Compound Characterization and Metabolic Profile Elucidation after In Vitro Gastrointestinal and Hepatic Biotransformation of an *Herniaria hirsuta* Extract Using Unbiased Dynamic Metabolomic Data Analysis

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Abstract: *Herniaria hirsuta* L. (Caryophyllaceae) is used for treatment of urinary stones and as a diuretic. Little is known about the active compounds and the mechanism of action. The phytochemical composition of *H. hirsuta* was comprehensively characterized using UHPLC-UV-HRMS (Ultrahigh-Performance Liquid Chromatography-Ultraviolet-High Resolution Mass Spectrometry) data. An in vitro gastrointestinal model was used to simulate biotransformation, which allowed the monitoring of the relative abundances of individual compounds over time. To analyze the longitudinal multiclass LC–MS data, XCMS, a platform that enables online metabolomics data processing and interpretation, and EDGE, a statistical method for time series data, were used to extract significant differential profiles from the raw data. An interactive Shiny app in R was used to rate the quality of the resulting features. These ratings were used to train a random forest model. The most abundant aglycone after gastrointestinal biotransformation was subjected to hepatic biotransformation using human S9 fractions. A diversity of compounds was detected, mainly saponins and flavonoids. Besides the known saponins, 15 new saponins were tentatively identified as glycosides of medicagenic acid, acetylated medicagenic acid and zanhic acid. It is suggested that metabolites of phytochemicals present in *H. hirsuta*, most likely saponins, are responsible for the pharmaceutical effects. It was observed that the relative abundance of saponin aglycones increased, indicating loss of sugar moieties during colonic biotransformation, with medicagenic acid as the most abundant aglycone. Hepatic biotransformation of this aglycone resulted in different metabolites formed by phase I and II reactions.

Keywords: *Herniaria hirsuta*; Caryophyllaceae; saponins; urolithiasis; machine learning; dynamic metabolomics

1. Introduction

Urinary stone disease is considered as an important healthcare problem that affects 10–15% of the population in the developed world, but the incidence can be as high as 20–25% in the Middle East, with a peak at ages 20 to 40 years. In addition, the disease is characterized by its high recurrence rate, about 50% in 10 years and 75% in 20 years [1–6].
Over the years, many remedies and surgical treatments have been described, varying from dietary recommendations to interventional procedures. Endoscopic management, both ureteroscopic and percutaneous, offers an efficient and efficacious way to treat stones [6]. Nowadays, extracorporeal shock wave lithotripsy (ESWL) is widely used for stone fragmentation without invasive instrumentation of the body. However, the aforementioned therapies show some significant side effects such as residual stone fragments that can act as a nidus for new stone formation [5,7]. Existing oral treatments with evidence that supports their long-term efficacy to prevent kidney stones only include a handful of drugs. Thiazide diuretics, potassium citrate and allopurinol have been used as therapy for more than 30 years. However, their adverse effects prevent long term consumption [2,8].

Phytotherapy might be useful as an alternative treatment. Some herbal remedies have been used for centuries and have been shown to be effective against urolithiasis, although the mechanism of action is often not well established through systematic pharmacological and clinical studies. Many plants have been and are still used to treat urinary stones, such as *Phyllanthus niruri*, *Zea mays*, *Agropyron repens* and *Herniaria hirsuta* [5,9].

An aqueous extract of the aerial parts of *H. hirsuta* (hairy rupturewort) is an herbal medicine widely used against urolithiasis and which contains diuretic properties. The Arabic name, herras lehjer, is translated as stonebreaker and refers to its use as a traditional herbal remedy for urinary stones. The beneficial effects of the extract have been demonstrated in several studies [9–14]. Even though a lot of research has been done to prove the activity of *H. hirsuta* against urolithiasis, little is known about the active compounds and the exact mechanism of action. Previous phytochemical research on *Herniaria* species revealed the presence of saponins, flavonoids and coumarins [15–18]. Four herniariasaponins have been reported in *H. hirsuta*: herniariasaponins E, F and G, three monodesmosidic derivatives of medicagenic acid and herniariasaponin H, a bidesmosidic derivative of medicagenic acid [17,19]. Literature suggests that the antilithiatic potential of *H. hirsuta* is attributed to saponins or metabolites thereof [17,20,21].

It is well known that many natural products are so-called prodrugs, which become active after biotransformation [22]. Nevertheless, this aspect is usually overlooked when searching for new therapeutic agents using classical approaches. After oral administration, an herbal extract is inevitably brought into contact with gastrointestinal enzymes and intestinal microflora, which might lead to biotransformation of the compounds. Saponins are high-molecular-weight glycosides, containing a triterpenoid or steroidal sapogenin aglycone covalently linked to one or two sugar chains via an ether or ester glycosidic bond, referred to as monodesmosidic or bidesmosidic saponins, respectively. Before absorption in the gastrointestinal tract, saponins are presumably hydrolyzed with loss of sugar chains [23]. The aglycones or metabolites thereof may be absorbed and may further be biotransformed in the liver to the ultimate active components [24]. In this study, the biotransformation of a well-characterized *H. hirsuta* extract was investigated in an in vitro gastrointestinal model followed by hepatic biotransformation. The longitudinal multiclass data were subjected to different data analysis workflows to screen and tentatively identify metabolites formed after in vitro biotransformation.

### 2. Results and Discussion

#### 2.1. Identification of Compounds

A dereplication workflow performed on the extract of *H. hirsuta* resulted in a molecular network containing different clusters, composed of different nodes connected by edges which define the degree of similarity between the MS/MS spectra. Compounds that are structurally related are linked to each other. Mass differences between the nodes were studied and provided information about hitherto unidentified compounds. Major clusters were analyzed to tentatively identify unknown compounds. As an example, the cluster containing herniariasaponin H is shown in Figure 1. Herniariasaponin H (Figure 2) is the most abundant saponin previously observed in *H. hirsuta* and served as starting point for identification of other compounds in this cluster.
Structures were assigned to peaks only when both the mass/charge (m/z) ratios and the molecular formulae of the precursor ion and product ions were in agreement. Additional information about the proposed structures was often provided by photodiode array (PDA) spectra and retention times. However, the information was not always sufficient for full elucidation of the structural composition at an acceptable confidence level. Additional information was obtained from published data.

Figure 1. Molecular network in negative ion mode with the cluster containing herniariasaponin H shown in detail.

Figure 2. Chemical structure of herniariasaponin H.

All amMS and PDA data used for peak identification are listed in Table 1. The table also specifies the literature consulted to obtain additional information in order to confirm compound identity. In accordance with previous research, mainly flavonoids and saponins were identified [15–21]. However, many of the identified compounds have not been reported previously in H. hirsuta.

Flavonoids were mainly observed as glycoconjugates, i.e., a flavonoid aglycone attached to one or more sugar moieties. Several flavonoid compounds were identified using analytical standards. Others were tentatively identified with ddMS². Collision-induced fragmentation resulted in product ions caused by elimination of sugar moieties, indicating glycosidic O-linkages [24]. Among others,
rutin, narcissin and quercetin-3-O-(2''-O-α-L-rhamnopyranosyl)-β-D-glucuronopyranoside were identified, in agreement with previous studies [17,20]. However, isorhamnetin-3-robinobioside and isorhamnetin-3-[3-feruloylrhamnosyl-(1→6)-galactoside], previously reported to be present in *H. fontanesii*, were not detected in *H. hirsuta*, which is probably due to variances in phytochemical profile between different plant species [18].

A rich diversity of saponins was detected, differing in the presence of various aglycones, a different composition of sugar chains and varied linkages of sugar moieties. Fragmentation led to the distinct presentation of product ions caused by elimination of sugar moieties, indicating glycosidic O-linkages. The product ions in the MS² spectra provide additional structural information about the sugar residues and the aglycones of the fragmentated saponins. Losses of *m/z* 132, 146, 162 and 176 were observed, corresponding to pentose, deoxyhexose, hexose and uronic acid moieties, respectively. Three different aglycone moieties were observed with fragment ions at *m/z* 543.33234, 541.31787 and 501.32074, corresponding to acetylated medicagenic acid, zanhic acid and medicagenic acid, respectively [25,26].

Several other compounds, such as free phenolic and hydroxycinnamic acids, were detected. Amongst others, caffeic, *p*-coumaric and chlorogenic acids were identified using analytical standards. The coumarins herniarin and umbelliferone, previously reported to be present in *Herniaria* species, were not detected. This variation in compounds is probably due to differences in origin or regional climate [17,20]. More than one chromatographic peak was often present for the same precursor ion, indicating the presence of structural isomers with different linkage positions and/or different sugar moieties.

Figure 3 illustrates the identification of a hitherto unknown saponin. As shown in Figure 3A, the unknown saponin has a retention time of 15.33 min. Figure 3B shows a molecular ion at *m/z* 1703.71838 [M-H]⁻. Since saponins contain the elements C, H and O, 19 molecular formulas are possible within an error range of 5 ppm.

The sugar residues and the aglycone were identified from the MS² spectra as shown in Figure 3C and Table 2. The product ion of *m/z* 1381.62830 [M-H-176-146]⁻ results from the loss of uronic acid (176 u) and deoxyhexose (146 u).

The ion at *m/z* 879.30701 originates from a sugar chain containing two deoxyhexosyl (146 u), two pentosyl (132 u) and two hexosyl (162 u) units. The ion at *m/z* 823.41089 [aglycone-H+146+176]⁻ represents the aglycone attached to a deoxyhexose unit (146 u) and a uronic acid moiety (176 u). The ion at *m/z* 501.32275 is characteristic for medicagenic acid as aglycone, resulting from the loss of sugar residues. The ion at *m/z* 439.31934 [aglycone-H-18-44]⁻ results from an additional loss of water (18 u) and CO₂ (44 u). This suggests a bidesmosidic saponin with R₁ comprising uronic acid and deoxyhexose and R₂ comprising two pentosyl, two deoxyhexosyl and two hexosyl moieties, with a hexosyl moiety in a terminal position. Combining all the information with the retention time and literature data, the saponin with *m/z* 1703.71734 supports a molecular formula of C₇₆H₁₂₀O₄₂ and was tentatively identified as medicagenic acid attached to three deoxyhexosyl, two hexosyl, two pentosyl and a uronic acid moiety [15–21]. However, MSⁿ does not provide enough information for absolute structural characterization. Nevertheless, the use of a hybrid orbitrap mass analyzer enabled the tentative identification of 15 saponins that have not been reported before in *H. hirsuta* (indicated with e in Table 2). Full structural characterization is not the main goal in this project, as the complete herbal extract is subjected to biotransformation studies.
Figure 3. LC–MS data for an unknown saponin. (A) Chromatogram obtained for the extract of Herniaria hirsuta. (B) MS scan of peak at 15.33 min in the heated electrospray ionization (HESI) negative mode. (C) MS² fragmentation spectrum of the ion at \( m/z \) 1703.71734.

2.2. Gastrointestinal Biotransformation

Herbal extracts comprise a mixture of compounds, covering a wide range of bioactive constituents, aside from active compounds or prodrugs. As the main compounds are classified as flavonoids and saponins, extensive biotransformation after oral intake can be expected. Therefore, gastrointestinal biotransformation of the lyophilized extract was simulated in vitro to monitor the levels of the identified compounds using amMS and UV data. However, gastrointestinal enzymes and fecal microflora present in the samples cause a lot of matrix interference, increasing the complexity of the data. The multiclass samples are measured as a function of time, adding a longitudinal aspect to the complex data and impeding the interpretation of the data. The concentration of precursor compounds and metabolites can increase, decrease or show any combination of these patterns during biotransformation.
Table 1. Chromatographic and Spectrometric Data of the Tentatively Identified Compounds in *H. hirsuta* Detected with a Generic LC-PDA-amMS Method for Moderately Polar Phytochemicals.

| Compound Number | Compound | Molecular Formula | RT (min) | HESI Full MS | HESI ddMS | Neutral Losses | Max Abs. (nm) | Literature |
|-----------------|----------|-------------------|----------|--------------|-----------|----------------|---------------|------------|
| 1               | quinic acid | C_{7}H_{12}O_{6} | 1.54     | reg          | pos       | 191.0561       | 169.0142      | 109.0290   | C_{6}H_{10}O_{2} | 259, 294 |
| 2               | gallic acid | C_{7}H_{6}O_{5}  | 4.06     | reg          | pos       | 169.0142       | 109.0290      | 109.0290   | C_{6}H_{10}O_{2} | 259, 294 |
| 3               | protocatechuic acid [3,4-dihydroxybenzoic acid] | C_{7}H_{6}O_{4} | 6.46     | reg          | pos       | 133.0193       | 109.0290      | 109.0290   | C_{6}H_{10}O_{2} | 259, 294 |
| 4               | caffeic hexose | C_{9}H_{8}O_{4} | 7.68     | reg          | pos       | 341.0881       | 365.08380     | 360.12839 | 181.04962; C_{6}H_{10}O_{2} - H_{2}O; 145.02861: C_{6}H_{10}O_{2} - 2H_{2}O, 153.04443: C_{8}H_{6}O_{2}; 259, 294 |
| 5               | asperulosidic acid | C_{18}H_{24}O_{12} | 8.01     | reg          | pos       | 431.11975, 499.10757: [M-H + NaFA]^{-}; 863.24756: [2M-H]^{-} | 367.10349, 707.18355: [M-H + NaFA]^{-}; 353.08685, 717.18355: [2M-H]^{-} | 329.09064: C_{6}H_{10}O_{2}, 161.04463: C_{6}H_{10}O_{2}, 271.06091: C_{6}H_{10}O_{2}, 145.04990: C_{6}H_{10}O_{2}, 127.03919: C_{6}H_{10}O_{2} | 369.11795 | 326 |
| 6               | dihydrocaffeic acid | C_{9}H_{8}O_{4} | 8.07     | reg          | pos       | 181.05041      | 181.05041     | 109.0290   | C_{6}H_{10}O_{2} | 259, 294 |
| 7               | 1-O-ferruloylquinic acid | C_{17}H_{20}O_{9} | 8.76     | reg          | pos       | 367.10349, 435.09108: [M-H + NaFA]^{-} | 353.10640, 717.18355: [M-H + NaFA]^{-} | 353.08685, 717.18355: [2M-H]^{-} | 367.11795 | 326 |
| 8               | chlorogenic acid [3,4-dihydroxycinnamoylquinic acid; 5-Caffeoylquinic acid] | C_{18}H_{24}O_{9} | 8.94     | reg          | pos       | 353.08685, 717.18355: [2M-H]^{-} | 353.10640, 717.18355: [M-H + NaFA]^{-} | 353.08685, 717.18355: [2M-H]^{-} | 367.11795 | 326 |
| 9               | isorhamnetin-3-O-rutinoside-7-O-glucoside | C_{34}H_{32}O_{21} | 9.27     | reg          | pos       | 785.21631, 853.20349: [M-H + NaFA]^{-} | 785.21631, 853.20349: [M-H + NaFA]^{-} | 785.21631, 853.20349: [M-H + NaFA]^{-} | 787.22931 | 479, 11838 |
| Compound a | Molecular Formula | RT (min) | HESI Full MS b | HESI ddMS c | Neutral Losses | Max Abs. (nm) | Literature |
|------------|-------------------|----------|----------------|-------------|----------------|----------------|------------|
| chlorogenic acid c | C_{16}H_{18}O_{9} | 9.36 | neg | $353.08781$ | 191.0561 | 191.0561: C_{7}H_{12}O_{3} |
| Catechin c | C_{15}H_{14}O_{6} | 11.21 | reg | $289.07176$ | - | - | 279 Mbark et al. |
| epicatechin c | C_{15}H_{14}O_{6} | 11.25 | reg | $289.07176$ | - | - | 280 |
| procyanidin B2 c | C_{26}H_{22}O_{12} | 9.68 | reg | $577.13515$ | 193.05003: C_{10}H_{9}O_{4} - CO_{2} |
| Caffeic acid c | C_{9}H_{8}O_{4} | 9.7 | reg | $179.03498$ | 135.04515: C_{9}H_{8}O_{4} - CO_{2} |
| 1-O-feruloylquinic acid d | C_{17}H_{22}O_{9} | 10.51 | reg | $367.10358$ | 193.05003: C_{10}H_{10}O_{4} - CO_{2} |
| Quercetin 3-rhamnosyl-(1->2)-rhamnosyl-(1->6)-glucoside or quercetin 3-rhamnoside | C_{33}H_{40}O_{20} | 11.05 | reg | $755.20471$ | 300.02771: C_{16}H_{16}O_{4} - CO_{2} |
| Matairesinoside | C_{26}H_{22}O_{11} | 11.3 | reg | $565.19336$ | 339.12378: C_{19}H_{20}O_{5} - CO_{2} |
| | | | | | | | 339.12378: C_{20}H_{20}O_{5} - CO_{2} |

Table 1. Cont.
| Compound | Molecular Formula | RT (min) | HESI Full MS | HESI ddiMS | Neutral Losses | Max Abs. (nm) | Literature |
|----------|------------------|----------|--------------|------------|----------------|---------------|------------|
| isorhamnetin-3-O-glucoside (one hexosyl and two deoxyhexosyl units) | C_{14}H_{22}O_{20} | 11.74 | neg 769.2204 | 1359.44527 | 2M-H^- | 632.32239 | C_{14}H_{20}O_{9}; 357.0610; C_{14}H_{18}O_{9}; 271.0252; 195.0651; 151.0223 | 355 |
| quercetin 3-O-β-(2"-O-α-L-rhamnopyranosyl)-glucopyranoside | C_{22}H_{25}O_{17} | 11.96 | neg 623.12537 | 1247.25657 | [M-H]- | 479.08180 | C_{14}H_{18}O_{8}; C_{14}H_{18}O_{7}; C_{14}H_{18}O_{6}; C_{14}H_{18}O_{5}; C_{14}H_{18}O_{4}; C_{14}H_{18}O_{3}; C_{14}H_{18}O_{2}; C_{14}H_{18}O_{1}; C_{14}H_{18}O_{0}; C_{14}H_{18}O_{−} | 255, 355 |
| rutin ^c | C_{20}H_{20}O_{16} | 12.08 | neg 609.14667 | 677.13349 | [M-H]+ | 479.08180 | van Dooren et al. (2016) |
| cynaroside ^c | C_{11}H_{11}O_{11} | 12.2 | reg 447.09364 | - | - | - | Wagner et al. (2013) |
| hesperidin ^c | C_{20}H_{20}O_{15} | 12.24 | reg 609.18153 | 661.19705 | - | - | - |
| ferulic acid ^c | C_{10}H_{9}O_{4} | 12.34 | reg 193.05083 | 177.0547; 145.02855; 135.04424; 117.03578 | 178.02463; 177.0547; C_{14}H_{18}O_{4}; C_{14}H_{18}O_{3}; C_{14}H_{18}O_{2}; C_{14}H_{18}O_{1}; C_{14}H_{18}O_{0}; C_{14}H_{18}O_{−} | - |
| hemiarasaponin 1 | C_{14}H_{20}O_{22} | 12.59 | reg 1215.2698 | 1239.26345 | [M+Na]^+ | - | - |
| isorqueretin ^c | C_{21}H_{22}O_{12} | 12.61 | reg 463.0882 | - | 300.02725; 271.0252; 255.0205; 243.01859; 151.01230 | 300.02725; C_{14}H_{18}O_{2}; C_{14}H_{18}O_{1}; C_{14}H_{18}O_{0}; C_{14}H_{18}O_{−} | - |
Table 1. Cont.

| Compound Number | Compound a | Molecular Formula | RT (min) | HESI Full MS b | HESI ddMS c | Neutral Losses | Max Abs. (nm) | Literature |
|-----------------|------------|-------------------|----------|----------------|--------------|----------------|--------------|------------|
| 26              | p-coumaric acid c | C_9H_8O_3 | 12.62 reg | 163.04007 147.04414 | 119.05024 147.04414 | 119.05024: C_9H_10O_2 | - | - |
| 27              | 2-(3,4-Dihydroxyphenyl)-5,7-dihydroxy-4-oxy-4H-chromen-3-yl-6-O(2-O-acetyl-a-L-arabinopyranosyl)-β-D-glucopyranoside | C_24H_32O_17 | 12.67 reg | 637.14125, 1275.28099 [2M+H] | 463.09000; 303.02724 | 463.09000: C_24H_30O_15 | - | - |
|                 |            |                   |          | 639.15543, 493.09749 | 465.10371; 303.04972 | 465.10371: C_21H_20O_12 | - | - |
| 28              | miquelianin [quercetin-3-O-glucuronide] e | C_21H_32O_15 | 12.89 reg | 477.06746 | 251.03828; 178.99860 | 251.03828: C_21H_30O_13 | - | - |
|                 |            |                   |          | 479.08202 | 303.04993; 257.04445 | 303.04993: C_21H_28O_15 | - | - |
| 29              | narcissin [isorhametin-3-O-rutinoside] | C_28H_32O_16 | 12.89 reg | 623.16111, 477.06749; 1247.33026 [2M+H] | 315.05072; 477.06749: C_28H_30O_14 | 284.03314; 255.03024; 227.03455 | 254; 355 | Wagner et al. (2013) |
|                 |            |                   |          | 625.14606 | 479.11804; 317.06586 | 479.11804: C_28H_30O_14 | - | - |
| 30              | avicularin [quercetin-3-O-arabinoside] e | C_20H_20O_11 | 13.16 reg | 433.0782 435.09219 | 301.0386 | 301.0386: C_15H_10O_6 | - | - |
| 31              | apigetrin e | C_21H_32O_9 | 13.27 reg | 431.0937 433.1192 | 268.03827 | 268.03827: C_15H_10O_6 | - | - |
| 32              | astragalin [kaempferol 3-O-glucoside] e | C_21H_32O_11 | 13.34 reg | 447.09381 | 253.02024; 227.03455 | 253.02024: C_15H_10O_6 | 265; 348 | - |
|                 |            |                   |          | 449.10814 | 284.03314; 255.03024 | 284.03314: C_15H_10O_6 | - | - |
| 33              | quercetin e | C_21H_32O_11 | 13.35 reg | 447.09328 | 253.02024; 227.03455 | 253.02024: C_15H_10O_6 | 265; 348 | - |

* a: Molecular formula
* b: HESI Full MS: Mass spectrum obtained using a high-sensitivity electrospray ionization (HESI) interface
* c: HESI ddMS: mass spectrum obtained using a high-sensitivity electrospray ionization (HESI) interface with double detection mode
* d: Neutral Losses: Mass losses observed in the mass spectrum
* e: Max Abs. (nm): Maximum absorbance wavelength
* Literature: References for the compounds
Table 1. Cont.

| Compound | Molecular Formula | RT (min) | HESI Full MS \(^a\) | HESI ddMS \(^b\) | Neutral Losses | Max Abs. (nm) | Literature |
|----------|------------------|----------|---------------------|----------------|----------------|--------------|------------|
| 34 salicylic acid \(^c\) | C\(_7\)H\(_5\)O\(_3\) | 14.07 | reg | 137.02442 | - | 93.03459 | C\(_6\)H\(_4\)O | 236; 302 |
| 35 herniariasaponin F | C\(_{30}\)H\(_{48}\)O\(_{28}\) | 14.17 | reg | 1249.58482, 647.29342 | 501.32283, 439.32258 | 647.29342: C\(_2\)H\(_4\)O\(_3\) + C\(_2\)H\(_2\)O\(_4\); 501.32283: C\(_2\)H\(_4\)O\(_3\); 439.32258: C\(_2\)H\(_4\)O\(_3\) - H\(_2\)O - CO\(_2\) | Mbark et al. (2000) |
| 36 acetylated medicagonic acid + two hexosyl + two deoxyhexosyl + one pentosyl moieties \(^c\) | C\(_{28}\)H\(_{46}\)O\(_{29}\) | 14.53 | reg | 1291.59644 | - | 705.39228: C\(_{20}\)H\(_{16}\)O\(_{10}\) + C\(_2\)H\(_2\)O\(_4\); 543.3337: C\(_{12}\)H\(_{4}\)O\(_{4}\); 501.32281: C\(_2\)H\(_4\)O\(_3\); 439.32205: C\(_2\)H\(_4\)O\(_3\) - H\(_2\)O - CO\(_2\) | Schröder et al. (1993) |
| 37 acetylated medicagonic acid + three hexosyl + one deoxyhexosyl moieties \(^c\) | C\(_{36}\)H\(_{50}\)O\(_{26}\) | 14.61 | reg | 1175.549 | - | 1411.60317: C\(_{20}\)H\(_{16}\)O\(_{10}\) + C\(_2\)H\(_2\)O\(_4\); 1235.56738: C\(_{12}\)H\(_{4}\)O\(_{4}\); 1102.56189: C\(_2\)H\(_4\)O\(_3\) | Mbark et al. (2000) |
| 38 medicagonic acid + two hexosyl + two deoxyhexosyl + two pentosyl + 1 glucuronic acid moiety \(^c\) | C\(_{20}\)H\(_{16}\)O\(_{30}\) | 14.89 | reg | 1557.6595, 1411.60317 | 1235.56738; 985.57922; 796.52348; 499.32043 | 1411.60317: C\(_{20}\)H\(_{16}\)O\(_{10}\) + C\(_2\)H\(_2\)O\(_4\); 1235.56738: C\(_{12}\)H\(_{4}\)O\(_{4}\); 1102.56189: C\(_2\)H\(_4\)O\(_3\) | |
| 39 zanhic acid + one hexose + four deoxyhexoses + two pentoses + one glucuronic acid moiety \(^c\) | C\(_{20}\)H\(_{12}\)O\(_{42}\) | 14.92 | reg | 1703.7106, 851.35632 [M-2H] \(^c\) | 1381.62800, 823.41089, 517.31879, 455.31766, 859.4945 | 1381.62800: C\(_{20}\)H\(_{12}\)O\(_{42}\) + C\(_2\)H\(_2\)O\(_4\) + C\(_2\)H\(_4\)O\(_3\) - H\(_2\)O - CO\(_2\); 859.4945: C\(_2\)H\(_4\)O\(_3\) + C\(_2\)H\(_4\)O\(_3\) | |

\(^a\) HESI: positive (pos), negative (neg) mass spectra.
\(^b\) ddMS: doubly charged fragment mass spectra.
\(^c\) Mass spectra of the compounds were obtained from Mbark et al. (2000).
Table 1. Cont.

| Compound Number | Compound * | Molecular Formula | RT (min) | HESI Full MS b | HESI ddMS 2 | Neutral Losses | Max Abs. (nm) | Literature |
|-----------------|------------|-------------------|----------|----------------|--------------|----------------|---------------|------------|
| 40              | herniariasaponin E | C_{35}H_{42}O_{25} | 15.03   | reg            | 1145.5387; 1213.5271 | 972.5673; 995.5634 | -             | -          |
|                 |            |                   |          |                | 543.33105; 483.31180 | C_{29}H_{30}O_{18}, H_{2}O, 439.32277 | C_{25}H_{23}O_{16}, 349.32277 | Mbark et al. (2000) |
|                 |            |                   |          |                | 439.32227; 439.32227 | 439.32208; 439.32208 | -             | -          |
|                 |            |                   |          |                | 601.19086; 601.19086 | 601.19086; 601.19086 | -             | -          |
|                 |            |                   |          |                | 543.33105; 543.33105 | 543.33105; 543.33105 | -             | -          |
| 41              | acetylated medicagenic acid + three hexosyl moieties * | C_{35}H_{35}O_{22} | 15.06   | reg            | 1029.49191; 1075.49754 | -             | -             | -          |
|                 |            |                   |          |                | 543.33105; 543.33105 | 543.33105; 543.33105 | -             | -          |
| 42              | medicagenic acid + three deoxyhexosyl + one pentosyl + one glucuronic acid moiety * | C_{35}H_{42}O_{28} | 15.12   | reg            | 1247.569 | 925.4815; 501.31976 | 1127.4972; 949.4744; 701.3485; 525.3280; 447.1437 | -          |
|                 |            |                   |          |                | 1048.53173; 1053.48612 | [M+Na]^+; [M+Na]^+ | 1398.65576; [M+Na]^+ | -          |
| 43              | zanhic acid + three deoxyhexosyl + one pentosyl + one glucuronic acid moiety * | C_{35}H_{42}O_{29} | 15.24   | reg            | 1263.5657 | 1163.15430; 1081.94128 | -             | -          |
|                 |            |                   |          |                | 1282.60583; [M+Na]^+ | 1163.15430; 1081.94128 | -             | -          |
| 44              | medicagenic acid + two hexosyl + three deoxyhexosyl + two pentosyl + one glucuronic acid moiety * | C_{35}H_{45}O_{42} | 15.33   | reg            | 1703.7174 | 1381.62830; 1219.22852; 879.30701; 843.52264; 823.41089; 501.32275 | 439.31954 | -          |
|                 |            |                   |          |                | 1727.71399; 1722.75891 | [M+Na]^+; [M+Na]^+ | -             | -          |
| 45              | medicagenic acid + one hexosyl + four deoxyhexosyl + two pentoses + one glucuronic acid moiety * | C_{35}H_{45}O_{41} | 15.52   | reg            | 1687.7229 | 1365.63477; 998.50337; 823.40276 | -             | -          |
|                 |            |                   |          |                | 1711.71956; [M+Na]^+ | 1711.71956; [M+Na]^+ | -             | -          |
| Compound Number | Compound * | Molecular Formula | RT (min) | HESI Full MS b | HESI ddMS c | Neutral Losses | Max Abs. (nm) | Literature |
|-----------------|------------|-------------------|----------|----------------|-------------|---------------|-------------|------------|
| 46              | medicagenic acid + three deoxyhexosyl + two pentosyl + one glucuronic acid moiety a,b | C_{34}H_{28}O_{22} | 15.12; 15.62 | reg            | 1379.6127   | 1057.52454; 925.48077; 823.41821; 501.32231; 439.32050; 1259.54272; 1081.57822; 847.40808; 579.18944 | pos 1403.60791 [M+Na] + | 1057.52454: C_{34}H_{28}O_{22} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 925.48077: C_{34}H_{28}O_{22} - C_{4}H_{2}O_{6}; 823.41821: C_{34}H_{28}O_{22}; 501.32231: C_{5}H_{4}O_{3}; 439.32050: C_{3}H_{4}O_{2} - H_{2}O - CO_{2}; 1259.54272: C_{34}H_{28}O_{22} - 2C_{4}H_{2}O_{6}; 1081.57822: C_{34}H_{28}O_{22} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 847.40808: C_{34}H_{28}O_{22} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 579.18944: C_{34}H_{28}O_{22} - C_{4}H_{2}O_{6}; | Charrouf et al. (1995) |
| 47              | medicagenic acid + one hexosyl + two deoxyhexosyl + two pentosyl + one glucuronic acid moiety a,b | C_{34}H_{28}O_{23} | 15.67 | reg            | 1395.6069   | 439.32248; 1219.58301; 439.32037: C_{34}H_{28}O_{23} | pos 1419.60039 [M+Na] + | 439.32248: C_{34}H_{28}O_{23} - C_{4}H_{2}O_{6} - H_{2}O - CO_{2}; 1219.58301: C_{34}H_{28}O_{23} - C_{4}H_{2}O_{6} | Charrouf et al. (1995) |
| 48              | herniariasaponin B | C_{30}H_{20}O_{13} | 15.83 | reg            | 1409.6244   | 1087.53479; 823.41272; 501.32300; 439.32343 | pos 1433.61982 [M+Na] +, 1428.66283 [M+NH4] +, 1411.63608 | 1087.53479: C_{30}H_{20}O_{13} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 823.41272: C_{30}H_{20}O_{13}; 501.32300: C_{30}H_{20}O_{13} - C_{4}H_{2}O_{6}; 439.32343: C_{30}H_{20}O_{13} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; | Charrouf et al. (1995) |
| 49              | acetylated medicagenic acid + one deoxyhexosyl + two pentosyl moieties a,b | C_{34}H_{24}O_{19} | 16    | reg            | 941.4756    | 501.32312; 821.43097; 543.53167 | pos 960.51893 [M+NH4] +, 965.47021 [M+Na] + | 501.32312: C_{34}H_{24}O_{19} - C_{4}H_{2}O_{6}; 821.43097: C_{34}H_{24}O_{19} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 543.53167: C_{34}H_{24}O_{19} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; | | |
| 50              | quercetin a,b | C_{15}H_{10}O_{7} | 16.4  | reg            | 301.0338    | 271.02539; 178.99774; 151.00273; 121.02832; 285.09694; 257.04440; 229.04947; 201.05476; 165.01811; 153.01846; 137.02357 | pos 303.04993 | 271.02539: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6}; 178.99774: C_{15}H_{10}O_{7}; 151.00273: C_{15}H_{10}O_{7}; 121.02832: C_{4}H_{2}O_{6}; 285.09694: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6}; 257.04440: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6}; 229.04947: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6}; 201.05476: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 165.01811: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 153.01846: C_{15}H_{10}O_{7} - C_{4}H_{2}O_{6} - C_{4}H_{8}O_{2}; 137.02357: C_{4}H_{2}O_{6}; | 255, 372 |
| Compound Number | Compound * | Molecular Formula | RT (min) | HESI Full MS $^b$ | HESI ddMS $^c$ | Neutral Losses | Max Abs. (nm) | Literature |
|-----------------|------------|-------------------|----------|-------------------|----------------|----------------|--------------|------------|
| 51              | acetylated medicagenic acid + one hexosyl + two deoxyhexosyl + one pentosyl + one glucuronic acid moiety$^d$ | C$_{63}$H$_{94}$O$_{30}$ | 16.76 | neg pos 652.28625 [M-2H]$^{-}$; 1305.57764; 1329.57288 [M+Na]$^{+}$; 1324.61719 [M+NH$_3$]$^{+}$ | 439.32321; 585.20691; 439.32321: C$_4$H$_8$O$_5$ + 2C$_4$H$_7$O$_9$ + C$_1$H$_2$O$_3$ | 439.32321: C$_4$H$_8$O$_5$ + 2C$_4$H$_7$O$_9$ + C$_1$H$_2$O$_3$ | - | 253.05069 | C$_{12}$H$_{10}$O$_4$ |
| 52              | naringenin $^e$ | C$_{15}$H$_{12}$O$_5$ | 16.8 | reg pos 271.06120 273.0755 | 253.05069 | 253.05069: C$_{15}$H$_{10}$O$_4$ | - | 253.05069 | C$_{12}$H$_{10}$O$_4$ |
| 53              | apigenin $^e$ | C$_{15}$H$_{10}$O$_5$ | 17.73 | reg pos 269.04555 271.06100 | 225.05510; 117.03339; 153.01845 | 225.05510: C$_4$H$_8$O$_5$; 151.00258: C$_2$H$_4$O$_3$ | 273.0755 | - | 153.01845 | C$_{12}$H$_{10}$O$_4$ |
| 54              | isorhamnetin $^e$ | C$_{16}$H$_{12}$O$_7$ | 17.87 | reg pos 315.05103 317.06555 | 300.02762: C$_{15}$H$_{10}$O$_7$ | 300.02762: C$_{15}$H$_{10}$O$_7$ | 153.01846 | - | - |
| 55              | luteolin $^e$ | C$_{15}$H$_{10}$O$_6$ | 18.09 | reg pos 285.04046 287.05001 | 199.03967; 151.00212; 153.01821; 153.01787; 155.04414; 153.01846 | 199.03967: C$_{15}$H$_{10}$O$_6$; 151.00212: C$_2$H$_4$O$_3$; 153.01821: C$_{14}$H$_7$O$_2$; 153.01787: C$_4$H$_8$O$_5$; 155.04414: C$_{13}$H$_{10}$O$_2$ | 258.05115; 231.06519; 165.01807; 153.01816; 121.02866 | - | 153.01787: C$_4$H$_8$O$_5$; 155.04414: C$_{13}$H$_{10}$O$_2$ |
| 56              | kaempferol $^e$ | C$_{15}$H$_{10}$O$_6$ | 18.09 | reg pos 285.04046 287.05001 | 258.05115; 231.06519; 165.01807; 153.01816; 121.02866 | 258.05115: C$_4$H$_8$O$_5$; 231.06519: C$_2$H$_4$O$_3$; 165.01807: C$_4$H$_7$O$_2$; 153.01816: C$_4$H$_8$O$_5$; 121.02866: C$_2$H$_4$O$_3$ | 265, 364 | - | - | C$_4$H$_8$O$_5$ |
| 57              | miquelianin [quercetin 3-O-glucuronide] $^e$ | C$_{22}$H$_{23}$O$_{13}$ | 12.89–13.49 | reg pos | 477.0675 | 301.03552; 271.02501; 255.03220; 178.99788; 151.00258 | 301.03552; 271.02501; 255.03220; 178.99788; 151.00258 | 257, 357 | - | C$_{12}$H$_{10}$O$_4$ |

* Com-Pound Number |
* Molecular Formula |
* RT (min) |
* HESI Full MS $^b$ |
* HESI ddMS $^c$ |
* Neutral Losses |
* Max Abs. (nm) |
* Literature |
| Com-Pound Number | Compound | Molecular Formula | RT (min) | HESI Full MS | HESI ddMS | Neutral Losses | Max Abs. (nm) | Literature |
|------------------|----------|-------------------|----------|-------------|-----------|----------------|---------------|------------|
| 58               | herniariasaponin G | C_{33}H_{44}O_{24} | 14.61 | neg | 1103.52828, 574.26304[M-2H+FA]^− | 821.43842, 501.32306 C_{30}H_{36}O_{8} 439.32156 C_{30}H_{40}O_{8} - H_{2}O - CO_{2} | 973.50128, 811.45524, 649.39624, 503.37704, 457.33189 | 1259.56189, 741.24219, 595.18250, 445.13116 | van Dooren et al. (2016) |
| 59               | medicagenic acid + two hexosyl + two deoxyhexosyl + one glucuronic acid moiety | C_{30}H_{115}O_{39} | 14.89−15.38 | reg | 1557.6595, 1411.60317 [M+Na]^+ | 1411.60317 C_{29}H_{117}O_{35} - C_{9}H_{15}O_{3} | | | |
| 60               | medicagenic acid + one hexosyl + two deoxyhexosyl + two pentosyl + one glucuronic acid moiety | C_{44}H_{116}O_{33} | 14.92−15.67 | reg | 1395.6069 | | | | |
| 61               | herniariasaponin H | C_{30}H_{115}O_{37} | 15.12; 15.66; 15.72 | reg | 1541.6657 | 1446.38293, 1219.58069 C_{30}H_{44}O_{8} - C_{9}H_{15}O_{3} | 823.41711 C_{30}H_{117}O_{35} - C_{9}H_{15}O_{3} - 2C_{2}H_{4}O_{6} | 501.32587: C_{30}H_{40}O_{8} | | van Dooren et al. (2016) |
| 62               | chlorogenic acid | C_{30}H_{15}O_{9} | 7.31; 8.88 | reg | 335.08665 | 191.05544 C_{9}H_{15}O_{4} | 179.03427 C_{9}H_{15}O_{4} | 173.04944 C_{9}H_{15}O_{4} - H_{2}O | 163.03915 C_{9}H_{15}O_{4} - H_{2}O, 145.02851 C_{9}H_{15}O_{4} - 2H_{2}O, 135.04443 C_{9}H_{15}O_{4} | | |
| 63               | 1-O-feruloylquinic acid | C_{17}H_{25}O_{8} | 9.20−10.62 | reg | 367.10349, 435.09108 [M+Na]^+ | 193.0003, 173.03485 | | | Jaiswal et al. (2011) |

^a If not specified otherwise: tentative identification based on accurate mass ^b Deprotonated and protonated molecules in negative and positive mode, respectively, unless stated otherwise ^c Identification with an analytical standard ^d Tentatively identified compound or an isomer ^e Tentative identification of saponins that have never been reported before in *H. hirsuta*.
Table 2. Most probable sum formula and ppm deviation for the fragment ions of m/z 1703.71734.

| m/z [M-H]− | Sum Formula | ∆ ppm |
|------------|-------------|-------|
| 1703.71838 | C₇₆H₁₂₀O₄₂ | 0.141 |
| 1381.62830 | C₆₄H₁₀₂O₃₂ | 0.113 |
| 879.30701  | C₃₃H₆₆O₂₆ | 9.445 |
| 823.41089  | C₄₂H₄₄O₁₆ | −1.541 |
| 501.32275  | C₃₀H₄₅O₆  | 1.172 |
| 439.31934  | C₂₉H₄₄O₃  | −5.528 |

Figure 4 shows the chromatogram of the H. hirsuta extract before and after biotransformation. Before biotransformation (t₀), peaks are mostly attributed to the tentatively identified compounds. After the colonic phase of gastrointestinal biotransformation, the chromatogram contains peaks of compounds, metabolites and matrix interferences, originating from enzymes and fecal microflora, which are absent at t₀.

Processing the chromatograms of every time point for every compound allows monitoring of the intensity over time. However, this manual approach is time-consuming and will only provide information about the abundance of previously identified compounds. Manual screening for metabolites
is comparable to looking for a needle in a haystack because of the complexity of the chromatograms and matrix interferences.

Generally, for most tentatively identified compounds, a decrease in relative abundance is observed mainly at the colonic phase, caused by bacterial degradation, suggesting formation of metabolites. However, some compounds, mainly saponins, show an increase in relative abundance over time. It is suggested that this increase is induced by conversion of one compound into another, caused by loss of one or more sugar moieties. For example, compound 46 (medicagenic acid + three deoxyhexosyl + two pentosyl + one glucuronyl moiety) shows an increase in relative abundance and reaches a maximum after 24 h colon phase, which is not observed in the negative control samples (NCHEX) (Figure 5). In addition, the signal was absent in the blank (MB), as expected. At the same time, a decrease of compound 45 (medicagenic acid + one hexosyl + four deoxyhexosyl + two pentosyl + one glucuronyl moiety) is observed. The increase of compound 46 can be related to the decrease of compound 45, caused by the loss of a hexosyl and a deoxyhexosyl moiety. The decrease is observed in both samples and negative control during the stomach phase and the small intestinal phase. At these time points, there is no difference in samples and negative control as the negative controls only lack fecal bacteria during the colonic phase. Since identification based on MS-data is not absolute, only tentative conclusions can be drawn from these observations. However, the data support the theory that gastrointestinal biotransformation leads to a loss of sugar moieties in saponins.

An automated data analysis workflow, previously described by Peeters et al. and Beirnaert et al., was used to screen for metabolites [23,27]. The workflow resulted in 11,072 features in negative ion mode, of which 4846 were annotated as isotopes. In positive mode, 5242 features were detected after exclusion of isotopes.

Firstly, the metabolic pathway of herniariasaponin H (m/z 1541.6675, rt: 17.3 min) was studied in detail to verify if the workflow was suitable for herbal extracts. The time profile of herniariasaponin H shows a decrease in intensity over time, especially during the colonic phase, which is not observed in the negative control samples (NCHEX) (Figure 6). In addition, the signal is absent in the method blank (MB). The feature is rated as notable based on the results of tinderesting, the interactive Shiny app developed in R, with a score of 1 and a ranking of 242 out of 6226 features. During biotransformation, stepwise elimination of sugar moieties is observed resulting in formation of the aglycone (Figure 7). All intermediate metabolites show a tinderesting score of at least 0.98, meaning that they are all rated
as interesting. Finally, the aglycone medicagenic acid (m/z 501.3213; rt: 21.68) is formed, showing an increase in signal over time in the samples, with a tinderesting score of 0.998 (Figure 6).

Features with a tinderesting score higher than 0.8 were selected for in-depth exploration with the following inclusion criteria: a reasonable intensity with good-quality MS/MS spectra and a retention time between 2 and 25 min.

The main challenge in the analysis of a large amount of analytical data remains in the identification and annotation of the features of interest. However, the data analysis workflow ranks features according to one single score taking the longitudinal aspect and the three sample classes into account, which facilitates selection of features of interest. On the other hand, determination of the molecular formula of compounds with a molecular mass above 1000 Dalton becomes more challenging. Therefore, this step in the data analysis process is still performed manually.

As expected, the features included in the in-depth exploration could be attributed to saponins, flavonoids and their metabolites. Saponins (compounds 35–41, 43–45, 47–48, 51, 58, 59 and 61) show the same biotransformation pathway as the studied example, herniariasaponin H. In brief, the intensity of the signal decreases over time due to stepwise elimination of sugar moieties, leading to formation of the aglycone. The observed biotransformation of flavonoids is similar. In nature, flavonoids mainly occur as glycosides. Gastrointestinal biotransformation causes deglycosylation, leading to formation of the aglycone. For example, quercetin glycosides are abundantly and ubiquitously present flavonoids in nature. The aglycone is formed after biotransformation of several precursors, such as isoquercetin and rutin. Isoquercetin (m/z 463.0871; rt: 11.67 min) has a tinderesting score of 0.86 and shows a decrease in intensity over time (Figure 8). The decrease is already observed in the stomach phase and continues in the small intestinal phase and colonic phases until the signal is completely absent. In the negative control samples, the signal stays stable during the colonic phase, because there are no bacteria present. The signal is absent in the method blank. For quercetin (m/z 301.0345; rt: 16.54 min), an increase in intensity is observed over time, which is not observed in the negative control or method blank (Figure 8). The feature is rated as interesting with a tinderesting score of 1. This suggests deglycosylation of isoquercetin leading to the formation of quercetin.

Figure 6. Time profile of gastrointestinal biotransformation of herniariasaponin H (m/z 1541.6675 [M-H]−) and medicagenic acid (m/z 501.3213 [M-H]−).
Figure 7. Metabolic pathway of herniarasaponin H by in vitro gastrointestinal biotransformation.
A feature with m/z 181.0497 and rt: 4.9 min was included in the in-depth exploration as it showed a interesting score of 0.992. The feature shows an increase in intensity over time, mainly during the colonic phase (Figure 8). The metabolite was tentatively identified as 3,4-dihydroxyphenylpropionic acid (A) or 3-methoxy-4-hydroxyphenylacetic acid (B). Both compounds were previously described as metabolites of quercetin [28,29]. Based on the obtained spectra, absolute structural characterization is not possible. Nonetheless, the product ion could be identified as a metabolite of quercetin. This shows that despite the increase in intensity of quercetin, the compound is further biotransformed (Figure 9).

![Figure 8](image1.png)

**Figure 8.** Time profiles of biotransformation of isoquercetin (m/z 463.0871 [M-H]⁻), quercetin (m/z 301.0345 [M-H]⁻) and a metabolite (m/z 181.0497 [M-H]⁻) tentatively identified as 3,4-dihydroxyphenylpropionic acid or 3-methoxy-4-hydroxyphenylacetic acid.

![Figure 9](image2.png)

**Figure 9.** Biotransformation pathway of isoquercetin showing quercetin as intermediate metabolite which is further transformed into 3,4-dihydroxyphenylpropionic acid (A) or 3-methoxy-4-hydroxyphenylacetic acid (B).

### 2.3. Liver Biotransformation

Figure 10 shows an overview of the proposed metabolic pathway of medicagenic acid after S9 biotransformation. For metabolites 1–3 (M1–3), three different chromatographically separated peaks were observed (rt 17.39; 17.60 and 18.45 min), suggesting the presence of three different structural isomers. This was confirmed by the prediction provided by Meteor. The software proposed hydroxylation at C-11, C-24, or C-29. Double hydroxylation (M4), oxidation of a hydroxy group (M5) and a combination of hydroxylation and oxidation (M9–10) were also predicted by Meteor. The non-target screening approach confirmed the formation of the predicted metabolites, and one additional metabolite was detected. Metabolite 8 (M8) shows a reduction of the carboxylic acid moiety. However, limited by the poor MS/MS fragmentation of the aglycons and lack of NMR.
data, identification remains tentative. Further elucidation of the molecular structures was not possible. The only phase II products observed were glucuronidated metabolites of medicagenic acid (M6–7). Sulfation reactions were not observed, although they were predicted by Meteor and Biotransformer. To confirm that the in vitro model was capable to form sulfated conjugates, biotransformation of testosterone was examined as positive control. Testosterone was analyzed after hepatic biotransformation and proved the capability of the in vitro model to form both sulfated and glucuronidated conjugates.

The phase II hydrophilic metabolites may be excreted via the kidneys and are often considered detoxifying as they usually possess lower activity compared to their parent aglycones. However, a number of glucuronide conjugates are known to be active. The 6-O-glucuronide of morphine, a widely used opioid analgesic, is the most well-known example of a glucuronidated metabolite possessing pharmacological activity greater than the parent aglycone [30]. Based on the observation of in vitro formation of glucuronidated hepatic metabolites, it is a possible hypothesis that these metabolites may be responsible for the urolithiatic activity [31]. A possible mode of action for prevention and dissolution of urinary stones could be the emulsifying properties of the glucuronidated metabolites, facilitating dissolution of crystals.

![Chemical structures of medicinal compounds](image)

**Figure 10.** B Metabolic pathway of medicagenic acid after in vitro hepatic biotransformation.

### 3. Materials and Methods

#### 3.1. Chemicals

Ultrapure water with a resistivity of 18.2 MΩ·cm at 25 °C was generated with a Millipore™-purification system. Ultrahigh-Performance Liquid Chromatography (UHPLC)-grade MeOH, acetonitrile and formic acid were purchased from Biosolve (Dieuze, France), dichloromethane by Merck (Darmstadt, Germany). The following analytical standards were provided by Sigma–Aldrich (St. Louis, MO, USA): apigenin, benzoic acid, caffeic acid, ±catechin, chlorogenic acid, cinnamic acid, coumarin, emodin, epicatechin, ferulic acid, isorhamnetin, naringenin, p-coumaric acid, protocatechuic acid, quercetin, quercitrin, rutin, salicylic acid, sinapic acid, β-sitosterol, stigmasterol, syringic acid, tannic acid, taxifolin, theophylline and vanillic acid. Luteolin and procyanidin B2 were provided by Santa Cruz Biotechnology (Santa Cruz, CA, USA). Gallic acid and p-hydroxybenzoic acid were provided...
by Carl Roth (Karlsruhe, Germany). Human liver S9 fraction and NADPH RS (Regenerating System) were purchased from XenoTech. Medicagenic acid was provided by Phytolab (Vestenbergsgreuth, Germany). All other chemicals and biochemicals were purchased from Sigma-Aldrich (St. Louis, MO, USA).

3.2. Preparation of Standard Solutions

Standard stock solutions of the analytical standards were prepared at a concentration of 1 mg mL$^{-1}$ in UHPLC-grade MeOH for each analyte separately and stored in the dark at −80 °C. Dilutions of these solutions were prepared in 60:40 (v:v) MeOH:ammonium formate buffer (40 mM, aqueous). Standard stock and working solutions were stored at −80 °C in the dark.

3.3. Sample Preparation

Aerial parts from *H. hirsuta* were collected in d’Oujda, Morocco. A voucher specimen of the plant is kept at the Muséum National d’Histoire Naturelle—Institut Scientifique-Université Mohamed V Agdal, Morocco—(Reference number: 5902). The air-dried plant material of *H. hirsuta* was ground prior to extraction with a PF 10 basic Microfine grinder drive (IKA-Werke GmbH & Co. KG, Staufen, Germany) using a sieve mesh size 0.5 mm.

To cover the full range of constituents, a comprehensive extraction protocol was applied, previously described by Bijttebier et al. with minor adaptations [32,33]. Combination of extraction with H$_2$O:EtOAc and CHCl$_3$:MeOH:H$_2$O afforded both polar and apolar compounds. Approximately 18 g of sample was subsequently wetted with 54 mL of water and 225 mL of EtOAc. The mixture was stirred for 2 h followed by 1 h of ultrasound-assisted extraction. The solvent was removed, and the residue was washed three times with 20 mL EtOAc. The extract was evaporated to dryness. CHCl$_3$:MeOH:H$_2$O extraction was performed by adding 18 g of sample to 180 mL CHCl$_3$:MeOH:H$_2$O (4:4:2). The mixture was stirred for 5 min, followed by 5 min of ultrasound-assisted extraction and stirred for 15 more min. The solvent was removed, and the residue was washed 3 times with 20 mL CHCl$_3$:MeOH:H$_2$O (4:4:2). The dried H$_2$O:EtOAc extract was redissolved with the extract from the CHCl$_3$:MeOH:H$_2$O extraction method, leading to an apolar and a polar phase. Both phases were separated and dried.

3.4. Gastrointestinal Biotransformation

Many natural products are pro-drugs that are biotransformed and activated after oral administration. A previously developed and validated in vitro gastrointestinal biotransformation model was used to mimic human biotransformation processes [34]. This model allows the study of biotransformation processes in the stomach, small intestine and colon using a culture of pooled human feces, avoiding extensive in vivo studies.

The digestive juices and fecal suspension were formulated corresponding to the human conditions, previously described by Breynaert et al. and Peeters et al. [23,34]. Small adaptations to the method were made [35]. Briefly, the pepsin solution was prepared by dissolving 16% (w/v) of pepsin powder in 0.1 M HCl (622,000 FIP-U 100 mL$^{-1}$). To obtain a pancreatin-bile mixture, 0.4% (w/v) pancreatin and 2.5% (w/v) bile were dissolved in 0.1 M NaHCO$_3$ (32,000 FIP-U lipase, 143,600 FIP-U amylase, 16,400 FIP-U protease and 58.4 mmol bile L$^{-1}$). A suspension of 10% (w/v) feces was prepared by homogenizing 3 human fecal samples with a sterile phosphate-buffer solution (0.1 M, pH 7.0). The phosphate-buffer solution consisted of Na$_2$HPO$_4$.2H$_2$O (1.03% w/v), sodium thioglycolate broth (3.45% v/v) and glycerol (17%). Prior to use, the fecal suspension was cultivated. To 10% (v/v) pooled fecal suspension in phosphate buffer 90% (v/v) phosphate buffer was added in an anaerobic glove-box (5% H$_2$, 5% CO$_2$ and 90% N$_2$) (Jacomex Glove Box T3, TCPS, Belgium). The bacteria were cultivated in an anaerobic environment for 1 h at 37 °C by continuously stirring.

Three groups were included in the experiment: (1) samples containing the Herniaria extract, which were treated with digestive enzymes and fecal microflora (HEX); (2) negative control samples, also containing the extract with addition of digestive enzymes but not of fecal bacteria (NCHEX);
and (3) method blanks, comprising an equal volume of solvent instead of extract and undergoing treatment with digestive enzymes and fecal bacteria (MB). For the samples, an amount of 300 mg lyophilized polar phase of the comprehensive extract of *H. hirsuta* was accurately weighed in triplicate.

Gastrointestinal biotransformation was simulated in vitro. An aliquot of each sample was taken at several time points: before biotransformation (t₀), after the gastric stage (S), after the small intestinal phase (SI) and at several time points during the colonic phase (after 2, 4, 6, 10, 14, 18, 22, 24, 32, 40, 48 and 72 h). Every sample was diluted with MeOH (1:2) and centrifuged for 5 min at 3500 rpm. The supernatant was collected, and samples were diluted 10 times with MeOH:H₂O (60:40) before analysis.

### 3.5. Liver Biotransformation

Liver biotransformation mimicking phase I and II reactions was performed using medicagenic acid, the most abundant aglycone after gastrointestinal biotransformation. Hepatic biotransformation was simulated in vitro by using pooled S9 fractions, previously described by Van den Eede et al. [36]. Small adaptations to the method were made. Briefly, a mixture of 65 mM TRIS buffer (pH adjusted to 7.4 at 37 °C), human liver S9 fraction (1 mg mL⁻¹ final protein concentration) and medicagenic acid (100 µM final concentration) was prepared in a total volume of 0.5 mL and preincubated in a shaking waterbath at 37 °C. The reaction was initiated by addition of an NADPH regenerating system (1 mM NADP, final concentration). Cofactors were added to expose the samples to phase II glucuronidation and sulfation: UDPGA (2 mM final concentration), GSH (2 mM final concentration) and PAPS (0.1 mM final concentration). The reaction was inhibited after 1 h by adding 0.5 mL of acetonitrile and storing the tubes on ice. Thereafter, the tubes were centrifuged for 5 min at 10,000 rpm (4 °C). The supernatant was collected and analyzed by LC–MS-MS. Blank samples were prepared as described above, replacing substrate by solvent. Negative control samples were prepared by immediately adding acetonitrile to quench the biotransformation reactions. A positive control was included by incubating testosterone (100 µM final concentration) [36].

### 3.6. Instrumental Analysis

For characterization of the *H. hirsuta* extract, an aliquot of 5 µL was injected with a CTC PAL™ autosampler (CTC Analytics) on a Waters Acquity UHPLC BEH SHIELD RP18 column (3.0 mm × 150 mm, 1.7 µm; Waters, Milford, MA, USA) and eluted with an Accela™ quaternary solvent manager. The temperature of the column was kept at 40 °C using a “Hot Pocket” column oven (Thermo Fisher Scientific, Waltham, MA, USA). The mobile phase solvents consisted of water + 0.1% formic acid (A) and acetonitrile + 0.1% formic acid (B). Compounds were eluted using the following gradient (min/B%): 0.00/0, 9.91/26, 18.51/65, 18.76/100, 20.76/100, 20.88/0, 23.00/0. For detection, an amMS (Q Exactive™, Thermo Fisher Scientific) was used with heated electrospray ionization (HESI). Spray voltage was set at −2.5 kV, sheath gas and auxiliary gas at 47 and 15 (adimensional), respectively, and capillary temperature at 350 °C. The PDA detector was set to scan from 190 to 800 nm. During the first analysis, full scan data were acquired in positive and negative ion mode over an m/z range of 120–1800 and resolving power was set at 70,000 at full width at half maximum (FWHM). Consecutively, a second analysis was performed to selectively fragment the generated ions by data-dependent fragmentation (ddMS²) in positive and negative ionization mode. Selective ion fragmentation was obtained by using a hybrid quadrupole-orbitrap MS analyzer (Q Exactive™, Thermo Fisher Scientific). The raw data were processed using XCalibur™ 3.0 software (Thermo Fisher Scientific).

For the qualitative UHPLC-UV-QTOF analyses of the biotransformation samples, an aliquot of 5 µL was injected on a Waters Acquity UHPLC BEH SHIELD RP18 column (3.0 mm × 150 mm, 1.7 µm; Waters). The temperature of the column was kept at 40 °C. The mobile phase solvents consisted of water + 0.1% formic acid (A) and acetonitrile + 0.1% formic acid (B), and the gradient was set as follows (min/B%): 0/2, 1/2, 14/26, 24/65, 26/100, 29/100, 31/2, 41/2. The flow rate was set at 0.4 mL min⁻¹. For detection, accurate mass measurements were done using a Xevo G2-XS QTof spectrometer (Waters,
Milford, MA, USA) coupled with an ACQUITY LC system equipped with MassLynx 4.1 software. During the first analysis, full scan data were recorded in ESI (+) and ESI (−) mode from $m/z$ 50 to 2000, and the analyzer was used in sensitivity mode (approximate resolution: 22,000 FWHM). The spray voltage was set at either +1.5 kV and −1.0 kV; cone gas flow and desolvation gas flow at 50.0 L h$^{-1}$ and 1000.0 L h$^{-1}$ respectively; source temperature and desolvation temperature at 120 °C and 500 °C respectively. Data were also recorded using MS$^E$ in positive and negative ionization modes, providing separate MS and MS$^E$ data. A ramp collision energy from 20 V to 30 V was applied to obtain additional structural information. Leucine encephalin was used as lock mass. To monitor analytical drift and assess precision, quality control (QC) samples were injected after every time point. QC samples were prepared using a dilution of the standard solution mix (39 ng mL$^{-1}$).

### 3.7. Data Analysis

Dereplication was used as an untargeted workflow for rapid identification of the major compounds present in the extract, regardless of their chemical class. Global Natural Products Social Molecular Networking (GNPS, a web-based mass spectrometry open-access knowledge base) was used to facilitate identification [37]. Raw files were uploaded into the METABOLOMICS-SNETS-V2 workflow from GNPS (Global Natural Products Social Molecular Networking) in order to cluster the data based on their MS/MS fragmentation pattern. The precursor ion mass tolerance and the fragment ion mass tolerance were set at 2.0 Da and 0.5 Da respectively, and the cosine score was set at 0.7. The generated molecular network was visualized using Cytoscape 3.7.1 software.

To encounter the dynamic nature of the biotransformation process, a data analysis workflow for in vitro gastrointestinal biotransformation was implemented, previously developed and validated using hederacoside C as a model compound [23,27]. Briefly, data were converted to the open-source mzData format to allow further processing [38]. XCMS was used to convert the raw data into features via peak-picking, using following parameters: ppm = 10, peakwidth = c(5, 25), snthresh = 10, noise = 1000, mzdiff = 0.01, prefilter = c(3, 5000), integrate = 1. XCMS was followed by grouping, and EDGE was used for the extraction of significant differential profiles [39,40]. Tinderesting, an interactive Shiny app developed in R, was used to rate the quality of the resulting features. These ratings were used to train a random forest model for predicting experts response. The machine learning model provided a single score for each feature, referred to as tinderesting score, which allowed ranking of all features based on the difference over time between the three groups. The maximal score of 1 corresponds to the model labeling this feature as interesting, and the minimal score of 0 defines an uninteresting feature [23,27].

Several peaks in the output correspond to the same type of sample molecule, reflecting its isotope pattern. Isotopes were removed, using following parameters: IsoMass = 1.003355, IsoMassMaxDiff = 0.05, RTdiff = 10, corrThreshold = 0.8, ppmIsoDiffThreshold = 5.

The metabolic pathway of herniariasaponin H was studied since it is the most abundant saponin present in the extract and it is structurally related to hederacoside C, which was used for optimization of the workflow [23]. Both compounds contain a sapogenin linked to sugar moieties via the hydroxy groups at C-3 and C-28 of the aglycone. Herniariasaponin H comprises medicagenic acid as aglycone, which only differs from hederagenin by an additional hydroxy group at C-2 and a C-24 hydroxycarbonyl group instead of a hydroxymethyl group. However, the challenge in revealing the metabolic pathway of herniariasaponin H was that these experiments were not conducted with a single saponin, but with an herbal extract, comprising a complex mixture of compounds of high structural variety.

A suspect screening method, based on in silico metabolite prediction, was combined with a nontarget screening workflow to enhance the identification of products formed by in vitro liver biotransformation [41].

A list of potential biotransformation products was generated using Meteor Nexus 2.1 (Lhasa Limited, Leeds, UK). For phase I biotransformation, all redox and non-redox biotransformations were selected. For phase II biotransformation, $O$-glucuronidation, $O$-sulfonylation, acetylation and...
conjugation with amino acids were selected. The maximum depth was set at 3, and the maximum number of biotransformation products at 1000.

Biotransformer was used as an additional software tool to predict phase II metabolites. The tool uses both a knowledge-based approach and a machine-learning-based approach to predict biotransformation [42]. The SMILES string of medicagenic acid was uploaded, and “Phase I Transformation” and “Phase II Transformation” were selected separately. The generated csv file contained InChIKey, synonyms, major isotope mass, molecular formula, type of biotransformation reaction and precursor ID.

A non-target screening workflow was conducted with MZmine 2.29 software (pluskal, mzmine 2) after converting the raw data files to the open-source mzXML format. Firstly, m/z features were detected using the centroid algorithm followed by a chromatogram building step. Resulting chromatograms were deconvoluted using the noise amplitude algorithm. Further data reduction occurred by deisotoping, keeping the lowest m/z value as the representative isotope. Meanwhile, chromatographic peaks were also filtered according to peak width: only peaks with a width between 0.05 and 1.00 min were retained. Next, peaks were aligned across samples using the random sample consensus (RANSAC) alignment algorithm correcting for nonlinear shifts in retention times. Finally, using the same rt and m/z range gap filler algorithm, any missing peaks were reiteratively extracted. The obtained m/z features were imported in R and processed as described by Vervliet et al. [43]. In brief, fold changes between samples and negative controls were calculated. A volcano plot was constructed to plot the p-value from a student t-test as a function of the calculated fold change for every feature. Features with a p-value lower than 0.05 and a log 10-fold change higher than 10 were selected for in depth investigation [41].

4. Conclusions

Since there is no prophylactic therapy for urinary stone disease, there is an urgent need for new drugs. An integrated strategy, based on natural prodrugs and their metabolites, is developed to characterize new lead compounds derived from *H. hirsuta*. The combination of a comprehensive extraction method and biotransformation via in vitro gastrointestinal and hepatic models, followed by metabolomics profiling, is an innovative concept to identify and further develop new lead compounds for drugs.

Comprehensive extraction led to the tentative identification of 15 saponins that have not been reported before in *H. hirsuta*. The automated data analysis workflow used for unbiased screening for metabolites revealed the gastrointestinal biotransformation pathway of several compounds. A decrease in relative abundance over time was observed for the majority of all identified compounds, especially during the colon phase. Additionally, the relative abundance of saponin aglycones increased, illustrating the biotransformation of possible saponin prodrugs to their respective aglycones, which was also observed for flavonoids. Further hepatic biotransformation of the most abundant saponin aglycone revealed further biotransformation of medicagenic acid to different phase I and phase II metabolites, which can be used as a possible target for in vitro activity studies against urinary stones.

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