Development, Validation and Application of a UHPLC-MS Method for the Quantification of Chios Mastic Gum Triterpenoids in Human Plasma*

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ABSTRACT
Chios mastic gum is the resinous secretion obtained from the barks of the shrub Pistacia lentiscus var. Chia, which is endemic to the Greek island of Chios. Since antiquity, Chios mastic gum has found several uses as a phytotherapeutic remedy, primarily for the treatment of gastrointestinal disorders while recently, Chios mastic gum was also recognized by EMA as an herbal medicinal product with specific indications. Chios mastic gum’s biological properties are attributed to triterpenes which comprise the major chemical group (approx. 70%) and notably isomasticadienonic acid and masticadienonic acid. However, due to their structural characteristics, the isolation thereof in high yield and purity is challenging and since they are not commercially available, pharmacological studies aiming to assess their biological properties are limited. In the present work, mastic’s phytochemical investigation by UPLC-HRMS is followed by the isolation and characterization of isomasticadienonic acid and masticadienonic acid to be used as analytical standards for their accurate and reliable quantification in human plasma. A UHPLC-Q-MS method that was developed and validated (in terms of specificity, linearity, limit of quantification, accuracy and precision), for the direct quantification of the targeted compounds in the low ng/mL range of concentration, was subsequently implemented on plasma samples of healthy volunteers thus demonstrating its fitness for purpose. The results presented herein might provide insight to the understanding of this traditional natural product consumed notably for its anti-inflammatory, antioxidant and lipid lowering properties. Moreover, this method might serve as a starting point for any study aiming to monitor bioactive triterpenes in biological fluids.

* Dedicated to Professor Arnold Vlietinck on the occasion of his 80th birthday.

* These two authors contributed equally to this work.

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Introduction

Chios mastic (mastiha or CMG) is the resinous secretion obtained from the wounds of the trunk and branches of *Pistacia lentiscus* L. var. *Chia*, a shrub of the Anacardiaceae family. Even if *P. lentiscus* shrubs are widespread in the Mediterranean basin, the exclusive source of mastic resin is only the trees of the Chia variety growing on the Greek island of Chios [1]. Mastiha is strongly linked to the insular tradition of Chios, and systematic cultivations dominate mainly the south part of the island due to the particular pedoclimatic and microclimatic conditions [2]. CMG is a product with protected designated origin and since 2014, the traditional cultivation of mastic resin in Chios is recognized as intangible cultural heritage by UNESCO [3]. Traditional usage of mastic as a spice and a therapeutic agent dates back more than 2500 years. The ancient Greek physicians mentioned its properties and recommended its use especially for stomach and liver problems. Mastic’s trade and value flourished during the Byzantine years and Medieval ages, when CMG occupied a special spot in folk medicine and, later on, in official Pharmacopoeias across Europe and Asia [4].

Several recent studies have attempted to investigate mastic’s pharmacological potential. In fact, CMG has been found to possess anti-ulcer [5], anti-dyspepsia [6], as well as antifungal and antibacterial activities [7–9]. In 2015, an assessment of mastiha was initiated by the committee on Herbal Medicinal Products of the European Medicines Agency as a traditional herbal medicinal product with the following indications: treatment of mild dyspeptic disorders (indication 1) and symptomatic treatment of minor inflammations of the skin and as an aid in the healing of minor wounds (indication 2) [10,11]. Mastiha was also found to be effective against *H. pylori* [12] and to demonstrate positive action against CD [13] as well as have antiproliferative activity in colon and prostate models in vitro and in vivo [14–18]. Back in 1975, Abdel-Rahman et al. [19] investigated the antioxidant activity of mastic, showing it was comparable to that of butylated hydroxyanisol, the most used synthetic antioxidant in the food industry. Later, other studies confirmed the potent antioxidant character of mastic against, notably, the oxidation of animal and vegetal oils [20] as well as of human low-density lipoprotein [21]. Furthermore, it was also shown that mastic displays anti-inflammatory activity in human aortic endothelial cells [22]. Anti-inflammatory action of mastiha was also demonstrated in active CD patients who presented a significant reduction of the CD activity index and of plasma interleukin-6 and C-reactive protein levels after a 4-week administration [13].

Interestingly, following these findings, focus was given to its hypolipidemic activity [23]. A clinical trial on 133 human subjects over 50 years old receiving a different daily dose of mastiha over 18 and 12 months was conducted [24]. Results showed that the high-dosage group displayed a decrease in serum total cholesterol, LDL, total cholesterol/HDL ratio, lipoprotein, apolipoprotein A-1, apolipoprotein B (apoB/apoA-1 ratio did not change), SGOT, SGPT, and gamma-GT levels. Potential in vivo hepatoprotective and cardioprotective effects of mastic were recently studied further in both animals [25] and humans [26]. A pilot study was carried out with over 156 healthy human volunteers receiving daily doses of mastic under different forms over an 8-week period. It was found that a daily dose of 1 g of crude mastiha has a significant lowering effect on total cholesterol and glucose levels, with excellent tolerance and no detectable side effects, especially in overweight and obese individuals [26]. Meanwhile, many products containing mastic can be found on the market, e.g., flavoring, liquor, natural chewing gum, dentistry products, and dietary supplements and is used in traditional therapies (relief of gastrointestinalgia, protection against peptic ulcer) [4].

Responsible for the biological and pharmacological properties of mastiha is its unique composition. From a chemical point of view, it is a considerably complex mixture of approximately 120 constituents that have been reported so far. Triterpenoids comprise the major chemical group corresponding to the 67–70% of the total resins’ weight. Terpenes have attracted particular attention from the scientific community and from the pharmaceutical industry for clinical studies and applications in the therapy of diseases due to their antioxidant [27] and anti-inflammatory properties, both associated with the biological activities reported in mastiha [28]. It is worth noting that many triterpenoids of natural origin and/or their synthetic analogues are in the clinical trials stage [29].

In more detail, the main constituents of mastiha are the triterpenic acid isomers, namely, 24Z-IMNA, MNA, and oleancolic acid as well as 24Z-IMLA, MLA, and moronic acid [30,31]. In addition to the triterpenic acids, mastic resin also contains neutral triterpenes [32] and significant quantities of the polymer cis-1,4-poly-β-myrcene [33].

Nevertheless, despite the scientific interest of mastiha triterpenoids directly related to its biological properties and phytothera-
phytochemical investigation of CMG soluble polymer greatly impedes sample handling and the overall bioactive natural products and their further development, since high yield and high purity are required to achieve meaningful results. To our knowledge, only two studies have been published dealing with this subject so far [34, 35].

Thus, in the framework of a study on the bioavailability of mastic resin terpenes and postprandial effect on human antioxidant potential [36], the scope of the present work was to address the need for analytical tools for the direct quantification of mastic characteristic constituents IMNA and MNA. Initially, IMNA and MNA were isolated in pure form and high yield from CMG in order to be used as reference standards for the development and validation of an LC-MS methodology in human plasma, suggesting a complete workflow. The method was also validated according to ICH [37] and might serve as a valuable tool to any future study aiming to examine the pharmacological effect of CMG’s constituents in human subjects.

Results and Discussion
As already mentioned, mastic resin is a very complex mixture of terpenes together with a natural polymer of poly-β-myrcene. Thus, the first and crucial step in CMG’s extraction and phytochemical investigation was the removal of the insoluble polymer, which due to its gum-like formation, hinders sample handling and the overall phytochemical investigation of CMG’s constituents. These facts lead to insufficient isolation procedures along with very low yields and purity, which inevitably results in the unavailability of reference compounds for further investigations [4]. Consequently, also similar is the lack of efficient and reliable analytical methods for the accurate identification of mastic triterpenoids and their quantitation, not only for quality control purposes (in resin) but also for the exploration of their pharmacological properties (in plasma, urine, tissues). However, this is a common obstacle in the field of bioactive natural products and their further development, since high yield and high purity are required to achieve meaningful results.

Since the current study focuses on the characteristic triterpenic acids of mastic and their quantitation in plasma, the AF was forwarded for separation, isolation, and purification. Thus, semipreparative SFC-UV/MS using a chiral stationary phase and MeOH as the cosolvent was used for this purpose, an approach which has been previously proposed, with minor modifications [39]. As shown in Fig. 2, this method offered a satisfactory separation between the two targeted compounds, which enabled their collection and isolation in high purity. Interestingly, the difference in response factors between UV and MS detection can be easily no-

![Fig. 1](structure.png) Structural formula of isolated compounds isomasticadienonic acid (IMNA) and masticadienonic acid (MNA).

![Fig. 2](chromatogram.png) SFC-UV/MS chromatograms of the enriched extract from the acidic fraction (AF) of the total mastic extract without polymer (TMEWP) [upper, UV chromatogram at 220 nm and lower, ESI (−)MS SIR chromatogram of m/z = 453.4]; peaks at Rt = 9.5 and 10.2 minutes collected as fraction 1 and fraction 2 corresponds to isomasticadienonic acid (IMNA) and masticadienonic acid (MNA) respectively.
ticed. In the latter, a minor compound of the extract, i.e., peak eluted at Rt = 9.0 min, displayed an intensity greater than those of the target compounds. This illustrates the necessity of using relevant analytical standards for the implementation of MS-based quantitative methodologies for the measurement of mastic gum triterpenes. A total of 50 mg of mastic’s Af was processed. Next, the collected fractions were merged and MeOH was evaporated. The remaining quantities were 18.7 mg for fraction 1 and 20.5 mg for fraction 2, which corresponds IMNA and MNA, respectively. Both fractions were transferred for further analysis, including NMR and HRMS for structure verification and purity evaluation to be used as reference standards, with a purity of >95% for IMNA and >99% for MNA (Figs. 25 and 35, Supporting Information).

Reference compounds of the analytical quality required is the most common issue that analytical chemists working in the field of natural products are facing when developing methodologies for the characterization of phytoconstituents in extracts or biological fluids. The most common way to overcome this issue is to use commercially available analytical standards of the same chemical family. The quantification of cinnamic acids as caffeic acid equivalents or polyphenols as gallic acid equivalents are well-known examples [40]. This strategy is very efficient when performing UV detection-based analysis, although it is limited when seeking to reach a lower LOQ implementing MS detection-based analysis. Indeed, ionization efficiency and matrix effects can possibly lead to a great difference in the response factor even for compounds that are structurally close. In that case, isolation of the targeted compounds appears to be the best option, even though it might be a challenging task, especially in regard to complex mixtures and structurally close compounds as in the case of triterpenic acids of mastic.

The next step in the experimental process was the development and validation of the analytical method for the quantitation of MNA and IMNA in plasma. To satisfy the sensitivity aspects of the analysis, a UHPLC system hyphenated to tQ was employed. Due to the particular chemical nature of the analytes, a pseudo-MRM (453.4 > 453.4) method was used. As illustrated in Fig. 3, where the reference compounds MNA (10.57 min) and IMNA (10.95 min) are compared with TMEWP, there are no coeluting compounds. The relative retention time of MNA and IMNA standards in relation to the TMEWP were 1.05 and 1.03, respectively, and the resolution between compounds was 1.1, thus demonstrating the effectiveness of the separation. It is useful to note that TMEWP displayed two other peaks eluting at 7.15 and 7.55 min identified as moronic acid and oleancolic acid, both isomers of MNA and IMNA [38]. Moreover, 3-O-acetyl ursolic acid was selected as an IS, demonstrating close structural features with MNA and IMNA (pseudo-molecular ion at m/z 497.3634, C32H49O4, Δm < 0.5 ppm). The IS was derived after acetylation of ursolic acid. Finally, the unequivocal assessment of the targeted compounds in the presence of the components that may be expected to be present was demonstrated.

A short-term stability study was initially conducted in order to demonstrate the extent to which the analyte’s response is stable in defined conditions, so as to guarantee the reliability of the analytical process. Stability of MNA and IMNA was evaluated by the recovery of the response (ratio of area analyte/IS) of standard solutions at different time points with that of zero time. Stability conditions were similar to those of the storage conditions of the samples to be further analyzed, i.e., 10°C. Three independent replicates of standard solutions were freshly prepared in MeOH at three different concentrations (20, 100, 300 ng/mL, n = 9) and were analyzed on 5 different days (days 1, 2, 3, 4, 5, and 7). The average recoveries of the replicate samples for every time point are displayed in Fig. 4. At the low level of concentration (20 ng/mL), recovery for the IMNA standard was of 100.65% after 7 days and of 100.33% for MNA, while respective RSDs were 6.11 and 4.70%. At 100 ng/mL, the IMNA standard displayed recoveries ranging from 100.09 to 106.40% with respective RSDs of 5.73 and 1.16% over the investigated period. In the same conditions, MNA standard minimum and maximum recoveries were 99.43 and 100.56% with an RSD of 1.58 and 4.99% respectively. Similar results were observed for both compounds at 300 ng/mL, with the average recovery for all investigated time points calculated at 102.27% for IMNA and 100.36% for MNA. Corresponding RSDs were 4.63 and of 5.06%, respectively. Finally, considering all analyzed samples and time points (n = 54), both solutions of IMNA and MNA in MeOH are stable during a period of at least 7 days at 10°C, reaching a maximum acceptable deviation of signal response of 10%. Consequently, all quantitative analysis took place within the characterized stability period.

Calibration curves were constructed as a function of area ratio (target compound/IS) against the concentration ratio (target compound/IS) for the 15 levels of concentration. Each level was randomly injected three times during the analytical sequence in-between all samples to be analyzed. Calibration curves for both targeted compounds in solvent are presented in Fig. 45, Supporting information. Coefficients of determination (R²) were 0.9995 for MNA and 0.9966 for IMNA. Moreover, a statistical evaluation of model linearity by ANOVA was performed. First, homogeneity of variances between each concentration level was verified using Cochrane’s C test with a significance level (α) of 0.05. In these condi-
tions (15 series of 3 data points each), the upper limit critical value was 0.33. The maximum observed C value was of 0.24 for MNA and 0.30 for IMNA, therefore, variances can be considered homogeneous for both compounds, hence repeatability is stable in the calibration model range. Then, F-tests were performed at the a level of 0.05 to compare variances of modelized values against residuals. Calibration models were shown to be significant for both compounds. Indeed, the explained data’s F values (mean square of modelized values/mean square of residuals) were greatly superior to F_5%, which implies that the tested variances are not equals; the calibration model explains the data. Also, F values of “nonlinearity’ (mean square of residuals/mean square of repeatability residues) were inferior to F_5%; linearity of the models is verified, and the defined calibration models are suitable. ANOVA tables for both compounds are detailed in Table 1S, Supporting Information.

LOQ is described as “the lowest amount of analyte in a sample that can be determined with acceptable precision and accuracy under the stated experimental conditions” [37], and therefore it should belong to a range of concentrations for which the linearity of the concentration against the measured signal has been demonstrated. Hence, the first level of the calibration curve for each analyte was defined as the instrumental LOQ. Taking into account the dilution factor from the preparation of plasma samples, the method LOQ was determined to be in plasma 6 ng/mL for IMNA and 1 ng/mL for MNA. Consequently, three independent pooled samples of control plasma (t = 0) were spiked accordingly to verify method performance at the LOQ. At this level, the RSD of the signal response was of 12% for IMNA and 5% for MNA and the difference of the average signal response with this of a standard solution at the same concentration was + 27% and −8% for IMNA and MNA, respectively.

Accuracy of an analytical method can be defined as the closeness of agreement between the measured value obtained by implementing this method and the true value [37]. This definition corresponds to what is designated as “trueness” in the International Vocabulary of Metrology and in ISO documents [41]. In the present work, the accuracy was evaluated by the preparation and analysis of three plasma samples artificially prepared so as to contain IMNA and MNA at low (20 ng/mL), medium (100 ng/mL), and high (300 ng/mL) levels of concentration. As described in the experimental section, the blank matrix consisting of pooled plasma samples of the zero time (pre-ingestion) were spiked accordingly and analyzed, following the whole analytical procedure including sample preparation. The experiment was then repeated after 1 week. The average measurement error in intermediate precision conditions (n = 6, 2 days × 3 independent replicates), that is to say the difference between the measured and the theoretical concentrations of the spiked samples, was determined for each investigated level of concentration.

Precision of an analytical method can be defined as “the closeness of agreement (degree of scatter) between a series of measurements” [37]. In the present study, precision of the method is expressed as the SD of the measurement bias of six independent samples analyzed in intermediate precision conditions. Results are presented in ▶ Fig. 5. At the low level of concentration (20 ng/mL), the measurement error of the IMNA standard was −6.88 and −1.53% for MNA, while associated SDs were 6.86 and of 6.09% respectively. At 100 ng/mL, the difference between the measured and theoretical concentrations was −1.25% for IMNA and 1.35% for MNA. At this level, corresponding SDs of the measurement error were 4.26 and 4.00%, respectively. Finally, at the high level of concentration (300 ng/mL), both analytes presented a slight negative measurement bias, which was −7.69% for IMNA and −7.45% for MNA, with associated SDs calculated at 6.47 and 4.90%, respectively. Considering all analyzed samples, accuracy

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▶ Fig. 4 Seven-day stability of standard solutions at low, medium, and high concentrations (i.e., 20, 100, and 300 ng/mL) kept at 10°C (error bars represent the relative SD between three independent replicates).
and precision of the present method were demonstrated for both targeted compounds within a maximum acceptable deviation of 20% as shown in Fig. 5.

After validation of the initial performances, the herein method was implemented for the measurements of IMNA and MNA in plasma samples of 17 healthy male adults after oral administration of 10 g of CMG powder dispersed in water [36]. Suitability of the analytical system was checked by investigation of the internal quality control samples. None of the targeted compounds were detected in the procedural blank nor in the injection blanks, which were performed every 10 injections. Pooled QC samples were injected 5 times in the beginning, 5 times at the end of the sequence, and every 40 injections in-between. The RSD of the MNA signal response in the QC samples was 6.24 and 11.87% for IMNA. These performances are in accordance with the those demonstrated during the method validation. The MNA and IMNA average concentrations in QC plasma samples were 19.10 ng/mL and 31.57 ng/mL, respectively. Plasma kinetic parameters are discussed and presented elsewhere [36] and will not be detailed here. Briefly, at zero time, both compounds were found to be inferior to the LOQ. The target compounds plasmatic concentrations then reached a maximum in the range of 30–767 ng/mL for MNA and 68–1381 ng/mL for IMNA before decreasing to a low ng/mL after 24 h. Maximum plasmatic concentrations were generally observed between 2 and 4 h after intake, even though this could vary depending on subjects.

To conclude, an analytical method using an LC-tQ platform was fully validated in order to establish for the first time reliable quantitative data on the bioavailability of active terpenic acids from CMG in human plasma. Moreover, a complete workflow was designed from the purification of the compounds of interest, MNA and IMNA, in the required purity and yield, to a full validation of a method for their quantitation in human samples as well as its application to real samples. A semipreparative SFC-UV/MS apparatus could serve as a method of choice for terpenoids in complex mixtures offering high purity (dual detection), high yield, as well as speed due to the high degree of automation. Furthermore, the methods described herein can be used as a valuable tool in future pharmacological evaluations of CMG’s constituents, since the detection and quantification of mastic’s bioactive triterpenic acids in plasma might provide further insight on the dose/response relationship of mastic’s supplementation.

Materials and Methods

Chemicals and reagents

For mastic’s extraction, all solvents were of analytical grade and purchased from Sigma-Aldrich. HCl (37%) and Na₂CO₃ (99.5%) were also purchased from Sigma. Based on previous works, 3-O-acetyl ursolic acid was used as an IS [34]. It was prepared by acetylation of commercial ursolic acid (p > 98.5%) with pyridine in diethyl ether, all acquired from Sigma-Aldrich. Food grade CO₂ was obtained from Revival A.E. HPLC and LC-MS grade solvents were purchased from Merck Chemicals and high-purity water was provided by a Millipore Direct-Q 3 UF purification system. Optima LC-MS grade formic acid was obtained from Thermo Fisher Scientific and NMR solvents were purchased from Eurisotop.

Chios mastic gum extraction

CMG was kindly provided by the Chios Mastic Growers Association. AF was obtained as previously described [12]. In brief, 76 g of crude mastic powder was dissolved in ethyl acetate/methanol 1:3 (600 mL in total). After 2 days at room temperature, the insoluble and decanted polymer was removed and the solution was filtered and condensed, giving a total of 66.8 g of TMEWP. TMEWP was thrice partitioned in a separatory funnel between aqueous 5% Na₂CO₃ and ether (1 L of each for all repetitions), which afforded 15.2 g of NF in the organic phase. The aqueous phase (3 L in total) was subsequently acidified with 1 N HCl. The acidic solution was reextracted with 1 L of ether, and the organic phase afforded 1.7 g of AF. All solvent evaporations were performed using a rotary evaporator (Buchi) with a water bath at 40 °C.

Isolation of isomasticadienonic acid and masticadienonic acid as analytical standards

MNA and IMNA were isolated by SFC instrumentation using a previously described method with some modifications [39]. In brief, instrumentation consisted of a Waters Prep 15 SFC Purification System coupled with a SQ Detector 2 mass spectrometer. The mass analyzer was interfaced with a Z-spray ion source operated in ESI negative. A ChiralPak IC column (5 μm, 4.6 × 250 mm) was used for the fractionation of the AF. The mobile phase was supercritical CO₂ with MeOH as an organic modifier. Chromatographic flow rate was set at 5 mL/min. Gradient elution started at 5% co-

![Fig. 5](image-url) Measurement error of blank matrix (pooled plasma samples t = 0) spiked at low, medium, and high concentrations (i.e., 20, 100, and 300 ng/mL) analyzed in intermediate precision conditions (n = 6: 2 days × 3 independent replicates, error bars represent the SD, dotted lines represent a maximum acceptable deviation 20%).
solvent for 2 min and increased to 30% in 10 min before going back to the initial condition in 30 s for a 2.5-min equilibration step. MS detection was carried out by simultaneously using an SIR function monitoring m/z = 453.40 as well as a full scan function with a mass range of 200 to 800 amu.

**Characterization of mastic’s fractions and the compounds used as analytical standards**

TMEWP, AF, and NF were analyzed by UPLC-HRMS. Liquid chromatography analysis was performed on an Acquity UPLC System. Detection was performed on a Thermo LTQ-Orbitrap XL hybrid mass spectrometer equipped with an ESI and an APCl source. Ten μL of each fraction at 100 μg/mL were injected into the system. Separation was achieved on a Supelco Ascentis C18 column (15 cm × 2.1 mm, 3 μm) using a water gradient containing 0.1% (v/v) formic acid (A) and ACN (B). Elution started at 5% B, which increased to 70% B in 3 min, reaching 100% B in a total of 20 min. These conditions were maintained for 4 min before returning to initial conditions in 1 min for a 3-min reequilibration (28 min in total). The column was maintained at 40 °C and the flow rate was set to 0.4 mL/min. HRMS data were acquired in the negative mode in the full scan m/z range of 115–1000 with a resolution of 30000. Data-dependent acquisition was simultaneously performed using a CID value of 35% and a mass resolution of 7500. For the ESI source, the capillary temperature was set to 350 °C and the source voltage was 2.7 kV. Tube lens and capillary voltage were respectively tuned at −100 V and −30 V. Nitrogen was used as the sheath gas (40 au) and auxiliary gas (10 au). For the APCl source, the capillary temperature was set to 200 °C, the APCl vaporizer temperature was set to 400 °C, and the source voltage was 6 kV. Tube lens and capillary voltage were respectively tuned at −125 and −30 V. Nitrogen was used as the sheath gas (70 au) and auxiliary gas (20 au). Spectral interpretation was performed using Xcalibur (Version 2.2) software.

Isolated compounds and the IS were characterized by HRMS and NMR. Direct infusion experiments were performed on an LTQ-Orbitrap XL hybrid mass spectrometer equipped with an ESI source. MS data were acquired in the negative mode in the full scan m/z range of 100–1000 with a resolution of 30000. 1D and 2D nuclear magnetic resonance spectra (1H, 13C, COSY, COSY-LR, HSQC-DEPT, HMBC) were acquired on a Bruker Avance 600 MHz spectrometer using CDCl3 as the solvent. Chemical shifts (δ) are expressed in ppm with reference to the solvent signals (δH 7.26, δC 77.0).

**Recruitment of the healthy volunteers and dosage information**

Apparently healthy men, aged 20–40 years old, were invited for screening (ClinicalTrials.gov Identifier: NCT02847117). The study protocol was reviewed and approved by the Harokopio University Ethics Committee (49/29–10–2015). It was conducted according to the principles of the Declaration of Helsinki and followed the Principles of Good Clinical Practice. Enrollment was based on certain inclusion and exclusion criteria. More specifically, a BMI >26 kg/m², history of alcohol or drug abuse, medication, vitamin or inorganic supplements intake as well as following a vegan or macrobiotic diet were considered exclusion criteria. A subject presenting gastrointestinal diseases such as IBD, peptic ulcer, or gastrointestinal cancer was also excluded from the study. Seventeen subjects were found to be eligible and followed a diet low in phytochemicals for 5 days. CMG intake (10 g) took place after overnight fasting. Blood samples were collected at specific time intervals (e.g., 0, 1, 2, 4, 8, and 24 h). Plasma was separated from serum by centrifugation immediately after blood collection. Finally, 1 mL of each sample was placed in microcentrifuge tubes and stored at −80 °C until further treatment.

**Preparation of plasma samples**

The protocol for the preparation of plasma samples was modified from Lemonakis et al. [34]. First, plasma samples were thawed in ice at 4 °C for 30 to 60 min. Then, 50 μL of each sample was transferred in a microcentrifuge tube of 1.5 mL in which 2 × 200 μL of cold ACN (−20 °C) were added for protein precipitation. Tubes were thoroughly mixed in vortex for 15 s each time ACN was added to the sample. Tubes containing precipitated proteins were centrifuged at 4 °C and 12 000 rpm for 10 min using a Mikro 200R centrifuge. After that, supernatants were transferred into microcentrifuge tubes and spiked with the IS in order to reach a final concentration of 40 ng/mL after reconstitution. Samples were further dried down in an Eppendorf vacufuge concentrator for 2.5 h. No heating was applied during the drying process. Once dried, samples were stored at −80 °C until the day of analysis when they were reconstituted in 100 μL of MeOH.

**Liquid chromatography mass spectrometry of plasma samples**

LC-MS analysis was carried out using an Advance UHPLC system coupled to a Bruker EVOQ Elite Triple Quadrupole Mass Spectrometer. Separation was achieved on an Acquity HSS T3 column (100 mm × 2.1 mm, 1.8 μm) heated at 40 °C with a gradient of water containing 0.1% (v/v) formic acid (A) and ACN (B) at a flowrate of 0.4 mL/min. Injection volume was set to 5 μL. Elution started at 70% B and reached 89% B in 12.5 min, then it increased to 100% in 0.5 min. These conditions were maintained for 3 min before going back to initial conditions for a 4-min reequilibration. The mass spectrometer was equipped with a heated ESI source operated in the negative mode. A tension of 5 kV was applied to the capillary. Heated probe and cone temperatures were set at 300 and 350 °C, respectively, while probe and cone desolvation gas (nitrogen) flows were set at 40 and 20 units, respectively. The nebulizer gas flow was set at 50 units and the exhaust gas was on. Argon under a vacuum of 1.5 mTorr was used as the collision gas. Finally, acquisition was performed in pseudo-MRM mode [42]. The deprotonated molecular ion was chosen as the precursor as well as quantifier and confirmation (453 > 453 for IMNA and MNA, 497 > 497 for 3-O-acetyl ursolic acid). Collision energies were set at 20 and 30 V for quantification and confirmation transitions, respectively. Q1 and Q3 were both operated with standard resolution settings and scan times were of 50 ms for each transition.

**Quality control**

Internal quality controls were systematically performed. LC-MS grade MeOH was used as injection blanks in order to control carryover. Moreover, procedural blanks made of high-purity water
were prepared to monitor eventual cross contamination during sample preparation. Pooled QC samples, which consisted of thoroughly mixed small aliquots of each plasma sample to be studied, were also prepared in order to check system performance before, during, and after analytical series. Finally, blank matrix, which was prepared from small aliquots of t = 0 plasma sample from each healthy volunteer participating in the study, was spiked for the verification of method performance.

**Method validation**

Specificity of the LC-MS method was investigated by the analysis of standard solutions in MeOH of MNA and IMNA as well as TMEWP. Concentrations were 10 ng/mL of MNA and IMNA, and 50 ng/mL for TMEWP. The resolution between the targeted compounds was determined and retention times of each standard in relation to the TMEWP were observed.

Short-term stability tests were performed over a one week period in order to guarantee the reliability of the analytical process. Fresh standard solutions were prepared in 3 independent replicates at 3 different concentrations (20, 100, 300 ng/mL, n = 9) and were analyzed on 5 different days (days 1, 2, 3, 4, 5, and 7). During this period, samples were left in the autosampler at 10 °C in similar storage conditions to those of the samples to be analyzed further.

Calibration curves were built upon 3 analytical replicates of standard solutions at 15 different levels of concentration (n = 45). Analyte concentrations ranged from 3 to 800 ng/mL for IMNA and from 0.5 to 500 ng/mL for MNA while IS concentrations were fixed at 40 ng/mL. Calibration curves were determined in intermediate precision conditions during 5 different days. Each sample was randomly injected three times. Calibration models were then analyzed by observation of R² and plots of residuals. Moreover, statistical evaluation of model linearity was conducted by ANOVA.

LOQs were determined by the concentration of the first level of the calibration curves. LOQs were then verified in matrix by spiking 3 sample replicates of blank matrix (pooled plasma samples t = 0) accordingly. Acceptance criteria were based on the observation of the RSD of the signal response with a maximum acceptable deviation of 20% and of its difference with a standard solution at t = 0 accordingly. Acceptance criteria were based on the observation of the RSD of the signal response with a maximum acceptable deviation of 20% and of its difference with a standard solution at t = 0 accordingly. Acceptance criteria were based on the observation of the RSD of the signal response with a maximum acceptable deviation of 20% and of its difference with a standard solution at t = 0 accordingly. Acceptance criteria were based on the observation of the RSD of the signal response with a maximum acceptable deviation of 20% and of its difference with a standard solution at t = 0 accordingly. 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Method accuracy was evaluated by the measurement error of a spiked blank matrix (pooled plasma samples t = 0) with a maximum acceptable deviation of 20%. All spiked samples were prepared in 3 independent replicates at 3 different concentrations (low, medium, and high: 20, 100, 300 ng/mL, respectively, for each analyte) while the IS concentration was 40 ng/mL. The experiment was repeated after one week. The SD of the measurement error between samples at each low, medium, and high levels of concentration was used to evaluate precision in intermediate precision conditions.

**Supporting information**

The following are available as supporting information: UPLC-HRMS profile of CMG AF with two different ionization sources (APCI/ESI) operated in the negative mode (Fig. 15), ESI(–)-HRMS spectra of fraction 1 (IMNA) and fraction 2 (MNA) (Fig. 25), 1H NMR spectra of IMNA and MNA (Fig. 35), calibration curves of both targeted compounds in solvent (Fig. 45), and variances analysis of the calibration models of the targeted compounds (Table 15).

**Contributors’ Statement**


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**Conflict of Interest**

The authors declare that they have no conflict of interest.

**References**


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