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Valorization of industrial wastes from French maritime pine bark by solvent free microwave extraction of volatiles

Hamid Mellouk ^a, Alice Meullemiestre ^{b, c}, Zoulikha Maache-Rezzoug ^c, Bouchra Bejjani ^a, Adil Dani ^a, Sid-Ahmed Rezzoug ^{c, *}

^a LGPE, FSTM, University of Hassan II, Mohammedia, Morocco

^b Université d'Avignon et des Pays de Vaucluse, INRA, UMR408, Sécurité et Qualité des Produits d'Origine Végétale, F-84000 Avignon, France

^c LaSIE, UMR CNRS 7356, Université de La Rochelle – Pôles Sciences et Technologie, Bâtiment Marie Curie, Avenue Michel Crépeau, 17042 La Rochelle, France

ABSTRACT

Solvent Free Microwave Extraction (SFME) of oil from French maritime pine bark waste and its antioxidant activity were investigated and compared to classical hydrodistillation (HD) method (Clevenger apparatus). A central composite design combined with response surface methodology was applied to evaluate the simultaneous influences of irradiation power and irradiation time. A maximal extraction yield of 3.48% (g/100 g dry bark) was achieved under optimal extraction time of 92.4 min and an irradiation power of 803.5 W compared 2.2% obtained for the conventional method (HD). Gas chromatography coupled to mass spectrometry (GC–MS) analysis showed that SFME extract is richer in oxygenated compounds (~40%) compared to HD extract with 26%. The results also showed that the two independent variables had a statistical significant effect on the considered responses namely oil yield, total phenolic compounds (TPC) and antioxidant activity assessed by 2,2-diphenyl-1-picrylhdrazyl (DPPH). Under optimized conditions, the amount of TPC was 139.15 mg GAE/g extract largely higher than that obtained for HD extraction (14.28 15 mg GAE/g extract) suggesting that SFME represents an interesting alternative technology for production of recoverable oil from waste of French Maritime pine bark. The percentage inhibition was also higher for SFME extraction technique (71.91%) than for HD (56.51%) indicating a higher antioxidant activity.

1. Introduction

Keywords:

Maritime pine bark

Hydrodistillation

Total phenolic compounds Antioxidant activity

Solvent free microwave extraction

The lignocellulosic waste materials represent an interesting source of chemicals. Their abundant and renewable origin as well as the qualities found in their components converts them in a promising alternative resource. Maritime pine (*Pinus pinaster*) is a conifer native to South-Western Europe and North-Western Africa, with major forests development on Atlantic coast of southern France, Spain and Portugal (Seabra et al., 2012). Extracts isolated from plants such as pines are used as fragrances in cosmetics, flavouring additives of foods and beverages, and scenting agents in a variety of household products including detergents, soaps or insect repellent. They are also used as intermediate in the synthesis of perfume chemicals and for unconventional medicinal purposes as well as in aromatherapy (Maimoonaa et al., 2011). In this field, the traditional synthetic antioxidants food additives such as butylated

* Corresponding author. Tel.: +33 546458615; fax: +33 546458241. *E-mail address:* sarezzou@univ-lr.fr (S.-A. Rezzoug). hydroxyanisole (BHA) and butylated hydroxytoluene (BHT) showed some negative side effects on health (Sarikurkcu et al., 2007) due to their instability and high volatility (Li et al., 2008). Therefore, increasing interest in natural antioxidant present in the diet has developed among consumers and the scientist community an interest to replace synthetic antioxidants (Inan et al., 2012). A lot of research work have been reported about antioxidant properties of different plant extracts (Ozkan and Erdogan, 2011; Zhang et al., 2014), and confirmed that the phenolic components in isolated oils were the main source of antioxidant activity (Karabegović et al., 2014).

The technologies used to extract essential oils from plant materials are enormous and usually have some disadvantages as high temperature, long processing time as in the conventional hydrodistillation or the solvent extraction in which a loss of volatile compounds occurs during solvent removal (Rodríguez-Rojo et al., 2012). For a few years, the change in attitude has evolved increasingly to a "green" tendency and there has been an increasing demand for new cleaner techniques for essential oil extraction. These new green processes will be more environmentally friendly

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with a shorter extraction times, lesser consumption of organic solvent and energy, and minor waste and CO₂ emissions, while maintaining a high quality of extract. Some intensified extraction methods were investigated including microwave (Gholivand et al., 2013; Fernández-Agulló et al., 2015) supercritical fluids (Herzi et al., 2013; Stashenko et al., 2013), D.I.C-assisted extraction (Mellouk et al., 2013; Rezzoug, 2009) or ultrasound assisted extraction (Meullemiestre et al., 2014: González-Centeno et al., 2015: Ghitescu et al., 2015). Currently, application of microwave technology-based methods such as solvent free microwave extraction (SFME) becomes highly desirable as a valid alternative to conventional methods and this extraction technology was the subject of several studies (Chen et al., 2011; Ma et al., 2012; Ranic et al., 2014). To the best of our knowledge, no work has been published on the antioxidant activity of oil extracted from industrial wood wastes by solvent free microwave extraction. In the present work, isolation of waste extracts from French maritime pin bark issued from timber industry was optimized using SFME method. A central composite design (CCD) has been developed to assess the effect of two independent variables namely microwave irradiation power and irradiation time on extraction yield, on the amount of total phenolic compounds and on the percentage inhibition of DPPH radical (2,2diphenyl-1-picrylhydrazyl). The results were compared with those of hydrodistillation (HD) as a conventional extraction method.

2. Materials and methods

2.1. Plant material and chemicals

French maritime pine bark (P. pinaster) provided by Archimbaud Company (Secondigné/Belle, France) in March 2014. The chips were coarsely crushed and dried for 24 h at 25 °C. Moisture content was measured using a halogen Moisture Analyzer (Ohaus - MB 35) at 105 °C and corresponded to 12.3% db (dry basis). After storage in a refrigerated room at 4 °C, fresh material was employed in all experiments after adding water to reach about 50% moisture content (for SFME). 2,2-Diphenyl-1-picrylhydrazyl hydrate (DPPH), anhydrous sodium carbonate, gallic acid, Folin-Ciocalteu's phenol reagent were purchased from Sigma-Aldrich and Methanol, Na₂CO₃ were from Fisher scientific.

2.2. Protocol

In the present study, the experimental design was achieved as illustrated in Fig. 1. Extraction of volatile molecules was performed by HD and SFME methods. Each HD operation was performed three times in order to test reproducible. SFME treatments were analysed and optimised through statistical study. For HD and for SFME in optimised conditions, the antioxidant activity and total phenolic compounds were evaluated.

2.3. SFME apparatus and procedure

Solvent-free microwave extraction has been performed on a Milestone NEOS microwave station (NEOS microwave laboratory oven) (Fig. 2). It is a multimode microwave reactor 2.45 GHz with a maximum delivered power of 1000 W variable in 10 W increments. Temperature was monitored by an external infrared sensor. In a typical procedure 100 g of moistened bark chips were subjected to microwave irradiations in oven cavity, initially at ambient temperature, during a fixed processing time. The microwave heating of the water contained inside the raw material allows releasing molecules constituting isolated oil. This oil was then driven by the generated vapour. A cooling system outside the microwave cavity permitted to condensate the distillate



Fig. 1. Protocol of extraction and analysis of oil from maritime pine bark waste.



Fig. 2. Solvent free microwave extraction apparatus (SFME): (1) refrigerant system; (2) aqueous fraction; (3) oil fraction; (4) maritime pine bark; (5) microwave oven.

continuously (5 °C). Condensed water was refluxed to the extraction vessel in order to provide uniform conditions of temperature and humidity. Isolated oil was dried with anhydrous sodium sulphate and stored at 4 °C in the dark until used. Extraction yield was calculated according to eq. 1

Extraction yield (%) =
$$\left(\frac{\text{mass of extracted essential oil}}{\text{mass of dry material}}\right) \times 100$$
(1)

2.4. Hydrodistillation apparatus and procedure

Conventional hydrodistillation apparatus (Clevenger-type apparatus) according to the European Pharmacopeia (2012) was employed. A quantity of 100 g of maritime pine bark for 1 L of

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distilled water was used to perform the hydrodistillation during 300 min from the first drop of distillate fell until the raw material has been completely exhausted. Isolated oil was collected, dried under anhydrous sodium sulphate, and stored at 4 °C for further analysis. Each extraction was performed at least three times, and a standard deviation was calculated. The extraction yield was calculated according to eq. (1).

2.5. Experimental design

The relationships between response functions and process variables have been established by using a central composite design (CCD) as well as the optimal conditions of the developed process. The independent variables were coded according to eq. (2):

$$\dot{x}_i = \frac{X_i - X_{i0}}{\Delta X_i} \quad i = 1, 2$$
⁽²⁾

where x_i and X_i are respectively the dimensionless and the actual values of the independent variable i, X_{i0} is the actual value of the independent variable *i* at the central point, and ΔX_i is the step change of X_i corresponding to a unit variation of the dimensionless value. Irradiation power (x_1) and processing time (x_2) were chosen as independent variables. The selected responses were the total yield of maritime pine bark oil (%), the total phenolic compounds (mg GAE/g extract) and the antioxidant activity expressed by the percentage inhibition of DPPH radical. For the two variables, the design yielded 14 experiments with four (2^2) factorial points, four axial points ($-\alpha$ and $+\alpha$) to form a central composite design and six center points for replications and estimation of the experimental error and to prove the suitability of the model. Both coded and actual values of the independent variables and their ranges of variations are listed in Table 1.

$$Y = \beta_0 + \sum_{i=1}^n \beta_i x_i + \sum_{i=1}^n \beta_{ii} x_i^2 + \sum_{i \neq j} \beta_{ij} x_i x_j$$
(3)

The responses Y are related to the coded independent variables x_i and x_j according to the second order polynomial expressed in eq. (3), with β_0 the interception coefficient, β_i the linear terms, β_{ii} the quadratic terms, β_{ij} the interaction terms. Fisher test for analysis of variance (ANOVA) performed on experimental data permitted to estimate the statistical significance of the proposed models. Response surfaces as represented by Fig. 3 were drawn by using the *analysis design* procedure of *Statgraphics Plus for Windows* software (Centurion version).

2.6. GC-MS identification

The volatile compounds were analyzed by gas chromatography coupled to mass spectrometry (GC–MS). Analyses were performed on a GC/MS Varian 3900 chromatograph coupled to a Saturn 2100T mass spectrometer using fused-silica-capillary column. The nonpolar column was Elite 5MS (30 m \times 0.25 mm \times 0.25 µm film

Table 1 Coded levels for independent variables used in developing experimental data.

	Coded 1	evel			
	$-\alpha$	-1	0	1	$+\alpha$
Irradiation power (W)	96.4	200	450	700	803.6
Processing time (min)	7.6	20	50	80	92.4

 α (axial distance) = $\sqrt[4]{N}$, N is the number of experiments of orthogonal design, i.e. of the factorial design. In this case $\alpha = 1.41$. The two variables were studied in five levels.



Fig. 3. Responses surfaces showing the simultaneous effect of irradiation power and irradiation time on the considered responses.

thickness). GC–MS spectra were obtained using the following conditions: He (helium) as carrier gas at flow rate of 1 mL/min; split mode1:20; 1 μ L as injected volume; 250 °C as injection temperature. The oven temperature programme was 60 °C for 5 min increasing at 2 °C/min towards 250 °C and held at 250 °C during 10 min. The ionization mode used was electronic impact at 70 eV. Most constituents were identified by comparison of their GC linear retention indices (RI), determined with reference to a homologous series of C5–C32 n-alkanes. The identification was confirmed by comparison of the mass spectral with those stored in the MS database (National Institute of Standards and Technology NIST08 and Wiley libraries) and also by comparison with mass spectra from literature data (Adams, 2007). The percentage composition was calculated from the summation of peak areas of the total oil.

2.7. Assay for total phenolics

Total phenols in maritime pine bark extracts was determined using spectrophotometric Folin—Ciocalteu method according to the

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literature methods (Orhan et al., 2013; Kim et al., 2010) with some modifications, using gallic acid as standard. Extract solution (0.5 mL) of diluted samples were added into test tubes followed by 2.5 mL of Folin–Ciocalteu reagent (20%, v/v). After 5 min, 2 mL of a solution of 7% Na₂CO₃ was added. All test tubes with the mixture were caped and shaken for 10 s and put on to incubation in a water bath at 45 °C for 5 min. Absorbance was measured after 30 min at 765 nm (Helios Omega UV/VIS Thermo Scientific Merck and Co. Spectrophotometer) against blank sample. The same procedure was repeated for all standard gallic acid solutions (2–200 µg/mL) and a standard curve was obtained with Eq. (4) (R² = 0.99):

Absorbance =
$$0.0109 \times \text{Gallic acid}(\mu g/mL) + 0.037$$
 (4)

The results were expressed as milligrams of Gallic acid equivalents (GAE) per g extract. The analyses were performed in triplicate and standard deviation was calculated.

2.8. DPPH assay for evaluation of antioxidant activity

The chemical compound 2,2-diphenyl-1-picrylhydrazyl (DPPH) is one of the first free radicals used for studying the structure—activity relationship antioxidant phenolic compounds (Brand-Williams et al., 1995). In this method, the antioxidant activity of oil extract is evaluated in term of the capacity to scavenging free radicals of DPPH formed, according to a method described by Lue et al. (2010). A solution of 4 mg of the radical DPPH dissolved in 100 mL of methanol was prepared. Then 3 mL of this solution was reacted with 1 mL of oil diluted bark extract (dissolved in methanol). The mixture was incubated in dark for 30 min at room temperature. The absorbance was measured at 517 nm with Helios Omega UV/VIS Thermo Scientific Merck and Co. Spectrophotometer. The percentage inhibition activity was calculated by eq. (5):

$$I\% = \left(\frac{A_0 - A_t}{A_0}\right) \times 100 \tag{5}$$

where A_0 is the absorbance of the control sample (without isolated oil) and A_t the absorbance of the extract with DPPH at 30 min. All analyses were run in triplicates and averaged.

2.9. Scanning electron microscopy

Microstructures were observed using an environmental SEM FEI/Philips Quanta 200 FEG (Field Effect Gun). Samples were observed under 1.00 mbar of water vapour pressure without any metallic coverage. The accelerating voltage used is 20 kV and the secondary electrons (SE) images were obtained with a Large-Field Detector (LFD).

3. Results and discussion

3.1. Regression coefficients and fitting the models

The complete design matrix together with the values of experimental yield, the total phenolic compounds and percentage inhibition are given in Table 2. A regression analysis was carried out to fit mathematical models to the experimental data aiming at an optimal region for the studied responses. The predicted models can be described by Table 3 in term of coded values. The significance of each coefficient was determined using Fisher test (*F-value*) and the probability p (*p-value*) in Table 4, which displays the variance analysis of the system (ANOVA). Corresponding variables would be more significant if absolute *F-value* becomes greater and *p-value*

Table 2

Experimental data and obtained responses with the different combinations of irradiation power (x_1) and irradiation time (x_2) used in the randomized central composite design.

Run	Coded variable level		Experimen	Experimental responses data		
	<i>x</i> ₁	<i>x</i> ₂	Yield ^a	TPC ^b	Inhibition ^c	
1	-1	-1	1.02	8.35	31.82	
2	1	-1	0.92	23.45	46.29	
3	$^{-1}$	1	1.72	20.31	40.75	
4	1	1	2.65	90.77	59.99	
5	$-\alpha$	0	1.23	1.19	27.34	
6	$+ \alpha$	0	2.67	60.20	64.31	
7	0	$-\alpha$	0.68	9.95	33.95	
8	0	$+ \alpha$	2.41	91.87	73.37	
9	0	0	1.73	69.29	69.79	
10	0	0	1.66	51.81	55.96	
11	0	0	1.47	46.52	47.90	
12	0	0	1.79	42.37	43.62	
13	0	0	1.88	50.52	55.19	
14	0	0	1.86	49.61	51.36	
Mean abso replicat	olute error ions	for	0.13	9.2	8.9	

^a g/100 g (d.m) or %.

^b mg GAE/g extract.

° (%).

Table 3	
Regression	coefficients of the second-order polynomial equations

Regression coefficients	Response variables			
	Yield (%)	TPC (mg GAE/g extract	Inhibition (%)	
βο	1.71	61.76	50.33	
β1	0.73	55.45	21.50	
β ₂	1.22	48.78	19.59	
β ₁₂	0.51	27.67	2.38	
β11	0.16	-19.60	-7.02	
β22	-0.25	-18.06	0.81	

The fitted models are given by the following eq. $Y = \beta_0 + \beta_1 x_1 + \beta_2 x_2 + \beta_{12} x_1 x_2 + \beta_{22} x_2^2 + \beta_{11} x_1^2$. Where x_1 and x_2 are the coded values of respectively irradiation power and irradiation time.

becomes smaller. For the yield of maritime pine bark isolated oil, it can be seen that the linear terms are strongly significant (p < 0.05). The interaction between irradiation power and processing time was also significant with a confidence level of approximately 97%. For total phenolic compounds, unlike the yield of extracted oil, all effects were found statistically significant with predominance for the linear effects of irradiation power and processing time. The significance of the two quadratic effects suggests a degradation of the phenolic compounds.

According to Abdelhadi et al. (2014), this degradation concerns the oxygenated compounds of the oil during a prolonged processing time or for high microwave intensity. The interaction between irradiation power and processing time was also statistically significant indicating an antagonistic effect of the two factors. Concerning the inhibition of DPPH radical, the linear effects of irradiation power and processing time were significant. In general way, the results suggest that changing in irradiation power and irradiation time had a highly significant effect on the yield of isolated oil, on its composition in phenolic compounds as well as on its capacity to scavenging DPPH radical. The quality of the models developed was evaluated based on the correlation coefficient R² and on the lack-of fit value. From ANOVA (Table 4), it can be seen that R² were systematically close to 90% and *p-value* of lack-of-fit systematically higher than 0.05 (non-significant) suggesting that

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Table 4

Analysis of variance showing the effect of the three independent variables as linear, quadratic and cross products terms on the studied responses.

	Source	DF	Sum of square	F-ratio	P-value
Yield (%)	<i>x</i> ₁	1	1.06	45.26	0.0025
	<i>x</i> ₂	1	3.00	127.99	0.0003
	$x_1 x_2$	1	0.26	11.26	0.0284
	x_{1}^{2}	1	0.03	1.78	0.2526
	x_{2}^{2}	1	0.14	4.50	0.1011
	Lack-of-fit	3	0.22	3.18	0.1462
	Pure error	4	0.094		
	R ²	0.93			
TPC	<i>x</i> ₁	1	6151.08	121.95	0.0004
	<i>x</i> ₂	1	4759.21	94.36	0.0006
	$x_1 x_2$	1	766.016	15.19	0.0176
	x_{1}^{2}	1	668.07	13.25	0.0220
	x_{2}^{2}	1	567.31	11.25	0.0285
	Lack-of-fit	3	904.48	5.98	0.0584
	Pure error	4	201.76		
	R ²	0.92			
Inhibition	<i>x</i> ₁	1	924.78	34.23	0.0043
	<i>x</i> ₂	1	768.08	28.43	0.0060
	$x_1 x_2$	1	5.69	0.21	0.6701
	x_{1}^{2}	1	85.82	3.18	0.1493
	x_{2}^{2}	1	1.14	0.04	0.8473
	Lack-of-fit	3	230.85	2.85	0.1690
	Pure error	4	108.07		
	R ²	0.84			

the predicted models reasonably represent the observed values. Thus the responses were sufficiently explained by the models.

3.2. Response surfaces analysis

Three-dimensional profiles of multiple non-linear regression models (Fig. 3) were used to illustrate the linear and quadratic effects of the two key variables as well as the interaction effects between irradiation power and irradiation time on extraction yield, total phenolic compounds and on the percentage inhibition obtained from DPPH test. In each plot, the third variable is fixed at its central value ("0"). The most influential effects on the yield of isolated oil are the linear terms of irradiation power (x_1) and irradiation time (x_2) . Fig. 3a clearly shows that the yield increased linearly as irradiation power and irradiation time increased. This increasing was also pointed out by Dai and Mumper (2010) and Al-Farsi and Lee (2008) for which a high temperature favoured the extraction process by enhancing the solubility of phenolic compounds and the mass transfer rate (Dai and Mumper, 2010) and by weakening the plant tissues. The interaction between irradiation power and processing time is also visible on Fig. 3a; when irradiation power increased from its low level (200 W) to its high level (700 W), a weak variation in the yield was observed, from 0.94% to 1.15%, for a processing time fixed at 20 min. Conversely, a strong evolution was observed, from 1.64% to 2.89% for a processing time fixed at 40 min. For the total phenolic compounds, the quadratic effect of irradiation power and in a lesser extent that associated to the processing time are visible on Fig. 3b on which a maximum of total phenolic compounds seems to be reached for the highest values of the two processing variables. On this figure, it can be also noticed a strong interaction; when irradiation power increase from 200 W to 700 W, TPC increased from 4.11 to 31.40 mg GAE eq/g extract with a maximum of 33.6 mg GAE eq/g extract for a processing time fixed at 20 min while for a same variation, the phenolic compounds increased continuously from 25.31 to 108.43 mg GAE eq/g extract for a processing time fixed at 40 min. Concerning the radical-scavenging properties, the linear effects of irradiation

power and irradiation time are clearly visible as testified by a flat growing surface (Fig. 3c).

3.3. SFME versus HD

3.3.1. Yield, total phenolics and antioxidant activity

For isolated oil yield the optimal conditions selected by the software were the highest values of irradiation power (803.6 W) and irradiation time (92.4 min). Under these conditions, the software indicates an optimum yield of 3.51% while the experimental yield was about 3.34%. This difference can be due to a loss of oil on the flask test walls. From kinetic point of view (Fig. 4), HD extraction results were compared with the results obtained for run 7, 8 and means of the four repetitions (runs from 10 to 13). The maximum HD extraction of 2.2% was reached after about 180 min while for SFME, a higher yield of 2.41% was obtained in a lesser time (92.4 min) for an intermediate irradiation power (450 W). It should be noted that in the optimal conditions of SFME extraction (803 W; 92.4 min) the yield was largely enhanced (by about 53%) compared to hydrodistillation method. In both HD and SFME extraction processes, the kinetics was reported to take place in two distinct phases (Pavićević et al., 2015). The first one is represented by an increasing line until the temperature reaches 100 °C and thus achieves the distillation of the first oil droplet. In the second part, the extraction temperature is approximately equal to water boiling temperature at atmospheric pressure. However, the most important differences were observed between the both extraction methods is the ability of SFME process to quickly raise the temperature of the sample to 100 °C. This rapid increasing of extraction temperature gives to acceleration of extraction rates under microwaves and could be due to a synergetic combination of heat and mass transfer phenomena. For HD, mass transfer occurs from inside to outside whilst heat transfer occurs from outside to inside. For microwave extraction, the two transport phenomena act in the same direction, from inside to outside, which facilitates oil diffusion (Farhat et al., 2011; Desai and Parikh, 2015). Fig. 4 seems to confirm this argumentation although the second phase is not really reached for SFME. The Folin-Ciocalteu method allowed a good discrimination between the SFME and HD extracts (Table 5). The results clearly shown that the application of microwaves leads to recover phenolic compounds with a higher concentration in comparison with methods based on hydrodistillation, respectively 139.15 and 14.28 mg GAE/g extract for SFME and HD. Thus SFME appears to be is an efficient technology to save the structure of phenolic groups



Fig. 4. Comparison of extraction kinetics of HD (\diamondsuit) and SFME (\square). (a): run 7, (b): means of runs 10–13 and (c) run 8.

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Table 5

Comparison of extraction yield, TPC amount and the percentage of DPPH inhibition for SFME (92.4 min, 803.5 W) and HD (180 min).

	Yield (g/100 g d.b)	TPC (mg GAE/g extract)	Inhibition (%)
SFME HD BHT	3.34 2.20	139.15 14.28	71.9 56.5 93.2

and avoid their contribution in side reactions. On the other hand, the two extract exhibited concentration-dependant DPPH radical scavenging activity. For a fixed value of extract concentration indicated in section 2.8, the reducing capacity of SFME isolated extract quantified as a percentage inhibition was higher than that of HD, respectively 71.9% and 56.5%. The findings above show the presence of natural antioxidant phenolic compounds in pine bark waste with a good performance since the SFME antioxidant activity can be favourably compared with that of BHT (butylated hydroxytoluene) (93.2%) known as a very efficient synthetic antioxidant activity of SFME extracts may be attributed to the presence of a higher quantity of oxygenated monoterpenes and in lesser extent monoterpene hydrocarbons> (Park and Lee, 2011).

3.3.2. Chemical composition

For both HD and SFME isolated oils, about 55 components were respectively identified among which monoterpene and sesquiterpene hydrocarbons as well as oxygenated monoterpenes and sesguiterpenes. From Table 6, it can be seen that isolated oils obtained by SFME and HD have equivalent percentages in non-oxygenated compounds, respectively 37.6% and 32.1%. Conversely for SFME the predominant group is that of oxygenated compounds (40.8%) compared to HD (26.8%). Orio et al. (2012) have linked this phenomenon to microwave energy effect which differs for oxygenated and non-oxygenated groups of compounds. For the more polar compounds (oxygenated compounds), the more easily the microwaves irradiations are absorbed, the better the interaction between electromagnetic wave and matter is established and more polar aromatic components are obtained, conversely to sesquiterpenes and monoterpenes hydrocarbons. Electromagnetic interactions are also cited as possible cause to presence of more oxygenated compounds in SFME extracts: organic compounds that have a high dipolar moment as many oxygenated compounds interact more vigorously with microwaves and can be extracted more easily. It should be noted that the oxygenated compounds are most valuable in plant isolated oils; they are characteristic of a strong odour and are known to play a key role in the antioxidant activity. Among the oxygenated compounds, α -terpineol seems to be predominant in SFME extracts and probably have also a part in antioxidant activity as it was remarked by Li et al. (2012).

3.4. Microstructural changes after extraction

Pine bark samples were examined by SEM for an evaluation of the microstructural alteration induced by SFME and HD extractions compared to untreated sample. As shown in Fig. 5A, the external surface of untreated sample was smooth and contains many folds. After extraction by HD (Fig. 5B) some ruptures and perforations appeared on the leaf surface and the folds were still present while after extraction by SFME (Fig. 5C), perforations and creation of canals was clearly observed and the surface appeared completely disrupted indicating that the strain induced by a rapid rise in temperature in SFME extraction and subsequent change in the surface tension of the glandular wall, causing it to crumble or rupture more readily.

Table	6
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Chemical composition of maritime pine sawdust waste oils obtained by GC-MS.

α-Pinene 11.51 2.35 2. 3-Carene 11.52 2.23 1. α-Thujene 11.57 0.33 0. Cis-ocymene 11.58 0.38 0. Trans-ocymene 11.6 0.57 0. Benzylalcohol 11 84 1.34 1	2.32 1.87 1.21 1.12 1.45 1.25 1.87 1.26 1.12
3-Carene 11.52 2.23 1. α-Thujene 11.57 0.33 0. Cis-ocymene 11.58 0.38 0. Trans-ocymene 11.6 0.57 0. Benzvlalcohol 11 84 1 34 1	1.87).21).12).45].25).87].26].12
α-Thujene 11.57 0.33 0. Cis-ocymene 11.58 0.38 0. Trans-ocymene 11.6 0.57 0. Benzylajcohol 11.84 1.34 1).21).12).45 .25).87 .26).12
Cis-ocymene 11.58 0.38 0. Trans-ocymene 11.6 0.57 0. Benzylalcohol 11.84 1.34 1).12).45].25).87].26].12
Trans-ocymene 11.6 0.57 0. Benzylalcohol 11.84 1.34 1).45 .25).87 .26).12
Benzvlalcohol 1184 134 1	.25).87 .26).12
).87 26).12
Tricyclo[2.2.1.0(2,6)]heptan 11.88 0.93 0.	.26).12
Tricyclo[3.2.1.0(2,4)]octane 11.92 0.95 1.).12
Bicyclo[3.1.1]hept-2-ene-2-e 11.98 0.42 0.	
1,3,6-Octatriene, 3,7-dimeth 12.04 1.09 0.).85
Cyclofenchene 12.1 4.36 3.	5.76
Camphene 12.92 0.31 0.).49
β-Pinene 13.97 5.46 5.	1.17
β-Cymene 16.85 0.37 0.	1.26
Limonene 16.97 2.13 1.	.93
Artemiseole 19.72 0.2 0.	1.18
(+)-4-Carelle 20.06 0.47 0.).4Z
rephiloiene 20.11 0.23 0.	1.13
α -4-Dimethylstyrelle 20.53 0.99 0. Exa Fonchol 22.24 1.02 0.	1.03
EXd-FEIICIIOI 22.24 1.05 0.	1.91
22.05 0.25 0.25 0.25 0.25	1.14
$\begin{array}{ccc} 0 - \text{Calliplicitor} & 22.77 & 0.26 & 0. \\ 0 + \text{trans ninesaryosl} & 22.41 & 1.07 & 0. \end{array}$	1.15
23.41 1.07 0.1	1.07
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	112
Cic_{β} termineol $24.11 + 0.52 = 0.$	1.15
$Trans_\beta$ -terpineol 24.25 0.55 0.75	126
Pinocaryone $24.62 0.49 0.1$	172
Perillaldehyde 24.81 0.37 0	162
Isoborneol 25.18 1.01 1	27
Borneol 25.10 1.01 1.	46
4-Terpineol 25.67 2.19 0)71
a-Terpineol 26.36 19.54 8	35
Valencene 36.85 2.7 2.	2.76
β-Carvophyllene 37.40 7.63 4.	1.92
α-Humulene 39.13 3.34 2.	2.19
α-Muurolene 41.32 0.16 0.).17
δ-Cadinene 42.20 0.61 1.	.06
α-Cadinene 42.38 0.24 0.).37
Guaiacol 43.79 0.54 0.).44
Eudesmol 44.08 0.99 0.).84
(+)-Longicamphenylone 44.56 0.16 0.).19
Caryophyllenyl alcohol 44.88 1.19 1.	.09
Caryophyllene oxide 45.15 1.42 1.	.51
Gamma-gurjunenepoxide-(2) 45.78 0.37 0.).18
Longiborneol 46.11 0.82 0.).63
3,7-Cycloundecadien-1-ol, 1,5,5,8-tetramethyl 46.31 0.47 0.47).42
Cubenol 47.24 0.81 0.).75
Calarene epoxide 47.59 0.14 0.).18
Isoaromadendrene epoxide 47.6 0.49 0.49).43
Aromadendrene oxide-(2) 49.29 0.36 0.).24
Biformene 58.81 0.68 1.	.34
α-Himachalene 60.32 0.2 0.).5
Verticiol 60.93 0.33 0.	1.77
Sclarene 61.16 0.36 0.).77
Manoyl oxide 61.88 1.28 1.	.95
Total oxygenated compounds 40.05 26	5.83
Monoterpenes hydrocarbons 21.27 18.	3.61
Sesquiterpenes hydrocarbons 16.34 14.	1.2

Rt: Retention time on varian Elite-5MS column.

%: Relative area percentage (peak area relative to the total peak).

3.5. Energy consumption and environmental impact

The reduced cost of oil extraction is clearly advantageous for SFME method in terms of energy and time saving. HD method required 180 min to reach a maximum yield of 2.2 mg/g d.b while 92 min were sufficient to reach a maximum yield 3.4 mg/g d.b for SFME. The amounts of water to evaporate were 1 kg for HD and about 50 g for SFME. Then, the energy required for performing HD

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Fig. 5. Scanning Electron microscopy for maritime pine bark waste. (A) Untreated, (B) after HD extraction and (C) after SFME extraction.

and SFME extraction methods were respectively 3,5 kWh and 1,2 kWh (in optimised conditions). Regarding the environmental impact, the calculated quantity of carbon dioxide rejected in the atmosphere for HD was from 2619 g to 3500 g CO₂, largely higher

than that of SFME (from 898 g to 1200 g CO₂). These calculations were preformed according to literature provided by the French Nuclear Energy Society (http://www.sfen.org/fr/lenergie-nucleaire/ nucleaire-et-environnement; downloaded on June 25, 2015): to obtain 1 kWh from coal and fuel, respectively 1000 g and 748 g of CO_2 is rejected in atmosphere during combustion. The GEEN extraction team of Avignon University (Filly et al., 2014; Petigny et al., 2014) recently tested the SFME technique at a pilot plant scale of 150 L capacity, for extraction of essential oil from various biomasses. The authors indicate that microwaves have wide-ranging large scale commercial applications as processing technology and can provide high returns on capital investment.

4. Conclusion

The sustainability of the wood processing industry can be improved through the use of by-products, by adoption of new technologies that maximize process profitability. This study focused on the optimization of SFME extraction process for isolation of oil from wastes of maritime pine wood. The maximum extraction yield was obtained for 803 W as irradiation power for 92 min as processing time was 3.34%, higher for that obtained with a conventional method (HD) for which the maximum extraction yield was 2.2% obtained for 3 h processing time. Moreover, the oil extracted under microwave irradiation was found to possess an improved antioxidant activity combined with a high quantity of phenolic compounds compared to HD method. Thus we can conclude that SFME is a green extraction method that offers important advantages as short extraction time, low energy input and no requirement of solvent since a comparison of SFME and hydrodistillation showed that it was possible to achieve a higher yield with a reduced extraction time and water volume. As a consequence, SFME exhibited a higher energy efficiency and a reduced environmental impact and thus may prove to be a better alternative to conventional methods. Further investigations are required on large scale SFME processing. Moreover it would be interesting to complete this work by a study on the mass transfer occurring during microwave extraction process as well as by testing the antimicrobial activity of isolated oils.

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