AN INSIGHT ON ALGAL CELL DISRUPTION FOR BIODIESEL PRODUCTION

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ABSTRACT

Objective: This review article deals with the effect that various cell disruption techniques have on the efficiency of lipid extraction. We have reviewed existing algal cell disruption techniques that aid the biodiesel production process.

Methods: Current rise in demand for energy has led the researcher to focus on the production of sustainable fuels, among which biodiesel has received greater attention. This is due to its larger lipid content, higher growth rate, larger biomass production, and lower land use. Extraction of lipid from algae (micro and macro) for the production of biodiesel involves numerous downstream processing steps, of which cell wall disruption is a crucial step. Bead milling, high-pressure homogenization, ultra-sonication, freeze-drying, acid treatment, and enzymatic lysis are some methods of cell disruption. The cell disruption technique needs to be optimized based on the structure and biochemical composition of algae.

Result: The lipid extraction efficiency varies depending on the algal species and the cell disruption technique used.

Conclusion: In-depth research and development of new techniques are required to further enhance the cell disruption of the algal cell wall for the enhanced recovery of lipids. In addition, the operating costs and energy consumption should also be optimized for the cost-effective recovery.

Keywords: Biodiesel, Cell disruption, Microalgae, Macroalgae.

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INTRODUCTION

Dependence on fossil fuels for energy production is now diminishing due to its rapid depletion and emission of environmentally toxic gases. For economic and environmental sustainability, it is necessary to explore alternative carbon neutral and renewable fuel [1]. Biodiesel is one among such sustainable source and carbon neutral fuel substitute for fossil fuels. Conventional sources for biodiesel production such as oil reserve crops, animal fat, and other sources cannot satisfy the increasing demand for fuels [2]. According to many reports, waste cooking oil and algae could be the promising feasible sources for the production of biodiesel to meet the worldwide energy demand [3].

Algae are capable of producing high quantity of lipid which can be used for biodiesel production [4]. Culturing of algae near to saline or brackish water region can minimize the use of land and water [5]. Algae help in reduction of carbon dioxide emissions by converting them into glucose. This glucose is converted to fatty acids for the synthesis of membrane [6]. Under stress conditions, this fatty acid is converted into lipids which can be used for production of biodiesel. Several algal species produce different types of lipids, hydrocarbons, and other complex oils. The total oil productivity, i.e., the total mass of oil that is produced per unit volume per day depends on the oil content in the biomass and growth rate of algae [7]. The growth rate of microalgae and macroalgae differs from each other. Extensive research is being carried out on macroalgae as they contain novel lipids and fatty acids. Jeong et al. have studied the efficiency of different pretreatment technique for the extraction of lipids from macroalgae. Effect of pH on disruption efficiency was studied on various macroalgae such as Ulva rigida, Polysiphonia strictissima, Enteromorpha intestinalis, and Porphyra species [8]. In general, the cell wall of algae is thick and composed of fibrillar matrix and crystalline polymers which cause hindrance in the extraction process of lipids. Its composition differs from each other and, hence, different disruption methods need to be developed. For example, the cell wall of Chlorella vulgaris predominantly consists of saccharides and hemicellulose for which grinding using liquid nitrogen has shown significant results [9]. The choice of the cell disruption technique depends on the type of algae, its cell wall structure, and composition. Hence, in this review, we have discussed about the algal cell wall structure, summarized on biodiesel production technique, and detailed on cell wall disruption methods.

ALGAL CELL WALL STRUCTURE

Algal cell wall is similar to that of plant cell wall and is generally trilaminar. An organized microfibrillar structure is embedded in a continuous matrix [10]. It consists of high protein content compared to plant cell wall, with major portion constituted of glycoprotein. 45% of the cell wall is made of cellulose, also containing other carbohydrates such as hemicellulose, and limited quantities of fructose, rhamnose, and glucose. It also has algaeenan, which is a resistant biopolymer. Figs. 1 and 2 represent algal spore cell wall and algal gelatinous cell wall composition, respectively. Based upon the composition of the cell wall, disruption technique is experimented. Algae produce a large quantity of oil, especially under stress and store it in between cell wall and membrane of the cell. The oil yield of algal cells varies from species to species.

PROCESS OF BIODIESEL PRODUCTION

Conversion of wet algal biomass into combustible fuel is a challenging process. Once the algal biomass is harvested, the processing of biomass takes place in various steps. Figs. 3 and 4 represent lipid content in various macroalgae and microalgae species. For extraction of energy-rich compounds (triglycerides), it is first dehydrated and then made to react with a solvent like hexane. The extracted triglyceride is reacted with methanol in the presence of catalyst in a reaction known as alcoholysis or transesterification to produce glycerol and biodiesel [11]. The following chemical reaction depicts transesterification reaction of lipids present in algae for biodiesel production [12,13].
The major steps involved in the generation of biodiesel using algae are the cultivation of algae, harvesting, lipid extraction by cell disruption, and lipid transesterification [14]. Fig. 5 represents the process of biodiesel production. Under optimal condition, green algae could double its biomass in a day with approximately 50% of lipid content [15,16]. The high-density biomass leads to an increase in the biodiesel production [13]. Although each one of these steps are important, cell wall disruption is particularly imperative, as the constituents of the extracted lipids are determined with respect to the disruption technique. Moreover, the challenge is microalgae are small in size and its surface is covered with a thick cell wall. The interested products are in general situated in globules or bound to cell membranes, making extraction more difficult. Hence, the use of relevant cell disruption strategy and method plays a major role in increasing the lipid extraction efficiency.

ALGAL CELL WALL DISRUPTION METHODS

A wide range of disruption methods are available for the disruption of the cell wall. They are classified into two main categories based on working mechanism which are mechanical and non-mechanical methods.

MECHANICAL DISRUPTION METHODS

Non-specific cell wall disruption is achievable by mechanical forces such as liquid-shear forces (employment of high-pressure homogenization and microfluidization), solid-shear forces (use of bead mill and high-speed homogenization), exchange of energy through waves (use of ultrasonication and microwave), electric current (application of pulsed electric field), or heat treatment/thermolysis [17]. Table 1 summarizes the effect of different techniques on lipid extraction from various algal species. Mechanical methods have higher efficiency compared to other methods as they do not depend on the species of algae to be processed and the chances of contamination of the lipid product extracted are low [18].

Bead milling

Bead mill method includes cell wall breakdown by agitated beads. This method leads to a direct damage that is induced by the highspeed spinning of fine beads along with the biomass slurry as represented in Fig. 6 [19,20]. Disruption depends mainly on the residence time of the beads in the system [21]. Other factors include bead size, cell size, and its strength [22]. Study on disruption of Chlorella vulgaris, Neochloris oleoabundans, and Tetraselmis suecica revealed that rate of release of intracellular carbohydrates and protein was higher with minimum energy consumption for smaller sized beads [23]. The rate of cell disruption is also directly proportional to volume ratio of the beads to that of cell suspension [24]. As the beads settle due to gravity, the extract could be easily removed by pipetting [25]. This technique is generally performed under laboratory scale. For large scale purposes, a dyno-mill is used, which has been successfully used for microalgal cell disruption. It utilizes rapidly rotating and notched discs for exciting the beads [26,27]. When biomass concentrations between 100 and 200 g/L are used, the method is effective with energy utilization [10,11]. The disadvantages are up scaling the process as it requires an extensive cooling system for the prevention of thermal degradation of the product [28]. Bead milling proved to be the most efficient cell disruption technique for C. protothecoides with lipid recovery of 18.8% [29].

High-pressure homogenization

In this method, cell suspension is pumped with a high pressure. In an accelerated cellular jet, suspension is impinged on the stationary valve surface causing shear stress due to the pressure drop [15]. The
Specific care has to be taken to minimize the damage that might be maximized as there is a large availability of the designs of valve seat. The cell disruption can be extent to which the disruption takes place depends on the pressure applied and the strength of cell wall [22]. The cell disruption can be maximized as there is a large availability of the designs of valve seat. Specific care has to be taken to minimize the damage that might be caused by the effects of cavitation [32]. This technique offers various advantages which includes lower cooling cost, lower heat formation, no dead volume in reactor, easy scale up, and low risk of thermal degradation. Although the homogenizers have many advantages, it makes the use of a large amount of energy [28]. This technique can rupture even the most hard-surfaced algae like Nannochloropsis [33] and can be utilized to process pretreated concentrated paste [34] but for application in large-scale processing, the energy consumption should be considerably low. The scale-up reduces processing capacity due to the increase in homogenizing pressure [35]. Successful disruption was achieved after the Chlorococcum cells were homogenized at a pressure of 850 bar [36]. Similarly, 8.5 times more oil was extracted using this technique from Nannochlorops oculata [37].

**High-speed homogenization**

A stirring device in this instrument works at a higher revolution per minute (rpm) and comprises a stator and a rotor assembly made of stainless steel. This technique incorporates hydrodynamic cavitation that is caused due to mixing at a higher rpm and shear stress formed at the solid–liquid interphase. At the point when the critical rpm (8500) value is reached by the impeller tip, hydrodynamic cavitation is caused, which diminishes the surrounding pressure to the level of vapor pressure of the fluid. This causes the fluid to move away from the impeller. The fluid pressure is then reestablished which bring the fluid toward the impeller and that collapses the cavities. It is an easy, effective, but an aggressive technique. This method incorporates the possibility to process the suspensions with a relatively higher dry cell weight concentration (2–6% w/w) with the short contact time, thereby lessening the water footprint and the cost of downstream processing. Wang et al. and Balasubramanian et al. reported lipid extraction from Nannochloropsis sp. up to 76 % and 38 ± 2%, respectively, using high-speed homogenization method [38,39].

**Ultrasonication**

Ultrasonication is an alternative method to overcome the problems faced in the conventional methods. It has simple working setup conditions which gives significant purity to the product. This technique

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**Table 1: Effect of different cell disruption techniques on lipid extraction**

| Algal species     | Techniques                  | Conditions                | Result                                      | Reference |
|-------------------|-----------------------------|---------------------------|---------------------------------------------|-----------|
| Scenedesmus dimorphus | Bead milling, Ultrasonication | 1 mm glass beads, processed for 2 min, 100 watt ultrasonic processor, runtime ~ 2 min | 20.5% of dry weight lipid content recovered | [29]      |
| Chlorella sp.      | Bead milling                | 3500 rpm                  | 0.15 g lipid content per 0.5 g/dry weight    | [55]      |
|                   | Microwave                   | 2450 MHz                  | 0.18 g lipid content per 0.5 g/dry weight    | [55]      |
|                   | Ultrasoundication            | 50 Hz, 15 min             | 0.2 g lipid content per 0.5 g/dry weight     | [55]      |
|                   | Enzymatic lysis - cellulase  | 5 mg/L, 55°C, 10 h        | Lipid concentration was 22% of dry weight    | [9]       |
|                   | Enzymatic lysis - lysozyme  |                          | Lipid concentration was 24% of dry weight    | [9]       |
| Nannochloropsis   | Microwave                   | 60°C                      | 40% biodiesel yield                         | [72]      |
| Botryococcus sp. MCC31 | Microwave, Sonication      | 20 kHz                    | 20% biodiesel yield                         | [73]      |
|                   |                              | More than 100°C, 6 min    | 48.33% of dry weight recovered              |           |
| Ankistrodesmus falcatus | Pulsed electric field treatment | 45 kV, 360 ns 1/e pulse duration | Doubles the extraction efficiency of lipids | [74]      |
requires lesser time to run with higher reproducibility, operational at lower temperature, and requires overall less energy input [40].

The mechanism of cell damage includes acoustic streaming and cavitation. Once the ultrasound is applied, there is the production of microbubbles known as cavitation, which generates pressure on the cells causing them to disintegrate [41]. The mixing of the sample is facilitated by acoustic streaming [42]. The energy that is released by the high frequency waves initiates the cavitation process, and propagation of shock waves. This causes the cell disruption. Sonicators are mainly of two types, as described by Lee et al. which include the bath and horn type. The latter makes the use of the piezoelectric generator while the former has transducers that create ultrasonic waves [43]. During the course of the treatment, there is a generation of rapid compression/decompression cycles leading to stable and transient cavitation. The transient cavitation caused by the unsteady oscillations implodes producing heat shock waves. This causes the disruption of microgalal cells, leading the cavitation to crack the membrane and the cell wall [44-46]. The collapse of bubble leads to increased mass transfer and microstreaming, aiding the increase of lipid extraction efficiency [47]. However, free radicals are produced when ultrasonication process is prolonged, decreasing the oil quality to be extracted [48]. The capability of bubble activity in cells disruption relies on ultrasound intensity, characteristics of the air bubble, and the relative nearness of the cells to the bubble [49]. Sheng et al. have reported 30% increase in yield of lipid after treatment of synechocystis PCC 6803 by ultrasonication method [50].

**Microwave treatment**

Microwave irradiation technique is one of the promising technologies being used in the extraction process of biodiesel production. This technique consumes less energy with high disruption efficiency and short processing time. The factors that affect the microwave-assisted extraction are dielectric properties of process mixture, process time, solid-liquid ratio, temperature, and type of solvent [51,52]. Efficient heating systems like microwave processing in combination with solvents such as hexane and ethanol can reduce the energy consumption in addition to the minimal use of solvent and is also economical. Continuous microwave systems for oil extraction from *Scenedesmus obliquus* were significantly affected by time and temperature. However, quality of the oil was reported high [53].

The cost of heating using a microwave is two-thirds in comparison to the conventional methods of heating. In addition, biodiesel production could also be increased using this technique. Radio frequency microwave energy provides various advantages like improvement in rate of reaction thereby leading to better separation. The use of non-contact heat source offers various advantages such as increase in energy transfer, selective heating, equipment size reduction, and quick startup. Evidently, microwave treatment could prove to be a promising technology to obtain maximum yield compared to conventional techniques [54]. It can be observed from experiments performed on *Chlorella sp.*, *Tolyphothrix* sp., and *Nostoc* sp. that microwave treatment is one of the most efficient cell disruption methods and unlike other methods, gives same efficiency of lipid extraction for different species of algae [55].

**Pulsed electric field treatment**

In pulsed electric field method, an external electric field is applied that initiates critical electrical potential charge along cell wall or membrane. Electromechanical compression and electroporation induces tension that leads to the formation of pores in the wall or membrane [56]. The number and size of pores are proportional to pulses and the electric field strength. The electroporation can be either reversible or irreversible [57]. In general, increase in conductivity leads to metabolites/compounds release from the disintegrated cells which in turn increases the temperature. This leads to decreases in the efficiency of cell disruption, and hence, this method is less preferable than other methods for extraction of lipid [43]. Eng et al. have demonstrated that the lipid yield increased 9 times when the algae *Auxenochlorella protothecoides* was pretreated by pulse electric field method before solvent extraction [58].

**Non-mechanical disruption methods**

Non-mechanical cell disruption techniques consume less energy when compared to mechanical methods [59]. The methods are more gentle and specific but are difficult to scale up to industrial levels. Non-mechanical methods include physical methods such as freeze-drying and osmotic shock, and chemical methods like enzymatic cell lysis and treatment with chemical agents which involve the permeabilization of cell wall by binding with specific cell wall components. Vogels et al. have demonstrated that combining non-mechanical pretreatment methods like enzyme lysis or heat treatment with other mechanical techniques increased the cell wall disruption efficiency [60].

**PHYSICAL DISRUPTION METHODS**

**Freeze-drying method**

Freeze-drying makes lipid extraction from algal biomass easier [27]. It ensures that there is no loss of lipids, as they are volatile due to evaporation [61]. In this method, the wet biomass is frozen at −84°C under vacuum to crystallize the intracellular water [62]. After freeze-drying, cell can be directly lysed by allowing the ice crystals to expand by thawing. It can be combined with other methods like grinding ultrasonication or microwave to increase the yield efficiency [63]. If the cells are freeze-dried before bead milling, lipid recovery is more due to enhanced specific area and reduced diffusion gradient [6-4]. Although this method helps in increasing the efficiency, it is very expensive and has high energy consumption.

**Manual grinding**

Manual grinding could be performed in different methods. The first method includes harvesting of microalgae sample into a ceramic mortar, followed by the addition of liquid nitrogen and allowing the sample to thaw and then grounded by a pestle. The second method includes addition of Quartz sand to the sample and grounded directly. The third method includes drying the sample at 60°C for 7-8 h and grounded with Quartz sand [9].

**CHEMICAL METHOD**

**Sulfuric acid treatment**

Pretreatment of algal biomass for lipid extraction can be performed by sulfuric acid treatment. It is carried out by mixing the concentrated sulfuric acid (3-8%) with the algal culture and autoclaving at high temperature (120°C-160°C) for 15-45 min. This type of acid treatment helps in the hydrolysis of polysaccharide carrageenan layer by chain depolymerization and hydrolysis of sulfate moiety [65]. Experiments conducted by Halm et al. suggests that the relative concentration of lysed cell is more when treated with high volume of concentrated sulfuric acid followed by thermal treatment at high temperature [66]. Although this technique has good efficiency and low energy requirement, it is not widely used as there are high chances of product degradation due to the harsh conditions.

**Enzymatic method**

One of the most commonly used alternatives for mechanical disruption method is enzymatic lysis. This technique is specific for each microorganism to be lysed and helps in release of specific product [67]. Algal cell walls are strong and stable due to the presence of polysaccharides such as cellulose and hemicellulose. Lysing these cells with mechanical methods is energy intensive and use of enzymes lowers energy requirement [68]. Membranes of the lipid sac are also made of phospholipids. Thus, the common enzymes used for algal cell wall disruption are cellulase and lipase [69].

![Cellulose](Cellulase) → Glucose

![Phospholipid](Lipase) → Glycerol

Treatment of *C. vulgaris* with lysozyme and cellulase resulted in lipid concentration of 22% and 24%, respectively, and had the highest
efficiency among all enzymatic treatment [Zheng et al] [9]. Algal cell wall is made of many complex layers and lysis of all these layers with just cellulase and lipase is not possible. Thus, a mixture of crude enzymes from sources like fungi is required for better efficiency. Like other non-mechanical method, this method can be used either independently or as a pretreatment step for mechanical methods to increase the efficiency and decrease the energy requirements. The major drawback of this method is the unavailability of the large quantity of enzymes required for industrial scale. Chong et al. reported that employment of alkaline pretreatment followed by enzymatic treatment resulted in 90% lipid extraction from Nannochloropsis sp. [70]. Lipid recovery of 92.6% was obtained when the enzymatic disruption technique was carried out in combination with mechanical methods (high-pressure homogenization) [71].

CONCLUSION

Different cell wall disruption techniques were found to be efficient for microalgae and macroalgae. Cell disruption efficiency toward extraction of lipid varies according to the method employed and species. Microwave method and pulsed electronic field method were found to be the most suitable techniques. For the techniques to be feasible, the operating costs and energy consumption should be optimized to ultimately fulfill the major goal of superior quality products and their easy recovery. In-depth research and development of new techniques are required to further improve the cell disruption of the algal cell wall for the enhanced recovery of lipids.

CONFLICTS OF INTERESTS

The authors declare that there is no conflict of interests regarding the publication of this paper.

AUTHORS CONTRIBUTION

The authors contributed equally to this work.

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