Article
Comparison of Bioactive Substances Content between Commercial and Wild-Type Isolates of Pleurotus eryngii

Antonella Calabretti 1, Stefania M. Mang 2*, Antonella Becce 2, Donato Castronuovo 2, Loriana Cardone 2, Vincenzo Candido 3 and Ippolito Camele 2

1 Department of Chemical and Pharmaceutical Sciences, Piazzale Europa 1-Via Giorgieri 1, University of Trieste, 34127 Trieste, Italy; antonella.calabretti@deams.units.it
2 School of Agricultural, Forestry, Food and Environmental Sciences (SAFE), University of Basilicata, Viale dell’Ateneo Lucano 10, 85100 Potenza, Italy; antonellabecce@ymail.com (A.B.); donato.castronuovo@unibas.it (D.C.); loriana.cardone@unibas.it (L.C.); ippolito.camele@unibas.it (I.C.)
3 Department of European and Mediterranean Cultures, Environment, and Cultural Heritage, University of Basilicata, Via Lanera 20, 75100 Matera, Italy; vincenzo.candido@unibas.it
* Correspondence: stefania.mang@unibas.it; Tel.: +39-097-1205-544

Abstract: Mushrooms belonging to Pleurotus genus have been demonstrated to have important nutritional and medicinal value and their regular intake prevent many diseases, reduce the infection probability and increase immunity. In order to investigate the bioactive compounds produced by seven commercial (‘142 F’, ‘142 E’, ‘D+', ‘V turbo’, ‘V 142’, ‘A 12’, ‘V 160’) and five wild-type (‘Albino 1107’, ‘Altamura 1603’, ‘Muro Lucano 139’, ‘Conversano 1250’, ‘Albino beige chiaro 1094’) Pleurotus eryngii isolates, the following qualitative analyses were performed: Total polyphenol content, antioxidant activity (EC50 of ABTS) and antiradical power (ARP) in fresh lyophilized and dry basidioma, and water content, β-glucans and phenolic compounds in fresh samples. Standard methods were employed for each of the above mentioned aims. Total polyphenol content was diverse among the P. eryngii isolates. In particular, an elevated polyphenolic content was found in fresh lyophilized P. eryngii samples of the commercial isolates ‘V 142’ followed by ‘A 12’. The highest antiradical activity (ARP) was obtained by ‘Muro Lucano 139’ isolate. Wild P. eryngii isolates were characterized by higher water and β-glucans contents compared to the commercial ones, and the highest values were registered for the ‘Albino beige chiaro 1094’ isolate. In conclusion, the present study allowed us to identify the commercial and wild-type P. eryngii isolates from the Basilicata region, with high nutritional and medicinal value based on their bioactive compounds.

Keywords: “King Oyster”; bioactive compounds; qualitative analyses; total polyphenols; glucans; antioxidants

1. Introduction

Mushrooms are important functional foods included in the recommended well-balanced diet [1–6] and in the “Third Food Kingdom” [7,8]. In recent years, many studies have been carried out to identify new compounds from edible mushrooms with biological activity [5,9–12] and pharmacological and medicinal properties [7,13,14]. Among the most cultivated edible fungi, Pleurotus is a particularly interesting genus due to its nutraceutical properties linked to the polysaccharide fraction [15,16], such as 1-3-β-D-glucans (pleuromutilin and pleurane). Other bioactive compounds present in Pleurotus spp. are proteins, peptides and lectins (with high molecular weight), terpenes, fatty acid esters and polyphenols (with low molecular weight) [17]. Many studies on the health effects of Pleurotus species have been reported, such as immune-stimulating activity [18,19], antimicrobial and antiviral [20], hypoglycemic [21–24], anticancer [19,25],...
neuroprotective activities [26,27], hypocholesterolemic [28–31], antineoplastic, hypotensive, anti-inflammatory, antioxidant and hepatoprotective effects [32–34] and anti-allergic properties [35].

Numerous molecules/classes of compounds (proteins, peptides, polysaccharides, lipid derivatives, glycoproteins, etc.) with therapeutic properties have been isolated so far from fungi including *Pleurotus* spp. Among the polysaccharides, the most abundant are chitin, hemicellulose, β and α-glucans, mannans, xylans and galactans, and the compounds that stimulated the greatest interest are the high molecular weight polysaccharides. Especially β-glucans (β-(1,3)→(1,6)-glucans), which interact with the immune system to increase/decrease specific aspects of host response and are; therefore, called biological response modifiers (BRMs) [36,37]. The antiviral activity is also due to the strengthening of the host’s immune system and to the production of interferons under the stimulus of β-glucans, and is inhibited by fungal glycoproteins, protein-bound polysaccharide K (PSK), as well as by the β-glucans themselves [38,39].

In particular, *Pleurotus eryngii* (DC.) Quél., 1872, known as the “King Oyster” mushroom or “Cardoncello”, is considered to be a medicinal mushroom due to its content of high value proteins, vitamins and minerals and the low amount of carbohydrates and cholesterol [20]. A hemolysin called “Eryngeolysin” was isolated from *P. eryngii* and demonstrated to have particularly good cytotoxicity against leukemic cells and antibacterial activity [40]. “Cardoncello” also produces lovastatin (mevicolin, monacolin K), an important molecule that acts by competitively inhibiting the enzyme 3-hydroxy-3-methylglutaryl coenzyme A reductase (HMG-CoA) [41] and is one of the most potent drugs for lowering blood cholesterol employed so far.

Bioactive compounds are influenced by type of mushroom and some information regarding the chemical characterization of wild and cultivated mushrooms is available in literature. Nutritional composition and positive properties of wild mushrooms from Portugal [11], Khasi Hills of Meghalaya [42], northwest Spain [43] and southern Italy [44] have been reported. Furthermore, very recently, Obodai et al. [45], evaluated the antioxidant properties, composition in minerals, lipophilic and hydrophilic molecules of four cultivated (*Lentinus squarrosulus* strains SQW and LSF, *P. ostreatus* strain EM-1, *P. sajor-caju* strain PScW and *Auricularia auricula*) and two wild mushrooms (*P. tuber-regium* and *Termitomyces robustus*) from Ghana, and they found significant differences between wild and cultivated mushrooms, reporting their relevant nutritional aspect, high antioxidant potential, carbohydrates, proteins and macro- and micro-elements diverse content and generally low fat levels.

Although the glucans and phenolic composition of extracts from *Pleurotus* species has been quite well documented over the years [11,46–48], there are no studies on the comparison of bioactive substances between commercial and wild *P. eryngii* in the Basilicata region (Southern Italy). Therefore, the aim of this study was to determine in twelve *P. eryngii* isolates (seven commercial and five wild) the following parameters: (i) Total Polyphenol Content (TPC) and antioxidant activity in lyophilized fresh and dried samples and, (ii) total amount of β-glucans in fresh lyophilized samples, to point out fungal isolates with superior nutritional and medicinal value.

### 2. Materials and Methods

#### 2.1. Fungal Isolation and Culture

Twelve *P. eryngii* isolates, commercial and wild-type, originating from the Basilicata region (Southern Italy) were used in this study. The commercial isolates investigated were: ‘142 F’; ‘142 E’; ‘D+’; ‘V turbo’; ‘V 142’; ‘A 12’; ‘V160’ and the wild-type fungal isolates were: ‘Albino 1107’; ‘Altamura 1603’; ‘Muro Lucano 139’; ‘Conversano 1250’ and ‘Albino beige chiaro 1094’. Fungal isolates belonged to the Fungal Culture Collection (FCC) of the Mycology laboratory (University of Basilicata, Potenza, Italy) and were stored at 4 °C as pure cultures, in tubes containing potato dextrose agar (PDA) media. After revitalization on PDA added with 0.5 g L⁻¹ streptomycin sulfate, the Petri plates were incubated at
22 °C in the dark. Well-grown mycelia were inoculated on wooden sticks in glass cylinders and placed again in an incubator at 22 °C. Subsequently, standard substrate bags were inoculated with the wooden sticks and positioned in a greenhouse at 25 °C at about 90% RH. A randomized block experimental design with three replicates was applied, and each experimental unit included two substrate bags well colonized by mycelium [6]. Mushroom cultivation was performed following the standard technique usually used by farmers. Finally, qualitative analyses were carried out on P. eryngii samples of different isolates.

2.2. Qualitative Analysis

Before proceeding with the qualitative analyses, each sample (fresh/dry) was subjected to lyophilization and, subsequently, the extract was obtained from each dry residue. To obtain the extract, 20 mL of an 80% methanolic solution was added to 2 g of mushroom powder and the mixture was left to stir for 30 min at room temperature (RT). After, the sample was centrifuged, the supernatant was discarded, and the residual pellet was extracted again with another 20 mL of methanol solution. The liquid phases were then combined and reduced in volume (20 mL) by means of a Heidolph Hei Vap G1 rotary evaporator (BUCHI, Germany). Finally, the fungal extracts were used in several analyzes to identify: (a) The total polyphenol content (TPC); (b) the antioxidant activity; (c) the phenolic profile; and (d) the content of β-glucans.

2.2.1. Determination of Polyphenols by the Folin-Ciocâlteu Method

To determine the polyphenols, the spectrophotometry method with Folin-Ciocâlteu reagent [49–51] was used and the reagent consisted of an aqueous solution of phosphomolybdate and phosphotungstate. For these analyses, the filtrate obtained previously was diluted at 1:10. Fifty hundred µl of Folin-Ciocâlteu reagent was added to 1 mL of each sample and after 3 min 1 mL of 10% sodium carbonate (Na₂CO₃) was added to the mix in order allow the reaction of the compounds. Then, the total volume was brought to 10 mL with distilled water and the sample was kept for 1 h in the dark. After, the absorbance was measured using a UV spectrophotometer (Cary 60, Agilent Technologies, Santa Clara, CA, USA) at 765 nm wavelength. The results were expressed as the mean value of three repetitions ± standard error of mean (SEM).

2.2.2. Analysis of Phenolic Compounds

The determination of the phenolic profile was performed by high performance liquid chromatography (HPLC) analysis [52,53]. An InfinityLab Poroshell 120 EC-C18 chromatographic column, 4.6 × 150 mm, 4 µm (Agilent Technologies, Santa Clara, CA, USA) was used. For the elution, a binary mixture of solvents A and B was used, whose compositional and elution characteristics are reported below: Solvent A was 0.1% formic acid in Milli-Q water; Solvent B was 75% acetonitrile; flow rate was 1 mL min⁻¹; the temperature of the column was kept at 40 °C and the wavelength was 260 nm. The elution profile is shown in Table 1. To determine the analytes, the calibration straight lines were prepared using pure reference standards: ferulic acid, gallic acid, epicatechin (ECG) and epigallocatechallallate (EGCG).

| Time (min) | 0.1% Formic Acid (%) | 75% Acetonitrile (%) |
|-----------|----------------------|---------------------|
| 0.00      | 98.00                | 2.00                |
| 15.00     | 85.00                | 15.00               |
| 25.00     | 85.00                | 15.00               |
| 35.00     | 75.00                | 25.00               |
| 45.00     | 0.00                 | 100.00              |
| 50.00     | 0.00                 | 100.00              |
2.2.3. Antioxidant Activity

Among the various methods commonly used to determine antioxidant activity, based on the deactivation of a radical whose concentration is known, (ORAC, ABTS, DPPH, reduction of thiobarbituric acid, reduction of Fe, etc.), the ABTS method as described by Miller et al. [54] and usually employed to study *P. eryngii* antioxidant activity [55–57] was used in this study. This generates the radical starting from the oxidation of the ammonium salt of 2,2-azobis-(3-ethylbenzothiazolin-6-sulphonic acid-ABTS) by means of a solution of potassium persulphate $K_2S_2O_8$. To 5 mL of ABTS with 7 mM concentration, 88 $\mu$L of $K_2S_2O_8$, 140 mM, was added. Then, the solution was allowed to react for 16 h in the dark in order to obtain and stabilize the radical cation (ABTS••). To proceed with the analysis, the solution measured at a wavelength of 734 nm was diluted with ethanol up to an absorbance of 0.70 ± 0.02. When the radical solution, with its intense green-blue color, is put into contact with the active extracts, the antioxidant compounds present cause its discoloration, whose intensity reduction is then measured with a spectrophotometer at 734 nm. The determinations were made at a steady-state (tss) time of 30 min. This discoloration was proportional to the antioxidant charge present in the sample and was calculated as a percentage of absorbance decrease, called “inhibition percentage” (I%). This parameter was calculated using the following formula (1):

$$I(\%) = \frac{\text{Abs ABTS}}{\text{Abs ABTS-Abs sample}} \times 100$$

where: Abs ABTS = the absorbance of the ABTS and Abs = the absorbance of the sample. To determine the I(%) of each extract, four dilutions were prepared in ethanol, in a concentration range 0.4–20.4. For each solution, 100 $\mu$L was taken and 900 $\mu$L of ABTS•• was added. Then, the absorbance was measured at the tss and the I(%) calculated. A straight line was used to calculate the percentage of inhibition, from which it was possible to extrapolate the value of EC$_{50}$ (the value at which the inhibition of 50% of the concentration observed of the radical ABTS).

2.2.4. $\beta$-Glucans Extraction

For the extraction of $\beta$-glucans, the method of Lee et al. [58] with minor modifications was used. To 2 g of dry mushroom extract, 20 mL of a 2% NaOH solution was added and then heated at 90 °C for 5 h. Subsequently, the suspension was cooled and centrifuged at 3200 × g rpm at RT for 15 min. The alkaline supernatant was first neutralized with 2 M acetic acid, and then $\beta$-glucans was precipitated with 45 mL of ethanol. Proteins removal was achieved by re-dissolving the alcoholic extract in a 3% acetic acid solution followed by centrifugation at 4000 × g rpm for 20 min at RT. The supernatant was neutralized with 2 M NaOH and subsequently dried. Finally, the dry residue was used to determine the total $\beta$-glucans content.

Analysis of $\beta$-Glucans

For the determination of $\beta$-glucans, the specific kit of Megazyme International Ireland Ltd. was used, based on the McCleary and Codd [59] method, in which an enzymatic demolition of the $\beta$-glucan up to D-glucose was carried out, measured by a spectrophotometer after addition of reagents that develop the color, of which the absorbance was measured at 510 nm wavelength.

An aliquot of 200 mg of dry $\beta$-glucan extract was first heated at 40 °C for 1 h with 200 $\mu$L of lichenase enzyme. Then, 40 mL of double-distilled water was added to the suspension followed by centrifugation at 3000 × g rpm for 10 min. An aliquot of the extract (1 mL) was treated with the $\beta$-glucosidase enzyme and incubated at 40 °C for 15 min. Subsequently, the GOPOD reagent (a mixture of glucose-peroxidase and 4-aminoantipyrin), was added to the sample, and the colorimetric reaction was allowed to take place at 40 °C for 20 min. The absorbance reading was made at 510 nm wavelength against the blank
(acetate buffer solution with GOPOD). Quantitative determination was made using the pure \(\beta\)-glucan as a standard [59].

### 2.3. Statistical Analysis

Values of three observations were considered for each *P. eryngii* isolate, all the assays were carried out in triplicate and the results were expressed as mean values (±standard error of mean, SEM). The outcomes were evaluated by using the two-way analysis of variance (ANOVA) with type of isolates as one factor and status of isolates (dry and fresh) as the second factor. Means were separated by least significant differences (LSD) test with a probability level \((p \leq 0.05)\), using SAS software (SAS Institute, Cary, NC, USA, 2015) [60]. Water content, \(\beta\)-glucans and phenolic compounds, determined on only fresh samples, were subjected to a one-way ANOVA (types of isolates as main factor) using the same software mentioned earlier. An unpaired \(t\)-test (when the criteria of normality and equal variance are satisfied) was conducted to compare the means of all studied parameters between commercial and wild isolates. When the variances of the \(t\)-test samples were not equal, the means were compared with a Welch’s \(t\)-test analysis [61].

### 3. Results

The results of the ANOVA test on all studied parameters are reported in Table 2. A significant effect of the type of isolate was found for all parameters. No significant effects of status of the isolate on TPC, antioxidant activity and antiradical power were found, even if samples of fresh mushrooms, subjected to pre-lyophilization, showed a higher polyphenolic content compared to dried ones (Table 2). In addition, a high significant interactive effect between the type of isolate and the status of isolate was found for the TPC (Table 2). This parameter was determined in all *P. eryngii* samples and showed quite different amounts, which varied from 0.24 ± 0.01–5.34 ± 0.05 mg g\(^{-1}\). The maximum and minimum values were reached by fresh sample of the ‘V142’ commercial isolate and by dry sample of the ‘Albino beige chiaro 1094’ wild isolate, respectively (Table 3).

Table 2. Analysis of variance of different sources on all studied parameters.

| Source               | d.f. | Total Polyphenol Content | Antioxidant Activity (ABTS) | Antiradical Power (ARP) | Water Content | \(\beta\)-Glucans | Ferulic Acid | Gallic Acid | ECG | EGCG |
|----------------------|------|--------------------------|-----------------------------|-------------------------|---------------|------------------|--------------|-------------|-----|------|
| Type of isolate (T)  | 11   | *** 1                    | ***                         | ***                     | ***           | ***              | ***          | ***         | *** | ***  |
| Status of isolate (S)| 1    | NS                       | NS                          | NS                      | -             | -                | -            | -           | -   | -    |
| T \(\times\) S       | 11   | ***                      | *                           | *                       | *             | *                | *            | *           | *   | *    |

1 Significance level indicated by the asterisks: *** \(p < 0.001\), * \(p < 0.05\), NS—not significant.

Commercial isolates showed higher values of the TPC than wild ones \((p< 0.001\) for Welch’s \(t\)-test) (Table 3). Among commercial mushrooms, the lowest value was registered in the dry ‘V 160’ isolate. Regarding the wild mushrooms, the fresh ‘Altamura 1063’ and ‘Conversano 1250’ isolates showed the highest value of TPC (Table 3).

The average values of antioxidant activity (concentration required to obtain a 50% antioxidant effect, EC\(_{50}\)) and antiradical power (ARP values compared to 100 g of dehydrated product) ranged between 2.12 –73.25 ± 0.72 mg mL\(^{-1}\) and from 1.4 ± 0.03–53.5 ± 0.37%, respectively. In particular, fresh and dry samples of ‘Muro Lucano 139’ exhibited the best antioxidant potential, reaching values of 53.5 ± 0.37% and 46.9 ± 0.39%, respectively (Table 3). Among the commercial type isolates of *P. eryngii*, ‘V160’ obtained values similar to those recorded for ‘Muro Lucano 139’ (Table 3). Whereas the lowest values of ARP were reached by fresh and dry *P. eryngii* samples of ‘D+’ followed by ‘V142’, ‘142 E’ and ‘142 F’ isolates.
Table 3. Interactive effect of isolate type and status on polyphenolic content, antioxidant activity (EC50 of ABTS) and antiradical power (ARP) in twelve P. eryngii isolates from the Basilicata region (Southern Italy).

| Type of Isolate | Status of Isolate | Total Polyphenol Content (mg g−1) | Antioxidant Activity EC50 of ABTS (mg mL−1) | Antiradical Power (ARP) (%) |
|-----------------|-------------------|----------------------------------|------------------------------------------|---------------------------|
| Commercial      |                   |                                  |                                          |                           |
| ‘142 F’         | Dry               | 2.55 ± 0.02 i<sup>1</sup>         | 23.81 ± 0.00 h                          | 4.2 ± 0.06 hi             |
|                 | Fresh             | 3.35 ± 0.04 f                     | 28.37 ± 0.37 f                          | 3.5 ± 0.05 hi             |
| ‘142 E’         | Dry               | 2.85 ± 0.03 g                     | 25.77 ± 0.01 g                          | 3.9 ± 0.04 hi             |
|                 | Fresh             | 3.68 ± 0.03 e                     | 31.78 ± 0.35 e                          | 3.2 ± 0.06 hi             |
| ‘D+’            | Dry               | 2.70 ± 0.09 h                     | 60.53 ± 0.02 b                          | 1.7 ± 0.03 i              |
|                 | Fresh             | 3.56 ± 0.09 e                     | 73.25 ± 0.72 a                          | 1.4 ± 0.03 i              |
| ‘V turbo’       | Dry               | 3.30 ± 0.15 f                     | 7.67 ± 0.00 n                           | 13.0 ± 0.12 def           |
|                 | Fresh             | 4.39 ± 0.05 f                     | 9.82 ± 0.38 m                           | 10.2 ± 0.09 fg            |
| ‘V142’          | Dry               | 4.05 ± 0.02 d                     | 43.18 ± 0.01 d                          | 2.3 ± 0.05 hi             |
|                 | Fresh             | 5.34 ± 0.05 a                     | 52.44 ± 0.74 c                          | 1.9 ± 0.03 hi             |
| ‘A12’           | Dry               | 3.65 ± 0.02 e                     | 16.70 ± 0.00 j                          | 6.0 ± 0.08 gbi            |
|                 | Fresh             | 4.91 ± 0.05 b                     | 20.35 ± 0.40 i                          | 4.9 ± 0.11 hi             |
| ‘V160’          | Dry               | 0.40 ± 0.01 op                    | 2.12 ± 0.00 q                           | 47.2 ± 0.38 b             |
|                 | Fresh             | 0.53 ± 0.03 mno                   | 2.63 ± 0.08 q                           | 38.0 ± 0.33 c             |
| Wild            |                   |                                  |                                          |                           |
| ‘Albino 1107’   | Dry               | 0.50 ± 0.00 no                    | 6.42 ± 0.00 op                          | 15.6 ± 0.23 de            |
|                 | Fresh             | 0.64 ± 0.01 lm                    | 6.01 ± 0.18 p                           | 16.6 ± 0.16 d             |
| ‘Altamura 1603’ | Dry               | 0.86 ± 0.01 k                     | 15.40 ± 0.00 k                          | 6.3 ± 0.13 ghi            |
|                 | Fresh             | 1.14 ± 0.03 j                     | 14.25 ± 0.35 l                          | 7.0 ± 0.14 gh             |
| ‘Muro Lucano 139’| Dry              | 0.50 ± 0.01 no                    | 2.13 ± 0.01 q                           | 46.9 ± 0.39 b             |
|                 | Fresh             | 0.62 ± 0.03 l m                   | 1.87 ± 0.14 q                           | 53.5 ± 0.37 a             |
| ‘Conversano 1250’| Dry             | 0.72 ± 0.01 l                     | 7.74 ± 0.00 n                           | 12.9 ± 0.17 def           |
|                 | Fresh             | 0.97 ± 0.04 k                     | 7.13 ± 0.06 no                          | 14.0 ± 0.18 def           |
| ‘Albino beige chiaro 1094’ | Dry         | 0.24 ± 0.01 q                     | 9.67 ± 0.00 m                           | 10.4 ± 0.15 efg           |
|                 | Fresh             | 0.33 ± 0.02 pq                    | 9.06 ± 0.12 m                           | 11.0 ± 0.17 efg           |

Table 4 shows the effects of type of isolate on water content, β-glucans and phenolic compounds, which were determined only in fresh samples. The wild P. eryngii isolates were characterized by a higher content of water (+3.4%) compared to commercial ones. In detail, ‘Albino beige chiaro 1094’ and ‘Albino 1107’ isolates of P. eryngii were the richest in water (Table 4).

The outcomes of β-glucans analysis emphasized the richness of these compounds within the matrix of different P. eryngii isolates. However, it was observed that the wild-type P. eryngii isolates had a higher β-glucan content compared to the commercial ones. The highest content of β-glucans was obtained by the ‘Albino beige chiaro 1094’ wild isolate followed by the ‘Muro Lucano 139’ one, while the lowest value was reached by the ‘V160’ commercial isolate (Table 4).

The phenolic profile results showed that the main phenols present were ferulic acid, gallic acid, epicatechingallate (ECG) and epigallocatechigallate (EGCG). The presence of gallic acid and EGCG, founded in all isolates both commercial and wild, could; therefore, motivate the important antioxidant properties exhibited by the P. eryngii isolates studied.

In general, as shown in Table 4, the concentration of the investigated phenolic compounds was superior in the P. eryngii commercial isolates compared to the wild-type ones. Besides, considerable variations were observed between all the P. eryngii isolates investigated. Ferulic acid, gallic acid, ECG and EGCG varied from 0.08 ± 0.03–1.31 ± 0.03 mg g−1,
from 0.14 ± 0.05–2.43 ± 0.05 mg g\(^{-1}\), from 0.04 ± 0.02–0.60 ± 0.03 mg g\(^{-1}\), and from 0.04 ± 0.01–0.83 ± 0.05 mg g\(^{-1}\), respectively. The ‘V 142’ commercial isolate recorded the highest values of ferulic acid, gallic acid, epicatechin and epigallocatechigallate. ‘Albino beige chiaro 1094’ wild-type isolate of *P. eryngii* resulted as being the poorest in phenolic compounds but the richest in \(\beta\)-glucans and water content (Table 4).

Table 4. Effect of isolate type on water content, \(\beta\)-glucans and phenolic compounds in twelve *P. eryngii* isolates from the Basilicata region (Southern Italy).

| Type of Isolate       | Water Content (%) | \(\beta\)-Glucans (g 100 g\(^{-1}\)) | Ferulic Acid (mg g\(^{-1}\)) | Gallic Acid (mg g\(^{-1}\)) | ECG (mg g\(^{-1}\)) | EGCG (mg g\(^{-1}\)) |
|-----------------------|-------------------|------------------------------------|-------------------------------|-------------------------------|---------------------|---------------------|
| **Commercial**        |                   |                                    |                               |                               |                     |                     |
| ‘142F’                | 83.2 ± 0.11 f \(^1\)  | 23.4 ± 0.72 d                      | 0.77 ± 0.04 c                 | 1.32 ± 0.02 c                 | 0.34 ± 0.05 cd      | 0.42 ± 0.09 c       |
| ‘142 E’               | 85.2 ± 0.06 d      | 21.6 ± 0.40 de                     | 0.87 ± 0.05 c                 | 1.54 ± 0.01 bc                | 0.47 ± 0.06 abc     | 0.54 ± 0.06 bc      |
| ‘D+’                  | 85.6 ± 0.17 cd     | 19.3 ± 0.70 fg                     | 0.86 ± 0.03 c                 | 1.41 ± 0.03 c                 | 0.27 ± 0.05 de      | 0.54 ± 0.05 bc      |
| ‘V turbo’             | 81.9 ± 0.23 g      | 20.8 ± 0.77 ef                     | 1.04 ± 0.07 b                 | 1.75 ± 0.03 b                 | 0.45 ± 0.07 bc      | 0.52 ± 0.10 bc      |
| ‘V142’                | 82.3 ± 0.17 g      | 18.5 ± 0.86 g                      | 1.31 ± 0.03 a                 | 2.43 ± 0.05 a                 | 0.54 ± 0.02 ab      | 0.83 ± 0.05 a       |
| ‘A12’                 | 83.1 ± 0.12 f      | 22.6 ± 0.63 de                     | 1.20 ± 0.08 a                 | 2.14 ± 0.33 a                 | 0.60 ± 0.03 a       | 0.67 ± 0.09 ab      |
| ‘V160’                | 84.7 ± 0.06 e      | 12.8 ± 0.43 h                      | 0.10 ± 0.03 de                | 0.14 ± 0.05 e                 | 0.07 ± 0.03 f       | 0.08 ± 0.01 d       |
| **Wild**              |                   |                                    |                               |                               |                     |                     |
| ‘Albino 1107’         | 87.4 ± 0.11 a      | 29.7 ± 0.82 c                      | 0.12 ± 0.03 de                | 0.26 ± 0.08 de                | 0.07 ± 0.04 f       | 0.11 ± 0.05 d       |
| ‘Altamura 1603’       | 85.8 ± 0.17 c      | 31.6 ± 0.53 bc                     | 0.12 ± 0.05 de                | 0.25 ± 0.04 de                | 0.08 ± 0.01 f       | 0.09 ± 0.03 d       |
| ‘Muro Lucano 139’     | 85.7 ± 0.06 c      | 32.5 ± 0.84 ab                     | 0.20 ± 0.05 de                | 0.51 ± 0.06 d                 | 0.11 ± 0.05 f       | 0.16 ± 0.06 d       |
| ‘Conversano 1250’     | 86.6 ± 0.12 b      | 21.3 ± 0.60 e                      | 0.23 ± 0.03 d                 | 0.41 ± 0.06 de                | 0.16 ± 0.05 ef      | 0.13 ± 0.04 d       |
| ‘Albino beige chiaro 1094’ | 87.5 ± 0.17 a      | 33.8 ± 0.70 a                      | 0.08 ± 0.03 e                 | 0.15 ± 0.07 e                 | 0.04 ± 0.02 f       | 0.04 ± 0.00 d       |

\(^1\) Each value expressed as mean ± SE (\(n = 3\) for type of isolate effect; \(n = 21\) for commercial isolates; \(n = 15\) for wild isolates) in columns not sharing the same letters are significantly different according to least significant difference test (\(p \leq 0.05\)).\(^2\) **, significance at \(p < 0.01\);

ECG = epicatechin; EGCG = epigallocatechillallate.

### 4. Discussion

Edible mushrooms (EMs) are important sources of nutrients for the human diet, being utilized worldwide, and contain many chemical compounds with several pharmacological, medicinal, cosmetic and biocontrol properties [5,13,16,62]. Results from our study, on various *P. eryngii* isolates, both wild and cultivated types, from the Basilicata region, surely added further knowledge on this fungi and also generally agree with the outcomes of several studies previously performed on EMs. Such as the earlier work of Cheung et al. [63], who analyzed the antioxidant activity and the total phenols in two EMs Shitake and straw mushrooms and showed that they may have potential as antioxidants. In addition, the present work furnished similar results as the studies of Oke and Aslim [64], who examined the phenolic compositions and antioxidative, protective and cytotoxic effects of *P. eryngii* and *Auricularia auricula-judae* and revealed that this fungi are able, based on their compounds, to protect against oxidative cell damage and can be used as a potential source of natural antioxidants. Furthermore, the results very much agree with the very recent studies of Bahadori et al. [65] who investigated the metal concentration, phenols and antioxidant activity in two negligible wild mushrooms *Melanoleuca cognata* and *M. stridula* and demonstrated that these types of EMs can have good nutritional value and also provide important health benefits.

Particularly, new and useful knowledge of the medicinal value of the *P. eryngii* mushroom has arisen from the outcomes of this study, showing that the polyphenols reached, on average, higher concentrations in the commercial isolates. Since the phenolic content has always been higher in the fresh mushroom samples compared to the dry ones, it can
be hypothesized that the heat could determine a partial degradation/loss of the phenolic compounds during the drying process. From the analysis of the phenolic profile it has emerged that ferulic acid, gallic acid, epicatechingallate (ECG) and epigallocatechingallate (EGCG) are the most present compounds. Of particular interest is the presence of gallic acid and EGCG, which confirms the remarkable antioxidant properties of all *P. eryngii* isolates studied. Especially, the wild-type ‘Muro Lucano 139’, ‘Albino 1107’ and ‘Conversano 1250’ *P. eryngii* isolates showed a remarkably high antioxidant activity.

Apart from polyphenols, other important functional components had been isolated from EMs, like β-glucans, which have demonstrated them to have hypocholesterolemic, antidiabetic and anticarcinogenic properties as well as positive effects on the immune system [66–68]. The mushroom species most rich in β-glucan were found to be *Lentinus edodes* and some members of the *Pleurotus* genus, such as *P. ostreatus* and *P. eryngii* [69].

*P. eryngii* contain, in general, 85–90% moisture, 11–12 g\_100 g\(^{-1}\) dried mushroom protein content, 39–40 g\_100 g\(^{-1}\) carbohydrates, 7–8 g\_100 g\(^{-1}\) lipids, 28–29 g\_100 g\(^{-1}\) fibers and 4–5 g\_100 g\(^{-1}\) minerals [70]. The water content of the *P. eryngii* isolates from this study was similar compared to what has been reported by other authors [70,71] and also match with the outcomes of the investigation of Gezer et al. [72], which showed that wild mushrooms had a low dry matter content but a high value of water, fiber and beta-glucans content.

Moreover, the present work integrates well among the only few studies reported so far that dealt with comparisons of bioactive compounds (mainly glucans) between cultivated and wild mushrooms. For example, the work of Sari et al. [71], who showed that many wild species, like *Trametes versicolor* and *Boletus edulis*, had a higher β-glucans content and also revealed that cultivated mushrooms had a range of β-glucans content between 8.60–26.93 g\_100 g\(^{-1}\), while *P. eryngii* had a β-glucans value of 15.32 g\_100 g\(^{-1}\) [71]. In addition, the work of Butkup et al. [73], who evaluated the β-glucan content of twenty-five wild edible mushrooms from Northeastern Thailand, showed that *Volvariella volvacea*, a gilled mushroom, reached the highest content (34.4 g\_100 g\(^{-1}\)), while *Alpova trappi*, a puffball mushroom, obtained the lowest one (0.62 g\_100 g\(^{-1}\)). In the present study, the β-glucans content, known for its hypocholesterolemic activity, was also of great significance and its concentration was, on average, elevated in wild-type isolates, being the highest for ‘Albino beige chiaro 1094’ than the commercial ones.

Research on wild edible fungi (WEF) compared to commercial types were earlier performed by Barros et al. [11], who showed that the wild mushroom species contained higher amounts of protein and lower fat concentrations. Besides, the α-Tocopherol was detected in higher amounts in the wild species along with a higher content of phenols, but a lower content of ascorbic acid, than commercial mushrooms. Wang et al. [74] indicated that wild mushrooms accumulate more macronutrients and micronutrients and they are rich mainly in potassium, phosphorus, calcium, magnesium, sodium and iron. Similarly, in our study an elevated bioactive substances concentration was found especially in the WEF investigated, identical to what was reported by Palazzolo et al. [44], who analyzed five WEF collected from Sicily (*Fistulina hepatica*, *Infundibulicybe geotropa*, *Laetiporus sulphureus*, *Macrolepiota procera* var. *procera* and *Suillus granulatus*) and reported a high content of vitamin B12 and riboflavin presence. These latter results suggested that the WEF could be very valuable natural products and; therefore, including them in many diets is certainly highly beneficial [44,75–77].

Some investigations focusing on *P. eryngii* and *P. nebrodensis* (Quél., 1886) furnished more information about these highly appreciated mushrooms and revealed their important nutritional and medicinal value [78–80]. Thus, further studies examining some *P. nebrodensis* isolates along with a few more of the *P. eryngii* ones could be useful and; therefore, will be planned.
5. Conclusions

The “Cardoncello” mushroom, an excellent product in gastronomy, is considered a food of high nutraceutical value, thanks to its strong antioxidant properties and elevated content in bioactive compounds, such as total phenolic and beta-glucan content.

On the basis of the results from this study, it can be concluded that type of mushrooms (commercial and wild) influenced both the phenolic compounds and β-glucans content. In particular, cultivated P. eryngii isolates showed a higher content of TPC, ferulic acid, gallic acid, epicatechingallate and epigallocatechigallate, while wild mushrooms were characterized by a superior content of β-glucans. This latter outcome suggested that wild P. eryngii isolates represent a very good source of beneficial compounds to human health.

However, further investigations, taking into account a greater number of “Cardoncello” isolates, are needed to generalize better and apply the results of the present study.

The results from future studies on WEF can furnish important and useful data for the production of novel active foods with high nutritional value and wide and diverse medicinal applications in mycotherapy.

Author Contributions: Conceptualization: V.C., I.C., S.M.M., L.C.; methodology: A.C.; software: L.C., D.C.; validation: all authors contributed equally to the validation of the work; formal analysis: A.B.; investigation: A.B., A.C., V.C., S.M.M., L.C.; data curation: V.C., A.C., I.C., S.M.M., L.C.; writing—original draft preparation: S.M.M., A.C., V.C., I.C.; writing—review and editing: S.M.M., V.C., I.C., A.C., L.C., D.C.; supervision: I.C., V.C., A.C. All authors have read and agreed to the published version of the manuscript.

Funding: This research received no external funding.

Institutional Review Board Statement: Not applicable.

Informed Consent Statement: Not applicable.

Data Availability Statement: The data presented in this study are available on request from the corresponding author.

Acknowledgments: The authors thank Donato Aiace of Cardopan Company, Tolve—Potenza (Italy) for his contribution to the research by providing the commercial “Cardoncello” isolates.

Conflicts of Interest: The authors declare no conflict of interest.

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