Introduction

Environmental pollution is one of the major ecological difficulties, rapid growth of industrial and explosion of population are the primary reasons for the dwindling water quality [1]. In last few decades, requiring urgent attention has been paid to the industrial wastes, which are usually discharged harmful and toxic substances into on land or into different natural water sources such as rivers and canals and creating health hazards [2]. Milk processing requires high specific water consumption and considerable raw material waste effluents [3]. Dairy industry is one of the major industries causing water pollution because it contains large quantities of milk constituents such as casein, lactose, inorganic salt, detergents, dissolved sugars and proteins, fats, rich in fats, oils and greases and can have negative impacts on wastewater treatment systems [4]. The dairy industries produce effluents as often cause foul odours, blockage of pipes and sewer lines [5,6]. When dairy effluents discharged into water bodies, these effluents rapidly degraded causing depletion of dissolved oxygen levels of receiving water bodies. These receiving water bodies then become a propagation place for flies and mosquitoes carrying malaria and other perilous disease like dengue fever, yellow fever and chicken [5]. The disposal of large quantities of this wastewater untreated or partially treated and domestic source rapidly causes deterioration of the environment [7]. Environmentalists and governments are looking for efficient, cheap and long lasting solutions for waste treatment and recycling. Chemical methods of wastewater treatment are inevitably cost intensive and cannot be employed in all industries, and these methods creates waste disposal problems by using neutralization and detoxification of the discharged wastewater is necessary which also causes further contaminations to the environment [8,9]. To overcome the shortage of chemical treatment, studies have been conducted using microorganisms for the biological industrial waste biological treatment offers an economical alternative to chemical treatment method as they are highly selective to the range of pollutants removed and are expensive. Microbial treatment systems have advantage of being simple in design and low in cost [10]. Fungi and bacteria are concerned in the biodegradation of undesirable materials or compounds and convert them into beneficial products [11]. The development of applications for a chitosan has expanded rapidly in recent years. Chitosan is a naturally available biopolymer coagulant, and it considers safe and more economical alternative for developing countries. Extraction of chitosan from fish waste is gaining significant importance in the world of industrial. Chitosan is a naturally available biopolymer of glucosamine and N-acetyl glucosamine. It has been produced by alkaline N-deacetylation process of shrimp (Crangon) and fish (Labeorohita). Chitosan, being polycationic, nontoxic, coagulation creates better quality flocks, have a faster settling velocity, biodegradable as well as antimicrobial, has many applications especially in the agriculture, food and pharmaceutical industries and wastewater treatment [12-14]. Use eco-friendly such as chitosan belonging to the polysaccharide to use as reducing and stabilizing agent to obtain metal nanostructures [15]. Chitosan nano silver generally showed stronger bactericidal effects for gram positive bacteria (E. faecalis and Staphylococcus aureus) than for a gram-negative bacteria (E. coli, Salmonella typhimurium and K. pneumoniae) [16,17]. One of the most eminent renewable energy resources is biodiesel and considered to be a potential alternative to petro-diesel fuel in many developed and developing countries worldwide [18]. The global biodiesel production has increased to 18.2 billion liters per year from 2000 to 2010 [19]. Oleaginous microbes, known for their ability to yield 20% to 87% of their total biomass as lipids and fatty acids [20,21]. Microbial oils when compared to plant oils were found to be more economical and easier for this processed.

Comparative Study of Biodegradation and Coagulation of Dairy Effluent by Using Immobilized Microbial Isolates and Chitosane Silver Nano Particles and Production Biodiesel

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Currently, much attention is being paid to the development of microbial lipids can represent a valuable alternative feedstock for biodiesel production, for many advantages such as short life cycle, microorganisms grows much faster and are easier to handle, less work required, are less affected by the location, season, climate, as well as faster growth and a potential solution for a bio-based economy [22]. In the future, production of microbial lipid will become a major oil source in the transportation sector [23,24]. Oleaginous micro-organisms such as yeasts, fungi and micro algae can accumulate high amounts of reserved lipids in the form of triacylglycerols under appropriate cultivation condition [25,26]. The biodiesel quality depends upon the fatty acid composition of the oil feedstock [27]. Biodiesel constitutes a renewable fuel that is convenient with current commercial diesel engines and has clear benefits relative to diesel fuel including enhanced biodgradation, decreased toxicity, and a lower emission profile [28]. To reduce the cost of microbial oils, exploring other carbon sources instead of glucose is very important especially for such oils applied to biodiesel production. It was reported that xylose, corn straw, molasses, whey and other agricultural and industrial wastes could be used as the carbon sources for microbial oils accumulation [19]. Oleaginous fungi can synthesized and accumulate high amounts of Triacylglycerols (TAG) within their cells. This TAG can be effectively changed over to biodiesel through a method know as transesterification via the conversion of TAG and methanol with potassium hydroxide as a catalyst [29]. Many researches such as yeast strains **Curvularia sp.** [23] and **Trichoderma reessei** has a wide spectrum in production of biodiesel [21], also **Trichoderma spp.** and **Gliocladium roseum** [30].

**Objective**

- The purpose of the study is to compare the reduction rate of BOD, COD and turbidity for dairy effluent after the addition of fungal and bacterial to units and using the chitosan as an adsorbent to treat dairy wastewater
- Application of chitosan silver nanocomposites efficiency as a biomaterial for dairy wastewater treatment by coagulation process
- Isolation and identification of fungal isolate capable of yielding high amount of storage lipids.
- Characterization and comparative analysis of fatty acid profiles of novel isolated fungus and ability of the selected isolate on the growth on molasses as a sole carbon source and test their potential utilities as biodiesel feedstock.

**Materials and Methods**

**Study area**

For the present study the dairy effluent samples were collected from dairy factories Panda milk in Kaliuob City industrial zones, Egypt. Samples were collected in sterile 5L plastic container. Samples were stored at 4 °C inside an ice box then transferred immediately to the laboratory of Central Laboratory for Environmental Quality Monitoring for further experiments.

**Characterization physico-chemical analysis of dairy effluent and treated effluent**

All the samples were analyzed for untreated dairy effluent and final filtrated after treatment, according to standard methods for the examination of water and wastewater (APHA, 2012). These parameters include; pH, turbidity, total suspended solids (TSS), biological oxygen demand (BOD), COD and Bacteriology analysis (Total coliform, fecal coliform, fecal streptococci and *pseudomonas aeruginosa*).

**Coagulation test**

In this research, it is very important to determine the optimum dosage of coagulant and to establish optimum chemical conditions as pH value. Then, the different efficiencies were chosen for a coagulant and pH dose for the different coagulants. For chitosane, coagulant dose was selected at 0.5 to 3 g/700 ml and pH from 6.39 to 6.58. A conventional jar test apparatus was employed for the test. All tests were carried out with 500 ml of wastewater at room temperature. Optimum pH was determined by adjusting of 700 ml aliquot of wastewater at different pH using sulfuric acid and sodium hydroxide. A fast stirring of 100 rpm was applied for 3 minutes. At the end of the speed period, the mixing was stopped and the formed flocs were left to settle down for 30 minutes. The samples to be tested were taken from the top of the water level in the flask.

**Isolation of microorganisms**

To count the number of bacteria and fungal colonies in the Wastewater, serial dilution followed with pour plating was done. Three samples were prepared after serial dilution to 1/10, 1/100, 1/1000, 1/10000 and from each sample one ml sample is poured into flask contain Sabouraud’s Glucose broth medium and is well mixed and incubated at 28 °C for 72 hours. Similar in bacteria one ml sample is poured into flask of Nutrient broth and incubated at 35 °C for 24 h. One loopful enriched sample from N.B. flask was streaked on Nutrient Agar plates and skim milk agar. One loopful enriched sample from S.B. was streaked on Sabouraud’s Agar plates. Nutrient Agar plates were incubated at 35 °C for 24 h, while the Sabouraud’s Agar plates were incubated for 3 days at 28 °C. Plating was done in triplicate for each medium. Isolated colonies were further studied for Gram’s staining and identified on the basis of Gram nature.

**Identification and phylogenetic analysis**

Lactobacillus isolate was identified using Biolog’s microbial identification system (Biolog® Gen III,USA). While *M. circinelloides* was identified depending on colony morphology and microscopic examination using lactophenol cotton blue staining method.

Identification of the *M. circinelloides* was performed based on molecular genetic analysis using the internal transcribed spacer region using ITS1-5.8S ITS2 regions by specific primers ITS1 (5’TCC GTA GGT GAA CCT TGC GG 3’) and ITS4 (5’TCC GCT TAT TGA TAT GC 3’). Partial sequences of the isolated 18s rDNA were obtained using a strategy based on [31].
The nearest fungus with similar sequence was aligned with GenBank database using BLASTn as the query sequence [32]. Alignment with the similar sequences was carried out using ClustalW [15]. Phylogenetic tree was structured applying neighbor-joining algorithm with MEGA 7 software and bootstrap for 1000 re-sampling to ensure robustness and reliability of trees constructed [12].

Experimental procedure for the extraction of chitosan

Preparation of chitosan from tilapia fish scales was done following three major steps, demineralization, deproteination, and deacetylation. For demineralization, 10 g of raw fish scales were washed thoroughly with water than was treated with in 1% HCL solution for 36 hours at room temperature. After demineralization, the sample was washed with tap water till the sample reaches neutral pH. For deproteination, fish scales powder was treated with 2N NaOH for 36h. The residue was then collected and washed with distilled water until the pH became neutral, followed by thorough washing and drying as mentioned above. After this step, the end product was chitin. For deacetylation, chitin was treated with strong alkali, 0.4 g of chitin was added to 50% NaOH for 5h at 90 °C followed by washing till it reaches neutral pH. After drying, the final product recovered was chitosan (Figure 1) [33].

Immobilization of bacteria in alginate beads

Sodium alginate was used as the immobilizing agent for bead preparation. Exponentially growing cells were harvested by centrifugation (4500 rpm for 10 min) and resuspended in 50 ml of sterile water. To this, 50 ml of 4% alginate solution was mixed thoroughly to get a final alginate concentration of 2%. The alginate-bacterial mixture with chitosane then added drop wise into CaCl₂ (0.1 M) solution. The beads were kept in the same solution for 30 mins at 4 °C for hardening.

Immobilization of fungi in alginate beads

Commercial grade sodium alginate was used as the immobilizing agent for bead preparation. Exponentially growing M. circinelloides, was harvested by centrifugation (4500 rpm for 10 min) and resuspended in 50 ml of sterile water. To this, 50 ml of 4% alginate solution was mixed thoroughly to get a final alginate concentration of 2%. The alginate-bacterial mixture with chitosane then added drop wise into CaCl₂ (0.1 M) solution. The beads were kept in the same solution for 30 min at 4 °C for hardening.

Experimental and Design of integrated treatment system

The sample water which has to be treated will undergo different treatment steps in specially designed parts of the plant is presented in Figure 2. The construction of wastewater treatment plant consists of six stages, filtration tank, Fungi unite, bacteria unite, coagulation tank and chitosane silver nano particles tank. The first tank was considered as the primary tank capacity 5L with 50 cm height, contain two layers of filtration media were placed, 15 cm height of black sand on the top, while 15 cm height of Zeolite was on the bottom than...
Determination of the chitosan-Ag nanoparticle

The one-step production of chitosan-Ag nanoparticle spheres is shown in Figure 3. Chitosan (0.3g, dissolved in 10 mL of 2% v/v CH$_3$COOH solution) and 2% 50 mL AgNO$_3$ was mixed by constant stirring for 30 minutes, and then an AgNO$_3$-chitosan mixture solution was obtained. The AgNO$_3$-chitosan mixture solution was then dropped into a 30%, 25 mL NaOH solution. After 10 minutes, yellow-brown color spheres were yielded. Spheres were collected and washed twice with distilled deionized water to remove residual alkali [34].

Characterization of chitosan-Ag nanoparticle

After preparation of chitosan-Ag nanoparticle, the stock solution was measured by TEM (JEOL-JEM-2100), a UV-Vis spectrophotometer model: Thermo Scientific Orion AQ 8000...
Aqua Mate 8000 was used to record silver plasmon band and Fourier transform infrared spectroscopy (FT-IR-jasco 4600).

**Bio-oil extraction**

Preparation of fungal cultures: Oleaginous fungus *M. circinelloides* was grown in the medium with the following components (in gl-1): Glucose, 20.0; (NH$_4$)$_2$SO$_4$, 1.4; KH$_2$PO$_4$, 2.0; MgSO$_4$.7H$_2$O, 0.3; 0.7KH$_2$PO$_4$; 0.6 yeast extract and sterilized by autoclaving. The broth was inoculated with the isolated fungal strains and incubated at 28 °C in rotary shaker at 120 rpm for 5 days. The fungal biomass determination *M. circinelloides* gl$^{-1}$. The fungal biomass of screened isolates, mycelia were harvested from the incubated flask by suction filtration through Whatman No 1 filter paper and thoroughly washed with distilled water. Then filtered mycelia dried at 65 °C in an oven for 2 hrs. The weight of the dried sample was calculated and dry biomass was expressed in gl$^{-1}$. In the above mentioned media the carbon source was replaced by various low cost substrates such as molasses, Banana peel, sugarcane and water hyacinth to test the ability of oleaginous fungi to accumulate lipids when grown on agro-industrial wastes. These low cost carbon substrates were sun dried and grounded into fine powder. The fine powdered carbon source raw materials were then sieved with 0.2 mm in diameter sieve and kept in desiccators until required for use. We have used concentration of dilute H$_2$SO$_4$, (5% wt/wt) for retention time 120 min at temperature (120 °C) for pretreatment of raw material. The fungal was inoculated in the production medium and kept in shaker at 150 rpm.

Extraction of lipid compounds: Lipids were extracted following modified the method of Bligh and Dyer [35]. The obtained biomass (dry weight) was transferred to 50 ml centrifugal tube and 20 ml of HCl (4 M) was added and incubated at 60 °C for 1 h, then kept at-80 °C for 2 hours and subsequently in boiling water for 10 min. This freezing process was repeated 2 times in order to break up the cells. The extraction was performed with 1.0 g dried biomass in a glass vessel containing 30 ml methanol and toluene [36]. The extraction was assisted by avortex in 10 min. After vortexing, further extraction with 30 ml chloroform/methanol (1:1) were added into the tube, shaken vigorously with a vortex and then centrifuged at 3000 rpm for 10 min. The lipid containing chloroform layer (the lower layer) lower chloroform layer was filtered on filter paper containing 1.0 g of anhydrous sodium sulfate and collected in vials pre-weighed glass. This procedure was repeated for the extraction of lipids remaining in the sample. All the organic phases were pooled and the solvent removed in atmosphere of nitrogen. Lipids content was expressed as gram lipid per liter.

**Determination of fatty acids profile by GC with FID:** A known amount (50 µl) of each extracted lipid was added into a glass tube and reacted with 0.6 ml of KOH (0.5N) in methanol at 80 °C for 10 min, then 1 ml of Boron trifluoride methanol complex was added to each tube and incubated at 80 °C for 10 min. Two milliliters of distilled water was added to stop the reaction, the suspension was vortexed vigorously. Equal volume of hexane was added to each tube and mixed thoroughly. After 30 min the upper phase containing the fatty acids methyl esters (FAMEs) were taken out and then transferred into a new clean tube for gas chromatography (GC) analysis. The gas chromatography (7890A, Agilent Technologies, USA) was equipped with a flame ionization detector (FID) and a capillary column DB225 (30 m × 0.25 m × 0.25 µm film thickness). The injector temperature was 250 °C; the initial column temperature was 160 °C and rose to the final temperature of 250 °C at a rate of 3 °C per min; and the detector temperature was 250 °C. The carrier gas was helium at a flow rate of 450 ml/min.

**Statistical analysis**

One way-ANOVA was performed to calculate significant differences in treatment means. SPSS version 20.0 software was used for interpretation of data. Data with a p value < 0.05 were considered significantly different. All experiments were done in triplicate.

**Results and Discussion**

**Characterization of dairy effluent**

In the present investigation, Table 1 summarizes the obtained various of the physicochemical analysis for untreated dairy wastewater.

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**Table 1:** The characteristics of studied of dairy wastewater before and after treatment.

| Parameter | Dairy effluent | Fungal stage | Bacterial stage | Concentration chitosan 3g/700 ml |
|-----------|----------------|--------------|----------------|----------------------------------|
|           |                |              |                | 0.5 | 1 | 1.5 | 3 |
| pH        | 5.73           | 6.48         | 5.38           | 6.39 | 6.42 | 6.55 | 6.58 |
| Colour    | Milky White    | Light green  | Faint green    | Semi clear | Semi clear | Semi clear | Semi clear |
| COD (mg/l)| 3900 ± 317.4   | 1200 ± 1.89  | 1500 ± 2.54    | 716 ± 0.15 × 102 | 630 ± 0.2 × 102 | 490 ± 0.36 × 102 | 423 ± 0.251 × 102 |
| Turbidity | 350 ± 25.1     | 120 ± 0.15   | 97 ± 0.107     | 76 ± 0.035 × 102 | 66 ± 0.035 × 102 | 45 ± 0.035 × 102 | 27 ± 0.025 × 102 |
| BOD (mg/l)| 1300 ± 80.8    | 136 ± 0.32   | 200 ± 0.36     | 89 ± 0.015 × 102 | 76 ± 0.047 × 102 | 57 ± 0.025 × 102 | 44 ± 0.04 × 102 |
| TSS (mg/l)| 791 ± 9.0      | 400 ± 1 × 102| 490 ± 0.95     | 195 ± 0.05 × 102 | 150 ± 0.036 × 102 | 132 ± 0.025 × 102 | 109 ± 0.01 × 102 |
| NH$_4$ (mg/l)| 37 ± 2.08   | -            | -              | 13.8 ± 0.076 × 102 | 11 ± 0.01 × 102 | 8.6 ± 0.76 × 102 | 2.7 ± 0.25 × 102 |
pH: The influent dairy wastewater was slightly acidic (5.86). The acidic pH is attributed to the breakdown of milk lactose into lactic acid [37]. It was variable even after treatment by using bacterial isolate pH 5.38, while in fungi unite pH observed to be slightly acidic 6.48.

Total suspended solids (TSS): The concentration of suspended solids in dairy effluent 791 mg/L. The percent reduction in total suspended solids of treated dairy effluent is recorded in Figure 3. It is depicted in the (Table 1) that after treatment by fungi and Lactobacillus units for 7 days the percent reductions shown by fungal was 49.2%. Reduction efficiency of bacteria Lactobacillus was 46.3%. It is clear that fungal caused the highest reduction where as bacterial caused least reduction in TSS. The efficiency of chitosan in the removal of TSS for dairy wastewater is shown in Figure 1. It can be observed that the performance of the chitosan is influenced by the pH of the effluent. The optimum pH value of the dairy wastewater was found to be in the pH 6.58. An interaction between NH\(^+\)_3 functional groups and anionic mineral particles which attract colloidal particles in the medium which get neutralized and settle down as flocs. On the other hand, in Figure 4 shows, the results indicates the effects of varying the dosages of chitosan-Ag nanoparticle, the clarity of water increase with increase concentration doses from 68.3% at 1 g/l, to 90% at 4 g/l. The reduction in TSS after aeration might be due to use of suspended organics by microorganisms for their growth and development. The overall reduction after filtration is directly associated with the uses of fungus, bacteria and chitosan-Ag nanoparticle.

Turbidity: Turbidity is a critical parameter for public water supplies and disinfection processes. The efficiency of turbidity removal is an important parameter to check the efficiency of coagulation. It can be seen that turbidity removal efficiency was 72.2% for M. circinelloides and 65.7% for Lactobacillus at 7 days, it can be seen that fungi caused highest reduction in turbidity than bacteria. Turbidity decreased due to consumption of organic materials and suspended particles by M. circinelloides/Lactobacillus through growth and survival. Under the condition controlling solution pH (6.58), the residual turbidities after mixing with various dosages of chitosan (0.3 to 2 g/700 ml) of dairy wastewater were shows in (Table 1). It says that the residual turbidity reduced as the coagulation dosage increased. Indeed, in this, it is shown that the percentage removal of turbidity of dairy wastewater was enhanced by increasing the chitosan concentration up to 3 g/l. It’s clear that effective coagulation was achieved with much lower doses of chitosan than would be required for complete charge neutralization of particles, and this process was guided by the combined effects of electrostatic patch and bridging mechanisms. This coagulation process was fast; turbidity removal was achieved within 30 min at natural pH. This is because the chitosan is a bio multi polymer, has a positive charge and contains free amine groups which give high capability for chemical relevance with molecules that have negative charges such as proteins, fats and mineral ions [38]. As can be observed from Figure 4, the turbidity was decreased with increasing chitosan-Ag nanoparticle concentration. When chitosan-Ag nanoparticle concentration increased to 4 g/l increases the water clarity up to 93.75%. The turbidity had decreased from 350 to 20 NTU for 24h.

Biochemical oxygen demand (BOD): Biochemical oxygen demand is widely used as an indication of water quality. It can be seen that after reduction by fungal and bacterial units as seen in (Table 1), the reductions in BOD values for bacteria
Chemical oxygen demand (COD): The graphical representation (Table 1) show the COD values that are observed during at 7 days treatment of the effluent after reduction by fungal and bacterial units were significantly reduced by fungal and Lactobacillus 74.3% and 69.2% respectively. From the graphs it is very clear that the COD of the effluent is continuously decreasing and it indicates that the strains used for treating the effluent are helpful in COD reduction. The best reduction efficiency of the effluent, whereas; fungal showed highest reduction in COD. The surface charge of the chitosan effectively varies with varying the pH of the wastewater. So, the experiments were performed at pH 6.58. chitosan dosage of 0.3 to 3 g/700 ml, settling time of 1h as constant and the observation is depicted in Table 1. From the Table 1, it was observed that the percentage of COD removal increased with increasing initial pH up to 6.58. This is mainly due to the formation -NH₃⁺ ions in acidic conditions, which could adsorbs the negatively charged organic matters present in the wastewater significantly [39] and increased the removal efficiency of COD (88.4%) up to the pH of 6.58. It can be seen from Figure 6 that COD concentration was decreased with increase of chitosan-Ag nanoparticle concentration (g/l). When chitosan-Ag nanoparticle concentration was 1 to 4 g/l, the COD reached 76.9% to 90% at 24h. Our results are in accordance with the reduction in COD seen by [40] had also reported 67.1% and 48.3% reduction in COD of dairy wastewater with use of two bacterial strains namely Neisseria sp. and Citrobacter sp.

Effects of Chitosane AgNP on Bacteria of Dairy Wastewater

As given in (Table 2), illustrated that the results for the total colifrom in influent was recorded (70000 CFU/100 ml) and after 40 and 60 minutes were recorded 3800 and 200 CFU/100 ml with removal 94.5% and 97.5% respectively, while at 2h the concentration bacteria was reduced to nil. As the results indicate, removal percentage is significantly increased by the increase of time.

The results are presented in (Table 2). Revealed that the results for the fecal coliform in influent was recorded (20000 CFU/100 ml) and after 40 and 60 minutes were recorded 1700 and 60 CFU/100 ml with removal 94.5% and 97.5% respectively, while at 2h the concentration bacteria was reduced to nil. As the results indicate, removal percentage is
significantly increased by the increase of time.

Given the analytical results in (Table 2), it could be noticed that, the fecal streptococci counts were relatively high at influent with (1800 CFU/100 ml). These counts were remarkably decreased in effluent water after 40 minute was recorded 500 CFU/100 ml with removal 72% while at 1h was rescored 5 CFU/100 ml with removal 99.0%, while at 2h the concentration bacteria was reduced to nil.

It can be seen from (Table 2), illustrated that the results for the *Pseudomonas aeruginosa* in influent was recorded (20000 CFU/100 ml) and after 40, 60 and 120 minutes were recorded 4000, 2800, 2500 CFU/100 ml with removal 80%,
The possible mechanisms are as follows, the electrostatic interaction between positive charged silver metal ions with the negative charged DNA and protein molecules could collapse the structure and function of DNA and protein [44]. Kim, et al. 2007 [45] suggested that the free radicals might be involved in the membrane damage. They have depicted that the antioxidant (N-acetyl cysteine) could influence the antibacterial activity of free radicals which may be released from the surface of silver nanoparticles.

Silver Chitosan Nanocomposites Analysis by FT-IR Spectrophotometry

Figure 8 Shows the FTIR spectra of the chitosan-Ag nanoparticle spheres. The bands between 3,500 cm⁻¹ and 3294 cm⁻¹ due to overlapping of O-H and amine N-H stretching bands. The bands between 2,925 cm⁻¹ corresponded to the alkane C-H-stretching lipids. The bands between 1,659 and 1586 cm⁻¹ corresponded to the amino groups of amide. 1715 (C = O stretching). The bands between 1450, 1376, and 86% and 87.5% respectively, while at 4h the concentration bacteria was reduced to nil. As the results indicate, removal percentage is significantly increased by the increase of time.

Removal percentage of total coliform, fecal coliform, fecal streptococci and pseudomonas aeruginosa have a significant difference with four other times (p < 0.05). Comparing the removal percentage in 40, 60 and 120 minutes there is no significant difference (p < 0.05).

Nano silver chitosan molecule has the ability to interact with bacterial surface and is adsorbed on the surface of the cells and stacks on the microbial cell surface and for mingan impervious layer around the cell, leading to the block of the channels [42]. The chitosan inhibits the bacteria during electrostatic reactions between anime group (NH⁺) of chitosan and phosphoryl groups of phospholipids, which are found in the cell walls of bacteria [43]. Generally, cell walls of G⁻ bacteria contain a highest fat ratio than cell walls of G+ bacteria.

Table 2: Effect of different contact times of chitosan-Ag nanoparticle on microbial contamination.

| Bacterial tests (CFU/100 ml) | chitosan-Ag nanoparticle (4 g/700 ml) |
|-----------------------------|--------------------------------------|
|                             | Influent | 40 minute | 1 hour | 2 hours | 4 hours |
| **Total coliform** (CFU/100 ml) (mean ± S.D/100 ml) | 100000 ± 15 × 10¹ | 3500 ± 4 × 10² | 600 ± 0.5 × 10² | 0 | 0 |
| **Fecal coliform** (CFU/100 ml) (mean ± S.D/100 ml) | 20000 ± 2 × 10¹ | 1700 ± 2.5 × 10² | 60 ± 7.6 | 0 | 0 |
| **Fecal streptococci** (CFU/100 ml) (mean ± S.D/100 ml) | 1850 ± 1.3 × 10² | 500 ± 1 × 10² | 7 ± 2.5 | 0 | 0 |
| **pseudomonas aeruginosa** (CFU/100 ml) (mean ± S.D/100 ml) | 200000 ± 104 × 10² | 4000 ± 2.5 × 10² | 2800 ± 2 × 10² | 330 ± 3.6 | 0 |

Figure 8: FTIR spectrum of chitosan silver nanoparticles.
1318 cm$^{-1}$ for C-H bending. Of 1000 - 1157 cm$^{-1}$ (CO stretching) were common in spectra due to the chitosan backbone. The peaks observed at 845, 751, 665 and 604 cm$^{-1}$ correspond to C-H stretching of alkenes. The FTIR spectrum of chitosan shows O-H stretching at 3,243 cm$^{-1}$, C-H and C-N stretching at 2,925 cm$^{-1}$, N-H bending at 1,637 cm$^{-1}$, N-H angular deformation in CO NH plane at 1,544 cm$^{-1}$ and C-O-C band stretching at 1,141 cm$^{-1}$.

**TEM Analysis**

The natural polymer chitosan nucleates, as well as stabilizes the silver nanoparticles formed and, thus, prevents aggregation of silver nanoparticles. The reason is the excellent chelating property of chitosan backbone, which binds the silver to it. Here it is reduced by NaOH and stabilized by the chitosan chains [15]. TEM micrographs shown in Figure 9 clearly depict the uniformly distributed silver nanoparticles embedded in chitosan. All particles exhibits poly dispersed and semi spherical morphology, the silver nanoparticles are visible as dark spots inside the chitosan-Ag nanoparticle with size approximately between 8 and 30 nm.

![Figure 9: TEM image of the chitosan-Ag nanoparticle.](image)

![Figure 10: UV-Vis spectra of chitosan silver nanoparticles.](image)
UV-Vis Spectroscopy Analysis

UV-Vis spectroscopic analysis is commonly used method to identify the formation of metal nanoparticles by analyzing the unique optical properties which depends on the size and the shape of the nanoparticles [46]. When AgNO₃ was mixed with chitosan solution, Ag⁺ ions could be bound to chitosan macromolecules via electrostatic interactions, because the electron-rich oxygen atoms of polar hydroxyl and ether groups of chitosan are expected to interact with electropositive transition metal cations [47]. Ag nanoparticles were formed by reduction of Ag⁺ using chitosan as both reducing as well as stabilizing agent and the process was accelerated by the use of NaOH. The colour of the solution changed from yellow to brown by NaOH indicating the formation of Ag nanoparticles [48]. The characteristic brown colour of colloidal silver solution is due to the excitation of surface plasmon vibrations in the nanoparticle and provides a convenient spectroscopic signature of their formation [39]. The UV-Vis spectrum of chitosan-Ag nanoparticle exhibited a strong absorption band at 400 nm (Figure 10). This is due to the strong surface plasmon resonance (SPR) shown by silver nanoparticles. Similar results were 400 nm obtained by [15].

Determination of Fatty Acids Profile by GC-FID

Fatty acids composition FAMEs of the test oleaginous fungal isolates *mucor circinelloides* in liquid medium molasses of date which were extracted by acid methanalysis during transesterification process of fungal lipids extract to Fatty Acid Methyl Esters (FAMEs) and were determined by GC/MS. As shown in Figure 11, fatty acid profiles by GC/MS showed the presence of a high fraction of saturated, and monounsaturated fatty acids which is considered a potential feature to indicate the fuel quality of fungal based diesel. Among saturated fatty acids, high amount of palmitic acid (C16:0) and significant amount of stearic acid (C18:0) were observed in the lipid profile of *mucor circinelloides*. Among monounsaturated fatty acids, oleic acid (C18:1) was shown in the Table 3. Few amounts of capric, undecanoic acid and vaccenic acid exhibiting percentage values of 1% respectively. Fungal oleaginous usually differ from most vegetable oils in being rich in PUFAs (polyunsaturated fatty acids) and hence are mainly exploited for PUFA production [49]. Fatty acids indicated for good biodiesel properties include C14:0, C16:0, C16:1, C18:0, C18:1 and C18:2 [50].

Fatty acids composition of lipid extracted

The growth and lipid production of oleaginous fungi *mucor circinelloides* in various molasses waste, Sugarcane bagasse and water hyacinth were studied. By accumulating lipids within a short period of time and growing well on a variety of wastes and inexpensive materials, such as nutritional residues molasses from industry and Sugarcane bagasse and water hyacinth, the oleaginous microorganisms plays a vital role in lowering the cost of fuels [51]. It was found that all the carbon sources used supported the growth and production of lipid to a considerable extent. Various low cost substrate were used which includes molasses, Sugarcane bagasse and water hyacinth. Figures 12 and Figure 13 show the biomass concentration and lipid concentration of fungi grown in different low cost substrate respectively. It was observed that there was an increase in biomass concentration and lipid concentration when the fungi were cultured in media containing molasses as carbon source. The lipid concentration for carbon sources molasses, Sugarcane bagasse and water hyacinth were 34.62%, 14.78% and 9.68% respectively. It was reported that the increased fungal biomass was due to the increased of reducing sugar content in the culture media [52].

Identification of Fungal Species

The colony morphology of mycelium by light microscope identified fungus isolate was *Mucor coccinellids* as shown in Table 3: Fatty acid composition of extracted total lipids from *M. circinelloides* grown in media containing molasses as carbon sources.

| Fatty acid               | Concentration (%) |
|--------------------------|-------------------|
| CAPRIC ACID (C10:0)      | 1.57%             |
| UNDECANOIC ACID (C11:0)  | 1.40%             |
| LAURIC ACID (C12:0)      | 1.96%             |
| MYRISTIC ACID (C14:0)    | 3.95%             |
| PALMITOLEIC ACID (C16:1 Ω 7) | 4.09%        |
| HEPTADECENOIC ACID (C17:0) | 1.04%        |
| STEARIC ACID (C 18:0)    | 4.86%             |
| OLEIC ACID (C18:1 Ω 9)   | 35.08%            |
| LINOLEIC ACID (C 18:2)   | 9.09%             |
| PALMITIC ACID (C 16 : 0) | 13.78%            |
| VACCINIC ACID (C18:2 Ω 7) | 1.04%        |
| GAMMA LINOLENIC ACID (C18:3 Ω 7) | 18.85%       |
| RUMENIC ACID (C18:2 Ω 7) | 2.48%             |

Table 3: Fatty acid composition of extracted total lipids from *M. circinelloides* grown in media containing molasses as carbon sources.

![Figure 11: GC chromatogram of fatty acid methyl esters extracted from of *mucor circinelloides*](image-url)
Figure 14. As well as, the 18S rDNA gene sequence was applied and confirmed that fungal isolate as *Mucor coccinellids* (1382 bp) using BLASTn on NCBI and showing the homology sequences with different percent. The nucleotide sequence of *Mucor coccinellids* containing internal transcribed spacer 1; 5.8S ribosomal RNA gene and internal transcribed spacer 2, complete sequence; and large subunit ribosomal RNA gene, partial sequence. The fungal isolate was submitted on the international NCBI/EMBL/DDBJ/Genbank databases with accession number MF078484.1 as *Mucor coccinellids* strain MC7001 (www.ncbi.nlm.nih.gov/nuccore/MF078484.1). The neighbor-joining phylogenetic tree, Figure 15, was constructed with these homologous sequences. The filamentous fungus was identified as *Mucor coccinellids* belonged to family Mucoraceae.

**Conclusion**

The dairy industry is the one of the leading industry in the
world. It can be concluded that the integrated treatment system designed here was effective for treatment of dairy wastewater by microbial isolates and natural coagulant chitosane. According to the results, fungal isolate is the most effective organisms than bacterial isolate to reduce the COD and BOD concentration more 82% and 98.6%. These microbial isolates can be efficiently applied for the biological treatment of the dairy effluent treatments. Microbial isolates uses the organics in Wastewater as a food supply so that water pollution threats due to dairy effluents can be reduced through biological methods and this will enable the recycling of water. On using chitosane silver nanoparticles in the pilot unites, it gives 90.3%, 82%, 98.6% and 76% removal efficiencies for TSS, COD, BOD and Ammonia, respectively. Antibacterial effect of chitosan-Ag nanoparticle materials increased with increasing time. *M. circinelloides* can be grown on agricultural wastes it can be used for producing cheap microbial oil and can control environmental pollution. Higher biomass and lipid content was obtained when the filamentous fungi was cultured in the media containing molasses. The present investigation supported that *M. circinelloides* MC7001 can be cultured in molasses as carbon source for biodiesel production. Presence of palmitic (16:0), oleic (18:1), stearic (18:0) and linoleic (18:2) acids strongly supports that the fungal can be used for the biofuel production. Finally, production of biodiesel at low cost opens the door for replacing fossil fuel and avoiding its environmental, health, and economic problems.

**Conflict of Interest**

The authors declare no conflict of interest.

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