Design of Natural Food Antioxidant Ingredients through a Chemometric Approach

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In the present work, an environmentally friendly extraction process using subcritical conditions has been tested to obtain potential natural food ingredients from natural sources such as plants, fruits, spirulina, propolis, and tuber, with the scope of substituting synthetic antioxidants, which are subject to regulation restrictions and might be harmful for human health. A full characterization has been undertaken from the chemical and biochemical point of view to be able to understand their mechanism of action. Thus, an analytical method for profiling the compounds responsible for the antioxidant activity has been used, allowing the simultaneous determination of water-soluble vitamins, fat-soluble vitamins, phenolic compounds, carotenoids, and chlorophylls in a single run. This information has been integrated and analyzed using a chemometrical approach to correlate the bioactive compounds profile with the antioxidant activity and thus to be able to predict antioxidant activities of complex formulations. As a further step, a simplex centroid mixture design has been tested to find the optimal formulation and to calculate the effect of the interaction among individual extracts in the mixture.

KEYWORDS: Antioxidant activity; simplex centroid mixture design; polyphenols; vitamins; MLR; PLS; FS; HPLC-DAD profile

INTRODUCTION

Food additives play an important role in today’s complex food supply; nevertheless, they represent one of the most misunderstood topics in food safety raising consumers’ concerns. Food control focuses mainly on chemical additives, which although often present are usually only in minor or trace amounts. They are intentionally added to food in order to produce a desired positive effect, although their level has to be maintained within regulated limits. Natural antioxidants are receiving increasing attention in food science because of the reports stating that diets rich in plant antioxidants derived from fruits and vegetables are associated with lower risks of coronary heart disease and cancer (1, 2). For example, recent reports have been published suggesting that naturally occurring homologous mixtures have a more beneficial effect than the intake of synthetically produced vitamin E (3). This fact is probably related to the absence of nonactive stereoisomers (4) in synthetic products. Therefore, the importance of finding natural sources of antioxidants is increasing. Renewed attention in the food industry is being focused on CO2 as a clean technology for ingredients and additives in manufacturing. CO2 extraction has been recognized as one of the new technologies able to extract higher quality natural ingredients, such as food aroma compounds, colorants, antioxidants, and even antimicrobial agents. Several advantages have been demonstrated compared to traditional extraction methods using organic solvents or steam-distillation to extract different compounds from natural sources. For instance, the potential to process natural products at mild temperatures under chemically inert conditions, using CO2 as an extraction fluid, also results in a low environmental impact. Moreover, due to these mild conditions, the functional, sensorial, and nutritional properties of the products are kept unaltered (5, 6).

An important challenge in the food industry when dealing with natural additives is to know their exact chemical composition and to be able to correlate it with their biological activity. Therefore, in the present work, a full characterization of the different extracts obtained under subcritical conditions has been undertaken from a chemical and biochemical point of view. The selected method for chemical characterization was based on a previous work carried out in our laboratory (7) for profiling different bioactive compounds such as water-soluble vitamins (ascorbic acid, thiamine, folate, pyridoxine, nicotinamide, cobalamin), fat-soluble vitamins (α-tocopherol, retinol acetate, cholecalciferol), phenolic compounds (phenolic acids, cinnamic acids, flavanones, isoflavones, anthocyanins), carotenoids (β-carotene and lutein), and chlorophylls in a single run, based on an HPLC-DAD analysis. The complete information obtained using this method can be

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combined with data corresponding to antioxidant capacity (DPPH radical scavenging method) and, once integrated and analyzed using different chemometric approaches, can provide a useful tool to correlate composition and bioactivity, with the added advantage of offering a method to predict antioxidant activities of similar or even more complex formulations.

Experimental designs are highly useful when it comes to optimize several parameters at once or when trying to find the best composition of a mixture of more than two components. A mixture experiment is an special type of experimental design in which the factors are the ingredients or components of a mixture. Among the different mixtures experiments, simplex centroid designs allow one to estimate not only main effects but also significant interactions on a mixture (8).

The present work is focused in the development of novel class of food additives based on substances extracted from natural sources by using environmentally friendly extraction processes, such as near critical carbon dioxide extraction. The use of chemometrics allows one to obtain information about the correlation between chemical composition and bioactivity considering, in a first step, all the compounds involved in the extract, meaning all the compounds giving a characteristic profile of the sample. This is a more comprehensive approach, since the compounds are not discriminated in advance for their supposed (described in the literature), or not, bioactivity. On the contrary, the hypothesis is related to the effectiveness of a profile to provide a specific biological activity. The optimization of the model is carried out by means of multiple linear regression methods whose validity is lately assured by using a simplex centroid design.

MATERIALS AND METHODS

Subcritical Fluid Extraction. In a first step, dried raw materials (Spirulina pacifica, Citrus compositum, Raphanus niger (radish), Rosmarinus officinalis (rosemary), Propolis, Medicago compositum (alfalfa), Carica papaya (papaya)) were extracted using a hydroethanolic mixture (30:70) and near critical CO2. Extraction conditions ranged between 68 and 75 atm and 37 and 45 °C for each raw material. A total of 10 extraction conditions were evaluated. The equipment used to obtain bioactive extracts was SFT-100 of Supercritical Fluid Technology Inc. (Newark, DE) with adaptations with a 100 mL extraction vessel. This adaptation has been designed and operated by Bioma Agro Ecology CO (Quartino, Switzerland).

The novelty of this approach is the extraction rationale which is the opposite of the traditional extraction processes. Instead of heating the extraction system in order to shorten the extraction time, the process is carried out near room temperature using a rising pressure on the extraction liquid that interacts with the solid matrix. Extraction at low temperatures is a relevant issue, since it is possible to avoid a thermal stress on thermolabile substances. Various experimental conditions, as described below, were used to determine the conditions that maximized the extraction. The SFE procedures were performed for 60 min at a flow rate of 5 mL/min of liquid CO2.

Chemical Characterization. A HPLC method previously developed in our laboratory (7) was used; under the selected conditions, water-soluble vitamins (ascorbic acid, thiamine, folic, pyridoxine, nicotinamide, cobalamin), fat-soluble vitamins (vitamin A, carotenoids), phenolic compounds (phenolic acids, cinnamic acids, flavonoids, anthocyanins), and carotenoids were monitored. The method consisted of measuring the change in absorbance that occurred at 280 nm by mixing 195 mL of DPPH solution (23.5 mg/L in ethanol) with 5 mL of extract. Since reaction time depends on the extract, data were collected every 15 min for 5 h. The stabilization time occurred when a plateau was reached, the end of the reaction for each sample.

The antioxidant activity of the extracts was measured by the DPPH radical scavenging method. This method is based on that previously developed by Brand-Williams et al. (6) adapted to use 96-microwell plates. The method consisted of measuring the change in absorbance that occurred at 517 nm by mixing 195 mL of DPPH solution (23.5 mg/L in ethanol) with 5 mL of extract. Since reaction time depends on the extract, data were collected every 15 min for 5 h. The stabilization time occurred when a plateau was reached, the end of the reaction for each sample.

Statistical Analysis. In order to find the main components of the extracts that contribute to its antioxidant activity, data related to the measured antioxidant activity and vitamin-phenolic HPLC profiles were submitted to multiple linear regression (MLR) analysis using forward stepwise (fs) and partial least squares (PLS). An equation of the form $y = b_0 + \sum_{i=1}^{p} b_i x_i$ is assumed, where $b_0$ is the intercept of the model, $b_i$ is the regression coefficient for the $i$th compound ($X_i$), $p$ is the number of compounds in the model, and $x_i$ is the antioxidant activity calculated using the model. All calculations were done with the STATISTICA software for Windows, version 7.1 (StatSoft, Tulsa, OK, 2006, http://www.statsoft.com), using the forward stepwise procedure in the Statistics, Multiple Regression module (with values of 4.0 and 3.9 for F-to-enter and F-to-remove, respectively, and fixing a limit of 10 steps) and the partial least squares regression procedure in the Statistics, Advanced Linear/Nonlinear Model module of STATISTICA program.

Mixture Experimental Design. The design of the antioxidant additive and its composition was studied by means of a modified simplex centroid design (12). The components of the mixture were selected among those with higher contribution to the antioxidant activity. This kind of modified simplex centroid with face experiments design is a mixture design for the full quadratic model with more runs than the classic design.

RESULTS AND DISCUSSION

Extraction conditions ranged between 68 and 75 atm and 37 and 45 °C. As can be seen, this is a relatively small area of experimental conditions; in fact, these ranges could be a normal variation in an industrial scale extraction plant. Our aim was to cover this area to appreciate the effect of normal variation, where pressure and temperature cannot be controlled as easily as on a laboratory scale. By using these extraction conditions, CO2 expands its liquid phase (water/ethanolic) and reduces its viscosity (13), increasing the mass transfer rate.

When the antioxidant activity of vegetable extracts was measured, three levels of activity were found in the DPPH radical scavenging test: low (citrus, propolis, and raphanus), medium (spirulina, alfalfa, and papaya), and high (rosmarin) antioxidant. On the other hand, the activity of the extracts showed the same trend when plotting antioxidant activity (EC50) versus total polyphenolic content, as can be seen in Figure 1.

Chemical characterization was done using a method previously developed in our laboratory (7). This screening method allows one to quantify simultaneously several kinds of bioactive compounds, namely, vitamins (hydro- and fat-soluble), phenolic compounds, and certain pigments such as chlorophylls and carotenoids. This method proved its efficacy in juices, fortified juices, beers, and milky drinks (7).
In general terms, three zones were easily differentiated in the chromatograms: water-soluble vitamins (2–12 min), phenolic compounds (10–20 min), and, finally, fat-soluble vitamins and pigments (20–40 min). Representative chromatograms of the different raw materials are shown in Figure 2. Table 1 shows the mean composition of the different raw materials considering the different families of compounds detected in the samples. As a matter of fact, individual components were identified tentatively and classified among the different groups or families of compounds, such as phenolic acids, cinnamic acids, flavanones, isoflavones, and anthocyanins and numbered according to their retention time; in order to simplify the table, the quantification is presented as the total content of the different groups of compounds, but the statistical analysis was applied considering the individual contribution of each compound corresponding to the different families. The quantification of individual compounds of each family was performed.

A first analysis of the data obtained showed the absence of fat-soluble vitamins. Only a small amount of α-tocopherol could be detected in some rosemary extract, but below the quantification level. The main vitamin found in the extracts was thiamine followed by nicotinamide and ascorbic acid. The HPLC analysis of the different extracts revealed, as expected, the presence of rosmarinic acid and carnosic acid and their derivatives in rosemary extracts, providing a high antioxidant activity. Through the study of the chromatographic profiles of the different extracts, it can be inferred that the medium antioxidant activity of Spirulina, alfalfa (Medicago composita) and papaya can be attributed to the presence of carotenoids and phenolic compounds and their synergistic effects. Also, many flavonoids have been detected in extracts from propolis, alfalfa, rosemary, raphanus, and fruits, and most of them have been related to the antioxidant and antimicrobial activity of the CO2-alcoholic and near-critical extracts (14–17).

Statistical Analysis. In order to identify the main components of the extracts that contribute to their antioxidant activity, statistical analysis was performed considering antioxidant activities and the vitamin-phenolic HPLC profiles allowing for the individual contribution of each compound in the whole profile. Two different statistical methods were used: FSMLR (forward stepwise multiple linear regression) and PLS (partial least squares).

When MLR using the FS procedure was applied to find the main components of the extracts that contribute to their antioxidant activity, the selected variables in the vitamin-phenolic HPLC profile were Flavonol15, Flavanone18, Cinnamic acid21, Flavanone15, Flavonol19, Cinnamic acid28, Flavanone13, Flavanone14, Flavonol23, and Flavonol13. These compounds have been numbered according to their retention time, as mentioned above. Results of the regression provide the following equation:

$$\text{antioxidant activity} = 77.80 + 227.25 \\ \times (\text{Cinnamic21}) + 3.17 \\ \times (\text{Flavanone13}) + 11.04 \\ \times (\text{Flavanone14}) - 7.77 \\ \times (\text{Flavonol13}) + 42.96 \\ \times (\text{Cinnamic28}) + 5.71 \\ \times (\text{Flavanone15}) - 4.71 \\ \times (\text{Flavonol15}) + 7.46 \\ \times (\text{Flavonol19}) + 0.96 \\ \times (\text{Flavanone18}) + 6.05 \\ \times (\text{Flavonol23})$$

The value of determination coefficient was $R^2 = 0.917$, and the value of the root-mean-square error of calibration (RMSEC) was 10.59. RMSEC is defined by the equation: \[\text{RMSEC} = \sqrt{\frac{\sum(y_i - \hat{y}_i)^2}{n}}\], where $n$ is the number of samples (in this study $n = 60$), $y_i$ is the observed antioxidant activity, and $\hat{y}_i$ is the antioxidant activity calculated using eq 1. Figure 3A shows the calculated values versus the observed values for antioxidant activity. As can be seen, the fit for the predictions of antioxidant activity can be considered appropriate. Despite the proven antioxidant activity of some identified compounds such as ascorbic acid (vitamin C) (18), these compounds are not indicated by the FS-MLR test as being 10 main contributors to the antioxidant activity; neither to increase EC50 (positive coefficient) nor to decrease it...
The same procedure was followed but grouping the compounds in families, but the adjustment obtained was very poor ($R^2 < 0.7$).

On the other hand, when MLR using the PLS procedure was applied to the prediction of antioxidant activity from all the compounds involved in the complete HPLC profile, three main components were selected by cross-validation, and values of $R^2 = 0.857$ and RMSEC = 16 were obtained. Figure 3B shows the predicted values for antioxidant activity from MLR using the PLS procedure versus the observed values. As can be seen, the fit can be considered as also appropriate.

**Figure 2.** HPLC-DAD chromatograms showing the different regions of bioactive compounds.

**Figure 3.** Scatter plot of predicted values for EC$_{50}$ from MLR using FS (A) and PLS (B) procedures versus observed values.
287 experiments are conducted to determine if mixed blends (or for-
288 mulations) are more desirable than additives obtained using a
289 single raw material. An important property of the mixture
290 experiment is that the change in the response depends on the
291 proportionality of the individual components present in the
292 mixture and not on the amount of the mixture (8).

293 The analysis of variance of the extraction conditions (con-
294 sidering the different experiments) versus antioxidant activity
295 only showed statistically significant differences among different
296 batches of Spirulina and alfalfa. On the other hand, although such
297 differences can be found in some values of antioxidant activity
298 of some extracts, the real values of antioxidant activity are
299 close enough (±7 μg/mL of the difference in EC50 values around
300 70–80 μg/mL) to consider that extraction conditions do not
301 strongly affect the results obtained in terms of antioxidant
302 activity. Therefore, by using this approach, each product (raw
303 material) was considered independent of the extraction condi-
304 tions (in the assayed range), and thus, 10 values were considered
305 and a mean and standard deviation were obtained. To be able to
306 design a formulation considering the different raw materials, a
307 PLS methodology was used, and eq 2 was obtained, as follows:

\[
antioxidant activity = 102.78 \times \text{Citrus} + 70.91 \\
\times \text{Papaya} + 67.67 \\
\times \text{Alfalfa} + 140.54 \\
\times \text{Propolis} + 118.81 \\
\times \text{Raphanus} + 10.07 \\
\times \text{Rosmarinus} + 80.23 \times \text{Spirulina} \quad (2)
\]

In this equation, a linear model of composition (7 factor
310 simplex lattice) was considered, and therefore, no synergies were
311 taken into account; each term is composed by the relative amount
312 of the raw material (expressed in %) and a coefficient that gives
313 an estimation of the relative contribution of each raw material to
314 the total antioxidant activity (expressed as EC50). As can be seen
315 in Figure 4, the best results (lowest EC50 values) are those with
316 high proportions of rosemary (100% and 50%), followed by
317 medicago (50%) and papaya (50%). The main disadvantage of
318 using this model is the lack of prediction in case of synergies or
319 inhibitions among different compounds coming from different
320 raw materials.

In order to demonstrate the ability of our approach to predict
321 the antioxidant activity of a mixture of different raw materials, a
322 systematic study was performed using a mixture experiment.
323 The final goal was, with this model, to be able to find the
324 optimum formulation, that is, the one providing the best anti-
325 oxidant activity/profile. Therefore, a modified simplex centroid
326 design for mixtures, with three components at different concen-
327 trations (prepared on a dry basis), was performed; the graphical
328 planning of the design can be seen in Figure 5, and the levels of the
329 factors in Table 2. Experiments 1, 2, and 3 were vertex points of
330 the design corresponding to pure components. Points 4, 5, and 6
331 on the sides were 1:1 binary mixtures, while points 7, 9, and 9
332 corresponded to mixtures of 1:1:4, 1:4:1, and 4:1:1 proportions,
333 respectively. Points 10, 11, and 12 were ternary mixtures with
334 equal parts of all three components. The design was run by using
335 three more antioxidant materials, namely, rosemary, alfalfa, and
336 papaya. All the mixtures were prepared in percentages of dry
337 weight and redissolved in ethanol/water (1:1). The batch selected
338 to run the design was batch 2 of each raw material, and the
339 extraction conditions used were 71.5 atm and 41 °C. A linear
340 predictive model including interactions (n = 12 runs) was
341 considered.
chemometric approach, the design of the antioxidant ingredient (additive-formulation) requires minimum experimental effort, which is ensured by a well-planned experimental design.

LITERATURE CITED


