Bio-enriched Pleurotus mushrooms for deficiency control and improved antioxidative protection of human platelets?

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Abstract The study investigated effect of substrate supplementation with Se alone or in combination with Cu or/and Zn Se on (1) the growth of Pleurotus ostreatus and Pleurotus eryngii; (2) elements accumulation in mushrooms; (3) the antioxidant activities of bio-enriched mushroom extracts in human platelets. The accumulation of elements generally increased over concentration gradient reaching its maximum at 1.2 mM for P. ostreatus and P. eryngii: (1) over 100 and 80 mg kg\(^{-1}\) of Se, respectively (Se supplementation); (2) over 15 and 30 mg kg\(^{-1}\) of Cu, respectively (Se+Cu); (3) over 30 and 85 mg kg\(^{-1}\) of Zn, respectively. Se was predominantly accumulated as an organic fraction. Contrary to P. eryngii, the P. ostreatus biomass decreased with substrate elements concentration but was satisfactory up to 0.9 mM of Se, Se+Cu and Se+Zn. The Se+Cu+Zn model yielded low biomass and elements accumulation.

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Extracts from mushrooms bio-enriched with Se and Se+Zn (0.6–1.2 mM) revealed significant antioxidant activities in human platelets by ameliorating reactive oxygen species (ROS) and preventing lipid peroxidation. The study demonstrated the potential application of Pleurotus mushrooms as functional food products bio-enriched with essential elements. ROS inhibition by extracts of these mushrooms may be useful in control of platelets activation cascade.

Keywords Pleurotus · Bio-enrichment · Selenium · Copper · Zinc · Human platelets

Introduction

Mushrooms are increasingly popular as foodstuffs in different world locations owing to their nutritional value and potential medicinal use, extensively evidenced over recent decades. Mushrooms belonging to the genus Pleurotus (higher Basidiomycetes) are, next to Agaricus sp., one of the most important cultivated species worldwide [1] and account for nearly 27% of global production [2, 3]. Considered as a delicacy, they are rich in B vitamins, polysaccharides and minerals, e.g. potassium, phosphorus, calcium, magnesium and iron [4]. Various pharmacological effects exerted by compounds of Pleurotus, particularly derived from P. ostreatus, have been reported and include antiviral antibacterial, antidiabetic, antihypercholesterolic and antiarthritic activities [5–8]. In all, it appears that Pleurotus mushrooms have particularly high potential as functional foods. Economically-wise, this is also supported by their short growth time, the inexpensive and readily available materials required for their cultivation (e.g. wheat, rice straw, cotton waste or sawdust) and their documented resistance to pathogens and pests [9].
Bio-enrichment (bio-fortification) of food is an interesting strategy for delivering nutrients and preventing deficiencies [10, 11], and it may be attractive for consumers as it applies products of natural origin, increasingly preferred over synthetic food supplements [12]. However, most attention in this regard is focused on plant-based products [13]. Mushrooms, on the other hand, also have great potential for such use if one considers their ability to uptake and accumulate various elements in their fruiting bodies efficiently [14, 15], and their already established place in various national cuisines [16]. Some previous studies have already demonstrated that Agaricus bisporus and Ganoderma lucidum grown on substrates intentionally supplemented with trace elements may reveal significantly increased nutritional value [17, 18].

Selenium (Se), zinc (Zn) and copper (Cu) are all known to play pivotal metabolic roles in various biological processes, and their altered status may exert numerous adverse effects in humans. A decreased intake of Se can lead, inter alia, to hair loss, blotchy nails, diarrhoea, garlic breath, gastrointestinal issues and decreased thyroid function [19, 20]. Zn deficiency, on the other hand, has been related to immunosuppression, retardation of physical growth and cognitive function as well as various dermatological problems [21]. Epidemiologically, the worldwide frequency of Se and Cu deficiency is not high but can be observed in regions characterized by extremely low content of these elements in the soil and subsequently plant products [22, 23]. This is particularly the case in some parts of Asia, a continent with a long tradition of mushrooms use, both as food and natural medicines [24]. Zn deficiency is becoming a worldwide problem with approximately 17% of the human population being at risk [25]. It is therefore essential to maintain the optimal status of these trace metals in humans in order to avoid any manifestation of the above-mentioned harmful effects and to support normal human growth, development and function.

Mushrooms bio-fortified with certain elements may also exert novel properties, e.g. their enrichment with lithium may have a potential use as a medical nutrition therapy in mood modification [26] while enrichment with selenium (Se) enhances the antitumor activities of mushroom polysaccharides [27] and significant immunomodulatory properties of proteins [28]. It has also been shown that P. ostreatus and P. eryngii enriched simultaneously with Se and Zn reveal a greater content of phenolic compounds and that methanolic extracts of these mushrooms exhibited an increased scavenging of radicals in assay with 2,2-diphenyl-1-picrylhydrazyl (DPPH) [29]. This would indicate that bio-fortified mushrooms may have greater antioxidative properties, yet it remained unclear whether they could have any use in the amelioration of oxidative stress in human cells.

The aim of the present study was to investigate whether two edible mushroom species of worldwide popularity, P. ostreatus and P. eryngii, can be used in bio-enrichment with Se, Cu, Zn in a single-, double- and triple-element cultivation model, and whether such fortified mushrooms may exert some beneficial antioxidative effects. Specifically, the preventive effect of hot water extracts of fruiting bodies on tert-butyl hydroperoxide (tBHP)-induced oxidative stress in human platelets isolated from healthy donors. It has been evidenced that reactive oxygen species (ROS) play a role in triggering the activation cascade in platelets that involves tethering to the endothelium, rolling adhesion, aggregation and eventually formation of thrombus [30]. Therefore, any antiplatelet effect of bio-enriched mushrooms could play a potentially positive role in the prevention of cardiovascular complications.

Materials and methods

Experimental design

The mushrooms of two species P. ostreatus and P. eryngii were grown on substrates supplemented in four different configurations of trace elements: (1) Se; (2) Se+Cu; (3) Se+Zn and (4) Se+Cu+Zn, and final concentrations of 0.1, 0.3, 0.6, 0.9 and 1.2 mM. The control was constituted of mushrooms grown without the addition of any element. The yield included fruiting bodies harvested successively as they matured. Collected material was subjected to analytical determinations of Se, Cu and Zn content in the fruiting bodies. Additionally, speciation analyses were applied to distinguish between the inorganic and organic Se fraction due to distinctive differences in their biological activity. These experiments were designed to choose those cultivation models at which the most efficient bioaccumulation and growth of fruiting bodies was observed. The fruiting bodies from the chosen model were then extracted at high temperature, and extracts were later used to test as regards their ability to attenuate tBHP-induced generation of ROS and lipid peroxidation in human platelets isolated from healthy donors.

Mushroom cultivation and harvesting

All experimental cultivations were performed in three independent replicates. The substrate for P. ostreatus was prepared from wheat straw cut into chaff 4–5 cm long. The straw for the experiment was moistened to a moisture content of 60% using distilled water and then pasteurized at 60 °C for 24 h. The following salts were used in the experiment: sodium selenite [Na2SeO3(IV)], sodium selenate [Na2SeO4(VI)] by Acros Organics (USA), copper(II) selenate (CuSeO4·2H2O) and zinc nitrate hexahydrate [Zn(NO3)2·6H2O] by Sigma-Aldrich. These salts were dissolved in such an amount of sterile distilled water sufficient to obtain their appropriate concentration (0.1, 0.3, 0.6, 0.9
and 1.2 mM for all elements) in the substrate after which they were added to the substrate. Following the addition of the salt solution, the substrate water content reached 70%. The substrate was mixed with grain spawn (on wheat grain) which constituted 3% in relation to the wet weight of the substrate and placed in bags of perforated foil. Each bag contained 1 kg of the substrate. Mycelium incubation was conducted at a temperature of 25 °C and relative humidity of air at 85–90%. Once the substrate was totally covered with the mycelium, it was transferred to the cultivation chamber in which the temperature was maintained at 15–16 °C and relative humidity of air at 85–90%. The cultivation was additionally irradiated with fluorescent light of 500 lx intensity for 10 h a day. The growth facility was aerated in such a way as to maintain CO₂ concentration below 1000 ppm.

The substrate for *P. eryngii* was prepared from a mixture of beech sawdust and flax shives (3:1 vol.) which was additionally supplemented with wheat bran to the amount of 20%, corn flour 5% and gypsum 1% in relation to the substrate dry matter. The mixture was moistened with distilled water to a moisture content of 45%. The substrate prepared as described above was placed in PP bags and sterilized at a temperature of 121 °C for 1 h and was then cooled down to a temperature of 25 °C. Cu, Se and Zn salts were dissolved in sterile water, and the solution was added to the substrate with intensive blending using a POLYMIX PX-SR 90 D stirrer (Kinematica AG, Switzerland) in an amount that allowed their appropriate concentration in the substrate and, at the same time, making sure that the substrate water content reached 60%. The same concentrations of Se and Zn salts were added in the substrate as for *P. ostreatus*. The substrates with Se and Zn addition were mixed with grain spawn (on wheat grain) of the examined mushroom species (5% of substrate weight) and placed in polypropylene bottles of 1 L volume. Each bottle was filled with 350 g of the substrate and closed with a cover consisting of a cellulose filter type 338 with a typical retention of 12–15 µm of the substrate and closed with a cover consisting of a cellulose filter type 338 with a typical retention of 12–15 µm. The fruiting bodies were then weighed, dried using an electric drier SLW 53 STD (Pol-Eko, Poland) at 50 ± 2 °C for 48 h, weighed again for analysis of dry weight and ground in a Cutting Boll Mill 200 (Retsch GmbH, Germany) for 1 min.

**Determination of trace elements**

The powdered mushroom samples were sieved through a 0.02 mm sieve. 1.000 ± 0.001 g of sample was extracted by 1 mol L⁻¹ phosphoric acid in an ultrasonic bath at an ambient temperature (30 min). Samples were then filtered through a paper filter (washed by 200 mL of water and 20 mL of phosphoric buffer). Sample pH was adjusted at 6.0–6.5 by the addition of 10 mol L⁻¹ sodium hydroxide solution, and finally the samples were diluted to 20.0 mL by a phosphate buffer.

For Cu and Zn determination, flame atomic absorption spectrometry (FAAS) SpectraAA 22FS (Varian, Australia) was applied with stoichiometric flame acetylene (2.0 L min⁻¹) and air (13.5 L min⁻¹). Hollow cathode lamps (Varian) were used with the following parameters for Cu: wavelength 324.8 nm, slit 0.5, lamp current 10 mA and for Zn: wavelength 213.9 nm, slit 1.0, lamp current 5 mA, both with background correction with a deuterium lamp. The determination limits were 0.1 mg kg⁻¹ with uncertainty about 5.0% (measured as RSD) for both elements determined.

For Se determination, electrothermal atomic absorption spectrometry (ETAAS) SpectraAA 280Z (Agilent, USA) with Zeeman background correction was applied. A selenium hollow cathode lamp (wavelength 196.0 nm, slit 1.0 nm, current 10 mA) was used, and a temperature programme was optimized as follows: drying 85–120 °C during 55 s; ashing 1000 °C during 8 s; atomization 2600 °C. Pyrolytic graphite tubes and palladium solution as a chemical modifier (10 µL of 500 mg L⁻¹ for 20 µL of sample) were used. The limit of detection 0.01 mg kg⁻¹ and the uncertainty of results (measured as RSD) at the level of 5.0% were obtained.

In the case of Se species analysis, hyphenated high-performance liquid chromatography with hydride generation atomic absorption spectrometry detection (HPLC-HG-AAS) systems was applied as described previously in detail [31]. The hyphenated analytical system consisted of a Shimadzu liquid chromatograph (LC-10A) equipped with an HPLC pump (LC-10AT), a vacuum degasser unit (GT-104), Rheodyne PEEK valve (IDEX, USA) and an anion-exchange column Supelco LC-SAX1 (250 mm, 4.6 mm i.d., resin particle size 5 µm) thermostatted by a column oven (CTO-10ASvp). The chromatographic run was isocratic at 3 mL min⁻¹ with an injection volume of 200 µL. The measurements were performed with a Model SpectraAA 220FS spectrometer (Varian, Australia) equipped with an UltrAA selenium hollow cathode lamp.
The selenium species Se(VI) does not form volatile hydride, so in order to receive an analytical signal in the HG-AAS system it was necessary to perform a preliminary reduction of Se(VI) to Se(IV). The reduction was carried out on-line by heating the sample (90–100 °C) with the reducing agent: 0.5 mol L$^{-1}$ thiourea solution in 10 mol L$^{-1}$ hydrochloric acid (Kozak 2012). The eluate from the chromatographic column was joined to the stream of the reducing agent (flow rate 1 mL min$^{-1}$ from the peristaltic pump) through a T-shape coupling and directed to the Tygon capillary loop (inner diameter 0.82 mm) heated in a water bath. The capillary loop outlet was connected to the hydride generation system. For both analytical systems, PEEK transfer tubing of the eluent from the LC column to the hydride generation unit was inserted into a Tygon sleeve. The continuous hydride generation system (VGA-77, Varian) consisted of a manually controlled, four-channel peristaltic pump with Tygon tubing (0.6 mm i.d.), one reaction coil (PTFE tubing 0.8 mm i.d., 75 cm length) and three-way connectors. The gas–liquid separator was made of glass, and the interior dead volume was 3 mL. For the atomization of the selenium hydrides (detected at 196.0 nm), a heating controller, an electrothermally heating mantle and a quartz tube (ETC-60, Varian) heated to 900 °C were used.

A number of validation parameters characterising the analytical method were determined. The limits of detection 0.01 mg kg$^{-1}$ for both Se(IV) and Se(VI) and the uncertainty of results (measured as RSD) at the level of 10% for both selenium forms were obtained. As it was impossible to estimate the measurement traceability owing to the lack of any certified reference materials for determination of inorganic selenium species, the recovery of each selenium species was measured upon the addition of a standard to the sample. A recovery of 96–105% of each species was considered as satisfactory.

**Mushroom extract preparation**

Homogenized samples of fruiting bodies were extracted in distilled water at the ratio of 1:10 (w/v) at 95 °C for 30 min. The samples were then cooled to room temperature and centrifuged, and supernatants were sterilized by filtration on 0.22 μm syringeless filter devices (Roth). The extracts were kept at −20 °C until use.

**Human platelet isolation**

Human platelets were isolated from whole blood samples (6 mL) collected in acid citrate dextrose (ACD) from three healthy donors (screened by physical examination, medical history and initial blood tests), with normal platelet count (150,000–450,000 cells μL$^{-1}$). The donors (aged 19–22 years old; three females) were non-smoking and normal weighted (BMI 18.5–24.9). Samples were collected at the Regional Centre of Blood and Blood Treatment in Poznan, Poland, according to accepted safeguard standards and legal requirements. Within 20 min of collection, platelet-rich plasma (PRP) was obtained by centrifugation at 200 g for 12 min. PRP was transferred into a polypropylene tube, and 1/10 volume ACD and 100 ng mL$^{-1}$ prostaglandin E1 was added to prevent platelet activation during isolation. PRP was centrifuged at 900g for 15 min, and plasma was aspirated. Platelets were then suspended in 6 mL of Hepes–NaCl$_2$ buffer (10 mM Hepes, 0.85% NaCl, pH 7.4) and layered on a discontinuous 10–17% iodixanol gradient in Hepes/NaCl buffer. The gradient was centrifuged at 300g for 20 min, and the platelet fraction was collected. Platelets were centrifuged at 900g for 15 min, and the resulting pellet was washed and resuspended with Ca$^{2+}$-free Tyrode–Hepes buffer (137 mM NaCl, 0.3 mM NaH$_2$PO$_4$, 3.5 mM Hepes, 5.5 mM [β]-glucose, pH 7.35).

**Intracellular reactive oxygen species assay**

Isolated platelets were loaded for 30 min at 37 °C in darkness with 20 μM of 2′,7′-dichlorofluorescin diacetate (DCFDA; Abcam, UK), a fluorogenic dye that measures hydroxyl, peroxyl and other reactive oxygen species (ROS) activity within the cell. Cells were then washed, pretreated with 10 μL mushroom extracts for 30 min, washed again, seeded in black clear bottom 96-well plate and exposed to 10 μM tBHP (Abcam, UK) for another 60 min. The negative/positive controls consisted of DCFDA-loaded platelets pretreated with 10 μL of Tyrode–Hepes buffer and not exposed/exposed to 10 μM tBHP. Fluorescence of DCFDA in all samples was measured kinetically after 15, 30 and 60 min of incubation using a Synergy HTX multi-mode plate reader (BioTek, USA) at an excitation of 495 nm and emission of 528 nm. The final results were presented as a percentage of the negative control.

**Lipid peroxidation assay**

Lipid peroxidation was analysed using a Lipid Peroxidation (MDA) Colorimetric/Fluorometric Assay Kit (BioVision, UK) by means of malondialdehyde (MDA) equivalents. Isolated platelets were pretreated with 10 μL mushroom extracts for 30 min, washed and exposed to 10 μM tBHP for another 60 min. The negative/positive controls constituted of platelets pretreated with 10 μL of Tyrode–Hepes buffer and not exposed/exposed to 10 μM tBHP. After the experiments, cells were harvested from each well and homogenized on ice in 300 μL of provided lysis buffer and centrifuged
to remove insoluble material. The resulting 200 μL of supernatants was transferred to a microcentrifuge tube and supplemented with 600 μL of thiobarbituric acid (TBA) to generate an MDA–TBA adduct. To accelerate the reaction, samples were incubated at 95 °C for 60 min and the final product was measured colorimetrically at 532 nm. The calculated values were compared to a calibration curve prepared using MDA standard (BioVision, UK). The coefficient of variation ($r^2$) for the calibration curve was 0.99. The final results were presented as a percentage of the negative control.

**Statistical analysis and calculations**

The results were analysed using STATISTICA 10.0 software (StatSoft, USA). Because the element accumulation and mushroom biomass data met the assumption on Gaussian distribution (analysed with the Shapiro–Wilk test), comparison of element accumulation and yielded biomass between cultivation models was assessed with multivariate analyses of variance (MANOVA) with the Tukey HSD method as a post hoc test. Since the data of platelet ROS concentration and lipid peroxidation had no normal distribution, the Wilcoxon signed-rank test was employed to compare the studied samples with the positive control. $p < 0.05$ was considered as statistically significant.

**Results and discussion**

**Accumulation of trace elements in bio-enriched mushrooms**

The addition of Se to the growth substrate increased its accumulation in the fruiting bodies of both investigated species in a concentration-dependent manner. Greater concentrations were, however, found for *P. ostreatus*. Nevertheless, Se concentrations were 40-fold higher compared to the control once the substrate overgrown by both species was supplemented with 1.2 mM of Se. It is worth noting that an increase in Se content was over 10-fold for *P. ostreatus* and 20-fold for *P. eryngii* when the lowest assayed concentration, 0.1 mM, was added (Fig. 1). For both species, organic Se had the greatest share for nearly all the studied substrate concentrations (Table 1). *P. ostreatus*, however, revealed higher concentrations, reaching maximally over 100 mg kg$^{-1}$ after the substrate had been supplemented with 1.2 mM of Se (Table 1). These findings are in line with previous studies demonstrating the potential of various mushrooms from the *Pleurotus* genus in the production of food bio-fortified with Se [31, 32].

Compared to mushrooms cultivated on substrates enriched only with Se, the simultaneous addition of Se and Cu resulted in decreased Se accumulation in both *Pleurotus* species although this effect was more evident for *P. eryngii*. For this species, Se concentrations did not exceed 20 mg kg$^{-1}$ regardless of the initial substrate content of trace elements. To compare, for *P. ostreatus* the minimum Se level did not fall below 40 mg kg$^{-1}$ once the initial concentrations were in the 0.6–1.2 mM range (Fig. 1). Moreover, greater organic Se levels were observed in *P. ostreatus*. In turn, *P. eryngii* was more effective in Cu bio-enrichment, particularly at the 0.6–1.2 mM concentration range at which the Cu concentrations in the fruiting bodies exceeded at least 20 mg kg$^{-1}$ (Fig. 2).

Compared to exclusive enrichment with Se, mushroom cultivation on substrates supplemented with Se in combination with Zn decreased the Se accumulation only 0.1 and 0.3 mM (*P. ostreatus*) or 1.2 mM (*P. eryngii*) concentration of elements had been applied. The greatest organic Se levels in fruiting bodies of both species were found in mushrooms growing on the substrate supplemented with

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**Fig. 1** Concentrations of Se accumulated in the fruiting bodies of *Pleurotus ostreatus* (a) and *P. eryngii* (b) growing on variously supplemented substrates. Bars represent mean with standard deviation ($n = 3$)
In this combination, the organic Se content reached nearly 80 mg kg\(^{-1}\) for \textit{P. ostreatus} and exceeded 30 mg kg\(^{-1}\) for \textit{P. eryngii} (Table 1). The latter species was, in turn, more effective in Zn accumulation; the concentrations in the fruiting bodies exceeded at least 45 mg kg\(^{-1}\) after the substrate was supplemented in the 0.6–1.2 mM element concentration range (Fig. 2).

The observed concentrations of elements in Se, Se+Cu and Se+Zn cultivation combinations revealed relatively low variability (with some exceptions), demonstrated by low values of standard deviation (Figs. 1, 2), especially that our experiment was performed in three independent replicates. This, in turn, allows repetitive levels of trace elements to be obtained.

Simultaneous substrate supplementation with Se, Cu and Zn resulted only in a slight increase in Se levels in the fruiting bodies of both, \textit{P. ostreatus} and \textit{P. eryngii}, and usually significantly lower Cu and Zn levels compared to the control (Figs. 1, 2). This is most likely due to high competitive sorption of metals from the substrate, an effect observed in various organisms including mushrooms [33, 34]. Due to unsatisfactory Se concentrations, speciation analysis was not performed for this cultivation model.

Considering that the recommended daily allowance (RDA) values were set at 55 µg for Se, 900 µg for Cu and 11 mg (male) or 8 mg (female) for Zn, for adult individuals [35, 36], the fruiting bodies of \textit{P. ostreatus} and \textit{P. eryngii} bio-enriched with Se alone or in combination with Cu or Zn demonstrate a potentially high nutritional value. However, the partial loss of trace elements during mushroom processing (e.g. washing, drying, cooking) has to be taken into account [14]. The Se in \textit{Pleurotus} mushrooms has been previously shown to be highly bioaccessible through in vitro studies, particularly in organic form [32]. In the present study, this form accounted for up to 89% for \textit{P. ostreatus} and 74% for \textit{P. eryngii} once the inorganic Se salts were added to the substrate at 1.2 mM concentration (Table 1). This is an important finding because the retention of organic forms of Se (selenocysteine and selenomethionine) is higher than that of inorganic Se (selenate—SeO\(_4^{2-}\) and selenite—SeO\(_3^{2-}\)) although both were shown to be absorbed in the gastrointestinal tract [37]. Further studies would be required to establish the bioaccessibility of Cu and Zn from mushrooms.

The results of the present study indicate that bio-enriched \textit{Pleurotus} mushrooms may help in deficiency prevention. It should, however, be highlighted that an excessive intake of Se, Zn or Cu may also bring about toxic effects in humans [38, 39]. On the other hand, mushrooms are not usually consumed on a daily basis and, moreover, the bio-fortified fruiting bodies of \textit{Pleurotus} species may be used in adequate amounts (e.g. in form of powder) as a natural ingredient of various food supplements.

### Table 1

| Cultivation model | Species | Control | 0.1 | 0.3 | 0.6 | 0.9 | 1.2 |
|-------------------|---------|---------|-----|-----|-----|-----|-----|
| Se                | \textit{P. ostreatus} | 0.6 (0.08) | 25.1 (2.7) | 9.1 (1.6) | 5.9 (0.6) | 30.6 (1.1) | 23.3 (0.4) |
| Se                | \textit{P. eryngii} | 1.1 (0.1) | 8.7 (1.1) | 28.0 (6.0) | 18.5 (1.9) | 16.9 (1.9) | 7.9 (0.5) |

| p value | * | ns | *** | *** | *** | *** | *** |

| Se+Cu            | \textit{P. ostreatus} | 0.6 (0.2) | 23.6 (5.5) | 1.6 (0.7) | 27.3 (3.8) | 17.9 (5.9) | 6.7 (2.0) |
| Se+Cu            | \textit{P. eryngii} | 1.4 (0.2) | 1.4 (0.1) | 9.6 (0.9) | 23.3 (0.5) | 9.2 (2.0) | 5.7 (2.0) |

| p value | * | ns | *** | *** | *** | *** | *** |

| Se+Zn            | \textit{P. ostreatus} | 0.6 (0.2) | 23.6 (5.5) | 1.6 (0.7) | 27.3 (3.8) | 17.9 (5.9) | 6.7 (2.0) |
| Se+Zn            | \textit{P. eryngii} | 1.4 (0.2) | 1.4 (0.1) | 9.6 (0.9) | 23.3 (0.5) | 9.2 (2.0) | 5.7 (2.0) |

| p value | * | ns | *** | *** | *** | *** | *** |

0.9 mM element concentration. In this combination, the organic Se content reached nearly 80 mg kg\(^{-1}\) for \textit{P. ostreatus} and exceeded 30 mg kg\(^{-1}\) for \textit{P. eryngii} (Table 1). The latter species was, in turn, more effective in Zn accumulation; the concentrations in the fruiting bodies exceeded at least 45 mg kg\(^{-1}\) after the substrate was supplemented in the 0.6–1.2 mM element concentration range (Fig. 2).
Growth of bio-enriched mushrooms

The growth of the studied mushroom species revealed distinctly different patterns in response to the investigated combinations and concentrations of trace elements. In the case of *P. ostreatus*, the biomass generally decreased with the increasing initial element concentration in the substrate (Fig. 3). This effect was observed on substrates supplemented exclusively with Se but also in combination with Cu and Zn. Regardless of the combination of metals in the substrate, the lowest yield was found after supplementation of 1.2 mM. Nevertheless, with the exception of the Se+Cu+Zn model, the biomass of fruiting bodies did not fall below 100 g in samples with an initial substrate element concentration up to 0.9 mM (Fig. 3).

*Pleurotus eryngii*, in turn, exhibited a tendency to increase the biomass with initial concentrations of elements, with the exception of the Se+Cu+Zn model. This was particularly observed for the Se+Cu combination for which the greatest biomass production was found after the substrate had been supplemented with 1.2 mM of each element (Fig. 4).
However, no sensory or instrumental measurement was applied; no change in colour or discolouration of the fruiting bodies of \textit{P. ostreatus} and \textit{P. eryngii} was noticeable after the mushrooms were grown on substrates supplemented with Se alone or in combination with Cu and/or Zn (Figs. S1, S2). This finding is important from the consumer point of view as it has been shown that changes in the appearance of the food, e.g. the yellowish colour of maize enhanced with vitamin A [40], can be discouraging and lead to a decrease in the commercial value of such food products.

**Antioxidant activity of bio-enriched mushrooms**

Antioxidant activities were tested on hot water extracts obtained from fruiting bodies collected from substrates supplemented with 0.6, 0.9 and 1.2 mM concentration of Se, Se and Cu, and Se and Zn, as these cultivation models yielded the most satisfactory results of element bioaccumulation, organic selenium fraction and mushroom growth.

As observed, extracts of both species grown on substrates not supplemented with any element slightly decreased the intracellular ROS content (Figs. 5, 6) and level of lipid peroxidation (Fig. 6) generated by tBHP in human platelets (Fig. 7). It has been previously shown using in vivo rodent studies that both mushroom species possess antioxidant properties, a phenomenon associated with mushroom-derived polysaccharides [41, 42]. The present study further demonstrates that extracts obtained from mushrooms bio-enriched with Se alone or in combination with Zn in the tested concentration range (0.6–1.2 mM) enhance their antioxidative effect exerted in platelets. The most significant amelioration of radicals was observed in cells pretreated with extracts from 1.2 mM of the Se and Se+Zn cultivation model with ROS levels decreased by 65.8 and 91.1% for \textit{P. ostreatus}, respectively (Fig. 5) and by 54.3 and 79.6% for \textit{P. eryngii}, respectively (Fig. 6). The counteraction of mushrooms extracts against tBHP-generated ROS resulted in significantly prevented lipid peroxidation in platelets, measured by means of MDA equivalents. The most beneficial effect was observed for extracts obtained from 1.2 mM Se+Zn supplementation—compared to the positive control the lipid peroxidation was decreased by 83.5% for \textit{P. ostreatus} (Fig. 5) and 67.8% for \textit{P. eryngii} (Fig. 6). Peroxidation of lipids is a chain reaction initiated by the hydrogen abstraction or addition of an oxygen radical, resulting in the oxidative damage of polyunsaturated fatty acids. If not terminated fast enough, a decrease in membrane fluidity and in the barrier functions of membranes is induced, final products of peroxidation (predominantly MDA and 4-hydroxy-2-nonenal) can induce genotoxicity, and cell death is promoted [43, 44]. Its prevention is thus essential for cell survival and homeostasis.

The observed in vitro effects of mushroom extracts are not only beneficial in view of the prevention of oxidative stress and its potential consequences (e.g. DNA damage,
Fig. 5 The intracellular ROS concentrations in human platelets exposed for 1 h to 10 µM tBHP and not pretreated with any mushroom extracts (tBHP; positive control), pretreated with extract of unenriched *P. ostreatus* (not enriched + tBHP) and pretreated with extracts of *P. ostreatus* bio-enriched with 0.6–1.2 mM of Se (Se + tBHP), Se and Cu (Se&Cu + tBHP), Se and Zn (Se&Zn + tBHP). Bars represent mean ± SD from three independent experiments corresponding to different donors. Asterisks represent statistically significant difference to the positive control (*p* < 0.05; Wilcoxon signed-rank test).

Fig. 6 The intracellular ROS concentrations in human platelets exposed for 1 h to 10 µM tBHP and not pretreated with any mushroom extracts (tBHP; positive control), pretreated with extract of unenriched *P. eryngii* (not enriched + tBHP) and pretreated with extracts of *P. eryngii* bio-enriched with 0.6–1.2 mM of Se (Se + tBHP), Se and Cu (Se&Cu + tBHP), Se and Zn (Se&Zn + tBHP). Bars represent mean ± SD from three independent experiments corresponding to different donors. Asterisks represent statistically significant difference to the positive control (*p* < 0.05; Wilcoxon signed-rank test).

Fig. 7 The level of lipid peroxidation (measured by means of MDA concentration) in human platelets exposed to 10 µM tBHP for 1 h and not pretreated with any mushroom extracts (tBHP; positive control), pretreated with extract of unenriched mushrooms (not enriched + tBHP) and pretreated with extracts of mushrooms bio-enriched with 0.6–1.2 mM of Se (Se + tBHP), Se and Cu (Se&Cu + tBHP), Se and Zn (Se&Zn + tBHP). *A* Data for *P. ostreatus*, *B* data for *P. eryngii*. Bars represent mean ± SD from three independent experiments corresponding to different donors. Asterisks represent statistically significant difference to the positive control (*p* < 0.05; Wilcoxon signed-rank test).
protein modification, lipid peroxidation) but also in terms of the potential inhibition of platelet activation. Although the present study did not investigate the effect of extracts on platelet aggregation, it is known that ROS may play, along with nitric oxide, adenosine and prostacyclin, a profound role in platelet activation by scavenging nitric oxide (which has a preventive effect on platelet aggregation), triggering tyrosine phosphorylation of β3 or supporting the metabolism of collagen and arachidonic acid [23]. Considering that primary mechanism of tBHP action is generation of oxidative stress [45] and that tBHP has been previously demonstrated to support platelet activation cascade [46], it is plausible that the antioxidant properties of mushroom extracts may also prevent platelets from aggregation. Moreover, a recent study demonstrated that an extract of P. eous inhibited human platelet activation in vitro, the effect linked to the high mushroom content of phenolic compounds and flavonoids [47]. Their concentrations in P. ostreatus and P. eryngii have been previously shown to be significantly increased by substrate supplementation with Se alone and particularly in combination with Zn [22]. Therefore, it might be interesting to investigate a potential application of bio-enriched mushrooms as antiplatelet agents. There is a clear evidence that dual antiplatelet treatment with aspirin and P2Y12 receptor inhibitors is essential in pharmacological treatment of acute coronary syndromes; it reduces the risk of ischaemic events and improves patients outcome [48, 49]. Further studies, preferably employing assays such as VerifyNowP2Y12, Multiplate Aggregometry or Light Transmission Aggregometry, are required to evaluate their possible use in prevention of major adverse cardio- and cerebro-vascular ischaemic events such as myocardial infarction, stent thrombosis and stroke [50].

Importantly, extracts obtained from mushrooms cultivated on substrates supplemented with 0.6 and 0.9 mM of Se and Cu did not prevent generation of ROS (Figs. 5, 6) nor peroxidation of lipids (Fig. 7). This indicates that Cu can ameliorate the beneficial antioxidative properties of Se, as well as other compounds extractable from Pleurotus mushrooms. However, Cu (along with Zn) is a constituent of cytoplasmic superoxide dismutase (Cu, Zn-SOD), an important enzyme of the cellular antioxidant system [51], and at higher concentrations it reveals significant pro-oxidant properties. It was shown that cultivation of Pleurotus sp. in a medium with increased Cu²⁺ concentration results in enhanced activity of laccases [52]; enzymes performing mono-electronic oxidation of phenols and aromatic or aliphatic amines to corresponding reactive radicals [53]. In summary, the findings of the present study indicate that P. ostreatus and P. eryngii bio-enriched with Se in combination with Cu do not reveal any promising antioxidative action, contrary to the other combinations tested.

Conclusions

The present study highlighted the potential of P. ostreatus and P. eryngii in the production of food bio-enriched with Se alone or in combination with Cu or Zn. Both species were able to grow on supplemented substrates and accumulate significant levels of elements (with a higher share of organic Se fraction) in their edible fruiting bodies. The only unsatisfactory results were obtained in the triple-element cultivation model—the biomass and accumulated content of elements were low. The study further demonstrated that bio-enrichment with Se alone and in combination with Zn largely enhances the antioxidant properties of these mushrooms in human platelets as observed by ameliorated ROS generation and prevented lipid peroxidation. These findings indicate that bio-enrichment of Pleurotus mushrooms with the studied elements may not only be a promising strategy in deficiency control but could also have a potential biomedical application, particularly in antiplatelet therapy.

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Compliance with ethical standards

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Conflict of interest All authors declare that they have no conflict of interest.

Compliance with ethics requirements This article does not contain any studies with human participants or animals performed by any of the authors.

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