

1 Article

2 Assessment, Validation and Application to Real 3 Samples of a RP-HPLC Method for the Determination 4 of Guayulins A, B, C and D in Guayule Shrub

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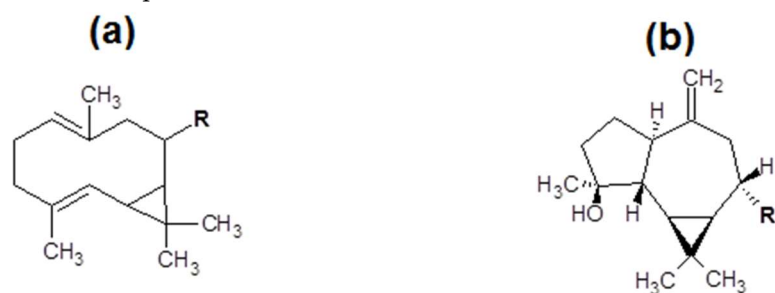
20 **Abstract:** Guayule (*Parthenium argentatum* Gray) is a shrub native of the arid regions of Mexico. In
21 the last decades, significant attention for its cultivation has risen because it is the raw material for
22 the production of hypoallergenic natural rubber. Guayule biomass contains also high amounts of
23 resin, which is not normally exploited in any way. Among other sesquiterpenic esters, guayulins
24 (i.e. the parteniol esters of cinnamic acid, guayulin A, or of anisic acid, guayulin B) are contained in
25 resin. In addition, minor amounts of guayulin C and guayulin D are formed by
26 degradation/oxidation of guayulins A and B, respectively. Guayulins likely act as cinnamate and
27 p-anisate reservoirs for Guayule shrub, in addition, it has been postulated that they might have a
28 key role in the chemical defense system of Guayule. Furthermore, it seems reasonable that
29 guayulins may possess significant biological properties (e.g. antibacterial and anticancer activities),
30 in close analogy with those shown by sesquiterpene lactones contained in many other species of
31 *Parthenium* genus. As a matter of fact, guayulins A and B play an important role in the synthesis of
32 antineoplastics used in breast cancer treatment. In this contribution we propose an original and
33 validated RP-HPLC approach to the simultaneous quantification of guayulins A, B, C and D. The
34 procedure of resin extraction from Guayule biomass has been optimized in terms of both extraction
35 method and solvent. RP-HPLC separation has been accomplished by an Ascentis® C18 column
36 under isocratic elution with a 80:20 (v:v) acetonitrile:water mixture. Validation was carried out in
37 terms of limits of detection and quantification, linearity, precision, and trueness. Finally, the
38 method was tested with a number of fresh and seasoned samples of spontaneous Guayule shrub
39 from Mexico.

40 **Keywords:** Guayule; *Parthenium argentatum* Gray; Resin; Guayulins; RP-HPLC.

1 Introduction

Guayule (*Parthenium argentatum* Gray) is a perennial and low-growing shrub belonging to the Asteraceae family, native of arid regions of Mexico and the southwestern United States. Since the early years of the last century it has gained considerable industrial and scientific interest for the production of a kind of natural rubber exhibiting properties similar to that obtained from the *Hevea brasiliensis* tree [1,2]. In the last decades, the discovery that guayule latex is free from the allergenic proteins contained in the *Hevea* latex led to a reemphasis and expansion of guayule research, mainly aimed to the production of hygiene and medical products compliant with the strictest standards of quality for hypoallergenic rubber [3]. Together with rubber, Guayule plant also contains a resinous material (the so-called *resin*), which has to be separated from rubber in order not to compromise its physical and technological properties. Rubber (8-26% of dry weight) and resin (5-25% of dry weight) are present in amounts roughly equivalent in Guayule shrub [1]. Since the resin fraction is soluble in polar solvents (like acetone), whereas the rubber one can be dissolved only in non-polar solvents (like hexane), sequential or simultaneous extractions have to be made in order to separate these fractions from plant tissue and, eventually, the resin from the latex [1]. The high manufacturing costs of the process of rubber extraction from Guayule make the success of its exploitation dependent in large extent by the possibility of using as much as possible the residual plant fractions, like resin or bagasse, as raw materials to obtain high-value coproducts [4].

Guayule resin is usually obtained from the extraction by the ground whole shrub or the coagulated latex with polar solvents, usually acetone [5]. Despite the fact that it is often not valorized and hence discarded, resin is a very interesting fraction from both an academic and an industrial point of view. Indeed, it is not only rich in sesquiterpenoids and triterpenes of potential commercial value [6], but it also contains a variety of interesting secondary metabolites of the shrub, like guayulins [6-8]. Guayulins A ((1R,2S,4Z,8Z,10S)-4,8,11,11-tetramethylbicyclo[8.1.0]undeca-4,8-dien-2-yl(2E)-3-phenylprop-2-enoate) and B ((4Z,8Z)-4,8,11,11-tetramethylbicyclo[8.1.0]undeca-4,8-dien-2-yl 4-methoxybenzoate) are respectively the cinnamic and anisic esters of the partheniol ((1S,2E,6E,9S,10R)-3,7,11,11-tetramethylbicyclo[8.1.0]undeca-2,6-dien-9-ol) [7], whereas the guayulin C ((1AR,1BR,2R,4AR,7S,7AR)-2-hydroxy-1,1,2-trimethyl-5-methylidene-decahydro-1H-cyclopropa[E]azulen-7-yl (2E)-3-phenylprop-2-enoate) and guayulin D ((1AR,1BR,2R,4AR,7S,7AR)-2-hydroxy-1,1,2-trimethyl-5-methylidene-decahydro-1H-cyclopropa[E]azulen-7-yl 4-methoxybenzoate) are likely formed by the oxidation of the guayulines A and B, respectively [6,8]. Scheme 1 reports the structures of these molecules.



R = *t*-cinnamate, guayuline A
R = *p*-anisate, guayuline B

R = *t*-cinnamate, guayuline C
R = *p*-anisate, guayuline D

35 **Scheme 1.** Structures of guayulins A and B (a) and guayulins C and D (b).

36 Guayulins A and B are abundant in the resin. Namely, up to 10% of the resin mass is constituted
37 of guayulin A [9-11], while the relevant amount of guayulin B in the resin from India is between 0.1
38 and 3% [10]. The wide variability in the amounts of guayulins A and B in the resin depends on a
39 number of factors, such as: soil nature, climate, phenological phases, cultivar, age and – mainly – the
40 nature of the different parts of the shrub [10,11]. On the other hand, the concentrations of guayulins

1 C and D in resin seems to be much lower than those of guayulins A and B. Schloman et al. estimated
2 that the amounts of guayulins C and D in resin are between 50% and less than 10% of the amount of
3 the guayulin A and B, respectively [6]. Guayulins likely act as cinnamate and p-anisate reservoirs for
4 Guayule shrub: when it is required by the plant, a metabolic turnover releases the corresponding
5 free acid [4]. The hypothesis that guayulins and rubber may have a common biochemical link [12]
6 suggested Teetor et al. [11] to use the amounts of guayulins A and B as possible predictive tools in
7 order to evaluate the rubber content in Guayule shrub, but the results obtained were absolutely
8 unsatisfactory. It is noteworthy that, in *Heliantheae* tribe and *Parthenium* genus, only *Parthenium*
9 *argentatum* contains both guayulins and rubber, whereas it does not contain any sesquiterpene
10 lactone (i.e. one of the principal families of compounds easily found in almost all the other species of
11 *Parthenium* genus) [13]. Since the sesquiterpene lactones are synthesized in the *Parthenium* genus
12 plants as chemical defense agents against animal attacks (these molecules are toxic for livestock, a
13 deterrent for insects and cause severe contact allergies in mammals) [14], it seems reasonable that
14 guayulins might play a similar physiological role also in *Parthenium argentatum*. Results of
15 pioneering studies conducted by Rodriguez et al. supported this insight [15]. As a matter of fact,
16 guayulin A has been found to be a powerful contact allergen for guinea pigs [8] while the same
17 behavior has not been observed for guayulin B. For humans, the sensitizing power of guayulin A is
18 much smaller than for laboratory animals. Hence, the possibility that traces of guayulin A may be
19 present in the guayule rubber at a sensitizing level seems to be remote at the moment [16]. In
20 analogy to what observed for sesquiterpene lactones in the most of the species of *Parthenium* genus, it
21 is likely that guayulins may also exhibit other significant biological properties (i.e. antibacterial and
22 anticancer activities) [15]. Indeed, guayulines A and B act as biological triggers in the synthesis of
23 lychnostatine and paclitaxel, which are antineoplastic agents used in breast cancer treatment.

24 It is evident that the continuous rise of interest in the properties of guayulins needs the
25 development of reliable, accurate and sensitive analytical methods for their identification and
26 quantification. Until now, the analytical methods reported in literature for the measurement of the
27 amount of guayulins in Guayule resin [3,4,8,11,13,17,18,19] or latex [16] are scarce and all lacking of
28 any validation protocol. Obviously, chromatographic methods are dominant in the analytical
29 characterization of such analytes and, among them, HPLC is the preferred approach. While few
30 papers reported the presence of guayulins C and D in extracts from resin [6,8,11], the quantification
31 of these analytes has been never simultaneously carried out to that of guayulins A and B [6,8].
32 Hence, it is evident that at present a validated HPLC procedure aimed to simultaneously quantify
33 the amounts of all known guayulins is not reported in literature. For this reason, the principal aim of
34 this contribution is to develop, validate and test with real samples an original RP-HPLC method
35 devoted to the quantification of guayulins A, B, C and D in different parts of Guayule shrub from
36 Mexico. As a side result of this study, different approaches of extraction of analytes from the matrix
37 were also compared in order to increase efficiency and make the procedure more eco-friendly.

38 2. Materials and Methods

39 2.1. Samples and Sample preparation

40 Spontaneous Guayule was collected in the province of San Pedro del Gallo, Durango, Mexico,
41 along the highways where it naturally grows. All the harvested Guayule shrubs were about 60 cm
42 high, thus estimating to be between 15 and 18 year old [20]. These plants were harvested in 2015 in
43 two sites located in the Durango desert. The first one (sampling of January 2015) is in the Mexican
44 Federal Highway 30 (road section Bermejillo-El Palmito, GPS coordinates: 25°52'20.68"N,
45 104°06'26.07"W), whereas the second one (sampling of September 2015) is along the highway Paso
46 Nacional-San Pedro del Gallo (GPS coordinates: 25°40'50.41"N, 104°16'33.53"W). A total of twenty
47 plants (ten for each sampling) were harvested following the recommendations specified by the
48 Official Mexican Rule [21], which establishes the procedures for the sampling, transportation and
49 storage. Shrubs collected in the harvest of January 2015 were air-dried at room temperature in open
50 air for eight months, whereas shrubs collected in the harvest of September 2015 did not undergo any

1 drying process. Only a few millimeters of rain fell during the first harvest. Just arrived in the
2 laboratory, all shrubs were cut up to constitute three fractions: leaves (L), fine stems (less than 10
3 mm in diameter, FS) and coarse stems (more than 10 mm of diameter, CS). All fractions were stored
4 in a freezer at -18°C until sample preparation. Before extraction, samples were allowed to reach the
5 room temperature. Later, all samples were dried at 50°C until reaching constant weight, roughly
6 pieced and then ground.

7 2.2. Chemicals and reagents

8 All reagents were at least of analytical grade. Ethanol (assay $\geq 99.8\%$), acetone ($\geq 99.8\%$) and
9 acetonitrile ($\geq 99.9\%$) were purchased from Sigma–Aldrich (Milan, Italy), whereas ultrapure (Type 1)
10 water (specific resistance 18 M Ω) was used throughout the analyses. Guayulins A, B, C and D were
11 from 1717 CheMall (Mundelein, IL, USA).

12 2.3. Instrumentation

13 Ultrapure water was from a New Human Power II Scholar UV apparatus (Human Corporation,
14 Seoul, Korea). A Retsch Grindomix GM 200 knife mill (Verder Scientific, Torre Boldone, Italy) was
15 used to grind the samples before extraction. ASE extractions were performed by an Accelerated
16 Solvent Extractor Dionex ASE 200, equipped with a Dionex Solvent Controller System (Dionex
17 Corporation, California, USA), whereas ultrasonic assisted extractions were performed by a
18 Bandelin-Sonorex model RK 255 H ultrasonic bath. Solvent was evaporated by extracted samples by
19 a Büchi Rotovapor model EL130 (Büchi, Cornaredo, Italy). The HPLC equipment consisted of a
20 Series 200 binary pump, a sampling valve, a 20 μ L sample loop, a Series 200 LC column oven, and a
21 Series 200 UV–vis variable wavelength detector, all from PerkinElmer, Milan, Italy. Data were
22 processed using a Turbochrom Workstation Software (PerkinElmer, Milan, Italy). Before use, the
23 mobile phase was filtered through a 0.45 μ m membrane from Millipore (Bedford, MA, USA) to
24 remove any particulates.

25 2.4. Validation

26 Validation of the proposed method was accomplished on the basis of limit of detection (LoD),
27 limit of quantification (LoQ), linearity, precision (measured as both repeatability and intermediate
28 precision) and trueness (measured by recovery tests of each analyte on real Guayule samples). LoD
29 was calculated according to the Upper Limit Approach (ULA1) approved by IUPAC [22]. For each
30 analyte, four different solutions at increasing concentrations not far from the expected LoD (i.e.
31 between 0.02 and 0.1 mg L⁻¹ for each analyte) were prepared and analyzed. Each measurement was
32 performed in triplicate. In addition, the ULA1 approach recommends that the LoQ value is three
33 times the relevant LoD value. Linearity was checked on at least three orders of magnitude of
34 concentration, as a function of the relative abundance of each guayulin in the different parts of the
35 Guayule shrub. Precision was evaluated in terms of both repeatability (i.e. the CV measured in five
36 consecutive replicates of the same sample in the same analytical session) and intermediate precision
37 (i.e. the CV obtained in five consecutive replicates of the same sample in different analytical sessions
38 along two weeks). The acceptability of these precision scores was checked in terms of HorRat ratio
39 values (i.e., the ratio between experimental and theoretical CV measured on the basis of the
40 Horwitz's theory) [23]. Due to the lack of any certified reference materials or of reliable independent
41 analytical methods, trueness was estimated through recovery tests. Four aliquots of the acetone (or
42 ethanol) extract of the stems were submitted to the overall analytical procedure after the addition of
43 increasing amounts of each guayulins to three of them. For each analyte, a plot of analytical
44 concentration versus the added amounts of mass was obtained: the recovery is represented by the
45 percent slope value of the regression line. The estimation of bias was made in duplicate.
46 Acceptability of the trueness values has been established according to the guidelines described in the
47 AOAC manual for peer-verified methods [24].

1 3. Results

2 3.1. Choice of the extraction procedure.

3 The choice of extraction technique of resin by Guayule biomass (i.e. leaves, fine stems, coarse
4 stems) was accomplished by means of a preliminary comparison among three well-known
5 procedures: the traditional extraction with warm solvents (40 °C) (WSE), the ultrasonic assisted
6 extraction (UAE) and the accelerated solvent extraction (ASE). Acetone (i.e. the most used solvent
7 for the extraction of the resin from Guayule biomass) and ethanol, one of the most important green
8 polar solvent, were the pure solvents used in this phase. Quantification of resin in organic extracts
9 has been performed by means of UV-Vis spectroscopy according to literature methods [25].

10 3.1.1. Procedures of extraction

11 3.1.1.1. Extraction with warm solvents (WSE)

12 2.5 g of selected Guayule biomass were suspended in 15 mL of acetone (or ethanol). The
13 mixture was stirred and heated up to 40°C in a thermostatic bath for 20 min. After the separation of
14 the solvent from the biomass and its replacement with a new 15 mL aliquot, the extraction procedure
15 was repeated for additional two times. The three extracts were collected and made up to the final
16 volume of 50 mL with pure solvent.

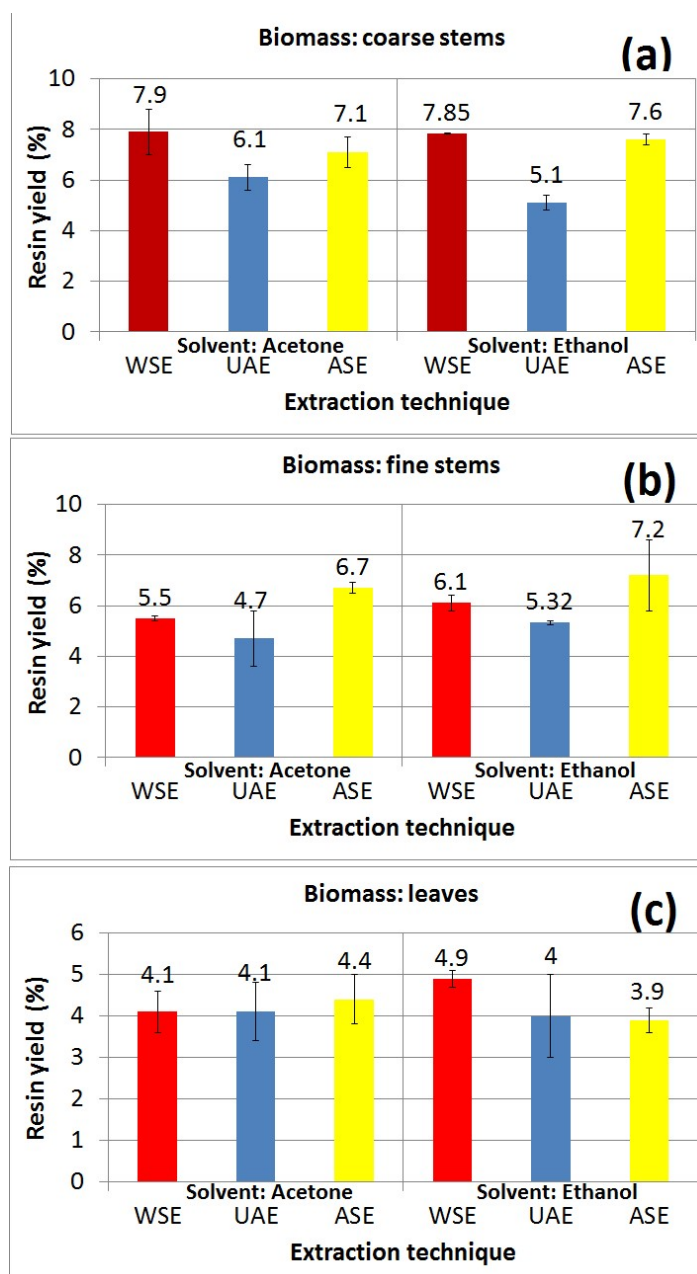
17 3.1.1.2. Ultrasonic assisted extraction (UAE)

18 2.5 g of selected Guayule biomass were suspended in 15 mL of acetone (or ethanol). The
19 mixture was stirred at room temperature in an ultrasonic bath for 20 min. After the separation of the
20 solvent from the biomass and its replacement with a new 15 mL aliquot, the extraction procedure
21 was repeated for additional two times. The three extracts were collected and made up to the final
22 volume of 50 mL with pure solvent.

23 3.1.1.3. Accelerated solvent extraction (ASE)

24 5.0 g of selected Guayule biomass were transferred in a 22 mL thimble. Extraction was
25 performed using 20 mL of the chosen solvent (acetone or ethanol). The extraction temperature was
26 set at 40°C. Three consecutive extraction cycles have been performed for each biomass aliquot. The
27 organic extracts were joined and made up to 100 mL with pure solvent.

28 Figure 1 reports a comparison of resin yields (% on the initial biomass amount) measured for
29 each extraction method and each biomass analyzed.



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Figure 1. Comparison of average solvent extraction yields in resin (% on the biomass amount), n = 3. Solvents: acetone or ethanol. **(a)** biomass: coarse stems; **(b)** biomass: fine stems; **(c)** biomass: leaves.

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Data obtained substantiate that the UAE method is characterized by the least recovery in resin, whereas the performances of WSE and ASE methods are roughly comparable as a function of the nature of Guayule biomass and the extraction solvent. However, it is possible to note that the net amount of resin extracted from the fine stems by means of ASE method is statistically higher than that obtained by WSE, whereas the yields in resin measured in the extraction from coarse stems are not statistically different among them from both WSE and ASE methods. Keeping into account the better intrinsic reproducibility of the ASE approach as compared with the WSE method, the ASE extraction method with acetone and ethanol was chosen as the technique used of the present analytical method.

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3.2. Assessment of the chromatographic method

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The first RP-HPLC methods aimed to quantify guayulins in Guayule resin were developed more than thirtyfive years ago, and were only aimed to the determination of guayulins A and B. On the other hand, reliable RP-HPLC methods devoted to measure the concentration of guayulins C and

1 D are still absent in the literature. For the reader's convenience, Table 1 summarizes the key features
2 of different literature RP-HPLC methods for the determination of guayulins.

3 **Table 1.** Key features of selected literature RP-HPLC methods of analysis for guayulins in resin
4 extracts from Guayule.

Stationary phase (mm x mm x μm)	Mobile phase (v:v)	Flow rate (ml min^{-1})	λ of quantification (nm)	Guayulins quantified	Reference
MicroPak MCH-10 (300 x 4.6 x 10)	from $\text{CH}_3\text{CN}:\text{H}_2\text{O}$ 50:50 to $\text{CH}_3\text{CN}:\text{H}_2\text{O}$ 75:25	n.r.	262	A and B ¹	[6]
C18 column ²	$\text{MeOH}:\text{H}_2\text{O}$ gradient elution ²	n.r.	215	C and D	[8]
RP-C18 column ²	$\text{MeOH}:\text{H}_2\text{O}$ 89:11	1	254	A and B	[10]
Microsorb-MV (250 x 4.6 x 5)	from $\text{CH}_3\text{CN}:\text{H}_2\text{O}$ 80:20 to pure CH_3CN	1.5	262	A and B ³	[11,16,17,19]
Altex Ultrasphere ODS type (150 x 4.6 x 5)	$\text{MeOH}:\text{H}_2\text{O}$ 93:7	1	254	A and B	[13]
Lichrosorb RP-18 (250 x 4.6 x 10)	$\text{CH}_3\text{CN}:\text{H}_2\text{O}$ 75:25	2.5	254	A and B	[18]

5 ¹ Guayulins C and D were quantified using a GC method; ² no further details were provided in the
6 paper; ³ in ref. [11], guayulins C and D were only identified, but not quantified.

7 Furthermore, it is of utmost significance to underline that no analytical method proposed until
8 now has been validated, and this poses serious doubts about the overall reliability of the data
9 reported. The results of preliminary tests performed on selected literature methods suggested the
10 assessment of an original procedure, based on a more recent C18 stationary phase than those
11 reported in Table 1. Ascentis[®] C18 is a monomeric-type, fourth-generation,
12 octadecylsiloxane-bonded silica stationary phase, introduced in the market in the early 2000s, and
13 characterized by unusually high surface area ($450 \text{ m}^2 \text{ g}^{-1}$) if compared with other C18 phases having
14 similar porosity ($10 \mu\text{m}$) and surface coverage ($3.7 \mu\text{mol m}^{-2}$). Since Ascentis[®] stationary phases have
15 already been successfully used for the HPLC determination of terpenoids in vegetal matrices [26],
16 we considered useful to check the performances of an Ascentis[®] C18 (25 cm x 4.6 mm x 5 μm)
17 column in the separation of guayulins A, B, C and D. Further steps of the method assessment have
18 been devoted to optimize the composition of the mobile phase and to the choice of the best
19 wavelength of UV detection. Starting from the literature methods, different methanol:water and
20 acetonitrile:water mixtures have been tested, working both in isocratic and in gradient elution. An
21 acetonitrile:water 80:20 (v:v) solution provided the best results. In addition, the acquisition of
22 chromatograms in the wavelength range between 250 and 280 nm demonstrates that the best
23 compromise among different UV maximum absorptions of the analytes is the λ at 262 nm, as
24 previously reported in the literature [4,6,10,11]. Table 2 summarizes the operating conditions of the
25 chromatographic method proposed.

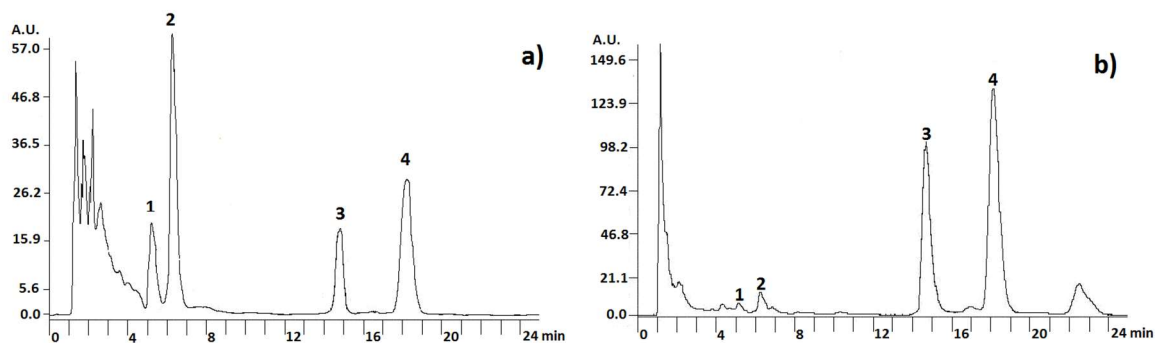
1 **Table 2.** Chromatographic conditions proposed for the contemporary RP-HPLC separation and
 2 quantification of guayulins A, B, C and D in resin extracts from Guayule biomass.

Stationary phase (length, mm x diameter, mm x particle size, μm)	Ascentis® C18 (250 x 4.6 x 5)
Mobile phase (v:v)	CH ₃ CN:H ₂ O 80:20
Elution mode	Isocratic
Volume (μL)	20
Flux of mobile phase (mL min^{-1})	1.5
UV wavelength (nm)	262
Column temperature ($^{\circ}\text{C}$)	25
Length of the chromatographic run (min)	25

3 The adoption of an Ascentis® C18 RP-HPLC column, more recent in comparison to those reported in
 4 literature (see Table 1) have allowed us to develop an isocratic method achieving the complete
 5 resolution at baseline level of all analytes in less than 20 minutes, thus with large time savings of the
 6 chromatographic run as compared with the most recent literature method [11].

7 3.3. Analysis of the resin extracts

8 2.5 mL of acetone (or ethanol) ASE extract of the resin from guayule biomass were evaporated,
 9 at room temperature and under reduced pressure, to dryness. The resin residue was dissolved in 5
 10 mL of acetonitrile, and then filtered through a 0.45 μm nylon filter before HPLC analysis. Figure 2
 11 shows typical chromatograms of ASE resin extracts from Guayule leaves (Figure 2a) and coarse
 12 stems (Figure 2b).



13 **Figure 2.** Guayulins A, B, C and D in resin extracts from a) Guayule leaves (sample L1, extraction
 14 solvent: acetone) and b) Guayule coarse stems (sample CS2, extraction solvent: ethanol). Peak
 15 attribution: 1; guayulin D, 2; guayulin C; 3, guayulin B, 4, guayulin A. A.U. = arbitrary units.
 16

17 Guayulins peaks were identified by comparing retention times with those of standard
 18 solutions, and the attribution of each analyte was confirmed by spiking each peak in the real sample
 19 with a standard solution containing known amounts of pure guayulins. In order to determine the
 20 retention times, the reference standards were injected both individually and as a mixture.
 21 Quantification was accomplished by external linear calibration on three different concentration
 22 levels in the relevant linearity interval of each analyte. Each sample was analyzed three times, and
 23 each analytical datum is reported as the average value \pm the relevant standard deviation.

1 3.3. Validation

2 Table 3 lists the validation parameters for the method proposed. Low LoD values (always
3 below 0.1 mg L⁻¹) supported a good sensitivity of the method proposed, which is able to quantify
4 analytes at concentration levels between 0.1 and 0.2 mg L⁻¹, as a function of the analyte. Also linearity
5 is more than satisfactory. It has been checked only within the three orders of magnitude of
6 concentration (i.e. inside the range of variability of the guayulins in the different parts of Guayule
7 shrub), obtaining values of the determination coefficients R² ranging between 0.9994 and 0.9984, but
8 it is likely that the linear dynamic interval for each analyte may largely overcome these ranges. In
9 addition, the analysis of the residuals of the regression line has excluded any possible deviation from
10 linearity of each calibration plot. A preliminary evaluation of the samples has substantiated a very
11 wide scattering of the concentration of the analytes in them. Due the fact that the meaning of the
12 precision measurements crucially depends on analyte concentration [23], repeatability and
13 intermediate precision were evaluated on the Guayule extracts characterized, for each analyte, by a
14 concentration that was as close as possible to its average value in the relevant calibration interval.
15 For this reason the ethanol extract of sample FS1, the acetone extract of sample FS2, the acetone
16 extract of sample FS1 and the ethanol extracts of sample CS1 were chosen for the precision
17 measurements for guayulins A, B, C and D, respectively. The acceptability of the levels of the
18 precision data, ranging between 1.3 and 3.2% (guayulin A) and 3.6 and 10% (guayulin D) for
19 repeatability and intermediate precision, respectively, were evaluated by a fitness-for-purpose
20 methodology based on the Horwitz's theory [23]. Operatively, acceptable values of the HorRat ratio
21 (i.e. the ratio between the experimental CV and the theoretical CV calculated on the basis of the
22 Horwitz's theory) should be less than 1.5. Since the HorRat ratios for the procedure subject of this
23 work never exceeded 0.6, the precision levels obtained from all analytes can be considered
24 acceptable. Trueness evaluations have been accomplished with the same Guayule samples involved
25 in the precision measurements. Recoveries obtained ranged between 73±5% (guayulin D) and 91±2%
26 (guayulin B) for acetone extracts, whereas ethanol extracts gave recoveries between 77±8% (guayulin
27 D) and 89±6% (guayulin A). The evaluation of the recovery values obtained on the basis of the
28 AOAC guidelines [24] have allowed to substantiate – for all analytes - a slight underestimation bias.
29 Unfortunately, the complete absence in the literature of validated methods for the determination of
30 guayulins prevented us to make any comparison with results from previous studies.

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Table 3 - Validation data

Guayulin n	Sensitivity (n = 3)		Linearity	Repeatability (n = 5)			Intermediate precision (n = 5)		Bias (n = 2)	
	LoD (mg L ⁻¹)	LoQ (mg L ⁻¹)		Concentration range (mg L ⁻¹)	R ²	CV _{exp,r} ^a	HorRat _r ^b	CV _{exp,IP} ^c	HorRat _{exp,IP} ^d	Recovery (%±SD ^e) on acetone
A	0.032±0.004	0.10±0.01	0.2 - 500	0.9994	1.3	0.26	3.2	0.43	87±4	89±6
B	0.051±0.004	0.15±0.01	0.2 - 250	0.9991	1.7	0.29	3.5	0.41	91±2	88±8
C	0.034±0.009	0.10±0.03	0.1 - 200	0.9989	1.5	0.58	4.0	0.39	81±9	80±10
D	0.055±0.005	0.16±0.02	0.16 - 200	0.9984	3.6	0.39	10	0.26	73±5	77±8

32 ^a CV_{exp,r} is the experimental coefficient of variation of repeatability, measured in the ethanol extract of FS1 for guayulin A, in the acetone extract of FS2 for guayulin
33 B, in the acetone extract of FS1 for guayulin C and in the ethanol extract of CS1 for guayulin D, respectively; ^b HorRat_r is the ratio between CV_{exp,r} and the theoretical
34 repeatability data (CV_{H,r}) according to Horwitz's theory; ^c CV_{exp,IP} is the experimental coefficient of variation of intermediate precision, measured in the ethanol
35 extract of FS1 for guayulin A, in the acetone extract of FS2 for guayulin B, in the acetone extract of FS1 for guayulin C and in the ethanol extract of CS1 for guayulin
36 D, respectively; ^d HorRat_{IP} is the ratio between CV_{exp,IP} and the theoretical intermediate precision data (CV_{H,IP}) according to Horwitz's theory; ^e standard deviation.

1 3.4. Application on real samples

2 Table 4 reports the analytical data obtained for the RP-HPLC determination of guayulins A, B,
3 C and D in acetone and ethanol ASE extracts from different Guayule biomass (leaves (L samples),
4 fine stems (FS samples) and coarse stems (CS samples)) harvested in January 2015 (dried samples, 1)
5 and in September 2015 (fresh samples, 2).

6 **Table 4.** Concentration of guayulins A, B, C and D in acetone or ethanol extracts of Guayule biomass
7 (leaves (L), fine stems (FS) and coarse stems (CS)) harvested in January 2015 (dried samples, 1) and in
8 September 2015 (fresh samples, 2).

Sample (solvent)	Guayulins (mg kg ⁻¹ ±SD)							
	A (A)	A (E)	B (A)	B (E)	C (A)	C (E)	D (A)	D (E)
L1	2080±40 ^a	2140±40 ^a	444±9 ^a	449±4 ^a	1638±8 ^a	1400±45 ^b	360±30 ^a	330±30 ^a
L2	2030±65 ^a	2190±20 ^a	428±6 ^a	460±9 ^a	1650±7 ^a	1560±45 ^b	370±20 ^a	330±20 ^a
FS1	3810±30 ^a	3230±40 ^a	615±9 ^a	500±20 ^a	372±6 ^a	204±4 ^a	8.4±0.7 ^c	6.52±0.07 ^a
FS2	4190±40 ^a	3640±10 ^a	1190±20 ^a	1150±9 ^a	61.5±0.1 ^a	26.4±0.7 ^a	<0.32 [*]	<0.32 [*]
CS1	8160±90 ^a	9830±20 ^a	700±10 ^d	884.8±0.7 ^a	1010±4 ^a	1044±7 ^a	25±2 ^a	55±2 ^a
CS2	8800±100 ^a	7800±100 ^a	3090±20 ^a	2810±40 ^a	249±4 ^a	199±4 ^a	23±2 ^a	16±2 ^a

9 SD = standard deviation; n = 3; *LoQ of guayulin D; paired values (i.e. same sample, same guayulins, different
10 extraction solvent) followed by the same letter are not significantly different according to ANOVA test at P =
11 0.05.

12 While data relative to guayulins A and B are roughly comparable with those previously
13 reported in literature [4,6,10,11], those relative to the amounts of guayulins C and D in acetone (or
14 ethanol) extracts by different parts of the Guayule shrub are - at the best of our knowledge -
15 unprecedented at all. Firstly, it is very important to highlight that these data are relative to a
16 spontaneous Guayule shrub, whereas almost all literature data are relative to specific and
17 well-known Guayule genotypes coming from cultivation.

18 3.4.1. Guayulins in Guayule leaves.

19 As a first remark, it is interesting to note that the relative composition in guayulins on leaves
20 from the first and the second harvest is roughly constant, and this is probably due to the fact that
21 also the leaves of the shrubs harvested in September 2015 arrived not entirely fresh in the laboratory,
22 like those harvested in January 2015. The relative composition of both guayulins is almost the same
23 as the nature of the solvent varies, with the only exception of a slight increase of the concentration of
24 guayulins C and D and a faint decrease of the guayulin A in the acetone extract in comparison to
25 those obtained by ethanol. The ratio between guayulin A and B in leaves is roughly close to 5:1. This
26 data is significantly higher than that (i.e., 1.3) measured by Sidhu et al. for a Gila cultivar [10], but is
27 within the range (between 2.5 and 7) measured by Teetor et al [11] in brown leaves of three Guayule
28 cultivars. On the other hand, leaves are very rich in guayulins C and D. In particular, the amount of
29 guayulin D in leaves is more than ten times higher than those found in the richest remaining part of
30 Guayule shrub (i.e. the coarse stems from the January harvest, CS1), whereas the concentration of
31 guayulin C in leaves is only ca. 50% higher than that measured in CS1. The significant increase of
32 concentration of guayulins C and D is likely a consequence of the drying process of the guayule
33 biomass. This fact confirms the hypotheses that these compounds could derive from
34 oxidation/degradation pathways from guayulins A and B, respectively [6,8]. The increase of
35 concentration is more evident for guayulin C rather than for guayulin D, but it is possible that this
36 should be caused by the difference between the relative abundance of guayulin A and guayulin B,
37 respectively.

1 3.4.2. Guayulins in Guayule stems.

2 The highest amounts of guayulins A and B were found in the coarse stems (CS samples).
3 Depending on the nature of the solvent used for the ASE extraction, amounts ranging between
4 7800 ± 100 and 9830 ± 20 mg kg⁻¹ were measured for guayulin A, whereas concentrations ranging
5 between 700 ± 10 and 3090 ± 20 mg kg⁻¹ were measured for guayulin B. These concentrations are in
6 partial agreement with those observed by Teetor et al. for guayulin A [11] and by Sidhu et al. for
7 guayulin B [10]. Indeed, the concentrations of guayulin B measured by Teetor et al. [11] and the
8 amounts of guayulin A found by Sidhu et al. [10] were not as high as those measured in this study. It
9 is interesting to note that the concentration ratio between guayulin A and guayulin B, which is
10 roughly constant at varying of the nature of the solvent, increases of more than 400% passing from
11 fresh to aged stems, and this is due to the loss of more than 70% in the average amount of guayulin B
12 during the storage. On the other hand, data show that changes of the concentration of the guayulin
13 A, albeit much lower than those shown by guayulin B, depend by both the nature of the extraction
14 solvent and the aging of the sample in non trivial way. Furthermore, the amount of guayulin C in
15 these samples tends to sharply increase (up to five times higher) with ageing, whereas the
16 concentration of this analyte is invariant towards the nature of the extraction solvent. Finally, only a
17 few tens of mg kg⁻¹ of guayulin D were found in the coarse stem samples. A moderate increase in
18 concentration of this analyte, likely due to aging effects, was observed only in ethanol extracts.

19 The amounts of guayulins in fine stems are intermediate between those measured in coarse
20 stems and leaves. The amounts of guayulins A and B in these samples are normally only 30-50% of
21 the relevant values measured for coarse stems, whereas the loss of guayulin B during the ageing is
22 only between 50 and 55% the amount measured in fresh samples. On the other hand, very high
23 increases of concentration of guayulin C were found (ranging between 600 and 770% in samples
24 extracted with acetone and ethanol, respectively) passing from fresh to aged samples, whereas just
25 few mg kg⁻¹ of guayulin D were measured only in FS1 samples, being the remaining samples below
26 the relevant LoQ for this analyte.

27 4. Conclusions

28 At the best of our knowledge, for the first time a RP-HPLC method specifically aimed to
29 simultaneously measure the concentration of the known guayulins in resin extracts from parts of
30 Guayule shrub has been developed, validated and applicated to real samples. The development of
31 the method has first involved the optimization of the resin extraction procedure from the Guayule
32 biomass among three different approaches (warm solvent extraction, ultrasonic assisted extraction,
33 accelerated solvent extraction). The latter technique provided best performances and was chosen to
34 be used in the analytical method. The HPLC separation was accomplished by means of an isocratic
35 elution on an Ascentis® C18 column with an 80:20 (v:v) mixture of acetonitrile and water. In this
36 way, the separation at the baseline level of all analytes in less than twenty minutes was obtained.
37 Low LoD and LoQ values, a very good linearity over more than three orders of magnitude and a
38 very good precision characterized the proposed method. On the other hand, a slight
39 underestimation bias was observed in all analytes. The method has been successfully tested to fresh
40 and aged samples of spontaneous Guayule shrub, harvested in the Chihuhaua desert (Laredo,
41 Mexico). The guayuline amount was measured as a function of the ageing level of the sample (fresh
42 or air-exposed for eight months), the parts of the plant (leaves, fine stems, coarse stems) and the
43 nature of the extraction solvent (acetone or ethanol). The data obtained are in substantial agreement
44 with those described in the literature for guayulins A and B, while there are no reference data for
45 guayulins C and D. The analysis of the stems revealed evident differences in the concentration of
46 guayulins as a function of their diameter, of the ageing level and - to a lesser extent - of the nature of
47 the extraction solvent. On the other hand, the analysis of the leaves showed no meaningful
48 differences among data obtained from samples obtained in the two crops or by variation of the
49 solvent nature. The leaves have been the richest fractions in guayulines C and D, while the coarse
50 stems have been the richest in guayulines A and B. The obtained data show that leaves constitute the
51 fraction of Guayule that is probably more is prone by the effect of oxidative degradation of

1 guayulins A (and B) and the consequent transformation in guayulins C (and D). This behavior is
2 much less evident, where it is present, on the stems. This analytical method is expected to be useful
3 in any further qualitative and quantitative evaluation of guayulins in different parts of Guayule
4 shrub as a function of the nature of the genotype, or the adoption of different cultivation practices.

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9 and N.S. wrote the paper.

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