41 (M. aquifolium herb) and shared several of the same non-goldenseal compounds (Fig. 6B), showing the supplement may contain M.
Fig. 6. Liquid chromatography-mass spectrometry (LC-MS) chromatograms from supplement samples GS-07 (A), GS-33 (B), and GS-20 (C) (suspected to be altered compared against reference material profiles for *H. canadensis* (GS-36), *C. chinensis* (GS-39), *B. vulgaris* (GS-43), and *M. Aquifolium* (GS-42). Highlighted alkaloid peaks are identified through comparison of accurate mass from high-resolution mass spectrometry in the positive ionization mode. GS-07 and GS-33 revealed peaks corresponding to berberine, palmatine, coptisine, and dihydrocoptisine, the last three of which are not found in the *H. canadensis* reference material. The *M. aquifolium* standard possessed berberine, palmatine, and jatrorrhizine; all of which were also present in GS-20 but the latter two were found in much lower concentrations than what was observed in the *M. aquifolium* reference material.
The marker ions in Hydrastis canadensis, Berberis vulgaris, Mahonia aquifolium, and Coptis chinensis with [M+H]⁺ values. Check marks represent whether or not the ions were present in the following goldenseal samples: commercial goldenseal supplements GS-07, GS-20, GS-33, GS-36 (Hydrastis canadensis root standard), GS-39 (Coptis chinensis rhizome standard), GS-42 (Mahonia aquifolium root material), and GS-43 (Berberis vulgaris root material).

<table>
<thead>
<tr>
<th>compound</th>
<th>GS-07</th>
<th>GS-20</th>
<th>GS-33</th>
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Table 2

There is a continued need to ascertain variability in complex botanical products, especially within the dietary supplement industry [Dietary Supplement Health and Education Act, 1994] to monitor quality control of products for adulteration [Tims, 2016]. Analyzing similarity and variation within commercial botanical supplement products remains a challenge due to their innate phytochemical complexity. Hydrastis canadensis supplements possess a variety of bioactive secondary metabolites, which vary depending on the plant portion used for formulation [Le et al., 2013; Leyte-Lugo et al., 2017]. This study illustrates the effectiveness of untargeted metabolomics methodologies to analyze this variability and differentiate outlier samples that could indicate possible adulteration in a large sample set. Multivariate analysis of the metabolomics dataset effectively modeled the variance between the goldenseal supplements based upon physiological origin of the product, as well as differentiating three potentially adulterated samples from the other supplements via the PCA scores plot (Fig. 1). Moreover, the distinction between the goldenseal supplements was achieved without any prior knowledge of the composition of the samples, nor of the identity of the possible adulterants. Subsequent investigation employing the PCA loadings plot (Fig. 2) and the stacked mass spectrometry chromatograms (Fig. 6C) yielded discriminative features (ions) that were responsible for the differentiation between sample groups, providing information that suggests the identities of the botanical adulterants. These marker ions were discovered from a large sample set comprised of commercial botanical dietary supplements (reported on the labels to contain only goldenseal). One potential disadvantage to untargeted metabolomics is the possibility of ion suppression and matrix effects, where co-eluting components can affect the ionization efficiency of one another [Jorge et al., 2016; Lei et al., 2011]. This predominantly impacts lower abundant compounds or compounds with poor ionization efficiency, but it can also enhance ionization. Both of these effects of ion suppression and matrix effects have the potential to compromise accurate quantitation of the analytes across a sample set [Jorge et al., 2016; Lei et al., 2011]. Steps can be taken to minimize the effect of ion suppression, including more refined chromatographic separation of analytes, an alteration to the ionization mode, or inclusion of an appropriate internal standard [Antignac et al., 2005]. However, targeted techniques would have required an a priori understanding of the identity of the adulterating botanicals and the relevant marker ions associated with each species, and untargeted metabolomics effectively provided a simultaneous comparison, which included more of the complex chemical profile and is much more efficient than a pair-wise comparison.

There is continued need to ascertain variability in complex botanical products, especially within the dietary supplement industry [Dietary Supplement Health and Education Act, 1994] to monitor quality control of products for adulteration [Tims, 2016]. The untargeted mass spectrometry-based metabolomics approach described herein has the potential to provide a versatile data acquisition tool for comparisons of multiple products with complex constituents. The collection of chromands of secondary metabolites represents a robust analytical technique that is capable of differentiating between closely-related samples [Dowlatabadi et al., 2017; Geng et al., 2015; Geng et al., 2017; Kang et al., 2008; Mncwangi et al., 2014; Simmler et al., 2017; Steuer et al., 2017], and employing untargeted follow up analyses to discern potential adulteration from multiple complex botanical matrices representing a potentially valuable application of this analytical methodology.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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Appendix A. Supplementary data

Supplementary data related to this article can be found at https://doi.org/10.1016/j.fct.2018.07.033.

Transparency document

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References


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