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Mathematical optimization of the green extraction of polyphenols from grape peels through a cyclic pressurization process

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Abstract

In the current era of high consumption and increasing waste, many products that are believed to be unusable can find a new purpose in the market. For example, the grape peel waste resulting from the production of wine contains numerous bioactive compounds. In reality, grape peels are by-products of winemaking that can be conveniently reused in many different ways, including agronomic use and cosmetic industry applications. Moreover, the by-products can also be used in the energy field as biomass for the production of biogas or in food plants for the production of energy. In this article, to extract polyphenols, grape peels were processed via a cyclically pressurized extraction method known as rapid solid-liquid dynamic extraction (RSLDE), which does not require the use of any

organic solvent or include heating or cooling processes that can cause the loss of substances of interest. To better understand the cyclically pressurized extraction process, a numerical simulation was performed to evaluate the exchange between the grape piece solid matrix and water during the extraction process. Furthermore, a finite element model was used to numerically determine the time-dependent concentration distribution at specific times.

Keywords: Nutrition, Food technology, Food analysis, Food science

1. Introduction

Polyphenols, particularly flavonoids, are ubiquitous in plant foods (fruits, vegetables, legumes, and some beverages) and are also widespread in many medicinal plants. Numerous epidemiological and experimental studies have suggested that polyphenols have multiple beneficial effects on human health [1, 2]. The biological activity of polyphenols is associated with their chemical and biochemical properties, including their ability to act as antioxidants, their antineoplastic effects, their ability to regulate gene expression in chronic degenerative diseases, and their anti-diabetic potential [3, 4, 5, 6]. However, the bioavailability of polyphenols varies, depending mainly on the food source and the forms of polyphenols that it contains [7]. The grapes used for wine production represent a good example of an agricultural product rich in nutritional value and health benefits due to their vitamins, minerals and antioxidants. In particular, the flavonoids in grapes are located mainly in the peels. Some types of flavonoids are also found in grape seeds and stalks, but they are generally absent in the flesh. Approximately 15% of the fruit, however, consists of the peel, and peels represent a wine-making process waste material that is reused in part for the production of brandy, ethanol, tartaric acid, dyes, and fertilizers. Today, the use of agro-food processing waste by-products represents a concrete action that addresses the problem of agricultural production sustainability and provides the growing food, nutraceutical, pharmaceutical and cosmetic industries with low-cost, high-quality raw materials that were destined for disposal for the production of natural ingredients. In particular, the target of some food industries is to develop snacks and drinks composed of grapes, which are known to contain high amounts of polyphenols. To extract these polyphenolic compounds from grapes, a type of liquid-liquid extraction that involves the use of organic solvents, such as ethanol, methanol, formic acid or acetone and water in different proportions, is usually used. However, this type of extraction can involve considerable health risks and can be damaging to the environment; consequently, extraction methods that reduce the use of organic solvents, including microwave extraction [8], supercritical fluid extraction (SFE) [9], and ultrasound-assisted extraction (UAE) [10, 11], have recently been developed. To reduce the environmental impact and create a greener

extraction technique, this work investigated a green extraction process for polyphenols using the peels of Aglianico grapes, a prestigious variety of grape grown in Campania (Italy) that is particularly rich in nutrients and antioxidants; this process uses a cyclically pressurized extraction method known as rapid solid-liquid dynamic extraction (RSLDE) with a Naviglio extractor at room temperature [12]. Literature data show that conventional and non-conventional methods have been applied for the extraction of high-added value compounds from winery-processed materials. Non-conventional technologies represent a promising tool for recovering high-added value compounds from winery wastes and by-products. However, several parameters influence the choice of technology used to recover these compounds, such as the matrix being processed, selectivity, energy consumption, equipment cost, and value of the extract [13]. In particular, compared to other known techniques, RSLDE is an innovative solid-liquid extraction technology that allows rapid extraction of solid matrices containing extractable substances with an organic or inorganic solvent and with their mixtures. Most current methods aim to heat the extraction system to increase yield and accelerate extraction times, while RSLDE performs extraction at room temperature, using an increase in extraction liquid pressure on the solid matrix for extraction. Extracting at low temperatures is important because the use of low temperature avoids applying thermal stress on thermolabile substances. In this way, it is possible to determine the composition of the substances present in the different matrices without inducing changes in the active principles contained therein [12]. In addition, the process was modelled and studied using a finite element method.

2. Materials and methods

2.1. Plant material

This study was conducted using the Aglianico grape cultivar, which is a red grape cultivar that is widespread in Southern Italy and is renowned for the quality of its wines. It is grown in several Italian regions, but it is mainly cultivated in Campania from the renowned "Feudi di San Gregorio" vineyard in Sorbo Serpico, Avellino (Campania, Italy) (41°00'35.39"N/14°57'34.29"E). Four types of grape samples were examined: Group I: directly harvested grapes; Group II: peels and seeds from grapes that had been subjected to the crushing process and subsequently sent to a tank for maceration; Group III: crushed and dried grapes produced by Imepa (Southern Industry Drying Agricultural Products, Salerno, Campania, Italy); and Group IV: crushed, dried and shredded grapes produced by Imepa.

Grape samples (200 g) were extracted using a Naviglio extractor, series LAB, model EXNA0101 (500 cc). This method is a solid-liquid extraction technique based on the

cyclic application of a variable pressure. The instrument uses pressure (approximately 8-10 bar) in a rapid and simplified manner to extract substances not chemically bound to a solid matrix in a liquid medium at room temperature using deionized water as the solvent.

2.2. Instrumentation and chemicals

The instruments used in this study included a UV-VIS spectrophotometer, model 1601, equipped with a 1-cm optical path cuvette (Shimadzu, Tokyo, Japan); a gas chromatograph equipped with a programmable split-splitless injector (PSS), a flame ionization detector (FID) (Auto System XL) (Perkin Elmer, Norwalk, CT, USA) and an Rtx-5 capillary column ($l = 30$ m, i.d. = 0.25 mm, and f.t. = 0.25 microns; Restek, Bellefonte, PA, USA) with a 5% diphenyl and 95% dimethylsilicone stationary phase; a gas chromatograph, model 17A (Shimadzu, Tokyo, Japan), equipped with a split-splitless injector and interfaced with a mass spectrometer, model QP-5000, with an SPB-5 capillary column ($l = 60$ m, i.d. = 0.25 mm, and f.t. = 0.25 micron; Supelco, Bellefonte, PA, USA); and a Karl Fisher titrator, model KF 2026 (Crisson Instruments, Baar, Switzerland). All High Performance Liquid Chromatography (HPLC) grade reagents and solvents were purchased from Merck (Darmstadt, Germany). Before the extraction process, the dried and crushed grape samples were weighed with a Gibertini Europe 1700 analytical balance (Novate Milanese, Milan, Italy). The grape samples had a uniform moisture content of 18%.

2.3. Experimental tests

Shredded and dried Aglianico cultivar grape peels were supplied by Imepa and were prepared via the following steps: 1) receipt of frozen raw material; 2) defrosting in a cell with a maximum temperature (T) of 8 °C; 3) washing in sterile water with a T of <50 °C; 4) drying on looms in an air-steam exchange oven at a T between 40 °C and 70 °C for 2 hours at a relative humidity (RH) of <18% 5) slow grinding; and 6) packaging. Before packaging, the samples were also subjected to microbiological analysis that included a visual macroscopic analysis of 1000 g to assess anything indicating foreign bodies in the product, an evaluation of any visible formations suggestive of surface moulds or signs of entomic infestation, and a direct microscopic and parasitological analysis on 10 g of sediment diluted 1:10 to evaluate normal non-pathogenic microbial flora, rare fungi, and pests and their eggs. The notes, comments and corrections pertaining to the hazard analysis critical control point (HACCP) guidelines provided by Imepa stated that the product is hygienically healthy for the above parameters and that the fruit is completely free from any visible signs of biological contamination.

2.4. Folin-Ciocalteu method for total polyphenols

The total amount of phenolics was determined using the Folin-Ciocalteu method [14]. A calibration curve of gallic acid was prepared, and the results were determined using a regression equation from the calibration curve, with each measurement expressed as mg of gallic acid equivalents per gram of sample (GAE/g). In this method, 1 ml of each extract, diluted appropriately with deionized water to obtain an absorbance in the range of the prepared calibration curve, was mixed with 1 ml of 3-fold-diluted Folin-Ciocalteu phenol reagent. Two millilitres of 35% sodium carbonate solution was added to the mixture, which was then shaken thoroughly and diluted to 6 ml by adding 2 ml of water. The mixture was allowed to stand for 30 min, and the blue colour that formed was measured at 700 nm using a spectrophotometer. As the samples from Group IV had the highest concentrations of polyphenols, the assays were focused on these samples (Table 1).

2.5. Experimental tests

A 200-g sample of grapes (Group IV) was stored at 10 °C in a hermetically sealed plastic container to prevent changes in its moisture content. The dried grapes were mixed several times for 2 min. Prior to extraction, the moisture content of the grapes was determined using an oven method at 105 °C for 24 hours, and the moisture content was 18% w/w [15].

2.6. Determination of the size

The size of the solid matrix of grape pieces (Group IV), including its length, width and thickness, was evaluated using a digital calliper before and during the process of solid-liquid extraction. From these values, the sphericity and geometric mean diameter (GMD) of the solid matrix of the grape pieces were determined [16]. The values obtained were used in numerical simulations to construct a mathematical model of the process.

2.7. Determination of total volume

To measure the volume without changing the moisture content, the grape solid matrix (Group IV) was sealed with food-grade polyethylene, which was very suitable for the volume used in this study. The initial total volume (V_{pt}) was evaluated using a 1000-ml graduated cylinder with a water volume of 500 ml by dipping in it into the volume to be measured.

The accessory mass volume (V_a) was evaluated and was subtracted from the V_{pt} value, providing the real volume of the considered sample (V_p), that is, $V_p = (V_{pt} - V_a)$. Subsequently, to evaluate the initial volume and weight, the

extraction process was performed for grape piece samples withdrawn from Group IV, and a rapid solid-liquid dynamic extractor (RSLDE) was used. The volume and weight values were calculated again during the extraction phase.

2.8. Determination of initial concentration

Experiments were performed to evaluate solute extraction from the solid matrix for the considered extraction process. A solid matrix of grapes was chosen. The samples (Group IV) were stored at $-20\text{ }^{\circ}\text{C}$ in tightly sealed plastic containers to prevent changes in their moisture content, and they were mixed several times for 2 min. Before the extraction process, the concentration of the grape piece solid matrix content was determined [17].

2.9. Determination of the dry residue

A 10-ml volume of extract was filtered and added to a calibrated dish, and the extract volume was evaporated to dryness using an oven at $50\text{ }^{\circ}\text{C}$. This process was continued until the remaining liquid volume was negligible, and the oven temperature was then increased to $105\text{ }^{\circ}\text{C}$. The sample was placed in a desiccator and was subsequently cooled to $20\text{ }^{\circ}\text{C}$ for weighing. This operation was repeated until a constant weight was attained. The weight of the dry matter was calculated by subtracting the tare weight. Each determination was repeated three times, and the measurement differences were lower than 7%; the average value of each sample was determined and reported. After determining their initial dry weight and volume, the samples were extracted.

2.10. Extraction process using a Naviglio extractor

Two hundred gram grape samples (Group IV) were subjected to a total of 30 programmed cycles of pressure (with a maximum pressure of 10 bars) applied to the liquid phase in contact with the grape pieces over a period of 2 hours. The number of hits in the dynamic phase (nd) was 12; the dynamic operative phase (td) and the static operative phase (ts) were performed for 2 min each. All of the experiments were performed in triplicate, and their averages were used for numerical modelling and analyses. Determinations of the weight and volume were performed as described below at the times reported in Table 1 for the included samples (see the Results And Discussion Section). The grape samples were then subjected to programmed cycles of pressure applied to the liquid phase in contact with the product, as described above, while maintaining the same solid–liquid ratio. All of the experiments were repeated three times, and the measurement differences were less than 5%; therefore, the averages were used for numerical modelling and analyses.

First phase. In the first phase of the procedure, using a Naviglio extractor (Atlas Filtri Ltd., Padua, Italy), the initially sampled raw material was split into three batches of 200 g each.

Second phase. In the second phase of the procedure, a 200-g sample of the raw material was removed from the Naviglio extractor at the end of the extraction process and transferred to a sterile bag that was labelled to indicate the Naviglio procedure, extraction ending time, initial time, final time, product, quality, batch and date of sampling [18, 19, 20, 21, 22].

2.11. Mass spectrometric analysis

Matrix-assisted laser desorption/ionization mass spectrometry (MALDI MS) experiments were performed on a 4800 ABSciex MALDI time-of-flight/time-of-flight (TOF/TOF) tandem mass spectrometer equipped with a nitrogen laser (337 nm). Sample extract (0.5 μ l) was mixed (1/1, v/v) with a 10 mg/ml solution of 2,5-dihydroxybenzoic acid in acetonitrile/water (90:10, v/v). Spectra were acquired using a mass (m/z) range of 100–4000 Da for polyphenol identification. Selected extracted samples were dissolved in methanol and subjected to the same MALDI-TOF/TOF analysis. The different species detected in the MALDI spectra were fragmented by MALDI-TOF/TOF and identified based on literature data MALDI MS/MS spectra [23]. The species identified in the different samples are summarized in Table 2 (see the Results And Discussion Section).

2.12. Analytical approach

The diffusion process at a constant temperature follows Fick's second law. Considering axisymmetric diffusion, Fick's three-dimensional (3D) equation is:

$$\frac{\partial M}{\partial t} = D \left(\frac{\partial^2 M}{\partial x^2} + \frac{\partial^2 M}{\partial y^2} + \frac{\partial^2 M}{\partial z^2} \right) \quad (1)$$

where M is the concentration of the extract from the considered matrix at time t and D is the diffusion parameter. Crank (1979) [24] proposed a suitable solution for Eq. (1) that considers a material with a spherical geometry having radius value r , described by:

$$MR = \frac{M - M_i}{M_e - M_i} = 1 - \left(\frac{6}{\pi^2} \right) \sum_{i=1}^{\infty} \left(\frac{1}{i^2} \right) \exp \left(-Di^2 \pi^2 \frac{t}{r^2} \right) \quad (2)$$

$$MR = \frac{M - M_i}{M_e - M_i} \quad (3)$$

where M is the instantaneous content concentration at a specified time t . Thus:

$$MR = 1 - \left(\frac{6}{\pi^2}\right) \left(\left(\exp\left(-D\pi^2 \frac{t}{r^2}\right) \right) + \left(\frac{1}{4}\right) \exp\left(-4D\pi^2 \frac{t}{r^2}\right) \right) \quad (4)$$

where MR is the concentration ratio and Me and Mi are the equilibrium and initial concentrations of the solid matrix during the process, respectively. Often, only a finite number of parameters of Eq. (2) are utilized to evaluate the MR parameter. Frequently, researchers exclusively utilize the first two terms of Eq. (4).

In this paper, the MR values were evaluated at a determined time t and utilized as input in a MATLAB program (R2016b). The diffusion parameter D for the considered matrix was evaluated at every instant t was examined. The geometry of the grape matrix is approximately ellipsoidal, so the diffusion parameter in Eq. (1) had to be modified. Gaston et al. (2002) [25] determined a method to evaluate the De parameter in cases of ellipsoidal geometry utilizing the following expression:

$$D_e = f_e^2 \times D \quad (5)$$

where f_e is the sphericity parameter for the ellipsoid. Subsequently, the diffusion parameter D must be replaced with the evaluated De parameter.

2.13. Finite element approach for extraction with the cyclically pressurized extraction process

To numerically simulate the extraction process, a mathematical simulation was set up using a COMSOL (Rel 5.1) program. Evaluating the exchange phenomena between the grape matrix and the liquid phase during the extraction process was possible with this program. A time-dependent concentration distribution as a function of time was determined by applying a finite element model.

The shape of the grape matrix pieces was approximated as an ellipsoid. To model the grape matrix pieces, only a quarter of an ellipsoid was considered in 3D (Fig. 1). The COMSOL (Rel.5.1) program allowed the evaluation of the concentration parameter for each node at each time considered.

A complete mesh consists of 10784 domain elements, 1406 boundary elements, and 103 edge elements with 8 layers near the external surface. The number of degrees of freedom was solved for 3341 (plus 1002 internal DOFs) with solution times of 8 s using an i7 processor at 3.3 GHz with 16 GB of RAM.

The diffusion parameter was evaluated by fitting Eq. (2), and the obtained data were input into the COMSOL program for computation of the concentration parameter for the considered nodes with a time range of 1 second. The total concentration of the grape solid matrix was also evaluated at 0.1, 2, 4, 10, 20, 32, and 48 hours by averaging the values of the considered nodes applied for the extraction process.

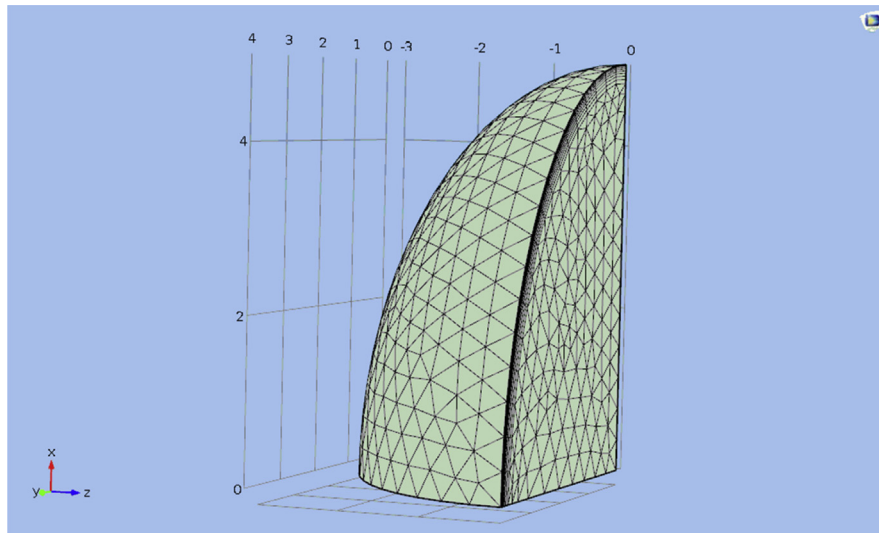


Fig. 1. Finite element grid for a single grape piece.

Moreover, the chemical and physical behaviour was defined by specifying the following parameters:

- effective diffusion coefficient for a spheroidal geometry
- initial essential oil content of the grape piece
- equilibrium essential oil content of the grape piece; and
- total time.

For a certain initial content concentration and for the proper diffusion coefficient of the material, the analysis evaluated the time-dependent concentration distribution at the nodes. An analogous approach has been performed to evaluate the temperature and moisture distribution during the cooling phase of the cryomaceration process for grapes [26]. For a finite element method, the calculated value of D is furnished in Eq. (1), and the MR parameters are computed for each node and for every instant t . The following hypotheses are considered when performing this approach for the considered matrix: (1) the diffusion parameter is independent of the concentration parameter; (2) the matrix is considered isothermal, and the heat exchange during the considered process is neglected; (3) the matrices are isotropic and homogeneous; and (4) the initial solute content of the grape matrix is uniformly distributed and constant inside the matrix.

3. Results and discussion

3.1. Extraction method

Table 1 shows the time period after which the extraction diagram was constant for each group and the corresponding value of the total extracted polyphenols. The

Table 1. Extractions performed and their results in terms of total polyphenols.

Group	Grape quantity used	Extraction process time	Total polyphenols
1	200 g grapes	72 h	264.8 ± 5.8 mg/L
2	200 g peels + seeds	48 h	674.5 ± 9.4 mg/L
3	200 g dried peels Imepa	48 h	4110.3 ± 20.8 mg/L
4	200 g dried and chopped peels Imepa	48 h	9210.4 ± 45.8 mg/L

(Mean value ± dxm = standard deviation of the mean).

kinetic curve for Group IV showed a higher content of extracted polyphenols than that observed for the other groups. Therefore, only the extraction model of Group IV was considered further, and only samples from this group were used to vary the geometry considered during the extraction process. Fig. 2 shows the extraction kinetics for Group IV during the cyclically pressurized process with a Naviglio extractor for polyphenol extraction. The trend of the rapid solid-liquid extraction kinetic dynamics was exponential, the initial extraction was very rapid, and the system reached a final equilibrium of polyphenols at approximately 48 hours, yielding a total extract of 9128 mg/L. The data used to evaluate the extraction kinetics diagram were computed in triplicate. Total polyphenol values were examined via a statistical study using multifactorial analysis of variance (ANOVA). The statistical importance of every parameter was evaluated utilizing the least significant differences (LSD) test. The data were statistically examined using Statgraphics Plus 5.1 software.

The total polyphenol content was determined spectrophotometrically in accordance with the Association of Official Analytical Chemists (AOAC) method [15], which is based on the Folin-Ciocalteu method and used the relevant reagent: gallic acid was

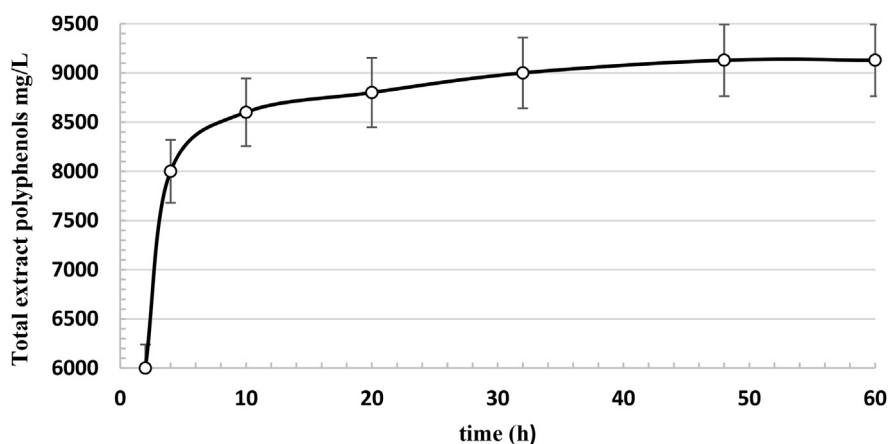


Fig. 2. Graph of the polyphenol extraction kinetics during a cyclically pressurized extraction process using a Naviglio extractor with the Group IV samples.

employed as the reference standard, and the results were expressed in mg of gallic acid equivalents per L of extract ($\text{GAE} \cdot \text{L}^{-1}$) (Fig. 2).

The compound name and MALDI-TOF/TOF product ions (m/z) for each identified polyphenol present in the Group IV extract are presented in Table 2.

3.2. Determination of the characteristic parameters

Table 3 shows the characteristic values of the parameters for the grape piece solid matrix in the Group IV grape samples as well as the diffusivity De calculated by Fick's law. The L_1 length along the x-axis of the considered particle (mm) is the largest principal dimension. The L_2 length along the y-axis of the considered particle (mm) is the second largest principal dimension, while the L_3 length along the z-axis of the considered particle (mm) is the smallest principal dimension.

Moreover, the sphericity value ϕ , which is an index of the roundness of the grain, was also calculated using the method described by Jain and Bal (1997) [16].

For non-spherical particles, the sphericity was calculated as the ratio of the surface area of an equivalent sphere to the surface area of the grain [27]. The results for non-spherical particles are reported in Table 3.

However, we detected a decrease in the width and thickness of the grape matrix, and its sphericity parameter changed from 0.9757 to 0.9770. fe was computed, and it changed from 0.970 to 0.972.

This decrease in dimensions was due to the migration of solute content in the grape peel cell solid matrix, causing overall deflation. Ahromrit et al. (2006) provides a detailed discussion on dimensional changes during the extraction process [28].

By applying the expression of Fick's law using Eqs. (2) and (3), De was precisely calculated for each time period examined. For the polyphenol components, the De values ranged from $0.942 \times 10^{-13} \text{ m}^2/\text{s}$ to $0.124 \times 10^{-10} \text{ m}^2/\text{s}$. These values were compared with those obtained in another study available in the literature, Madieta et al., (2008), in which De values ranging between $D_1 = 7 \times 10^{-9} \text{ m}^2\text{s}^{-1}$ and $D_2 = 5 \times 10^{-10} \text{ m}^2\text{s}^{-1}$ were used [29].

3.3. Chemical content

The results summarized in Table 1 show that the tests carried out using only deionized water with peels and seeds and no pulp yielded the highest polyphenol content at the maximum extraction time of 48 hours. After this time, the additional quantities obtained were negligible. Additionally, if a 48-hour process time is too long, it is possible to obtain an acceptable

Table 2. Compound names and molecular weights of polyphenols identified in the Group IV samples via MALDI-TOF/TOF analysis.

Grape peels dried	
Compound	m/z
Delphinidin	303.26
Petunidin	317.28
Malvidin	331.34
Peonidin-3-O-Monoglucoside	463.12
Delphinidin-3-O-Monoglucoside	465.12
Petunidin-3-O-Monoglucoside	479.13
Malvidin-3-O-Monoglucoside	493.16
A Type Vitisin of Pn-3-Glc/Acetone derivat of Malvidin-3-O-Glucoside	531
Malvidin-3-O-(6-O-Acetyl)Monoglucoside	535.17
Peonidin-3-O(6-O-P-Coumaroyl) Monoglucoside	609.21
Delphinidin-3-O-(6-O-Acetyl) Monoglicoside/Cyanidin -3,5-O Diglucoside	611
Petunidin-3-O-(6-O-Acetyl)Monoglicoside/ Peonidin-3,5-Diglucoside	625.20
Malvidin-3,5-O-Diglucoside	639.22
Malvidin-Petunidin	647.18
Mv-3-O-(6-O-Caffeoyl)Monoglucoside	655.22
Malvidin-Malvidin (Fragment)	661.21
A-Type Vitisin of Pn-3-P-Coumaryl- Glucoside	677.18
Delphinidin-3(6-O-P-Coumaroyl)-5-O- Diglucoside	773.24
Delphinidin-3-O-(6-O-Ferulyl)-5- Odiglucoside	803.26
Malvidin-Peonidin-Glucoside (Fragment)	823.28
Malvidin-Cyanidin-Diglucoside	941.32
Malvidin- <u>Peonidin</u> - Diglucoside	955.33
Malvidin -Delphinidin-Diglucoside	957.34
Malvidin- Malvidin -Diglucoside	985.36
Malvidin- <u>Peonidin</u> - Glucoside-P-Coumaroyl	1101.39
Malvidin -Delphinidin -P Coumaroyl Glucoside	1103.39
Malvidin- Malvidin -P Coumaroyl Glucoside	1131.42

Table 3. Some physical characteristics of the experimental mashed grape particles.

Time (h)	L1 Length (mm)	L2 Width (mm)	L3Thickness (mm)	GMD = D (mm)	Sphericity ϕ	Sphericity factor $-fe$
0	5.06 ± 0.92	4.22 ± 0.62	3.27 ± 0.31	3.71	0.9757	0.970839
2	5.01 ± 0.87	4.18 ± 0.43	3.23 ± 0.23	3.67	0.9754	0.970553
4	4.91 ± 0.78	4.09 ± 0.35	3.18 ± 0.08	3.60	0.9757	0.971199
10	4.86 ± 0.57	4.05 ± 0.63	3.15 ± 0.13	3.57	0.9758	0.971356
20	4.76 ± 0.71	4.01 ± 0.59	3.11 ± 0.05	3.53	0.9771	0.971997
32	4.71 ± 0.73	3.98 ± 0.34	3.08 ± 0.06	3.50	0.9775	0.972029
48	4.67 ± 0.72	3.93 ± 0.29	3.05 ± 0.04	3.46	0.9770	0.972019

Data are expressed as the mean ± SD. GMD: geometric mean diameter.

polyphenol content in the extract with shorter process times, causing a loss of approximately 30% compared to the 48-hour yield, as seen from the results reported in Table 1. Therefore, detecting a high rate of extraction using the experimental process was possible.

Food processing intended for conservation, transformation or extraction occupies an important place in various production processes. Therefore, green food processing represents a global strategy based on the discovery and design of processes to reduce energy and water consumption [30]. In particular, in the extraction sector, there has been a growing demand in recent years for new extraction techniques with reduced extraction times, reduced consumption of organic solvents and greater prevention of pollution, namely, so-called "green" methods. To improve the sensitivity and selectivity of analytical methods, ecological extraction techniques represent the sustainable alternative to the classic procedures of sample preparation used in the past. New extraction methods, such as UAE), microwave-assisted extraction (MAE), SFE and accelerated solvent extraction (ASE), are fast and efficient in the extraction of bioactive substances from different solid matrices [31]. These techniques can work at high temperatures and/or pressures, thereby greatly reducing extraction times. In the field of innovative techniques, a technology called rapid solid-liquid dynamic extraction (RSLDE) has been developed as a viable alternative to maceration. It is a "green" technique; in fact, compared to extraction with techniques such as SFE or ASE, it requires a minimum energy expenditure. It allows aqueous extraction to be performed in reduced times, which is impossible to achieve with maceration. The extraction is carried out using the formation of a negative pressure gradient from the inside to the outside of the solid matrix, so the solid-liquid extraction can be carried out at room temperature. Extracting at low temperatures is

important because the use of low temperature avoids applying thermal stress on thermolabile substances.

On the other hand, the RSLDE, because of its innovative technology [12], allows the extraction of plants and vegetables in a much shorter time than is allowed by current technologies such as maceration, percolation, Soxhlet, microwaves, supercritical fluids, and ASE, avoiding the thermal stress to heat-sensitive substances. As a result, its application fields range from the production of herbal products to the beverage industry, the cosmetics industry, and the food industry, among others [18, 19, 20, 21, 22, 32, 33, 34, 35, 36, 37, 38]. In this study, the cyclically pressurized extraction process was found to significantly increase the extraction rate of the grape pieces, and the equilibrium conditions were thus attained in much shorter times than those required by other methods. The juice thus obtained must subsequently be subjected to a pasteurization process to make it stable and suitable for human consumption. Given the complexity of the food matrix in question, as well as the different variables that contribute to the modification of phenolic composition, the results obtained herein provide

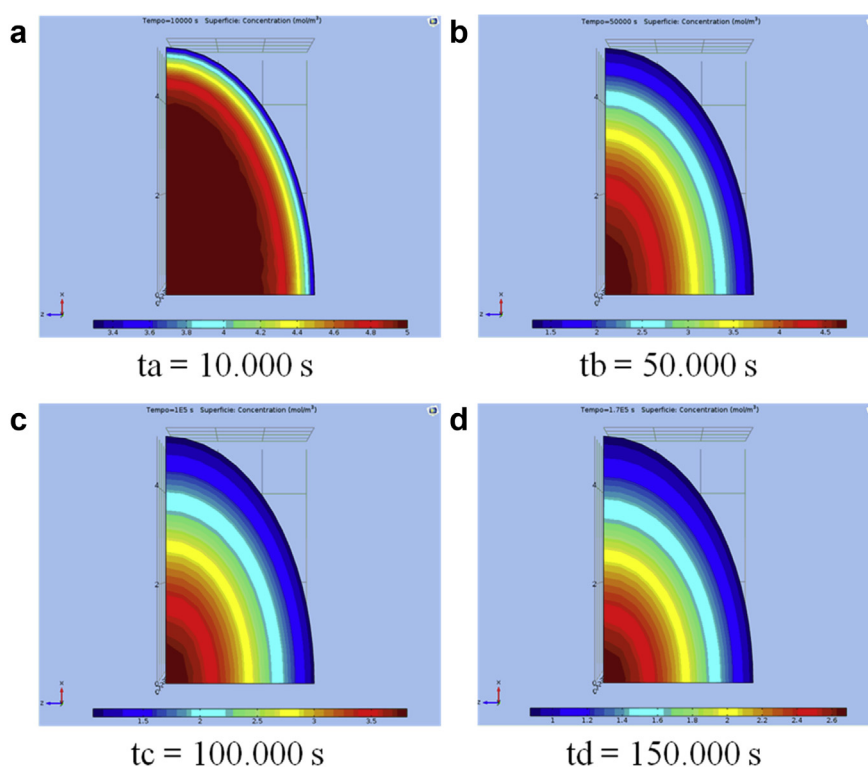


Fig. 3. a, b, c, and d show the typical distribution of the solute concentration (diffusion) in the solid matrix of grape pieces during extraction as determined by a finite element analysis after $t = 10\text{ s}$, $t = 50\text{ s}$, $t = 100\text{ s}$, or $t = 150\text{ s}$, respectively, for the component polyphenols. The bar below each shape represents the content concentration (%) within a particle region at a given time.

the first indications of the production of a grape-based beverage rich in polyphenols. In conclusion, the extraction and stabilization of grape polyphenols through the use of a Naviglio extractor provide an innovative approach for the use of grape pieces to develop new food ingredients. These results are of considerable importance because a diet rich in polyphenols provides significant protection against the development and progression of many chronic pathological conditions, including cancer, diabetes and cardiovascular problems. Furthermore, the intake of polyphenols, owing to their anti-radical action, contributes to the delay of ageing [39, 40, 41, 42, 43, 44].

3.4. Numerical analysis results

To perform a numerical simulation of the extraction process investigated in this study, a finite element model for grapes was used (Fig. 1).

The computed diffusion parameter was replaced in Eq. (1), and the equation was solved using the finite element method to evaluate the nodal flux-time diagrams for the polyphenols and their concentration distribution in the matrix at each instant considered. The data obtained for the predictions of the concentration values during the extraction process are shown in Fig. 3 a, b, c, and d.

Fig. 3 a, b, c, and d shows a gradual migration of the solute extract from the centre of the solid matrix to the surface. Theoretically, the finite element model, as shown in Fig. 3, indicates that even at the end of the extraction process, the concentration of the content throughout the solid matrix is not quite uniform. In Fig. 3 a, b, c, and d, we note that the extract solute gradient inside the matrix has high concentration values during the initial phase of the extraction and that these values decrease during the extraction process. At the end of the process, the concentration distribution is near constant for each node. These figures show the concentration at various points along a cross-section of a grape piece particle, from the surface to the centre of the solid matrix.

4. Conclusion

In recent years, interest has increased in specific compounds that can be obtained from the by-products of the wine industry. This interest, combined with the interest of the wine industry in reducing the impact of their products and reducing the volume and disposal costs of waste, has led to a greater interest in research on more efficient utilization of by-products rich in substances with high added value. In fact, in exhausted by-products of grapes, it is possible to find different bioactive compounds, including polyphenols. It is therefore important to develop methods that make it possible to use these by-products completely on a large scale and economically. The extraction of

chemical compounds with a high biomass added value already represents an important market in the pharmaceutical, cosmetic, nutraceutical and agricultural sectors; thus, extraction processes can be defined as sustainable, and it is important to use energy efficient technologies based on environmentally friendly solvents. Based on the results obtained, RSLDE is an efficient procedure for extracting polyphenols, in addition to demonstrating the exhaustion of solid matrices containing extractable substances in a short time, at low operating temperatures (environmental or sub-environmental) and with reproducibility of extraction. These features ensure the production of quality extracts, and compared to other techniques currently in use, the process has a low environmental impact. In addition, to better understand the cyclically pressurized extraction process, it has been modelled and studied using the finite element method. For added value, the exhausted peels that remain after the extractive process can be dried and used as manure and/or animal feed.

Declarations

Author contribution statement

Monica Gallo, Daniele Naviglio: Conceived and designed the experiments; Performed the experiments; Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data; Wrote the paper.

Rosalba Giacco, Gabriele Riccardi, Delia Lungo: Conceived and designed the experiments; Analyzed and interpreted the data; Wrote the paper.

Andrea Formato, Gaetano Formato: Performed the experiments; Analyzed and interpreted the data.

Angela Amoresano: Analyzed and interpreted the data; Contributed reagents, materials, analysis tools or data.

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Competing interest statement

The authors declare no conflict of interest.

Additional information

No additional information is available for this paper.

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