In many countries, mushrooms and dishes from them are considered to be delicacies due to their specific aroma, which is provided by hundreds of volatiles such as octane derivatives and octens, lower terpenes, benzaldehyde, sulfur compounds, and others. Fatty C8-compounds such as 1-octanol, 3-octanol, 3-octanone, 1-octen-3-ol, 2-octen-1-ol, 1-octen-3-one are formed in lipoxygenase pathway from polyunsaturated fatty acids.

Mushrooms have long been used as food or food flavoring materials because of their unique and flavor. Fresh and preserved mushrooms are consumed in a lot of countries as a delicacy, particularly for their specific aroma and texture [1]. Consumers highly appreciate the characteristic flavor of mushrooms, which consists of hundreds of odorous compounds such as derivatives of octane and octenes, lower terpenes, derivatives of benzaldehyde, sulphur compounds and others [1]. Volatile aroma substances of edible mushrooms may be divided into three groups: carbon-eight (C8) derivatives impart the “characteristic mushroom flavor”, terpenoid volatiles and sulfur-containing odor compounds [2].

Eight-carbon volatile compounds are a key contributor to mushroom flavor and have been described by many authors in fungi [3]. 1-Octanol, 3-octanol, 3-octanone, 1-octen-3-ol, 2-octen-1-ol, and 1-octen-3-one are the main of them. They account for 44,3–97,6% of the total volatile fraction. [4].

The aliphatic alcohol 1-octen-3-ol (first called “matsutake alcohol”) is the principal compound contributing to the unique mushroom aroma and flavor [3]. Isoprenoids (also known as terpenoids) belong to a group of secondary metabolites that are synthesized in mushrooms. Twenty-four isoprenoids have recently been reported in fruiting bodies of the white Tuber magnatum. Limonene and cedrol were the most abundant of them [5]. Aromadendrene, alpha-farnesene and other terpenoid compounds were detected in T. borchii fruit body [6].

**Key words:** *Pleurotus ostreatus*, volatile aroma compounds, sunflower oil, corn oil, sensory profile analysis, UV spectroscopy.
Sulfur volatile organic compounds (S-VOCs) are key contributors to truffle aroma. The diversity of sulfur volatiles in truffles is large, ranging from relatively small compounds, such as dimethyl mono-(DMS), di- (DMDS) and tri- (DMTS) sulfides, which are produced by most truffle species, to complex S-volatiles such as 2-methyl-4,5-dihydrothiophene, characteristic of the white truffle *T. borchii*, and bis(methylthio) methane, characteristic of the white truffle *T. magnatum*. The latter species contain further 27 sulfur volatiles [5].

The characteristic “sulfurous” note of Shiitake mushroom is composed from S-compounds. They include the straight chain compounds dimethyl disulfide, dimethyl trisulfide, 1-(methylthio)dimethyl disulfide and cyclic compounds lenthionine (1,2,3,5,6-pentathiepane, $C_2H_4S_5$), 1,2,4-trithiolane ($C_2H_4S_3$), 1,2,4,5-tetrathiane ($C_3H_6S_4$) and 1,2,3,4,5,6-hexathiepane ($CH_2S_6$) [7].

Eight-carbon volatiles are enzymatically formed by oxygenation of polyunsaturated fatty acids (PUFAs). This reaction is catalyzed by lipoxygenases (LOXs), thus starting the so-called lipoxygenase (LOX) pathway [8]. Metabolites originating from these pathways are collectively named oxylipins [9].

The purpose of the study was to analyze possible ways of synthesis of volatile flavor compounds in mushrooms and to determine the intensity of this synthesis by *Pl. ostreatus* (the oyster mushroom) in the process of intensive cultivation on substrates with the addition of vegetable oils as a potential source of unsaturated fatty acids, which are the main substrates for the synthesis of aliphatic aroma compounds.

Sunflower (*Helianthus annuus* L.) and corn (*Zea mays* L.) oils as vegetable oils were used in this work. Sunflower oil contains up to 90% unsaturated fatty acids (linoleic and oleic) and up to 10% saturated fatty acids (palmitic and stearic) [10]. Quantitatively the predominant components of the fatty acid composition of corn oil are palmitic, oleinic and linoleic acids [11].

The composition of fatty acids of sunflower [12] and corn [13] oils is given in Table 1.

### Materials and Methods

#### Mushroom strains. Three strains of the edible mushroom *Pleurotus ostreatus* (Jacq.:Fr.) Kumm.: IBK-549, IBK-551 and IBK-1535 from the mushroom collection of the Kholodny Institute of Botany of the National Academy of Sciences of Ukraine were objects of the study [14]. This mushroom belongs to the *Pleurotaceae* family of *Agaricales* of the *Agaricomycetes* class of the *Basidiomycota*, regnum *Fungi*.

#### Solid-phase cultivation. The substrate for the production of fruiting bodies was the agricultural waste: sunflower husk and barley straw. As additives to substrates, sunflower and corn oils were used at a concentration of 1% and 5% of the weight of the wet substrate. The sunflower oil for unrefined cold pressing of the first spin of the first grade [12] and the corn oil refined deodorized grade P [13] were used in the research.

Preparation and sterilization of substrates were carried out according to commonly accepted methods [15]. The substrate was evaporated for 2 hours, CaCO$_3$ was added in an amount of 1% to the mass of the substrate and sterilized twice autoclaving at 121 °C for 30 minutes with an interval of 24 hours. Straw was pre-minced to a size of 2–3 cm. The cooled substrate was inoculated with *Pl. ostreatus* mycelium in an amount of 5% by weight of the substrate. Seeding mycelium was obtained on barley grain. Cultivation was carried out in glass jars at 26–28 °C and 70–80% humidity to the full mycelial overgrowth of the substrate. The weight of the wet substrate on the basis of sunflower husk in one jar was 150 g, and on the basis of barley straw — 110 g. Containers with the substrate were transferred to a growth room with a temperature of 15–16 °C, humidity of 80–90% and 8-hour photoperiod after full growth of the substrate by mycelium. The 1st and 2nd flushes were harvested. Mushrooms were dried at 40–45 °C in a dry oven for 24–48 hours.

Growing strains of *Pl. ostreatus* on the substrates without the addition of oils was used as a control experiment.

| Kind of oil | Mass fraction of fatty acid,% to the sum of fatty acids |
|-------------|--------------------------------------------------------|
|             | Palmitic $C_{16:0}$ | Stearic $C_{18:0}$ | Oleic $C_{18:1}$ | Linoleic $C_{18:2}$ | Linolenic $C_{18:3}$ |
| Sunflower oil | 3.0–10.0 | 1.0–10.0 | 14.0–35.0 | 50.0–75.0 | <1.0 |
| Corn oil | 9.0–14.0 | 0.5–4.0 | 24.0–42.0 | 34.0–62.0 | <2.0 |
During the cultivation process, the following growth parameters of the *Pl. ostreatus* mycelium were determined: the time of the mycelial development on the substrate, the time of primordia formation, the number of formed bunches per 100 g of wet substrate, and the yield of the first and second flushes of fruitage per 100 g of wet substrate. Cultural and morphological mushroom features were studied in order to establish a relationship between them and the synthesis of aroma-forming substances.

**Sensory profile analysis.** The sensory profile of the aroma of dried mushroom samples was studied according to [16].

The panel consisted of 5 experts trained for organoleptic analysis. First, the characteristic attributes of the aroma were determined, and then the intensity of each of them on a 5-point scale: 0 — not present; 1 — just recognizable or threshold; 2 — weak; 3 — moderate; 4 — strong; 5 — very strong. The studied samples were evaluated three times.

The organoleptic evaluation of different strains of dried mushrooms, collected at the same stages of maturation, was carried out in specially prepared, well-ventilated rooms at the Department of Biotechnology of the Ukrainian State University of Chemical Technology.

Microsoft Office Excel 2007 software was used to construct the aroma profiles of dried mushroom samples.

**Spectrophotometric analysis.** For a spectrophotometric study, the dried fruiting bodies of the first flush were crushed on an electric mill to a powder. 1 g of the obtained material was placed in the extractor, then 100 cm$^3$ of solvent were added (the hydromodule was 1:100). Hexane was used as a solvent. Extraction was carried out at boiling point (69 °C) of the solvent for 30 minutes. The extracts were cooled in a fume hood, filtered through a paper filter on a Buchner funnel and transferred quantitatively into a volumetric flask of 250 cm$^3$. Then the solvent volume was adjusted to the mark. Absorption spectra were recorded using a spectrophotometer SF-2000 in the 200–350 nm wavelength range. Pure hexane was used as a comparative solution.

**Statistical analysis.** The obtained data were processed statistically using one-way analysis of variance [17]. All samples were carried out in triplicate. Values are presented as means ± standard error of the mean. Differences at $P \leq 0.05$ were considered to be significant.

**Results and Discussion**

**Possible ways of synthesis of volatile flavor compounds in mushrooms.**

Taking into account the data of other authors and our analysis, the most probable way of synthesis of 1-octen-3-ol is presented in Figure 1.

Fatty acids are precursors in the reaction of eight-carbon volatile synthesis and are key components in a variety of lipids [3].

Lipoxygenases (linoleate: oxygen oxido reductase, EC 1.13.11.12; LOXs) are
a family of enzymes found ubiquitously in plants and mammals, but have also been detected in coral, algae, fungi, yeast and a number of bacteria. In general LOxs are non-heme iron-containing dioxygenases [9]. This group of enzymes catalyzes the insertion of molecular oxygen into polyunsaturated fatty acids containing a (1Z,4Z)-pentadiene system, such as linoleic acid, linolenic acid, and arachidonic acid, yielding the corresponding hydroperoxides [18].

Although free fatty acids is the preferred substrate of LOX, it was recently reported that fatty acids acylated to phospholipids are slowly oxygenated by lipoxygenases [19].

A lot of researches conducted to study the structure and properties of plant and animal lipoxygenases.

Oxygen insertion into PUFAs by LOX is regio- and stereo-specific, and this specificity is used as the decisive criterion for LOX classification. In plants, linoleic acid and linolenic acid are the primary substrates of LOX, because they are the most abundant fatty acids. Plant LOxs are classified as 9- or 13-LOxs with respect to their positional specificity of linoleic acid oxygenation. Animal LOxs are classified as 5-, 8-, 9-, 11-, 12-, or 15-LOxs with respect to their positional specificity of the oxygenation of arachidonic acid, the predominant substrate of animal LOxs. Furthermore, LOxs are classified as S-or R-LOxs on the basis of the chirality of their hydroperoxide products [18].

Fungal LOxs, similar to plant LOxs, act mainly on C18 fatty acids. Linoleic acid (18:2, Δ9,12) is the most abundant in widely appreciated cultivated mushrooms, followed by palmitic and stearic acids. Most fungi convert C16 fatty acids into 9-hydroperoxy or 13-hydroperoxy fatty acids (i.e. they contain 9-LOX and 13-LOX, respectively). Interestingly, 10-hydroperoxy and 12-hydroperoxy fatty acids can also be formed as possible products of fungal polyunsaturated fatty acid metabolism [20].

Products formed by LOxs can be converted by a hydroperoxide lyase (HPL), an allene oxide synthase (AOS), a peroxygenase or a reductase.

Hydroperoxide lyases are said to perform either homolytic or heterolytic cleavage of hydroperoxides, yielding different short-chain volatiles, depending on their cleavage mode. The first cleavage mechanism, homolytic, involves the cleavage of the hydroperoxide between the carbon bearing the hydroperoxide group and the saturated carbon. This mechanism has been observed in algae and mushrooms. The second mechanism is called heterolytic. It is found in most plants: the enzyme cleaves the hydroperoxide between the carbon bearing the hydroperoxide group and the unsaturated carbon [3].

Wurzenberger and Grosch showed that after incubation of the 9-, 10-, 12- and 13-hydroperoxide isomers of linoleic with a protein fraction of mushroom Psalliota bispora only 10-hydroperoxide isomer was cleaved to 1-octen-3-ol and 10-oxo-trans-8-decenoic acid [21].

Results of Akakabe et al. indicated a stereochemical correlation between (R)-1-octen-3-ol and (S)-10-hydroperoxy-(8E,12Z)-8,12-octadecadienoic acid [(S)-10-HPODE] with homogenates of Lentinula edodes and Tricholoma matsutake [22].

Assaf et al. showed that 13-hydroperoxy-cis-9,trans-11-octadecadienoic acid (13-HPOD) and 10-oxo-trans-8-decenoic acid (10-oxo-acid) were found to be the major nonvolatile metabolites of Pl. pulmonarius submerged culture, associated with the enzymatic cleavage of linoleic acid to 1-octen-3-ol. But despite its accumulation, 13-HPOD was not to be the precursor of 1-octen-3-ol. These results suggest the involvement of two different lipoxygenases in 1-octen-3-ol and 13-HPOD formation [23].

But besides 1-octen-3-ol, other aliphatic saturated and unsaturated C6-C10 compounds, which are likely to be formed with the participation of other lipoxygenases and hydroperoxidases, are responsible for the formation of the characteristic aroma of mushrooms. Probably there are several metabolic pathways of the synthesis of volatile aroma compounds in fungal cells.

Terpenoids constitute the most abundant and structurally diverse group of plant secondary metabolites [24] that is why the metabolic pathways for the formation of volatile aroma compounds of terpenoid nature are sufficiently studied and highlighted in the literature on the example of plant organisms. The scheme of biosynthesis of terpenoid compounds is shown in Fig. 2.

Terpenoids are derived from the universal C5 precursor isopentyl diphosphate (IPP) and its allylic isomer dimethylallyldiphosphate (DMAPP), which in higher plants are generated from two independent pathways located in separate intracellular compartments. In cytosol, IPP is derived from the mevalonic acid (MVA) pathway that starts with the condensation of acetyl-CoA. In plastids, IPP is formed from pyruvate and glyceraldehydes-3-
phosphate. This MVA-independent pathway, also called MEP pathway after the key intermediate methylerythritol phosphate (MEP), was discovered [24–26].

Two main S-VOC biosynthetic pathways (Fig. 3), both relying on L-methionine (Met) catabolism, have been investigated on bacteria and ascomycetes: the one-step conversion of L-methionine to metanethiol (MTL) by methionine lyase (a typical bacterial rather than yeast enzyme) or by cystathionine lyase; and a two-step pathway, initiated by L-methionine transamination to 4-methylthio-2-oxobutyric acid, which is then converted to 3-(methylthio)propanal (also known as methional) via decarboxylation, with the ultimate formation of MTL [5].

Lenthionine (1,2,3,5,6-pentathiepane), a cyclic sulfur compound found in L. edodes, is derived from a γ-L-glutamyl-cysteine sulfoxide precursor (lentinic acid) in a two-step enzymatic reaction. Lentinic acid is first activated by the removal of its γ-glutamyl moiety catalyzed by γ-glutamyl transpeptidase (GGT) producing a L-cysteine sulfoxide derivative, which then undergoes α, β-elimination catalyzed by cysteine sulfoxide lyase, resulting in a highly reactive sulfenic acid intermediate. The sulfenic acid is then rapidly condensed to form thiosulfinate, and the thiosulfinate is often further transformed into other sulfur compounds including lenthionine (Fig. 4) [28].

The analysis of possible ways of synthesis of volatile flavor compounds in mushrooms showed that the precursors of this synthesis are organic acids, including those that are part of vegetable oils.

Culture and morphological characteristics of fungal growth depending on the type of investigated substrate

The term of overgrowing of the substrate by mycelium was on the studied substrates
from 6 to 7 days, that is, there was no significant difference between this growth rate in different experimental variants compared with the control. According to the morphological features, *Pl. ostreatus* mycelia of all examined strains were white, fluffy, and denser in sunflower husk.

*Pl. ostreatus* fruit bodies cultivated on substrates with the addition of sunflower oil (A) and corn oil (B) are shown in Fig. 5.

Fruit bodies, obtained on different substrate variants within the mushroom strain, did not differ morphologically.

The growth parameters of *Pl. ostreatus* IBK-549, IBK-551 and IBK-1535 on different substrates are given in Table 2.

The period of primordial emergence varied depending on the mushroom strain from 18 to 30 days and did not differ significantly in various variants of substrates within the strain.
Fig. 5. Fruiting bodies of various strains of *Pleurotus ostreatus* (Jacq.:Fr.) Kumm. cultivated on substrates with: the addition of sunflower oil (A) and corn oil (B) (the upper line — the substrate is sunflower husk; the lower line — the substrate is barley straw)

Table 2. The growth parameters of *Pleurotus ostreatus* strains on various substrates

| Substrate version | Time of primordial emergence, day | First flush of fruitage, day | Number of mushroom bunches per 100 g of substrate, pcs | Mushroom yield of the 1st flush, g/100 g | Time of primordial emergence, day | First flush of fruitage, day | Number of mushroom bunches per 100 g of substrate, pcs | Mushroom yield of the 1st flush, g/100 g |
|-------------------|----------------------------------|----------------------------|----------------------------------------------------------|----------------------------------------|----------------------------------|----------------------------|----------------------------------------------------------|----------------------------------------|
| Sunflower husk |                                  |                            |                                                          |                                        | Barley straw                     |                            |                                                          |                                        |
| **Pl. ostreatus** strain IBK–549 |                            |                            |                                                          |                                        | **Pl. ostreatus** strain IBK–551 |                            |                                                          |                                        |
| **Pl. ostreatus** strain IBK–1535 |                            |                            |                                                          |                                        | **Control**                      |                            |                                                          |                                        |
| SO 1%          | 18                               | 25                         | 11.6±1.4                                                 | 10.8±0.4*                             | 3.8±0.3                         | 17–18                      | 25                         | 12.7±1.1                                                 | 13.0±0.4*                             | 5.6±1.0                           |
| SO 5%          | 18–20                            | 25                         | 10.4±0.7                                                 | 10.9±0.4*                             | 4.2±0.5                         | 18                         | 25–27                      | 15.5±2.3                                                 | 14.9±0.9                             | 5.5±0.5                           |
| CO 1%          | 19–20                            | 25                         | 17.1±1.0                                                 | 12.7±0.5*                             | 3.4±0.4                         | 16–17                      | 24–25                      | 18.5±1.0                                                 | 13.9±0.7                             | 5.6±0.8                           |
| CO 5%          | 20                               | 25                         | 15.1±2.0                                                 | 10.5±0.3*                             | 3.9±0.4                         | 18–19                      | 24–28                      | 15.2±0.4                                                 | 12.4±0.5                             | 9.1±1.1*                           |
| Control        | 18–20                            | 25–27                      | 12.2±0.7                                                 | 9.2±0.4                               | 3.4±0.2                         | 17                         | 25–27                      | 14.8±1.6                                                 | 13.0±0.6                             | 4.5±0.2                           |
| SO 1%          | 18–20                            | 28                         | 13.8±1.8                                                 | 10.2±0.2*                             | 3.7±0.3                         | 18–20                      | 28–29                      | 13.0±2.1                                                 | 13.1±0.4*                             | 4.7±0.2                           |
| SO 5%          | 20–22                            | 28–32                      | 11.6±2.0                                                 | 11.9±0.6*                             | 3.4±0.1                         | 20–22                      | 27–29                      | 15.8±2.6                                                 | 13.2±0.4*                             | 5.9±1.2                           |
| CO 1%          | 19–21                            | 25                         | 14.2±1.0                                                 | 10.8±0.4*                             | 3.5±0.1                         | 19–20                      | 28                         | 14.8±0.4                                                 | 15.4±1.2*                             | 4.8±0.4                           |
| CO 5%          | 21–23                            | 25–29                      | 8.9±0.7                                                  | 11.4±0.5*                             | 4.0±0.4                         | 23                         | 30                         | 19.7±1.0                                                 | 16.8±1.8*                             | 6.1±0.4*                           |
| Control        | 20–22                            | 27–28                      | 10.4±0.5                                                 | 9.0±0.4                               | 3.1±0.3                         | 22–23                      | 29                         | 19.7±2.6                                                 | 11.6±0.3                             | 4.4±0.3                           |
| SO 1%          | 22                               | 29                         | 11.1±0.3                                                 | 9.7±0.3*                              | 3.3±0.1                         | 22                         | 32–36                      | 16.7±1.0                                                 | 14.9±1.6                             | 4.9±0.4                           |
| SO 5%          | 25–26                            | 32–36                      | 14.9±1.5                                                 | 10.6±0.5*                             | 3.5±0.4                         | 22                         | 29–36                      | 14.5±1.3                                                 | 14.2±0.5                             | 5.4±1.1                           |
| CO 1%          | 30                               | 39                         | 11.8±0.3                                                 | 11.3±0.4*                             | 4.1±0.4                         | 25                         | 30–34                      | 18.5±0.4                                                 | 12.8±0.5                             | 5.3±0.3                           |
| CO 5%          | 25–29                            | 30–36                      | 8.4±0.7                                                  | 10.8±0.1*                             | 4.0±0.4                         | 25–26                      | 33–36                      | 20.6±2.7                                                 | 17.1±0.8*                             | 4.9±0.3                           |
| Control        | 24–28                            | 30–36                      | 8.4±0.3                                                  | 7.8±0.5                               | 3.5±0.4                         | 22–24                      | 30–34                      | 8.8±0.7                                                  | 11.4±1.5                             | 5.4±0.4                           |

*Note:* * — *P* < 0.05 with compared to control; SO — sunflower oil; CO — corn oil.
of control. The primordia were formed first by
the strain IBK-549, 2–3 days later by IBK-551,
and 4–5 days later by IBK-1535.

The examined strains had significant
differences by fruitage time. Fruit bodies
were most rapidly formed by strain IBK-549,
2–3 days later by the strain IBK-551, and 4–5
days later by IBK-1535. Also, there was no
significant effect of additives to the substrate
on the terms of fruiting.

There was an increase in the formation of
bunches on sunflower husk with the addition
of corn oil at a concentration of 1% for all
strains. The strain IBK-1535 produced 1.3–
2.3 times more bunches on both substrates
with additives of vegetable oils in both
concentrations as compared to control.

Yield of the first flush was higher for all
strains grown on sunflower husk with both
vegetable oils. On barley straw the increase
of yield in the first flush was observed only for
the strain IBK-551.

Analysis of literary data on the application
of oils in the cultivation of mushrooms has
shown that the addition of soybean oil at a
concentration of 3 g/l in submerged cultivation
of Pl. mutilis contributed to an increase of
mushroom biomass yield and an increase in the
synthesis of fatty acids [27–30].

The stimulating effect on the growth of
mycelium A. bisporus was also revealed in
agar medium with sesame oil additives at 1%
concentration [31].

Profile analysis of the fungal aroma

During the sensory analysis by the panel,
the following attributes of the aroma of
dried mushroom samples were determined:
mushroom, sweet, woody, herbaceous, sour,
fish, meat, earthy, floral, and putrescent.

The results of sensory analysis of dried
samples of different strains of Pl. ostreatus are
presented in circle plots at Fig. 6.

From the provided data it is evident that
the aroma profile of mushroom samples varied
depending on the substrate, the strain of
mushroom and additives. For all strains with
both additives there was an increase in the intensity
of mushroom notes in 1.2–1.5 times.

Samples of all strains grown on sunflower
husk had a 1.4–2.4 times higher intensity of
herbaceous notes and in samples obtained on
barley straw, the intensity of the herbaceous
attribute was higher in 1.5–2.4 times when
only corn oil was added to the substrate.

As for meat notes, their higher intensity (in
1.4–1.8 times) was noted in comparison with
the control in samples of dried mushrooms
of all strains cultivated on barley straw with
additives of both oils. The higher intensity of
meat notes was recorded for the strain IBK-551
(in 1.3–1.5 times) on sunflower husk with the
addition of both oils, and for strain IBK-549 (in
1.5 times) with the additive of sunflower oil.

There was an increase in the intensity of
sweet (in 1.4 times) and floral (in 1.8–2.6
times) notes for the strain IBK-551 and earthy
(in 1.3–1.8 times) for strains IBK-549 and
IBK-1535 on both substrates with the addition
of corn oil.

For some samples there was a slight increase
in the sour and putrescent characteristics of
the aroma on the substrates with additives of
vegetable oils. And the nature and strength of
woody and fish notes were almost unchanged
compared to control for any of the samples.

Statistical processing of the sensory
analysis data showed that the standard error
does not exceed ±1 point, indicating the
statistical homogeneity of the set of expert
assessments [32].

Ultraviolet spectroscopy

The registered UV absorption spectra of
hexane mushroom extracts are presented in
Figures 7 and 8.

Hexane extracts of dried samples of
Pl. ostreatus fruit bodies had light absorption
maxima in ranges of 204–210 nm and
250–290 nm. Such spectral properties are
characteristic of solutions of unsaturated
compounds with unbound double bonds,
saturated and unsaturated aldehydes and
ketones, as well as derivatives of benzene
[33]. As it was found in previous studies,
the solution of 1-octen-3-ol in hexane has a
maximum absorption at \( \lambda = 207 \text{ nm} \) [34].

A 1.2–1.4 times higher light absorption
intensity was observed throughout the studied
range of wavelengths for samples of strain
IBK-549 obtained on sunflower husk, with the
addition of both oils at a concentration of 1%,
and in the range of 250–290 nm only for the
strain IBK-551.

Mushroom extracts of the strain IBK-1535
cultivated on husk with sunflower and corn oil
additives in both concentrations also showed
an increase in light absorption compared to
control. In addition, it is higher in 1.4–1.6
times as at 207 nm (the maximum is typical for
1-octen-3-ol), and 1.5–2.5 times higher in the
range of 250–300 nm (the maxima inherent in
aldehydes and ketones).

A similar dependence was observed for
the strain IBK-1535, cultivated on barley
straw with additives of oils. And also for the
strain IBK-549, whose extracts revealed more
intense (in 1.2 times) light-absorption maxima
**Fig. 6.** Sensory profile of aroma of dried samples of *Pleurotus ostreatus* strains

*Pl. ostreatus*, strain IBK-549
(substrate is sunflower husk)

*Pl. ostreatus*, strain IBK-551
(substrate is sunflower husk)

*Pl. ostreatus*, strain IBK-1535
(substrate is sunflower husk)
Experimental articles

for strains grown on straw with both oils at a concentration of 1%.

The intensity of light absorption was 1.1–1.2 times higher only in the range of 260–290 nm for samples of the strain IBK-551, collected from substrates with additives of both oils at a concentration of 1%.

Comparison of the intensity of light absorption of various strains of *Pl. ostreatus* showed the highest level for the strain IBK-549 on both substrates compared to control.

It should also be noted that the ratio of the intensity of the light-absorption maxima at $\lambda = 207$ nm (typical for 1-octen-3-ol) and at $\lambda = 260–280$ nm (typical for aldehydes, ketones and benzene derivatives) for different substrates and strains was dissimilar. Investigated extracts of the strain IBK-549 cultivated on sunflower husk had in 2.3 times higher optical density at 260–280 nm than at near ultraviolet light. And for samples of this strain grown on barley straw, the intensity of the maxima in these ranges is almost the same. Both other strains showed almost the same ratio of intensity of light absorption on both ranges.

Thus, as a result of the study, it was found that the addition of vegetable oils to the substrate, as precursors of the synthesis of flavor compounds by the *Pl. ostreatus* strains, promotes the formation of aroma compounds by mushrooms during solid phase cultivation.

The sensory profile analysis of dried samples of the fruit bodies obtained on substrates with the addition of sunflower and corn oils at concentrations of 1% and 5% showed an increase in the intensity of mushroom, meat and herbaceous notes of

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*Fig. 7. UV spectra of hexane extracts of Pleurotus ostreatus strains:*

the substrate is sunflower husk
aroma. Spectrophotometric study of hexane extracts of dried fruit bodies of studied strains showed an increase in the intensity of light absorption of samples cultivated on substrates with the addition of vegetable oils compared to control.

The results of the study can be used in mushroom cultivation to increase the organoleptic quality of *Pl. ostreatus* fruit bodies through enriching the composition of lignocellulose substrates with vegetable oils.

**Fig. 8.** UV spectra of hexane extracts of *Pleurotus ostreatus* strains: the substrate is barley straw

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БІОСИНТЕЗ ЛЕТУЧИХ СПОЛУК ГРИБАМИ Pleurotus ostreatus (Jacq.:Fr.) Kumm.
НА СУБСТРАТАХ, ЗБАГАЧЕННИХ РОСЛИННИМИ ОЛІЯМИ
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Метою дослідження було проведення аналізу можливих шляхів синтезу запашних сполук грибами Pleurotus ostreatus (Jacq.:Fr.) Kumm. за культивування на соняшниковій лузпинні та соломі ячменю з додаванням рослинних олій (соняшникової та кукурудзяної) як джерела ненасичених жирних кислот. Сенсорний профільний аналіз висушених зразків плодових тіл показав підвищення інтенсивності грибних, м'ясних та трав'янистих нот запаху на субстратах з додаванням рослинних олій у концентрації 1% і 5%. Для штама ІВК-551 відзначено зростання інтенсивності солодких і квіткових складових запаху на обох субстратах із додаванням кукурудзяного масла. УФ-спектроскопія висушених зразків плодових тіл виявляла максимуми світлопоглинання у діапазоні 200–210 нм та 260–300 нм. Спостерігали неоднакове збільшення інтенсивності світлопоглинання зразків різних штамів, культивованих на субстратах із додаванням рослинних олій.

Ключові слова: Pleurotus ostreatus, леткі запашні сполуки, соняшникова олія, кукурудзяна олія, сенсорний профільний аналіз, УФ-спектроскопія.

БІОСИНТЕЗ ЛЕТУЧИХ СОЕДИНЕНИЙ ГРИБАМИ Pleurotus ostreatus (Jacq.:Fr.) Kumm. НА СУБСТРАТАХ, ОБОГАЩЕННЫХ РАСТИТЕЛЬНЫМИ МАСЛАМИ
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Целью исследования было проанализировать возможные пути синтеза летучих душистых соединений Pleurotus ostreatus (Jacq.:Fr.) Kumm. при культивировании на подсолнечной лузге и соломе ячменя с добавлением растительных масел (подсолнечного и кукурузного) в качестве источника ненасыщенных жирных кислот. Сенсорный профильный анализ высушенных образцов плодовых тел показал повышение интенсивности грибных, мясных и травянистых нот запаха на субстратах с добавлением растительных масел в концентрации 1% и 5%. Для штамма ІВК-551 отмечен рост интенсивности сладких и цветочных составляющих запаха на обоих субстратах с добавлением кукурузного масла. УФ-спектроскопия гексановых экстрактов высушенных образцов плодовых тел обнаружила максимумы светопоглощения в диапазоне 200–210 нм и 260–300 нм. Наблюдалось неоднаковое увеличение интенсивности светопоглощения образцов различных штаммов, культивируемых на субстратах с добавлением растительных масел.

Ключевые слова: Pleurotus ostreatus, летучие душистые соединения, подсолнечное масло, кукурузное масло, сенсорный профильный анализ, УФ-спектроскопия.