Purification and Characterization of Milk Clotting Enzyme from Edible Mushroom (*Pleurotus florida*)

Ashraf Bakr 1, Osama Ibrahim 2*, Abd El-Sattar El-Ghandour 3, Noha El-Deeb 3

1 Food Science & Technology Department, Tanta University, Egypt; ashraf.bakr@agr.tanta.edu.eg (A.B.);
2 Dairy Science Department, National Research Centre, Cairo, Egypt; osama_nrc@hotmail.com (O.I.);
3 Dairy Technology Department, Animal Production Research Institute, Dokki, Egypt; dr.elghandour@yahoo.com (A.E.); nanosh66@yahoo.com (N.E.);
* Correspondence: osama_nrc@hotmail.com (O.I.);

Abstract: Oyster mushroom (*Pleurotus florida* 14 MICC) is one of the most widely cultivated edible fungi in the world. Milk clotting enzyme (MCE) derived from the mold was developed as a calf rennet alternative; thus, it was purified and characterized, and its effect on different milk species was investigated. The highest MCE activity (75.49 SU/ml) was observed in mushroom fruit bodies dissolved in 0.2 M sodium acetate pH 5.0; as well as the highest total MCE activity (367.85 SU) was recorded at 20% of ammonium sulfate with a specific activity of 343.79 SU mg⁻¹ protein while size exclusion column chromatography on Sephadex G-100 purified MCE 3.46-fold with 17.96% yield. Also, it could be capable of coagulating different milk species. Mushroom MCE exhibited their optimal activity at pH 5.0 for crude extract (CE) while at pH 6.0 for partial purified (PP) and purified (P) MCE fractions; as well as at 55 °C for CE and PP MCE fractions while 50 °C for P MCE. CaCl₂ concentration (0.01%) recorded the maximal activity for CE while (0.04%) and (0.02%) for PP and P fractions, respectively. It could be concluded that MCE from Oyster mushroom may be a good candidate as a calf rennet substitute in cheese production.

Keywords: milk clotting enzyme; oyster mushroom; calf rennet substitute; coagulant.

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1. Introduction

Calf rennet considers the oldest known application of enzymes in cheese making [1] as a traditional milk clotting enzyme (MCE) obtained from the fourth stomach of suckling calves, which consists of chymosin (EC 3.4.23.4) as the major coagulating component, as well as pepsin (EC 3.4.23.1) as a proteolytic enzyme [2]. Chymosin is known for its high specificity for releasing the caseinomacropeptide from κ-casein, which triggers the destabilization of the casein micelles, which induces milk clotting [3], whereas pepsin is much less specific and hydrolyses bonds with Phe, Tyr, Leu, or Val residues [4].

However, the use of calf rennet may be limited for religious reasons (e.g., Judaism and Islam), diet (vegetarianism), or consumer concern regarding genetically engineered foods (e.g., Germany, Netherlands, and France forbid the use of recombinant calf rennet) [5] Since cheese is a popular dairy product worldwide, the demand of milk coagulant has rapidly increased as the cheese production increased [6] Thus, much research interest has been directed towards discovering calf rennet alternative for cheese making from the microbial origin as an attempt to meet the demand for milk coagulants for cheese making.
Fungi belong to the microbial diversity, which is generally regarded as safe (GRAS), and their extracellular enzymes are easily recovered in bioprocess [7]. Among fungi, mushrooms are macrofungi which belong to basidiomycetes and ascomycetes. Oyster mushroom is one of the most widely cultivated edible fungi in the world [8]. Thus, MCE derived from mold has been developed as a calf rennet alternative such as Rhizomucor miehei, Rhizomucor pusillus, Cryphonectria parasitica [9], Mucor circinelloides [10], Mucor thermohyalospora, Rhizopus azygosporus [11], and Aspergillus oryzae [12] have been established. As regard mushroom species, Moharib 2007 [13] reported that Pleurotus sajorcaju STCPI-27 mushroom could be used as a source of protease enzymes with proteolytic and MCE activities. Also, Nakamura et al. (2014) [14] explored MCE-producing strains among mushrooms consumed routinely, including Hericium erinaceum MAFF 435060, MAFF 430234, and NBRC 100328, which were used in cheese making. Shamtsyan et al. (2014) [15] identified edible basidial mushroom Coprinus lagopides novel active producer of MCE as a calf rennet substitute in cheese production. Sato et al. (2016) [16] found that MCE from H. erinaceum MAFF 435060 coagulates heat-treated milk at both 66 °C for 30 min and 130 °C for 2 sec. Furthermore, Martim et al. (2017) [7] recorded that Pleurotus albidus as an edible mushroom is considered an available source of milk clotting protease.

Although the scientific interest focused on the nutritional value and medicinal properties of edible oyster mushroom which opened a new area to study their biologically active substances, where it's used as a source of MCE as well as the study of their biochemical characteristics is still limited to investigate their technological suitability in cheese production. Therefore, the present work aimed to extract and purify MCE from edible oyster mushroom (Pleurotus florida) available in Egypt as a calf rennet substitute in cheese production. Moreover, the biochemical characteristics of mushroom MCE, and its effect on different milk species were investigated.

2. Materials and Methods

2.1. Raw material and chemicals.

Dried fruiting bodies of oyster mushroom Pleurotus florida 14 MICC were obtained and identified from Central Laboratory for Agricultural Climate, Agricultural Research Center, Giza, Egypt. The dried samples were ground into a fine powder by laboratory mill and store in closed containers at 4 °C until use (Figure 1). Skim milk powder was purchased from BIEMLEK Spoldzielniamleczrska, Poland. All other reagents and chemicals were used in analytical grade.

2.2. Methods.

2.2.1. Extraction of MCE from the oyster mushroom.

MCE activity was detected in different extracts of oyster mushroom (Pleurotus florida 14 MICC) using 0.2 M acetate buffer pH 5.0, 0.2 M citrate buffer pH 5.0, and distilled water at the level of 10 (w/v) which was stored for 48 h at 5 °C. The recovered supernatants resulting from the cooling centrifugation of extracts (5000 rpm for 20 min at 5 °C) were considered crude enzymatic extract, and then MCE activity and protein content were determined.
2.2.2. Purification of oyster mushroom MCE.

2.2.2.1. Ammonium sulfate precipitation (ASP).

The selected crude enzymatic extract was partially purified using ammonium sulfate to 90% saturation at 5 °C, according to Colowick and Kaplan (1955) [17]. MCE activity and protein content of the recovered fractions were determined. The rich active fraction was dialyzed using 0.2 M acetate buffer pH 5.0 for 24 h at 5 °C for further purification.

2.2.2.2. Size exclusion column chromatography on Sephadex G-100.

The dialyzed fractions were further purified by applied on Sephadex G-100 (Sigma-Aldrich, Germany) column (2.5×20 cm, Pharmacia, Uppsala, Sweden), equilibrated with 0.2 M acetate buffer pH 5.0, and the sample eluted with the same buffer. The recovered fractions were assayed for MCE activity and protein detection at OD 280 nm. The rich fractions of MCE activity were pooled and considered as purified oyster mushroom MCE.

2.2.3. Electrophoresis of MCE fractions.

MCE molecular weight and purity were determined by Sodium Dodecyl Sulphate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) method according to Laemmli (1970) [18]. The protein samples were loaded for electrophoresis at 50 V, performed in a vertical slab gel apparatus in 12.5% polyacrylamide separating gel and 4% stacking gel. Protein samples were visualized by Coomassie brilliant blue R–250. The pre-stained standard proteins in the range of 10–250 kDa (Thermo scientific) were used as a molecular weight marker.

2.2.4. MCE activity assay.

The MCE activity of all resulted enzymatic extracts was measured according to the method of IDF (1992) [19]. Milk clotting activity is expressed in terms of the Soxhlet unit. One Soxhlet unit (SU) of milk clotting activity was defined as the amount of enzyme required to clot 1 ml of the substrate within 40 min at 35 °C. Soxhlet units were calculated using the following equation according to IDF (1992) [18], SU/ml = M×2400/E× t
Where M is the volume of substrate (ml); E, is the amount of enzyme extract (ml) and t, is the clotting time (sec).

2.2.5. Determination of proteolytic activity.

The proteolytic activity of crude, partially purified, and purified enzymatic extracts were determined according to Chopra and Mathur (1983) [20]. One unit of proteolytic activity is defined as the amount of enzyme required to release TCA-soluble fragments giving a blue color equivalent to one µg of tyrosine (Sigma-Aldrich, Germany) under the standard assay condition.

2.2.6. Protein content determination.

The protein content in all enzymatic extracts was determined as described by Bradford's (1976) [21] procedure using Coomassie brilliant blue G-250 dye (Bio-Rad, USA). The developed color was measured at 595 nm using UV 1201-vis spectrophotometer Shimadzu,
Japan. The protein concentration is calculated from a calibration curve of bovine serum albumin (Mallinckrodt, France).

2.2.7. Specific MCE activity calculation.

The specific MCE activity is calculated by divided the MCE activity (SU/ml) by the protein content (mg/ml).

2.2.8. Biochemical characterization of mushroom MCE.

2.2.8.1. Optimum pH.

The MCE activity was measured at different pH values ranging from 3-8 using 0.2 M Acetate buffer (pH 3.0-5.0), 0.2 M phosphate buffer (pH 6.0-7.0), and 0.2 M Tris-HCl buffer (pH 8.0) in order to define its optimum pH of MCE reaction.

2.2.8.2. Optimum temperature.

The MCE optimal reaction temperature was determined after the reaction was performed at different temperatures ranging from 35 to 65 °C.

2.2.8.3. Effect of CaCl$_2$ concentration on MCE activity.

The MCE activity was measured using skim milk dissolved in different concentrations of CaCl$_2$ ranging from 0.01-0.05 %.

2.2.8.4. Effect of NaCl concentration on MCE activity.

The presence of 1.0-5.0 % NaCl in all resulted MCE fractions was studied. The residual MCE activity was measured under standard assay conditions, while the sample without any addition served as a control (100%).

2.2.8.5. Thermal stability.

MCE was pre-incubated at different temperatures ranging from 40 to 70 °C for 15, 30, and 60 min, and then the residual MCA was determined.

2.2.8.6. Determination of MCE activity on different milk types.

MCE activity was determined using different milk types as a substrate, such as a cow, buffalo, goat, and sheep.

2.2.9. Statistical analysis.

SAS statistical software performed all statistical analysis of results (SAS, 1999) [22] using the ANOVA procedure to analyze variance. The results were expressed as mean ± standard error, and the differences between means were tested for significance using Duncan’s multiple ranges at $p \leq 0.05$. 

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3. Results and Discussion

The genus *Pleurotus* has been used to study the production of different enzymes and secondary metabolites due to it is an edible fungus. Also, research regarding the edible mushroom increased mainly due to integrating the information generated by genomics, proteomics, transcriptomics, and metabolomics [23]. Therefore, oyster (*Pleurotus floridana* 14 MICC) mushroom was used as dried fruiting bodies to produce MCE as a calf rennet substitute to meet the demand for milk coagulants for cheese production. Hence, MCE is extracted and purified from an oyster mushroom with successive purification processes such as ASP and gel filtration chromatography. Moreover, its biochemical characteristics, as well as their coagulating activity on different milk species, were investigated.

3.1. Extraction of MCE from the oyster mushroom.

MCE activity was determined on dried fruit bodies of oyster (*Pleurotus floridana* 14 MICC) mushroom (Figure 1) using different extraction buffers as presented in Table 1. It could be noticed that the highest MCE activity (75.49 SU/ml) and its specific activity (356.08 SU/mg protein) was observed in oyster mushroom dissolved in 0.2 M sodium acetate buffer pH 5.0, followed by citrate phosphate pH 5.0 and distilled water. Similar findings are reported with Martim et al. (2017) [7], who detected MCE on edible mushrooms, including *Pleurotus albida* DPUA 1692 with 73.39 U/ml. However, MCE activity detected in oyster (*Pleurotus floridana* 14 MICC) mushroom is slightly higher than *Thermomucor indicae-seudaticae* N31 with 60.5 U/ml [24] and *Aspergillus flavo furcatis* DPUA 1493 with 68.61 U/ml [25]; while it is lower than MCE activity detected in *Termitomyces clypeatus* with 333.33 U/ml [26].

The MCE from oyster (*Pleurotus floridana* 14 MICC) mushroom formed a strong clot with forming quite whey after adding the enzymatic extract in skim milk solution (Figure 1). A similar result of clot formation was observed by Alecrim et al. (2014) [25] and Martim et al. (2017) [7], who detected MCE in *Aspergillus flavo furcatis* DPUA 1493 and *Pleurotus albida* DPUA 1692, respectively.

### Table 1. Extraction of MCE from oyster (*Pleurotus floridana*) mushroom.

| Extraction buffers          | MCE activity (SU/ml) | MCE specific activity (SU/mg) |
|----------------------------|----------------------|------------------------------|
| Sodium acetate pH 5.0       | 75.49±2.28<sup>a</sup> | 356.08±9.95<sup>a</sup>    |
| Citrate phosphate pH 5.0   | 60.82±4.82<sup>ab</sup> | 291.00±21.83<sup>ab</sup> |
| Distilled water            | 55.50±5.38<sup>b</sup> | 263.03±25.99<sup>b</sup>   |

Figure 1. Oyster (*Pleurotus floridana* 14 MICC) mushroom and their milk clotting activity. (a) Dried fruiting bodies of mushrooms. (b) Dried mushroom powder. (c) Whey formed in the milk clotting test.

3.2. Purification of MCE from oyster mushroom.

MCE was partially purified from dried oyster mushroom (*Pleurotus floridana* 14 MICC) using ASP procedure. Table 2 shows that the highest total MCE activity (367.85 SU) was
recorded at 20% saturation of ammonium sulfate with specific MCE activity of 343.79 SU mg\(^{-1}\) protein as well as the MCE was partially purified 1.01-fold from the crude extract of oyster mushroom with a yield of 24.36%. Also, mushroom MCE activity gradually decreased with ammonium sulfate saturation increased, affecting the resulted yield and purification fold (data not shown). This result is in agreement with Nestor et al. (2012) [27], who reported that the highest MCA of Solanum elaegnifolium seeds extract was recorded at 20% of ammonium sulfate, as well as 20-40% saturation of ammonium sulfate precipitated MCE from prepared seed cake of Moringa oleifera [28]. However, low saturation of ammonium sulfate is superior to high concentrations usually used to purify MCE from different plants [29, 30].

Mushroom MCE was further purified using size exclusion column chromatography on Sephadex G-100, as shown in Figure 2. It could be noted that MCE activity was observed in one peak among 40 collected fractions with maximal activity in fraction no. 18 (67.8 SU/ml). Also, the purification of MCE using Sephadex G-100 column recovered 17.96% with 3.46 as a purification fold where the total MCE activity was 271.20 SU with 1232.73 SU/mg protein as specific activity (Table 2). The final MCE recovery is similar to that obtained from previously described laboratory-scale procedures by Lebedeva and Proskuryakov (2009) [31], which purified MCE from oyster mushroom (Pleurotus ostreatus (Fr.) Kumm).

### Table 2. Purification results of MCE from oyster (Pleurotus florida) mushroom.

| Purification steps       | Total Activity* (SU) | Total protein* (mg) | Specific activity (unit/mg protein) | Proteolytic activity (PA) (U/ml) | MCA/PA ratio | Yield (%) | Purification fold |
|-------------------------|----------------------|---------------------|-------------------------------------|----------------------------------|---------------|-----------|-------------------|
| Crude enzyme            | 1509.80              | 4.24                | 356.08                              | 2.39                             | 31.59         | 100       | 1.00              |
| ASP (20%)               | 367.85               | 1.02                | 360.64                              | 2.45                             | 30.03         | 24.36     | 1.01              |
| SEC on Sephadex G-100   | 271.20               | 0.22                | 1232.73                             | 2.82                             | 24.04         | 17.96     | 3.46              |

*Calculation based on 0.1 g/ml of dried oyster mushroom in 0.1 M sodium acetate buffer pH 5.0. **Specific activity** = Enzyme activity/Protein content; **Total activity** = Enzyme activity X Fraction volume; **Total protein** = Protein content X Fraction volume; **MCA** = milk clotting enzyme activity (SU/ml); **Yield** = Total activity of purified enzyme/Total activity of crude enzyme X 100; **Purification fold** = Specific activity of purified enzyme/Specific activity of crude enzyme; **ASP** = Ammonium sulfate precipitation; **SEC** = Size exclusion column chromatography.

![Figure 2](https://nanobioletters.com/

### 3.3. Electrophoretic profile of mushroom MCE fractions.

The electrophoretic profile of MCE fractions showed different bands in the crude fraction, which decreased in the partially purified fraction using ASP as presented in Figure 3; it could be due to the effect of the purification process. Moreover, SDS-PAGE of the purified...
MCE from oyster (*Pleurotus florida* 14 MICC) mushroom showed a single band with a molecular mass of about 45 kDa (Figure 3). This finding is similar to MCE aspartic peptidase from basidiomycete *Piptoporus soloniensis* [32], basidiomycetous yeast *Cryptococcus* sp. S-2 [33], and *Pleurotus ostreatus* [34].

**Figure 3.** Electrophoretic profile of MCE factions from the oyster mushroom. (c) Crude extract. (PP) Partial purified MCE fraction. (P) Purified MCE fraction. The black arrow indicates the protein corresponding to purified MCE.

### 3.4. Proteolytic activity of oyster mushrooms.

The proteolytic activity (PA) of crude MCE extracted from oyster (*Pleurotus florida* 14 MICC) mushroom was 2.39 U/ml with a ratio of MCA/PA of 31.59 as shown in Table 2. Moreover, PA of partial purified MCE was 2.45 U/ml, while the purified mushroom MCE recorded PA of 2.82 U/ml with a ratio of MCA/PA of 30.03 and 24.04, respectively. Hence, PA in the purified fractions was higher than a crude fraction; it could be due to the purification process since eliminating other components affects the overall PA of the resulting fractions (Figure 3). However, the relation of MCE activity and PA as a coagulation ratio is associated with the enzymatic specific in the clot formation, which influences the selection of MCE alternatives [25].

### 3.5. Biochemical characterization of oyster mushroom MCE.

Oyster (*Pleurotus florida* 14 MICC) mushroom MCE exhibited the optimal activity at 55 °C for the crude and partial purified extracts as shown in Figure 4, with a broad temperature range of 40-55 °C followed by gradually decreasing in their activity at 60 and 65 °C, respectively. Moreover, the purified mushroom MCE recorded their maximal activity at 50 °C. However, the optimum reaction temperature of mushroom MCE was similar to that found for MCE extracted from *Pleurotus sajor-caju* STCPI-27, and *Pleurotus eryngii* were 45 °C [13, 35]; while it was quite lower than of MCE extracted from *Pleurotus albidus* [7], *Rhizopus oryzae* [36], *Rhizopus microsporus* var. *rhizopodiformis* [37] which observed at 60 °C.

Figure 5 shows the optimal reaction pH for crude MCE of oyster mushroom was 5.0 while the purified and partial purified MCE was observed at pH 6.0. Also, neutral and alkaline pH reduces mushroom MCE for all resulted fractions. It could be due to the conformation changes in the protein structure caused by charge repulsion, where the distribution of charge in the protein surface and the conformations are modified, which affect the association between the MCE and milk protein as a substrate. Similar findings were reported by Sakovich et al. (2019) [34], who recorded that the highest MCE activity was shown at pH 5.0, meanwhile, Martim et al. (2017) [7] recorded the maximal activity of MCE from *Pleurotus albidus* at pH 6.0, and then the alkaline condition decreased its activity which only 19.37% was recorded at pH 10.0. Also, *Pediococcus acidilactici* SH exhibited high MCE activity at pH 6.0 [38], as well
as Vishwanatha et al. (2010) [39] recorded high MCE activity at pH 6.3 from Aspergillus oryzae MTCC 5341 followed by the activity reduction of in higher pH values.

![Figure 4](https://example.com/figure4.png) **Figure 4.** Temperature profile of MCE fractions from the oyster mushroom.

The CaCl₂ concentration (0.01%) recorded the maximal MCE activity for crude MCE extracted from oyster mushrooms, followed by a gradual decrease in their activity, as shown in Figure 6. Moreover, purified and partially purified mushroom MCE activity increased until 0.02% and 0.04%, respectively, after their activity decreased. However, it is well known that the non-enzymatic secondary phase, in which the aggregation of para-k-casein and other casein components occurs in association with Ca²⁺ ions and eventually results in the formation of a gel [40]. Also, Abd El-Salam et al. (2017) [41] reported that artichoke MCE required Ca²⁺ ions in the curd formation. Thus, clot formation is Ca²⁺ dependent [42]; it was confirmed with milk clotting enzyme extracted from mushroom, requiring suitable CaCl₂ concentration for optimal activity.

Sodium chloride is widely used as a preservative and flavoring agent in food and dairy products [43, 44]. Figure 7 shows that the mushroom MCE activity gradually decreased as the level of sodium chloride increased for all enzyme fractions. It could be due to inducing conformational changes in the enzyme molecule affected by a high sodium chloride level, which leads to a decrease in its MCE activity. Also, the addition of NaCl decreased the rate of MCE enzymatic reaction for coagulation [45]. These findings follow Ahmed et al. (2016) [46], who recorded the highest bacterial MCE activity extracted from Bacillus stearothermophilus at the lower level of NaCl.

The thermal behavior of the enzyme is a useful property in the cheese-making process, as it can be quickly inactivated by moderate heating [47]. The thermal stability of all mushroom MCE fractions was shown in Figure 8. The results revealed that the residual MCE activity decreased as both temperature and pre-incubation time increased, which could be attributed to the conformational changes in the protein structure under high temperatures during the pre-incubation time. Also, the results indicated that the highest reduced activity of mushroom MCE after pre-incubation for 60 min at 40 °C is ~12, 28, and 36% for crude, partially purified, and purified MCE fractions, respectively. Moreover, all mushroom MCE fractions activities were quickly diminished after incubation at 60-70 °C. MCE from Termitomyces clypeatus MTCC 5091 was stable between 35 and 50 °C which retaining more than 80% of its activity, followed by a fast reduction of activity as the incubation temperature increased [26]. Also, Alecrim et al. (2014) [25] reported that MCE from Aspergillus flavo furcatis DPUA 1493 had thermal stability higher than 70% between 40 and 60 °C, while in above temperature was decreased to 28%. Thus, mushroom MCE exhibit higher thermal stability at 40 °C for all enzymatic
fractions, while its stability at 50 °C is varied from 30 min for both crude and purified fractions. Hence, the thermal stability of mushroom MCE is similar to calf rennet since calf rennet reached its maximal activity at 45 °C, followed by a sharp decline when the temperature exceeded 50 °C [48].

Figure 6. Effect of calcium chloride on MCE activity from oyster mushroom

Figure 7. Effect of sodium chloride on MCE activity from oyster mushroom

Figure 8. Thermal stability of MCE fractions from the oyster mushroom.
(a) Crude extract. (b) Partial purified MCE. (c) Purified MCE.

The suitability of mushroom MCE fractions towards different milk types, including cow, buffalo, goat, and sheep milk, was shown in Figure 9. Mushroom MCE could be capable of coagulating all examined milk types with various coagulation times started with cow milk followed by buffalo, sheep, and goat milk for both crude and partially purified MCE fractions, while the purified MCE recorded the shortest coagulation time for buffalo milk among other milk species followed by cow, sheep and goat milk. It could be due to the different compositional components of milk types, as well as the difference of casein fractions content among the examined milk species, which affects their coagulation time [49]. Park et al. (2007) [50] mentioned that the casein micelles diameter, hydration, and mineralization generally lead to shorter coagulating time, weaker gel consistency, and decreased yield of the cheese curd. Similar findings were reported by Barbagallo et al. (2007) [51] and Abdeen et al. (2021) [28], who observed the caprine, ovine and bovine milk coagulated with (Cynara cardunculus L.) flowers and Moringa oleifera of seed cake, respectively. Hence, the capability of oyster (Pleurotus florida 14 MICC) mushroom MCE for coagulation of various milk casein types supports their suitability to be used as a calf substitute in cheese making.
4. Conclusions

MCE isolated and purified from edible oyster (*Pleurotus florida* 14 MICC) mushroom fruit bodies showed thermal stability, as well as it could be capable of coagulating different milk species, including cow milk followed by buffalo, sheep, and goat milk. Mushroom MCE exhibited their optimal activity at pH 5.0 for crude extract while at pH 6.0 for other purified MCE fractions, at 55 °C for crude and partial purified MCE fractions while 50 °C for the purified MCE. Moreover, CaCl₂ concentration (0.01%) recorded the maximal activity for crude MCE while the purified and partially purified mushroom MCE activity increased to 0.02% and 0.04%, respectively.

Thus, it could be concluded that MCE from Oyster (*Pleurotus florida* 14 MICC) mushroom be a good candidate as a calf rennet substitute in cheese production. Further studies will be examined its technological suitability for cheese production and its effect on the chemical, textural and sensory properties of the resulted cheese compared to commercial coagulants.

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**Conflicts of Interest**

The authors declare no conflict of interest.

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