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Furanolysis with menthofuran: A new depolymerization method for analyzing condensed tannins

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ABSTRACT

An improved analytical depolymerization method for characterizing condensed tannins was developed with menthofuran (3,6-dimethyl-4,5,6,7-tetrahydro-1-benzofuran) as the nucleophilic trapping reagent. Herein, menthofuran was compared with routinely used nucleophiles, phloroglucinol and 2-mercaptoethanol. At 30°C and in the presence of 0.1 M HCl, menthofuran displayed the outstanding ability to enable the fast and full depolymerization of procyanidin B2 using only a 1:1 molar ratio of both reactants. In the same conditions, phloroglucinol and 2-mercaptoethanol led to a reaction equilibrium with significantly lower conversion yields. Application to commercial tannin extracts showed that a menthofuran to extract weight ratio of 1 gave the same yields of procyanidin constitutive units as 10-fold higher mol. eq. phloroglucinol and 100-fold 2-mercaptoethanol. Finally, guidelines for implementing the menthofuran depolymerization method are proposed to assess the tannin content and composition of extracts as well as of plant materials without prior extraction.

KEYWORDS

Menthofuran, Furan derivatives, Condensed tannins, Depolymerization, Analytical method, Furylated flavonoids, UHPLC-DAD-MS
INTRODUCTION

Condensed tannins (proanthocyanidins) are polymers of flavan-3-ol units. The constitutive units are covalently linked by interflavan bonds between the phloroglucinol ring (C8 and/or C6 carbon atoms) and the benzylic C4 carbon atom of the extension units, thus resulting in B-type proanthocyanidins (Scheme 1 for carbon numbering). Additional linkages resulting from oxidation processes can lead to A-type proanthocyanidins, or biaryl and biaryl ether linked compounds. Condensed tannins, along with hydrolyzable tannins and phlorotannins, constitute the polyphenolic secondary metabolite class of tannins. These plant polyphenols represent the fourth most abundant organic polymer in the terrestrial biomass and the second one, after lignin, when considering only aromatic polymers.

The need to characterize and quantify the condensed tannin fraction from plant or food samples generally meets three objectives. First, characterization of condensed tannins can be used in taxonomy or structure-function relationship studies, including response to environmental stresses. The second objective is to qualify plant extracts or tannin-rich formulations eventually intended for commercial products, such as cosmetics and dietary supplements, in relation to health benefits including antioxidant activities. The third motivation is to develop specialty chemicals or polymer materials from renewable phenolics accordingly to the properties and specificities of the different types of tannins.

Non-degradative and degradative methods have been developed to characterize the condensed tannin fraction of plant materials. Among the former, colorimetric assays based on redox reactions are not specific to polyphenols and tannins and should be interpreted with caution. Methods based on $^1$H NMR and $^{13}$C NMR, 2D $^1$H–$^{13}$C HSQC NMR spectroscopy, as well as on mass spectrometry with electrospray ionization and MALDI-TOF systems, have been developed with the advances of technologies. These methods provide good
information on the nature of constitutive units, types of linkages and degrees of polymerization of tannin structures. The limitations of these methods mainly result from the dispersity of tannin polymers associated with a possible discrimination against highly polymerized structures. On the other hand, degradative methods are based on a chemical depolymerization reaction leading to the release of two kinds of constitutive units: the terminal units, with free C4 carbon atom, and the extension units, where the C4 position is linked to the next unit of the polymeric chains (Scheme 1). The depolymerization products can then be analyzed by chromatography to infer characteristics of the initial tannin structures, including the type and amounts of constitutive units, mean degree of polymerization and galloylation degree. Depolymerization-based methods are currently the most informative methods to characterize condensed tannins. They have recently been shown to give results consistent with NMR and MALDI analyses. Recent developments based on proanthocyanidin in-source fragmentation and mass spectrometry analysis also gave similar results in terms of composition and mean degree of polymerization of oligomeric and polymeric tannin fractions as chemical depolymerization. However, some information remains inaccessible to depolymerization-based methods, such as the molecular weight distribution of the tannin fractions or the sequences of constitutive units in the polymeric chains beyond hexamers, owing to their too low content in the samples. Moreover, some linkages are much more stable with respect to cleavage, such as interflavan linkages of 5-deoxy condensed tannins found in quebracho and acacia, while others are totally resistant, like the A-type patterns and biaryl or biaryl ether linkages resulting from oxidation. Since the first report by Betts et al., different depolymerization-based methods have been developed, compared, and applied to characterize the condensed tannin fraction in
plant extracts or food samples.\textsuperscript{21,33,34} Updates and improvements of these methods are still regularly published.\textsuperscript{35–37} The first nucleophiles used to trap the extension units released by the acid-catalyzed depolymerization of condensed tannins were mercaptans, such as thioglycolic acid and benzylmercaptan.\textsuperscript{22,28,34,38} Later works proposed to substitute mercaptans by analogs of the catechin A-ring (e.g., phloroglucinol, 2,4,6-trihydroxytoluene or resorcinol).\textsuperscript{30,39} Ever since, these methods have evolved with analytical techniques and useful optimizations were performed on reaction conditions, solvents and work-up, but no significant breakthrough was achieved regarding the reactants. Indeed, the typical smell of mercaptans has often been an obstacle to their use in the analysis of condensed tannins.\textsuperscript{30} The toxicity of the chemicals involved in the depolymerization methods is more generally questioned, owing to the fact that the trapping nucleophiles are used in large excess. This especially concerns mercaptans frequently used in the analysis of proanthocyanidins,\textsuperscript{40,41} while phloroglucinol seems to require higher doses to cause adverse effects.\textsuperscript{42} Recently, the possibility to use metalloles (five-membered heterocyclic aromatic compounds) in the depolymerization of condensed tannins to produce biobased chiral ligands or fully biobased aromatic building blocks for applications in specialty chemicals and materials was evidenced by Fu et al.\textsuperscript{43} with pyrrole derivatives and by Rouméas et al.\textsuperscript{44,45} with furan derivatives. In the framework of our studies on the depolymerization of condensed tannins in the presence of substituted metalloles, preliminary experiments have led us to identify menthofuran (3,6-dimethyl-4,5,6,7-tetrahydro-1-benzofuran), a tri-substituted furan, as both an efficient and commercially available nucleophilic trapping reagent. Menthofuran, a major component of essential oils such as pennyroyal oil, has been the subject of numerous toxicological studies.\textsuperscript{46,47} It is used as a flavoring agent (strong peppermint odor) in the food industry at a concentration up to 1000 ppm (i.e., in the same order of magnitude as in the
method described herein). This led us to evaluate it as a potential new reagent for the analytic depolymerization of condensed tannins. In the present work, menthofuran was compared to phloroglucinol and 2-mercaptoethanol, that are routinely used in standard depolymerization methods.

MATERIAL AND METHODS

Chemicals

The grape seed extract was purchased from Union des Distilleries de la Méditerranée (UDM, France). Pycnogenol, a commercial tannin bark extract from maritime pine (Pinus pinaster Aiton subsp. *atlantica* syn. *P. maritima*), was kindly offered by Horphag Research (Geneva, Switzerland). Samples of grape pericarp powder (*Vitis vinifera* L. subsp. *sativa* (DC.) Hegi, cultivar Savagnin), prepared as previously published were kindly supplied by Dr. Charles Romieu. Bark from Douglas fir (*Pseudotsuga menziesii*), kindly provided by Brassac Industrie sawmill (Brassac, France), was obtained from trees felled in March 2015 on a plot located at 43° 33’ 39.5” N, 2° 42’ 14.0” E (altitude 936 m) and debarked in April 2015 under batch reference “chantier Caraman n°UG 47109”. The bark was ground by knife milling using a Retsch SM 100 system operating at room temperature at a speed of 1500 rpm with a 2 mm size screen. Menthofuran (3,6-dimethyl-4,5,6,7-tetrahydro-1-benzofuran, ≥95%) and 2-mercaptoethanol (>99%) were purchased from Sigma-Aldrich (France). Phloroglucinol (>99%) was purchased from Merck (France). Procyanidin B2 (≥90%), (-)-epicatechin (≥99%), (+)-catechin (≥99%), (-)-epicatechin-3-O-gallate (≥97.5%), (-)-epigallocatechin (≥98%) and (-)-epigallocatechin-3-O-gallate (≥98%) were purchased from Extrasynthese (France).
Depolymerization experiments

All depolymerization experiments described in the following sections were performed in three independent replicates.

Depolymerization of procyanidin B2 with 1 molar equivalent of nucleophile. Owing to the purity of the commercial B2 sample (≥90%), a B2 solution was first prepared with approximate concentration. After determination of the concentration by measuring the peak area at 280 nm with the UHPLC-DAD-MS system, the B2 solution was then precisely adjusted to 1.05 mM by addition of methanol. For each depolymerization kinetics, equal volumes of methanolic solutions of procyanidin B2 (1.05 mM), nucleophile (1.05 mM) and hydrochloric acid (HCl, 0.3 M) were mixed and distributed in 8 vials, which were immediately sealed and incubated at 30°C. Vials were withdrawn at different times and directly analyzed by UHPLC-DAD-MS.

Depolymerization of procyanidins from a grape seed extract with optimized amounts of nucleophiles. Methanolic solutions of phloroglucinol (30 g·L⁻¹; 0.24 M), 2-mercaptoethanol (165 g·L⁻¹; 2.1 M, prepared as a 15:85 v/v 2-mercaptoethanol/methanol mixture) and menthofuran (3 g·L⁻¹; 0.020 M) were prepared. For each depolymerization kinetics, equal volumes of methanolic solutions of grape seed extract (3 g·L⁻¹), nucleophile and HCl (0.3 M) were mixed and distributed in 10 vials, which were immediately sealed and incubated at 30°C. Vials were withdrawn at different times and directly analyzed by UHPLC-DAD-MS.

Characterization of pycnogenol with the menthofuran method. For each depolymerization kinetics, equal volumes of methanolic solutions of pycnogenol (3 g·L⁻¹), menthofuran (3 g·L⁻¹; 0.020 M) and HCl (0.3 M) were mixed and distributed in 3 vials, one for each reaction time, which were immediately sealed and incubated at 30°C. Vials were withdrawn after 90, 120 and 150 min and directly analyzed by UHPLC-DAD-MS.
Characterization of pycnogenol with the phloroglucinol method. The analysis was based on the protocol proposed by Kennedy and Jones.\textsuperscript{30} Methanolic solutions of pycnogenol (15 g·L\textsuperscript{-1}), ascorbic acid (30 g·L\textsuperscript{-1}; 0.39 M) and HCl (0.3 M) were prepared. For each depolymerization experiment, the pycnogenol solution was used to solubilize phloroglucinol (150 g·L\textsuperscript{-1}; 1.2 M). Immediately after, equal volumes of ascorbic acid, HCl and pycnogenol/phloroglucinol solutions were mixed and the depolymerization solution was incubated at 50°C in a closed flask. The final pycnogenol concentration was thus 5 g·L\textsuperscript{-1}. At 20 min of reaction time, the depolymerization solution was mixed with five volumes of an aqueous solution of sodium acetate (40 mM). The final solution was directly analyzed by UHPLC-DAD-MS.

Characterization of grape pericarp powder with the menthofuran method. A sample of grape pericarp powder (18 mg) was suspended in 0.6 mL of methanol in a closed flask. Then, methanolic solutions of menthofuran (0.6 mL; 30 g·L\textsuperscript{-1}; 0.24 M) and HCl (0.6 mL; 0.3 M) were added. The closed flask containing the depolymerization solution was incubated at 30°C. At 2h of reaction time, a sample was withdrawn, centrifuged 1 min at 3000 \( x \) \( g \), filtrated and directly analyzed by UHPLC-DAD-MS.

Characterization of grape pericarp powder with the 2-mercaptoethanol method. The analysis was based on the protocol proposed by Tanaka et al.\textsuperscript{31} A sample of grape pericarp powder (18 mg) was suspended in 0.6 mL of methanol in a closed flask. Then, methanolic solutions of 2-mercaptoethanol (0.6 mL; 165 g·L\textsuperscript{-1}; 2.1 M) and HCl (0.6 mL; 0.3 M) were added. The closed flask containing the depolymerization solution was incubated at 40°C. At 2h of reaction time, a sample was withdrawn, centrifuged 1 min at 3000 \( x \) \( g \), filtrated and directly analyzed by UHPLC-DAD-MS.

Characterization of the procyanidins from Douglas fir barks with the menthofuran method. 500 mg of Douglas fir barks ground and sieved at 2 mm were suspended in 50 mL methanol
(10 g·L⁻¹). Then, menthofuran (94 µL; final concentration 2 g·L⁻¹; 0.013 M) and HCl (417 µL; final concentration 0.1 M) were added and the depolymerization medium was incubated at 30°C. At defined time intervals, a sample was withdrawn, centrifuged 1 min at 3000 x g, filtrated and directly analyzed by UHPLC-DAD-MS.

**Characterization of the procyanidins from Douglas fir barks with the 2-mercaptoethanol method.** Douglas fir barks ground and sieved at 2 mm (500 mg) were suspended in 47 mL methanol. Then, 2-mercaptoethanol (2.5 mL; final concentration 5:95 v/v) and HCl (417 µL; final concentration 0.1 M) were added and the depolymerization mixture obtained was incubated at 40°C. At defined time intervals, a sample was withdrawn, centrifuged 1 min at 3000 x g, filtrated and directly analyzed by UHPLC-DAD-MS.

**Preparation, isolation and characterization of epicatechin-menthofuran (EC-MF)**

The grape seed extract (20 g) was dissolved in methanol (280 mL). Menthofuran (21.0 mL; 0.136 mol) and HCl (4.17 mL of 37% HCl in 200 mL methanol) were added. The reaction was performed for 1h at 30°C under magnetic stirring. The medium was then neutralized with a solution of sodium hydrogenocarbonate (4.2 g) in water (700 mL). Methanol was evaporated under vacuum. The remaining aqueous phase was extracted with ethyl acetate (3 times 500 mL). The organic layers were gathered, dried with sodium sulfate, filtrated and evaporated under vacuum. The dark powder obtained (27 g) was triturated and sonicated for 5 min in diethyl ether (3 times 300 mL). The diethyl ether fractions were pooled, dried with sodium sulfate, filtrated and evaporated under vacuum. Remaining traces of menthofuran were eliminated by trituration in petroleum ether (3 times 100 mL). A purple powder was obtained (21 g) containing C, EC, ECG, C-MF, EC-MF and ECG-MF (see the abbreviation section and Scheme 1). A sample (1 g) was purified by flash chromatography on a PF430 system (Interchim, France) equipped with a silica gel column (120 g; granulometry 63-200 µm). The flow rate was
set to 40 mL·min⁻¹ and the gradient was: solvent A (CH₂Cl₂), solvent B (CH₂Cl₂–CH₃OH, 90:10, v/v); 0–2 min, 0% B (isocratic); 2–15 min, 0% to 50% B (linear gradient); 15–28 min, 50%–80% B (linear gradient); and 28–45 min, 80% B (isocratic). Fractions were analyzed by UHPLC-DAD-MS before being combined and evaporated under vacuum to yield a pale purple pulverulent solid (80 mg) containing EC-MF (> 90 % purity according to UHPLC analyses). For NMR characterization, EC-MF was dissolved in d₆-DMSO.

1D and 2D NMR spectra acquisitions were performed at 25°C with an Avance III HD NMR spectrometer (Bruker, Germany) at 500 MHz for ¹H and 126 MHz for ¹³C. HRMS spectrum was acquired on a MicroTof QII mass spectrometer (Bruker) using the TOF MS ES+ mode, with samples dissolved in MeOH. Spectra are provided as supporting information (Figures S1 to S5).

**Epicatechin-(4→5)-menthofuran.** ¹H NMR δ (ppm): 9.12 (1H, s, H₁₀), 9.04 (1H, s, H₁₁), 8.89 (1H, s, H₈'), 8.77 (1H, s, H₁₁'), 6.85 (1H, s, H₂'), 6.68 (1H, d, J = 8.3 Hz, H₅'), 6.58 (1H, d, J = 8.3 Hz, H₆'), 5.87 (1H, s, H₈), 5.78 (1H, s, H₆), 5.05 (1H, bs, H₉), 4.77 (1H, s, H₂), 4.05 (1H, s, H₄), 3.78 (1H, bs, H₃), 2.56 (1H, m, H₇''), 2.27 (2H, m, H₄''), 2.10 (1H, m, H₇''), 1.84 (1H, m, H₆''), 1.77 (1H, m, H₅''), 1.67 (3H, s, H₈''), 1.28 (1H, m, H₅''), 1.03 (3H, d, J = 6.6 Hz, H₉''). ¹³C NMR δ (ppm): 157.2 (C₇), 156.9 (C₅), 156.0 (C₈a), 148.3 (C₂'), 147.0 (C₇a'), 144.7 (C₃'), 144.6 (C₄'), 130.3 (C₁'), 118.0 (C₃a'), 117.7 (C₇'), 114.9 (C₅'), 114.8 (C₉'), 113.9 (C₃'), 98.3 (C₄a), 95.2 (C₈), 94.0 (C₆), 75.1 (C₂), 69.5 (C₃), 37.9 (C₄), 31.9 (C₅'), 30.8 (C₇'), 29.2 (C₉'), 21.5 (C₄'), 19.6 (C₆'), 7.8 (C₈').

HRMS (ESI): [M+H]⁺ found with m/z 439.1751 (calculated for C₂₅H₂₇O₇: 439.1751).

**Analytical method (UHPLC-DAD-MS system)**

The liquid chromatography system was an Acquity ultra-high-pressure liquid chromatography (UHPLC) equipped with a photodiode array detector (DAD, Waters, Milford, MA). The column (HSS T3, 100 × 2.1 mm, 1.8 mm) contained a Nucleosil 120-3 C18 endcapped phase (Macherey-Nagel, Sweden). The flow rate was 0.55 mL·min⁻¹ and the gradient conditions were as follows,
except for experiments with menthofuran: solvent A (H₂O–HCOOH, 99:1, v/v), solvent B
(\text{CH}_3\text{CN–H}_2\text{O–HCOOH, 80:19:1, v/v/v}); 0–5 min, 0.1% to 40% B (linear gradient); 5–7 min, 40%
to 99% B (linear); 7–8 min, 99% B (isocratic); and 8–9 min, 99% to 0.1% B (linear). For the
analyses involving menthofuran: 0-5 min, 0.1 to 60% B (linear gradient); 5–7 min, 60% to 99%
B (linear); 7–8 min, 99% B (isocratic); and 8–9 min, 99% to 0.1% B (linear). The Acquity UHPLC
system was coupled online with an amaZon X Ion-Trap mass spectrometer (Bruker Daltonics,
Germany), with electrospray ionization operating in the positive ion mode. In the source, the
nebulizer pressure was 44 psi, the temperature of dry gas was set at 200°C with a flow of
12 L·min⁻¹ and the capillary voltage was set at 4 kV. The mass spectra were acquired over a
m/z range of 90-1500. The speed of mass spectrum acquisition was set at 8100 m/z s⁻¹.

**Peak identification and quantification**

The peaks from the UV chromatograms (280 nm) were attributed to the corresponding
compounds by comparing the associated mass spectra and retention times to those obtained
with authentic standards of (+)-C, (-)-EC, (-)-ECG, (-)-EGC and (-)-EGCG. The products resulting
from the trapping of extension units by a nucleophile, i.e., (epi)catechin-(4→X)-nucleophile;
or (E)C-NU, are not commercially available. Their mass spectra and retention times were
determined by depolymerizing procyanidin B2 with the nucleophiles studied, as this reaction
yields mainly epicatechin and the targeted EC-NU. The procyanidin depolymerization products
obtained with 2-mercaptoethanol have also been characterized by NMR in a previous study.⁵¹
Figures 1 and S6 show examples of the UV chromatograms obtained along the kinetic
experiments.

The molar responses at 280 nm of C, EC, ECG, EGC and EGCG were determined by calibration
with the corresponding commercial standards. The molar responses at 280 nm of the (E)C-NU
products were assessed by depolymerizing a procyanidin B2 sample with a large excess of
nucleophile. As the amount of extension units trapped by the nucleophile (i.e., EC-NU) and EC produced from the terminal units were expected to be equal, the ratio of the corresponding peak area was attributed to the ratio of respective molar responses of the products.45,51 For ECG-PG, the molar response relative to EC of 3.7 given by Kennedy and Jones30 was applied. For ECG-MF and ECG-ME, a molar response relative to EC of 3.7 was used based on the coefficient experimentally determined for ECG, assuming that the nucleophile moiety did not alter the molar response of ECG. This assumption was consistent with the molar response of 1 relative to EC found for EC-MF and EC-ME. All values, expressed proportionally to EC molar response, are given in supporting information (Table S1).

RESULTS AND DISCUSSION

Experiments were first conducted on procyanidin B2 with 1 molar equivalent of nucleophile relatively to B2 (Scheme 2). Then, menthofuran, phloroglucinol and 2-mercaptoethanol were compared in the depolymerization of a grape seed commercial extract to characterize its tannin composition. Each nucleophile was used at the optimized concentration described earlier in standard literature procedures, while the other reaction conditions remained equal. The analytical method using menthofuran was then compared to the Kennedy and Jones30 phloroglucinolysis method for the characterization of the tannin composition of another commercial tannin extract, pycnogenol9,10 and with mercaptolysis for the direct analysis of a grape pericarp powder and of a Douglas fir bark sample without prior extraction.51

Depolymerization of procyanidin B2 with one molar equivalent of nucleophile

The B2 dimer was chosen as a model of condensed tannins because its depolymerization conveniently yields only two products: (2R, 3R)-epicatechin as the released terminal unit, and (2R, 3R)-epicatechin-(4→X)-nucleophile as the trapped extension unit, respectively referred
to as EC and EC-NU. To examine the difference of reactivity between the nucleophiles quantified by the corresponding amounts of EC-NU produced, the reactions were carried out with a stoichiometric amount of each nucleophile with respect to procyanidin B2. Reactions were performed in methanol in the presence of 0.1 M HCl, as in most of the standard methods. The depolymerization tests were performed at 30°C to limit epimerization at C2 carbon atom of the flavanol unit that may occur following the ring opening in the acidic conditions. The percentages of EC-NU produced and of residual B2 with respect to initial B2 concentration were determined from the depolymerization experiments for the three nucleophiles tested (Table 1). The time required to reach the plateau of maximum EC-NU concentration is also indicated.

Menthofuran exhibited the highest efficiency to promote procyanidin B2 depolymerization under the reaction conditions applied. Indeed, an almost full consumption (>98.8%) of the procyanidin B2 dimer was observed in 40 min, with a recovery of 92% of the extension units in the form of EC-MF. Menthofuran purity (≥95%) may have limited the calculated EC-MF yield as the nucleophile concentration was adjusted assuming 100% purity. The actual menthofuran to B2 dimer initial molar ratio was thus between 0.95:1 and 1:1. In contrast, the reactions performed with 2-mercaptoethanol (ME) and phloroglucinol (PG), prepared from >99% pure products, reached an equilibrium with much lower proportions of EC-NU (41% of EC-ME and 23% of EC-PG, respectively), a lower proportion of terminal units, and a sizeable proportion of remaining B2 (5% with ME and 14% with PG, see also Figure S7).

The differences between the initial B2 concentration and final of EC-NU, EC and residual B2 concentrations, likely correspond to oligomers that could not be accurately quantified in the UHPLC-DAD-MS analysis, even though dimers like EC-(4→8)-EC-(4→2)-PG and trimers were observed in the chromatograms. Indeed, depolymerization products including EC-NU and EC,
as well as procyanidin B2, are competitive nucleophiles that can add onto the cationic site of extension units after cleavage, resulting in a large diversity of products and decreasing the yields in EC and EC-NU. The relative amounts of these oligomers could be assessed based on stoechiometry, from the difference between the amounts expressed in EC equivalents, of products formed (EC-NU + EC) and B2 consumed at the considered reaction time. They were found to account for 39% and 53% of B2 consumption in the reactions with 2-mercaptoethanol and phloroglucinol, respectively, when the proportion of EC-NU reached its maximal value. In the case of menthofuran, they were estimated to account for 9% of B2 consumption.

The high initial reaction rate observed with phloroglucinol evidences its good reactivity as a trapping reagent, but the only partial depolymerization of the B2 dimer at reaction equilibrium shows that the EC-PG product is also cleaved at a high rate in a reverse reaction. Such an equilibrium between procyanidin B2 and EC-PG was predictable considering the structural similarity of these products. Indeed, they both consist in a phloroglucinol-like ring linked to the EC benzylic carbon at C2, and epicatechin-(4→2)-phloroglucinol also undergoes acid-catalyzed cleavage of the (4→2) bond in the reaction conditions applied. Other equilibria involving the new oligomers occurred at the same time, impacting the equilibrium between procyanidin B2 and EC-PG. The same phenomenon occurred with EC-ME, where the mercaptoethanol moiety can also be substituted in acidic conditions. Contrarily, the high yield of conversion of the B2 dimer into depolymerization products obtained with menthofuran in a 1:1 initial molar ratio indicate that EC-MF units were not significantly affected by this reversibility issue (Table 1). Menthofuran thus advantageously solves this concern by displacing strongly and rapidly the depolymerization equilibrium towards EC and EC-MF, even when this nucleophile is used in stoichiometric amount.
Depolymerization of procyanidins from a grape seed extract with optimized amounts of nucleophiles

The promising results shown with menthofuran for the depolymerization of procyanidin B2 motivated the development of an analytical method for characterizing more complex proanthocyanidin extracts. Preliminary experiments carried out on a grape seed extract (1 g·L\(^{-1}\)) showed that a 1:1 (w/w) menthofuran to extract weight ratio was sufficient to achieve maximal depolymerization yield of procyanidins. A comparison with phloroglucinol and 2-mercaptoethanol at the optimized ratios reported in literature was done by performing kinetic experiments at 30°C on 1 g·L\(^{-1}\) of the same grape seed extract in methanol containing 0.1 M HCl. In these experiments, 10:1 and 55:1 (w/w) nucleophile to extract ratios were used for phloroglucinol and 2-mercaptoethanol, respectively, corresponding to the 10:1 (w/w) weight ratio defined by Kennedy and Jones\(^{30}\) for phloroglucinol and to the 5% (v/v) volume ratio proposed by Tanaka and coworkers\(^{31,35}\) for 2-mercaptoethanol. In the extreme case where the tannin extract would consist only of EC extension monomers (molecular weight 290 g·mol\(^{-1}\)), these weight ratios corresponded to a molar excess of nucleophile of 2, 23 and 200 for menthofuran, phloroglucinol and 2-mercaptoethanol, respectively.

The depolymerization products were categorized in four types of units, considering on one hand, extension units versus terminal units and on the other hand, galloylated units versus non-galloylated units (Figure 2). For instance, the C-NU and EC-NU concentrations measured were summed to evaluate the amount of non-galloylated extension units. This enabled to infer the average composition in constitutive units of the polymers.

The release of non-galloylated units and of galloylated units followed different kinetics. The maximum concentration of non-galloylated units was reached faster (after 120-150 min) than that of galloylated units (after 200-300 min). It can also be noted that the amount of non-
galloylated units decreased over time, contrary to galloylated units, indicating higher stability of the latter. These results point to the importance of performing complete kinetic experiments when characterizing tannin composition instead of selecting an arbitrary time, because this optimal time may vary depending on the tannin extract. This is especially important when reactions are slow, which can lead to numerous side-reactions.

The concentrations reached at the plateau in the kinetic experiments were used to calculate the weight percentages of the four types of constitutive units obtained with each nucleophile for the grape seed extract (Table 2). In each case, depolymerizable units represented around 46% (w/w) of the grape seed extract and consisted of around 25% of extension units, 13% of terminal units, 5% of galloylated extension units and 2% of galloylated terminal units. The results were thus comparable despite the different amounts of nucleophile applied.

The analytical depolymerization involving menthofuran as the nucleophilic reagent thus demonstrated the same performance as with 10- and 100-times higher amounts of phloroglucinol and 2-mercaptoethanol, respectively. To our knowledge, this makes menthofuran the only nucleophilic trapping reagent described so far that enables the depolymerization of condensed tannins with maximal yield using a near to quantitative nucleophile to procyanidins molar ratio.

**Comparison of furanolysis with standard methods**

The high efficiency of menthofuran to trap the extension units released from tannin depolymerization led to evaluate its use in comparison with the phloroglucinolysis developed by Kennedy and Jones\textsuperscript{30} and with mercaptolysis based on the work of Tanaka et al.\textsuperscript{31} Figure 3A compares the results of the analysis of a maritime pine bark extract, commercially available under the name pycnogenol, using a 1:1 (w/w) weight ratio of menthofuran to procyanidin extract and a 10:1 (w/w) weight ratio of phloroglucinol to procyanidin extract as
optimized in the standard method. Both reactions were performed in methanol containing 0.1 M HCl. Reaction with menthofuran, performed at 30°C for 90 min, resulted in a very similar procyanidin composition profile as phloroglucinolysis (50°C, 20 min), with a good reproducibility and equivalent depolymerization yields, although a 10-fold lower nucleophile to extract weight ratio was used. This represented a non-negligible saving of reactants.

The menthofuran method was also compared to mercaptolysis for the characterization of the proanthocyanidin fractions of a grape pericarp powder. These methods were applied directly on the biomass sample (i.e., without prior extraction of tannins), in methanol containing 0.1 M HCl. Furanolysis was performed with a 1:1 weight ratio of menthofuran to grape pericarp powder at 30°C for 2h, while mercaptolysis was performed with a 5.5:1 weight ratio of 2-mercaptoethanol to grape pericarp powder at 40°C for 2h. Chromatograms are given as supporting information (Figures S8 and S9). Both methods showed similar results and good reproducibility (Figure 3B), despite using a 10-fold lower molar amount of nucleophile for furanolysis.

The menthofuran method was also applied to the direct analysis of Douglas fir bark powder, without a preliminary extraction step. The depolymerization products reached their maximal concentration after 20h, yielding a procyanidin content of 3.9% (w/w) of biomass dry weight, including 3.2 ± 0.2% of extension units and 0.7 ± 0.0% of terminal units. On the same sample, direct mercaptolysis (selected for reference since the standard phloroglucinolysis protocol first proceeds with tannin extraction) gave a similar procyanidin content (3.7% w/w), including 3.0 ± 0.1% extension units and 0.7 ± 0.0% terminal units. It should be noted that mercaptolysis was faster (the maximal concentration was reached in 4h) due to the higher temperature (40°C vs. 30°C) and the high nucleophile excess (60-times higher than menthofuran) used in this method. The differences in polarity and solvation of the
nucleophiles may also affect the depolymerization kinetics through mass transfer limitations within the solid bark sample.

Menthofuran thus displays interesting properties as a trapping reagent for the analytical depolymerization of procyanidins. Its high efficiency allows its use at low concentration, contrary to the large molar excesses required with the classical nucleophiles, and low temperature for a fast conversion of condensed tannins into monomeric units without significant reversal of the reaction. It is a readily available commercial chemical, and its use in near to stoichiometric amounts contributes to save cost and to lower the exposure to reactants. In the same way to the other nucleophiles, adjustments may be required in the protocol according to the sample to be analyzed. Since the amount of procyanidin in the sample is unknown, the quantity of menthofuran needs to be estimated to ensure it is higher than the quantity of extension units. Also, like with the other nucleophiles and methods, the minimum reaction time required to reach the maximum depolymerization yield may vary according to the sample (plant species, organ and physiological status affect tannin concentration and constitutive unit composition) and especially according to its type of preparation (ground raw biomass or more or less purified extracts). The good stability of the depolymerization products in the presence of menthofuran at 30°C enables to use a single, longer than required reaction time for samples of a same type when (recommended) systematic kinetics experiments cannot be performed. As a general guideline, the following conditions may be applied for unknown samples.

*For the characterization of tannins in soluble extracts*, the following reaction conditions are proposed as a standard setup: the depolymerization of a 1 g·L⁻¹ tannin extract in methanol in the presence of 0.1 M HCl is carried out with 1 g·L⁻¹ menthofuran at 30°C over 2h, either with end-point analysis or, preferably when possible, following the complete kinetics.
For the direct analysis of raw biomass, a concentration of 10 g·L⁻¹ of dry biomass sample may allow a good response in LC-DAD(-MS), using 2 g·L⁻¹ menthofuran. Compared to soluble extracts, mass transfer limitations may require a longer reaction time, as shown with the analysis of the bark sample, for which the reaction was complete in 20h. Alternatively, menthofuran concentration can also be increased to speed up the reaction. As the water content of raw biomass samples may affect the efficiency and rate of the depolymerization process, it is advisable to dry such samples to less than 15% water (w/w).

At appropriate times, the reaction medium can be directly analyzed by a LC-DAD(-MS) system according to a protocol similar as the one given in Material & Methods. When significant delays are expected between the reaction and the analysis of the reaction products, for example when large series of samples are scheduled in parallel, it is preferred to raise the pH of the reaction medium to pH 4-5 after the reaction is over, in order to avoid side-reactions.

On a more general note, the menthofuran method, referred to as furanolysis, demonstrates the potential of furans as nucleophilic trapping reagents in the depolymerization of condensed tannins. In a former work, furan and sylvan, two compounds that can be obtained by conversion of C5 sugars from wood biomass, were indeed proven to be efficient nucleophiles for quantitatively supplying fully biobased building blocks from condensed tannins.⁴⁴,⁴⁵ The superior efficiency of the menthofuran method needs to be tested against tannin structures known to be more recalcitrant to the usual depolymerization conditions, such as 5-deoxy tannins or A-type proanthocyanidins, using harsher conditions (e.g., higher temperature and/or acid concentration).

ABBREVIATIONS USED
Flavanols: B2, procyanidin B2 or B2 dimer; C, catechin; CG, catechin-3-O-gallate; EC, epicatechin; ECG, epicatechin-3-O-gallate; EGC, epigallocatechin; EGCG, epigallocatechin-3-O-gallate. Nucleophiles: ME, mercaptoethanol; MF, menthofuran; PG, phloroglucinol; NU, nucleophile. Flavanol derivatives (representative examples of the numerous combinations, see also Schemes 1 & 2): C-ME, catechin-(4→2S)-mercaptoethanol; EC-MF, epicatechin-(4→5)-menthofuran; ECG-PG, epicatechin-3-O-gallate-(4→2)-phloroglucinol; EGCG-NU, epigallocatechin-3-O-gallate-(4→X)-nucleophile; (E)CG-NU, (epi)catechin-3-O-gallate-(4→X)-nucleophile.

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SUPPORTING INFORMATION

UV chromatograms (280 nm) of the grape seed extract depolymerized with menthofuran throughout the kinetic experiment. Detailed kinetics of the depolymerization tests performed on procyanidin B2 with 1 molar equivalent. UV chromatograms (280 nm) of the products of depolymerization of the grape pericarp powder containing procyanidins and prodelphinidins with menthofuran and 2-mercaptoethanol.
REFERENCES


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FIGURE CAPTIONS

Scheme 1: Depolymerization of procyandinis leading to the release of terminal units and the trapping of extension units by a nucleophile (example of the products obtained with menthofuran).

Scheme 2: Other procyandin derivatives encountered in this study. Abbreviations: C, catechin; EC, epicatechin; CG, catechin-3-O-gallate; ECG, epicatechin-3-O-gallate; PG, phloroglucinol; ME, mercaptoethanol.

Figure 1: UV chromatograms (280 nm) of (A) the grape seed extract dissolved in methanol (1 g·L⁻¹) and (B) the same extract after acid-catalyzed depolymerization with menthofuran (1 g·L⁻¹; 65 min reaction). 1: C (291 m/z), 2: EC (291 m/z), 3: ECG (443 m/z), 4: C-MF (439 m/z), 5: EC-MF (439 m/z), 6: ECG-MF (591 m/z).

Figure 2: Depolymerization kinetics of the procyandinis contained in the grape seed extract according to the nucleophile used: menthofuran, phloroglucinol or 2-mercaptoethanol. Extension units (C-NU + EC-NU) are represented by red circles, terminal units (C + EC) by blue squares, galloylated extension units (ECG-NU) by purple triangles and galloylated terminal units (ECG) by green diamonds. Experimental points are means and error bars are standard deviations, calculated from three independent kinetic experiments.

Figure 3: Comparison of the menthofuran method (A) with a standard phloroglucinol method through the characterization of a pine bark extract (pycnogenol), (B) with mercaptolysis through the characterization of a grape pericarp powder containing procyandinis and
prodelphinidins. Mass contents are means and error bars are standard deviations calculated from three independent experiments.
### Table 1: Procyanidin B2 conversion yields in the presence of 1 molar equivalent of nucleophiles

<table>
<thead>
<tr>
<th>Nucleophile</th>
<th>Time to reach the plateau</th>
<th>EC-NU production yield at $t_{\text{max}}$</th>
<th>EC production yield at $t_{\text{max}}$</th>
<th>Remaining B2 dimer at $t_{\text{max}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$t_{\text{max}}$ (h)</td>
<td>mol% of theoretical maximum$^i$</td>
<td>mol% of theoretical maximum$^i$</td>
<td>mol% of initial B2 concentration</td>
</tr>
<tr>
<td>Menthofuran</td>
<td>0.7</td>
<td>91.6 ± 1.1</td>
<td>88.1 ± 1.9</td>
<td>1.2 ± 0.0</td>
</tr>
<tr>
<td>2-Mercaptoethanol</td>
<td>2.0</td>
<td>40.5 ± 0.4</td>
<td>74.9 ± 2.1</td>
<td>5.4 ± 0.7</td>
</tr>
<tr>
<td>Phloroglucinol</td>
<td>0.2</td>
<td>22.5 ± 0.4</td>
<td>65.8 ± 0.7</td>
<td>14.1 ± 0.1</td>
</tr>
</tbody>
</table>

$t_{\text{max}}$: time at which the plateau of maximum EC-NU concentration is reached

$^i$ the theoretical maximum corresponds to the conversion of 1 mol B2 dimer into 1 mol EC plus 1 mol EC-NU.

Values of mean and standard deviation were determined by performing three independent experiments.

### Table 2: Composition in procyanidin constitutive units of a grape seed extract according to the nucleophile used for its characterization.

<table>
<thead>
<tr>
<th>Nucleophile</th>
<th>Extension Units$^a$ % (w/w)$^i$</th>
<th>Terminal Units$^b$ % (w/w)$^i$</th>
<th>Galloylated Extension Units$^c$ % (w/w)$^{ii}$</th>
<th>Galloylated Terminal Units$^d$ % (w/w)$^i$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Menthofuran</td>
<td>25.9 ± 0.4</td>
<td>13.1 ± 0.3</td>
<td>5.2 ± 0.1</td>
<td>1.6 ± 0.0</td>
</tr>
<tr>
<td>2-Mercaptoethanol</td>
<td>25.0 ± 0.3</td>
<td>12.9 ± 0.2</td>
<td>5.1 ± 0.1</td>
<td>1.6 ± 0.0</td>
</tr>
<tr>
<td>Phloroglucinol</td>
<td>25.6 ± 0.3</td>
<td>12.5 ± 0.2</td>
<td>5.2 ± 0.2</td>
<td>1.6 ± 0.0</td>
</tr>
</tbody>
</table>

$^a$ Mean values and standard deviations were calculated for each nucleophile from three independent kinetic experiments using concentrations at (i) three reaction times (9 experimental points) and (ii) two reaction times (6 experimental points).
**Scheme 1**

2,3-trans : catechin-\((4\rightarrow 5)\)-menthofuran, C-MF

2,3-cis : epicatechin-\((4\rightarrow 5)\)-menthofuran, EC-MF

**Procyanidins**

Extension units

Terminal unit

R = H or Gal

Depolymerization

(+ Nucleophile)

2,3-trans : catechin, C

2,3-cis : epicatechin, EC

Nu = menthofuran

R = H : R = Gal :

2,3-trans : catechin-3-O-gallate, CG

2,3-cis : epicatechin-3-O-gallate, ECG
Scheme 2

Procyanidin B2

Epicatechin-(4→8)-epicatechin-(4→8)-phloroglucinol, EC-EC-PG

\[ \text{Procyanidin B2} \]

\[ \text{Epicatechin-(4→8)-epicatechin-(4→8)-phloroglucinol, EC-EC-PG} \]
Figure 1

A

B
Figure 2

Menthofuran (2 molar eq.)

Phloroglucinol (23 molar eq.)

2-Mercaptoethanol (200 molar eq.)

Concentrations (mmol/L)

Time (min)

- Extension units
- Terminal units
- Galloylated extension units
- Galloylated terminal units
Figure 3

A

Composition (g/100 g)

- Extension units
- Terminal units
- Taxifolin

Menthofuran method    Phloroglucinol method

B

Composition (g/100 g)

- C
- ECG
- EGC-NU
- CAU
- ESG-NU
- EC-NU
- ECG-NU

Menthofuran method    2-Mercaptoethanol method
For table of contents only.