INFLUENCE OF GLUCOSE AND PEPTONE ON THE MYCELIAL GROWTH OF OYSTER MUSHROOM (*Pleurotus ostreatus*)

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**ABSTRACT**
This present study was carried out to investigate the effect of glucose concentrations (5, 10, and 15) g.L⁻¹ as carbohydrate source and peptone (0, 2, 4, and 6) g.L⁻¹ as nitrogen source on mycelial growth, colony diameter, growth period, and mycelial morphology of *Pleurotus ostreatus*. The results indicated that the best mycelial extension after three and seven days from isolation was recorded at 5 g.L⁻¹ glucose and 6 g.L⁻¹ peptone whereas the lower growth of colony diameter recorded at 15 g.L⁻¹ glucose and in the control plates of peptone. Nevertheless, complete colonization of mycelial growth period was obtained after six days of incubation between lower concentration of glucose with all concentrations of peptone and between 10 g.L⁻¹ glucose with 6 g.L⁻¹ peptone, while concentration 15 g.L⁻¹ glucose plus control or 2 g.L⁻¹ peptone were fully colonized after ten days from colony growth. However, the best interactions between glucose at concentrations (5 and 10) g.L⁻¹ with 6 g.L⁻¹ peptone were showed white mycelium color, abundant colony growth, high density mycelium and cottony texture as mycelial morphology characters for oyster mushroom.

**KEYWORDS:** *Pleurotus ostreatus*, Oyster Mushroom, Potato Dextrose Agar, Glucose, Peptone, Mycelial Growth.

**INTRODUCTION**

Oyster mushroom (*Pleurotus ostreatus*) is one of the four important species based on their production and demand (Martinez and Lopez, 2010). *Pleurotus ostreatus* belongs to the Pleurotaceae family within the order Agaricales (Thorn *et al.*, 2000). *P. ostreatus* is distributed throughout temperate forests of the northern hemisphere including Asia, Europe, and North America. In addition, the presence of *P. ostreatus* has been confirmed in South America (Zervakis and Balis, 1996).

The nutritional benefits of mushrooms have long been appreciated in the traditional medicine of many cultures and have been studied for their anti-cancer effects (Tong *et al.*, 2009). In general, mushrooms can be used as a fresh food as well as for industrial purposes. The most commonly and easily cultivated mushrooms in Iraq and Kurdistan region are oyster mushrooms but their spawn exported from other countries, while the cultivation of this important mushroom has not been very common.

Many factors should be considered during spawn production particularly mycelial growth. Generally, there are several factors that minimizing the period of mycelial colonization and obtaining thick mushroom mycelia with fast growth rate. Those factors include culture kinds, carbon to nitrogen ratio, carbon source, nitrogen source, temperature, pH, plant growth regulators and some other factors (Lu, 2009). However, the effect of culture media on the mycelial growth of oyster mushrooms strains with the addition of glucose and peptone were the most suitable for mycelium growth (Pereima, 2017). Consequently, Nwokoye *et al.* (2010) investigated the mycelial growth requirements of *Pleurotus ostreatus*, they used different kinds of carbon and nitrogen sources including glucose and peptone, the results shown that glucose significantly enhanced mycelial growth and peptone improved the greatest mycelial growth as a nitrogen source.

Therefore, the present study is investigating the impact of different concentrations of glucose and peptone on the mycelial growth and morphology characters of oyster mushroom, this will provide basic knowledge and techniques required for mycelial growth. In addition, it is the first introduction of spawn culture and production in Iraqi Kurdistan in a simple way. Thus, the recent study will provide better

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opportunities for mushroom production in Iraqi Kurdistan with lower costs. Furthermore, it will provide extra job opportunists for farmers and graduated students as well as providing a substantial product when the price of other products decreases.

MATERIALS AND METHODS

This experiment was carried out at University of Duhok, College of Agricultural Engineering Sciences, Kurdistan Region-Iraq.

Preparation of culture media

The pure culture used as substrate for isolation of cultivated oyster mushrooms which is obtained on the table (1). This medium was used as a substrate for isolation and sub-culture or regeneration agar medium for study treatments.

| Content of pure medium | PDA |
|------------------------|-----|
| Potato                 | 250g |
| Dextrose               | 15g  |
| Bacteriological agar   | 20g  |
| Peptone                | 1g   |
| pH                     | 7    |
| Distilled water        | 1 litre |

* PDA: Potato Dextrose Agar

Procedure

The potato dextrose agar (PDA) medium was prepared by boiling 250g of sliced unpeeled potato in 600ml of distilled water for 20 minutes then added enough water to infusion of potato for dissolving 20g of bacteriological agar and 15g dextrose plus 1g peptone and completed the volume to 1000ml by distilled water, subsequently pH-7 was adjusted via electronic pH meter (Type BP3001) using HCl and NaOH. The medium was sterilized using autoclave at 15 lbs pressure 121°C for 20 minutes. After this process, 10ml of the sterile medium were poured into plastic petri dishes (70 x 15 mm) and allowed to solidified, then incubated for 24 hours at 25°C to ensure the purity of the media from any possible contaminants (Guadarrama-Mendoza et al., 2014).

For tissue culture, one day aged, fresh and a healthy mushroom was obtained from local distributor. However, under aseptic conditions mushroom was isolated according to (Singh et al., 2011), then incubated at optimal temperature 28°C for mycelium growth of oyster mushroom (Hoa and Wang, 2015). After ten days mycelium colonies regenerated from growing edges and carefully transferred to another PDA that prepared at the same way in the table (1) and again incubated for another 10 days to obtain pure cultures of mycelium for isolation to the experimental treatments.

Glucose and peptone concentrations

In this study three concentrations of glucose (5, 10, and 15) g.L⁻¹ and four concentrations of peptone (0, 2, 4, and 6) g.L⁻¹ were used with 3 replications and 36 experimental unit, under aseptic conditions the culture disc about 5 mm in diameter cut out from growing edges of pure mycelium that prepared before from the petri dishes (70 x 15 mm) and transferred into the larger dishes (90 x 15 mm) for each selective plate. The inoculated petri dishes were incubated aseptically at optimum temperature 28°C (Pereima, 2017).

Colony diameter

The diameter speed of mycelium growth (mm) in (90 x 15 mm) petri dishes was measured via electronic vernier and data were recorded at 3 and 7 days (Mbogoh et al., 2011). The experiment was arranged in RCB design and data were analysed using IBM SPSS Statistics software, V. 25. Then, the means were compared using Duncan multiple range test at p < 0.01 level of confidence.

Mycelial morphology

The nature of mycelia growth was recorded through visual observation after complete colonization of mycelium in petri dishes according to (Sobal et al., 2007), he states that the major characters of mycelia morphology as texture (cottony, velvety or floccose), density (high, regular or low), colour (off-white, white...
or pale pink), colony growth (scarce, regular or abundant) and growth period per days.

**RESULTS AND DISCUSSION**

**Colony diameter:** Table (2) shows that the growth of mycelium inoculation after three days, the impact of glucose at 5 g.l⁻¹ exhibited highest mycelium growth of *Pleurotus ostreatus* followed by 10 g.l⁻¹, and 15 g.l⁻¹ (Fig. 1) while there was no significantly observed between them, this is in accordance with the findings of (Neelam et al., 2013) who studied the carbon requirement on the growth of oyster mushroom, their results showed that glucose was the best carbon source, In addition, (Lisha and Lulu Das, 2015) recorded that glucose was the best carbon source among many types of sugar for mycelial growth of white button mushroom, but it is contradictory to the results of experiment conducted by (Memon et al., 2017), they shown that the effect of dextrose sugar on mycelial growth of oyster mushroom significantly recorded at 20 g.l⁻¹ glucose, concerning that increasing sugar concentrations more than 20 g.l⁻¹ produced fewer mycelial growth. One possible reason can be related to the unavailability of nitrogen sources in culture media. Another reason may be the inadequate amount of nitrogen sources in the media. this is confirmed by (Ma et al., 2014) who suggested that the optimum carbon to nitrogen ratio of culture media for white mushroom was 20:1 ~ 30:1.

However, the results show growth on all plates which tested and the mycelial growth diameter was seen significantly differ depending on the available of nitrogen sources in four concentrations of peptone even in control plates, this might be returned to the reasons of one gram peptone inside two subcultures of agar causes to improve growth in the control plates while only to a certain level (Fig. 1). In addition, that the best maximum diameter of mycelial growth was found in 6 g.l⁻¹ peptone but it is not significant differences from the other concentrations except control which recorded the lowest diameter of colony growth. On the other hands, (Nwokoye et al., 2010) who reported that peptone supported the greatest mycelial growth of oyster mushroom as a nitrogen source. However, the significant interactions between glucose and peptone obtained in the lower concentration of glucose with higher concentration of peptone.

| Table (2): Effect of glucose and peptone on the mycelial growth diameter (mm) of oyster mushroom after 3 days from isolation. |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Glucose g.l⁻¹   | Peptone g.l⁻¹   | Means of glucose|
|                 | 0               | 2               | 4               | 6               |
| 5               | 24.33bc         | 30.47abc        | 33.54abc        | 37.01abc        | 31.34a         |
| 10              | 22.29bc         | 25.05bc         | 26.04bc         | 27.87abc        | 25.31a         |
| 15              | 20.76c          | 23.67bc         | 23.79bc         | 27.12abc        | 23.84bc        |
| Means of peptone| 22.46c          | 26.40bc         | 27.79bc         | 30.67bc         |                |

* Means within the same column followed by the same letters are not significantly different at p < 0.01 according to Duncan’s test.
Colony diameter: Table (3) revealed that the growth of mycelium inoculation after seven days. Hence, the influences of glucose significantly recorded at (5 and 10) g.l\(^{-1}\) which has maximum mycelium growth of *Pleurotus ostreatus* (Fig. 2). Those results agree with the findings of (Thirumalvalavan et al., 2005) that recorded glucose as better carbon sources than others and petri dishes that contain 15 g.l\(^{-1}\) glucose obtained the minimum growth of colony diameter (Fig. 2). The effect of peptone significantly increased in the high concentration 6 g.l\(^{-1}\) which has higher mycelium growth, also (Ma et al., 2014) and (Lisha and Lulu Das, 2015) recorded the best sources of nitrogen was peptone after yeast and beef extract for the mycelial growth of white button mushroom. And plates that contain (0 and 2) g.l\(^{-1}\) peptone had the lowest growth of colony diameter. On the other hands, (Nwokoye et al., 2010) found that peptone gradually increased mycelial dry weight of *Pleurotus ostreatus* until 50 g.l\(^{-1}\). Consequently, after seven days from incubation the interaction between glucose and peptone significantly seen in many treatments (Fig. 2) but the better one was glucose at 5 g.l\(^{-1}\) with the entire levels of peptone which had fully colonized petri dishes. And the lesser amount of mycelium growth recorded at concentration 15 g.l\(^{-1}\) glucose + without peptone.

**Table (3): Effect of glucose and peptone on the mycelial growth diameter (mm) of oyster mushroom after 7 days from isolation.**

| Glucose g.l\(^{-1}\) | Peptone g.l\(^{-1}\) | Means of glucose |
|---------------------|---------------------|------------------|
|                     | 0       | 2     | 4     | 6     |
| 5                   | 87.00\(a\) | 87.00\(a\) | 87.00\(a\) | 87.00\(a\) |
| 10                  | 83.85\(a\) | 81.44\(ab\) | 83.16\(a\) | 86.00\(a\) | 83.61\(a\) |
| 15                  | 63.18\(c\) | 67.18\(cd\) | 72.4\(b\) | 79.32\(ab\) | 70.52\(b\) |
| Means of peptone    | 78.01\(a\) | 78.54\(ab\) | 80.85\(a\) | 84.11\(a\) |

* Means within the same column followed by the same letters are not significantly different at p < 0.01 according to Duncan’s test.
Fig. 2: Shows that the extend of mycelial growth at 7 days after inoculation in replication one for entire plates that starting from (5g glucose + control peptone and ended by 15g glucose + 6g peptone) l-1 respectively.

Mycelial morphology

Mycelial colonies after 10 days of entire plates for each replication were observed white colour (Table 4) and there were no recorded significant differences among them. This might be due to in this study two subcultures of PDA used to obtain pure mycelium as substrate for isolation.

On the other hands, after complete colony growth at 6 days the best treatments (5g or 10g glucose + 6g peptone) l-1 were recorded abundant colony growth, high density mycelium and cottony texture (Fig. 2) and the rest treatments were highly different. The result is in agreement with (Guadarrama-Mendoza et al., 2014) who suggested that cotty mycelium of Pleurotus spp. presented significantly higher growth rates in comparison with floccose mycelium and they confirmed that cotty mycelium resulted in two various types of growth. Firstly, cotty has high density and abundant growth. Secondly, cotty texture also has regular density and regular growth as shown in table (4).

Generally, complete colonization after 6 days that showed a shortest growth period of mycelium and significantly recorded in some plates including glucose at 5 g.l\(^{-1}\) plus all concentrations of peptone, and 10 g.l\(^{-1}\) glucose with 6 g.l\(^{-1}\) peptone. Whereas the interactions between 15 g.l\(^{-1}\) glucose + control and 2 g.l\(^{-1}\) peptone were fully colonized after 10 days from inoculation.


**Table (4): Effect of glucose and peptone on the mycelial morphology and growth period of oyster mushroom.**

| T. | Culture media | Colony growth | Mycelial morphology | Growth period (days) |
|----|---------------|---------------|---------------------|---------------------|
| T₁ | 5g glucose + 0g peptone | regular | low | cottony | white | 6 |
| T₂ | 5g glucose + 2g peptone | abundant | low | velvety | white | 6 |
| T₃ | 5g glucose + 4g peptone | regular | regular | floccose | white | 6 |
| T₄ | 5g glucose + 6g peptone | abundant | high | cottony | white | 6 |
| T₅ | 10g glucose + 0g peptone | regular | regular | velvety | white | 7 |
| T₆ | 10g glucose + 2g peptone | abundant | regular | floccose | white | 8 |
| T₇ | 10g glucose + 4g peptone | abundant | high | cottony | white | 8 |
| T₈ | 10g glucose + 6g peptone | abundant | high | cottony | white | 6 |
| T₉ | 15g glucose + 0g peptone | scarce | low | velvety | white | 10 |
| T₁₀| 15g glucose + 2g peptone | scarce | high | cottony | white | 10 |
| T₁₁| 15g glucose + 4g peptone | regular | regular | cottony | white | 9 |
| T₁₂| 15g glucose + 6g peptone | regular | high | cottony | white | 9 |

* Mycelial morphology recorded after complete colonization of mycelium in each plate.

**CONCLUSION**

In conclusion, the study shows that different levels of glucose and peptone sources can be used for mycelial growth of oyster mushroom. According to study results the mycelial growth, colony diameter and growth period were the best when decreased in the concentration of glucose, whereas increasing in the concentration of peptone causes to increase the growth rate of mycelial colony. Also improving mycelial morphology characters and complete colonization after six days for dual interaction significantly recorded between low and medium concentration of glucose with high concentration of peptone.

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(Pleurotus ostreatus)

تأثیر الگلوکوز و الیثیتون علی نمو الیسیله میلی لفطر المحاری

(الخلاصة)

آروعتت هذه الدراسة للتحقق من تراکیز مختلف من الالگلوکوز (5، 10، 15) غم/لتر و الیترویدات، والیثیتون (0، 2، 4 و 6) غم/لتر كمصدر لللیتروجین علی نمو الیسیله میلی، فطر المحاری . اثرات النتایج إلى أن افضل امتداد للیسیله بعد ثلاثة و سبعه أيام من العزل تم تسجيله عند 5 غم/لتر الالگلوکوز و 6 غم/لتر الیثیتون بينما سجل انخفاض معدل نمو فطر المستمرة عند 15 غم/لتر الگلوکوز وفي معاملة السيطرة من الیثیتون. ومع ذلك، تم الحصول على الاستعمار الكلي لفترة نمو الیسیله بعد ستة أيام من العزل بين أقل تركیز الکلوگواز مع جميع تركیزات الیثیتون وبین 10 غم/لتر الگلوکوز مع 6 غم/لتر الیثیتون، بينما 15 غم/لتر الگلوکوز مع معاملة السيطرة و 2 غم/لتر الیثیتون سجلت الاستعمار الكلي بعد عشرة أيام من نمو الیسیله. بالإضافة فإن أفضل النتایج بين الگلوکوز/ترکیزات (5 و 10) غم/لتر می شد، و کنترل نمو المستعمرات، و میلیلیم عالية الكافيه، و نسق قطبي كیفیتی مورفولوجیة لبیسیله فطر المحاری.

الکلمات المجازیة: نموز، Fleurots ostreatus، Fleurots ostreatus، نموز، مستعمرات، میلیلیم عالية الكافيه، و کنترل نمو، الجلوکوز، الیثیتون، نمو، Fleurots ostreatus.