Extracting Edible Oil from *Nannochloropsis oculata*: A Functional Food for Future

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Abstract Microalgae is known to grow at high rates and thrive in the non-arable land. It is also the most favorable feedstock for an affordable and sustainable supply of food and oil which can produce significant oil yield in less area. Its oil is considered as functional oil with great commercial potential, rich in beneficial fatty acids composition. Data from archaeological discoveries indicate that humans have developed on a diet that had a ratio of about 1/1 of omega-6, (ω-6) to omega-3, (ω-3) fatty acids. Western and Malaysian diets are deficient in ω-3 as both have greater ω-6/ω-3 ratio. In this study, Nannochloropsis oculata, (N. oculata) is used as an alternative source for edible oil application and the lipids are extracted via both conventional extraction (CE) and microwave-assisted extraction (MWE). The lipid yield was determined by the gravimetric technique and GC-FID was used to identify the fatty acid methyl ester profiles. Findings revealed the CE technique is more appropriate to be used when extracting lipids from N. oculata for edible oil production. Both techniques are successful in extracting the two most important types of fatty acids (C16 and C18). The lipids produced also comprise of specific valuable fatty acids such as arachidonic acid (ARA) and eicosapentaenoic acid (EPA) but do not produce trans-fatty acid. This edible oil obtained has a healthier ratio (~ 1) of ω-6/ω-3 for the human diet. Thus, N. oculata has the potential to be used as a source of edible oil.

Keywords Edible Oil, Fatty Acids, Lipid, Marine Microalgae, Microwave, *Nannochloropsis oculata*

1. Introduction

Several sources of evidence indicate that human beings developed on a diet that had a ratio of about 1/1 of ω-6 to ω-3 fatty acids; while today, Western diets have a ratio of 10/1 to 20-25/1 and Malaysian diets is about 3.5/0.35, i.e. 10 [1] suggesting that compared to the diet on which humans evolved and their genetic patterns were formed, Western and Malaysian diets are deficient in ω-3. The essential fatty acids such as ω-3 and ω-6 are not interconvertible in the human body. One of the ways that can remedy this is through the usage of cooking oil which comprises of balanced ω-6/ω-3 besides including fresh foods rich in ω-3 in the household food basket [2]. Moreover, deforestation making way for oil palm plantation has led to climate change. In view of this issue, many attempts were conducted to overcome these problems. In line with these efforts, this study proposes the use of microalgae as an alternative source for cooking oil. Microalgae is potentially a good source of food because it is fast growing, has high crop yield per area, high efficiency in carbon dioxide capture and solar energy conversion. Microalgae are not a food competitor with agricultural crops as they can be produced concomitantly.

with other harvest value co-products [3]. Microalgae can also be used in the diet of animals and humans. Microalgae are promising sources for novel products and applications [4]. Microalgae offers an interesting combination steps between cultivation and wastewater treatments [5] which are capable in generating valuable products such as high quality biodiesel feedstock [6]. Most of the conducted research showed microalgae as a promising resource feedstock for biofuel [7] and biodiesel [8]. Many species of microalgae comprise of lipid [9], protein [10], and carbohydrates [11]. Since lipid is important for oil production, attempts were conducted to extract lipid from microalgae. Lipids can be extracted by using methods such as mechanical, solvent, subcritical water extraction, supercritical fluid extraction (SFE) and hydrothermal liquefaction (HTL).

One of the well-known extraction methods for lipid is solvent extraction, also known as conventional extraction (CE). This method is very effective and can be up scaled easily giving high yield products. Moreover, it is also known as a simple and inexpensive extraction technique [12]. Solvent extraction technique has also been applied in large-scale vegetable oil production which is economical and efficient. Recently, several assisted techniques were introduced for solvent extraction such as ultrasonic and electric pulse [13] to enhance lipid yield. The microwave treatment is also capable of assisting solvent extraction. Microwave exposure helps to rapidly rupture microalgae cell wall which stored lipid, oil and other valuable components. Although microwave-assisted extraction (MWE) has been used on microalgae by many researchers, these works were mostly focused on biofuel and biodiesel application.

In a previous study, Wahidin, Idris & Shaleh [14] used a mixture of ionic liquid and methanol in their microalgae extraction and found that an increase in reaction time (15 mins) has a positive impact on the amount of FAMEs that can be obtained from the wet microalgae. However, it was reported that under the influence of mixture of ILs – methanol solvents and microwave, a complete absence of arachidic acid methyl ester (C20:0) and linolenic acid methyl ester (C18:3) were observed; the long-chain arachidic acid methyl ester (C20:0) tends to break down to shorter chain of FAMEs. Teo et al. [15] studied the influence of microwave heating in the extraction of lipids from marine microalgae for biodiesel and it was revealed that the microwave irradiation appears to generate a low percentage of the methyl ester with a C22:0 carbon chain relative to the traditional method. The absence of methyl esters with >19 carbon chains ensures superior biodiesel with low viscosity. In all of these studies on microwave exposure for biofuel and biodiesel application, it is found that time frame and extraction techniques used will affect the lipid yield and fatty acid profile. However, there are limited studies on the effect of microwave on the quality of lipids produced for functional oils from microalgae.

Microalgae also contain many important sources of dietary ω-3 such as eicosapentaenoic (EPA) and docosahexaenoic (DHA) as well as ω-6, i.e.: arachidonic, (ARA) which are favorable components for edible oil. Such types of fatty acids consist of long carbon chains which are normally found from marine microalgae species. *Schizochytrium limacinum* is one of the marine *thraustochytrid* (recently classified as marine microalgae) [16] that can synthesize lipids with a high content of DHA [17]. Meanwhile, *Nannochloropsis oculata*, *N. oculata* is capable of exhibiting high EPA content [18]. The EPA diet substantially reduced tumor multiplicity, incidence and tumor size on both the promotion and initiation of carcinogenesis [19]. All these three fatty acids (EPA, DHA and ARA) play important roles in regulating physiological processes impacting human health, as nonesterified fatty acids, esterified (membrane-associated) fatty acids, or oxidized fatty acids [20]. *N. oculata* is comprised of beneficial sources of ω-3 and its derived oil has been determined safe for dietary supplement application [21]. Moreover, *N. oculata* composition and its toxicological tests support the conclusion that this microalga shows the absence of toxicity [22]. The existence of long carbon chain such as fatty acids which contain ω-3 and ω-6 are also important and believed to give benefits for health performance. The balanced ratio of ω-6/ ω-3 is one of the significant dietary indicators to qualify as healthy edible oil. Besides that, the ratio of total polyunsaturated fatty acids to total saturated fatty acids (TPUFA/ TSFA) must be in the range of 0.8 – 1 as recommended by WHO [23]. A pooled analysis of eleven cohort studies of dietary fat and coronary disease confirmed that a diet higher in PUFA and lower in saturated fatty acids (SFA) reduced risk of fatal coronary heart disease (CHD) [24].

A study revealed that the *Nannochloropsis* genus microalgae, i.e., *N. oceanica*, contain TPUFA/TSFA ratio which almost meet the recommended ratio set by WHO and is capable of producing valuable fatty acids such as ARA and EPA [25]. However, the ω-6/ω-3 ratio of *N. oceanica* did not meet the WHO recommended requirement. Besides the work of Huang et al. [25] who studied the potential of microalgae for edible oil production, not much work has been done to further explore on the quantity and quality of lipids produced from microalgae for edible oil applications. Therefore, the objective of this study is to explore the potential of *N. oculata* which has never been used as an alternative source for edible oil application. The suitability of both CE and MWE techniques to produce lipid and fatty acids were assessed taking into consideration the time frame. Best practices of lipid and fatty acids production were also reported in this article.
2. Materials and Methods

2.1. Nannochloropsis Oculata, N. Oculata

*N. Oculata* powder was purchased from Xi’an Nate Biological Technology Co., Ltd. 1 g of *N. Oculata* powder was used for each sample in extraction and transesterification techniques.

2.2. Fourier Transfer Infrared (FTIR) Analysis

The *N. oculata* powder sample was characterized using a Fourier Transfer Infrared Spectrophotometer (Perkin Elmer). The sample was subjected to FTIR spectral measurement in the frequency range of 4000 – 370 cm⁻¹.

2.3. Conventional Extraction, CE – Modified Bligh-Dyer (Step 1)

1 g of *N. oculata* dried powder was mixed with 25 ml methanol, 12.5 ml chloroform and 5 ml of distilled water in a 100 ml of the conical flask. This mixture was mixed by placing it on the orbital shaker, (IKA® KS 130 basic) at 160 rpm for 5 minutes. Another 12.5 ml of chloroform and 12.5 ml 1.5 % w/v sodium sulphate were added to the mixture [26]. The conical flask was placed on the orbital shaker for another 5 minutes at 160 rpm of rotation frequency. Total mixing period was 10 minutes (5 minutes before and after addition of 12.5 ml of chloroform and 12.5 ml 1.5 % w/v sodium sulphate). This mixing period (10 minutes) will be used for MWE as well as 2 minutes for comparison study. The mixture was separated after 24 h in a separator funnel. The bottom layer which comprised of lipid and its solvent was extracted out into round flask. The solvent was evaporated by using a rotary evaporator and the remaining lipid was collected. The lipid yield was calculated using Equation (1).

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\text{Lipid yield (\%) = } \frac{M_f}{M_i} \times 100
\]

where \(M_i\) is mass of lipid whilst \(M_f\) mass of *N. oculata* powder. This sampling was triplicated at different mixture times to observe its effect towards lipid yield more accurately.

2.4. Microwave-assisted Solvent Extraction (Step 1)

1 g of *N. oculata* dried powder was mixed with 25 ml methanol, 25 ml chloroform, 12.5 ml of 1.5 % w/v sodium sulphate and 5 ml distilled water in 100 ml of the 3-necked mini round flask. This mixture was then subjected to the microwave irradiation at 250 W and was operated for 2, 4, 6, 8 & 10 minutes at 50 °C under 160 rpm of rotational frequency. The microwave irradiation was performed in MAS-II PLUS, Microwave Synthesis Workstation. The mixture was separated after 24 h in a separator funnel. The bottom layer comprised of lipid and it’s solvent. The solvent was evaporated by using rotary evaporator and the remaining lipid was collected and its yield was calculated using Equation (1).

2.5. Transesterification (Step 2)

The lipids extracted from Sections 2.3 and 2.4 were then allowed to undergo transesterification by mixing the lipid with sulphuric: methanol (1:100 v/v). The solution was then heated in a water bath, (WiseBath®) at 80 °C for 1 h at 50 rpm and then cooled at room temperature. n-hexane and distilled water were then added to the cooled solution and were subsequently vortexed, (IKA® VORTEX GENIUS 3) for 5 minutes before centrifugation (Hermle labotechnik GmbH, 2323 K) at 4500 rpm for 10 min [25]. The fatty acids methyl esters, FAMEs with solvent were finally collected from the upper layer. The solvent was evaporated by using a rotary evaporator.

2.6. Fatty Acids Methyl Ester Characterization

The collected FAMEs in the round flask from rotary evaporator was then diluted with dichloromethane and filtered through 0.22 μm PTFE syringe filter. The FAMEs profile identification was analyzed by gas chromatography with flame ionization detector (GC-FID) (Agilent Technologies, 7820A) and HP-88 capillary column (60 m x 0.25 mm x 0.2 μm). The gas chromatographic separation was performed with the following instrumental conditions: Injector temperature: 260°C, Carrier gas: Hydrogen at 0.8 mL/ min, Split ratio: 40: 1, Oven program: 50°C (hold 1 min), to 150°C (hold 5 min; 10°C/ min), to 215°C (2.8°C/ min) to 230°C (hold 8 min; 5°C/ min). Detector: Flame Ionization Detector, Temperature: 275°C, Hydrogen Flow: 40 mL/min, Air Flow: 450 mL/min and Injection Volume: 1 μL. The samples were quantified by area percentage calculation using Supelco 37 Component FAME Mixture.

2.7. Morphology Characterization

The *N. oculata* from MWE and CE were subjected to morphological analysis to examine changes of cell wall damage. A small amount of sample was taken and observed under a Field Emission Scanning Electron Microscope (FESEM), Hitachi SU8020.

2.8. Statistical Analysis

Single factor analysis followed by Turkey test was used to analyze data from lipid yield and fatty acids profile study. All the ANOVA were tested at 95% confidence level.
3. Results and Discussion

3.1. Nannochloropsis Oculata, N. Oculata

The FTIR was conducted to estimate the existence of lipid content in N. oculata powder. Figure 1 shows 14 distinct transmission bands over wavenumber range of 4000 - 370 cm⁻¹ for N. oculata powder. Based on numbers of the previous studies, most of FTIR analysis detected the presence of lipid at a wavenumber of 1723 – 2916 cm⁻¹ [27], [28]. Based on the obtained FTIR spectrum, the lipid existence was detected at 2918 – 2850 cm⁻¹. Thus, the FTIR result proved the existence of lipid in N. oculata.

![Figure 1. FTIR Spectrum of Nannochloropsis oculata, N. oculata](image)

3.2. N. oculata Characterization

Figure 2 shows the lipid yields of N. oculata obtained using the conventional and MWE technique at different mixing time. It was proven statistically that reaction time played a significant role in lipid yield (p < 0.05). Based on Figure 2, higher lipid yield (15.0 %) was obtained using CE compared to MWE (13.0 %). The Bligh-Dyer solvent extraction method was used for both techniques. In the MWE the extraction was assisted by microwave at 50°C, 250 W whereas the CE was performed at room temperature with no assistance. The results obtained contradicted to previous research findings where higher lipid yield was attained when MWE was used to be compared with CE [29]. It was believed that 10 minutes of microwave exposure onto N. oculata could be too long which possibly caused the decrement of its lipid yield. An additional MWE was conducted on N. oculata at reaction time of 2, 4, 6, 8 and 10 minutes. Based on this experiment, 2 minutes of microwave exposure shows the highest lipid yield, 48.0 %. It also shows that at eight minutes, lipid yield of N. oculata started to decrease from 24.0 % until 13.0 % for 10 minutes. Based on these results, it shows that less time of microwave exposure gives higher lipid yield for N. oculata. However, the findings also showed that MWE obtained higher lipid yields compared to CE at 2 minutes [30].

Microwave is believed to cause an increase in the heating rate of extraction process which caused higher lipid production compared to conventional extraction. Such phenomena have been observed by other researchers Teo & Idris [26] and Wahidin, Idris & Shaleh [31]. The presence of the polar compounds when performing the MWE also helps in increasing the heating rate of the extraction process. In this experiment, polar solvent and water were used. The combination effects from polar compounds and oscillating electric field from microwave generate vibration among molecules create inter- and intra-molecular friction. The effects of molecular oscillation, combination (inter-and intra-molecular) friction, collision as well as the movement of large quantities of charged ions lead to high heating rates within seconds of the N. oculata cell. Such intracellular heating is simultaneously capable of producing a pressure effect that ruptures the cell membranes. Furthermore, the electroporation effects result in the spilling out of the oil and lipids. Moreover, the energy carrier during MWE causes direct heat generation (volumetric heating) within microalgae cells. This volumetric heating creates a rapid temperature rise within the matrix, creating a pressure effect on the microalgae cell wall membrane structure. Thus, the lipids diffuse out rapidly from the microalgae cell wall into the solvent phase. Conversely, for CE minimal heat is transmitted to the substrate through convection and conduction [32]. Considering the mentioned results, reaction period and extraction technique play a significant role on lipid yield.

Similar results were obtained by one of the previous research studies by Prommuak, Pavasant, Quitain, Goto, & Shotipruck, [33] in which they conducted extraction of lipid from Chlorella vulgaris and Haematococcus pluvialis. Their findings also found that greater exposure time resulted in a decrease in lipid yield. This phenomenon is caused by lipid oxidation or lipid degradation in biomass treatment under drastic conditions [34]. Most of the previous findings on microalgae treated with microwave achieved higher lipid yield at lowest time. There was a study which achieved ideal condition at one minute of microwave at 300 W and 80 °C [35]. Another finding from Nogueira, da Silveira, Vidal, Ribeiro, & Veiga Burkert, [34] reported their lipid was efficiently extracted by microwave at 40 s. Lee, Yoo, Jun, Ahn, & Oh, [36] stated that the efficiency in lipid extraction from microalgae depends on many factors such as microalgae species and the extraction method. They have evaluated five methods of cell disruption and found the highest lipid yield contents were attained by microwave compared to osmotic shock, autoclave, ultrasound and bead milling. Ma et al. [37] also obtained higher lipid when applied microwave onto Chlorella sp. as to be compared with ultrasound. The authors attributed the findings not only to
the fast heating and humidity inside the cell but also to high pressure that was generated in the microwave process.

In this study single factor analysis of ANOVA with Turkey test was used in this study to analyze results. The lipid and FAME yields are the most important parameters as they are the indicators for healthier ω-6/ω-3 ratio for the human diet in edible cooking oil. The multivariate analysis like partial least square regression (PLSR) which can provide better analyzing results will be used in future work [38, 39].

Figure 2. Lipid yield of *Nannochloropsis oculata, N. Oculata* using conventional extraction, CE and microwave-assisted extraction, MWE at different mixing times

3.3. Fatty Acids Profile

Table 1 shows the types of fatty acid methyl ester which is detected from lipid samples from both processes CE and MWE for different time frame (2 and 10 minutes). The results revealed methyl butyrate, C4:0 methyl cis-5,8,11,14-Eicosatetraenoic (ARA) detected in CE samples at 2 minutes. However, C20:4, ω-6 was not detected using CE at 2 minutes. Some components such C14:0, C16:0, C16:1, C18:1 and C18:2 were detected in both samples of CE and MWE. It is also believed that C16 and C18 are fatty acids which are similar to vegetable oil [40] like palm oil [41]. However, it is observed that MWE attained a lowest quantity of methyl tetradecanoate, C14:0 at 2 minutes of reaction times. There was a study reported that low amounts of C14:0 is an advantage for those who are undergoing hypercholesterolemia treatment [42]. However, ARA, PUFA, ω-6, C20:4 and cis-13,16-docosadienoic acid, C22:2, ω-6 were not detected in CE samples with 2 minutes duration time. The very brief mixing time was insufficient to extract all the free fatty acids, FFAs but extending the duration to 10 minutes enables ARA, PUFA, ω-6, C20:4 to be extracted. The results surprisingly show that cis-13,16-docosadienoic acid, C22:2, ω-6 was also detected in MWE samples but not in CE samples. Fatty acids, C22:2 is a stronger inhibitor than docosahexaenoic acid, DHA and docosapentaenoic acid, DPA and it is said to have an anti-cancer effect among C22-fatty acids [43]. Unfortunately, methyl cis-5,8,11,14,17-Eicosapentaenoate, EPA, C20:5, ω-3, was not detected in MWE samples.

| Fatty acid methyl ester, FAME | CE, 2 min (%) | CE, 10 min (%) | MWE, 2 min (%) | MWE, 10 min (%) |
|-------------------------------|---------------|----------------|----------------|-----------------|
| Methyl butyrate, SFA, C4:0    | 3.0 ± 0.49    | ND⁴           | ND³            | ND³             |
| Methyl tetradecanoate, SFA, C14:0 | 9.0 ± 0.96 | 8.0 ± 0.37    | 7.0 ± 0.25     | 8.0 ± 1.14      |
| Methyl palmitate, SFA, C16:0  | 38.0 ± 0.61   | 38.0 ± 0.25   | 37.0 ± 0.83    | 36.0 ± 0.56     |
| Methyl palmitoleate, MUFA, (ω-7); C16:1 | 33.0 ± 0.85 | 34.0 ± 0.50 | 31.0 ± 0.86 | 30.0 ± 0.25 |
| cis-9-Oleic acid methyl ester, MUFA, (ω-9); C18:1 | 9.0 ± 0.87 | 9.0 ± 0.07 | 11.0 ± 0.33 | 11.0 ± 1.00 |
| Methyl Linoleate, PUFA, (ω-6); C18:2 | 3.0 ± 0.20 | 3.0 ± 0.06 | 4.0 ± 0.11 | 4.0 ± 0.10 |
| Methyl cis-5,8,11,14-Eicosatetraenoic (ARA), PUFA, (ω-6), C20:4 | ND³ | 2.0 ± 0.08 | 3.0 ± 0.34 | 4.0 ± 0.24 |
| Methyl cis-5,8,11,14,17-Eicosapentaenoate (EPA), PUFA, (ω-3), C20:5 | 6.0 ± 1.42 | 6.0 ± 0.26 | ND³ | ND³ |
| cis-13-16-Docosadienoic acid methyl ester, PUFA, (ω-6), C22:2 | ND³ | ND³ | 7.0 ± 0.87 | 8.0 ± 0.55 |

ND⁴: not detected
Apparently, CE can retain ω-3, unlike MWE where losses of ω-3 were observed. These results show that microwave radiation causes the oxidation of lipids. Methyl cis-5,8,11,14,17-Eicosapentaenoate, EPA, C20:5 is one type of ω-3 fatty acids which is very sensitive to oxidation [44]. The presence of a double bond is naturally susceptible to oxidation (electron can be ripped away to make the opportunity for additional hydrogen atoms). The more polyunsaturated fatty acids, PUFA (more double bonds), the more likely it is prone to oxidation. The hydrogen that is bound to methylene is located between two double bonds (“active methylene”), is the site of initial oxidation by an active oxygen species (see Figure 3). Consequently, PUFA with many double bonds comes with many “active methylene” groups; hence, it is easily oxidized. Compared to other PUFAs, EPA, C20:5 has the most number of double-bonds which is 5 double bonds. Therefore, EPA, C20:5 is most susceptible to oxidation among detected fatty acids in this study. This is depicted in the results obtained where ω-3 fatty acids were not detected when using MWE because they were oxidized regardless of the time frame due to the harshness of the technique. The presence of ω-3 is important to determine the ratio of ω-6/ω-3. The ratio of ω-6/ω-3 is one of the reliable indicators for healthy edible oil performance as well as the TPUFA/TSFA ratio. Hence, CE is a promising technique to be applied on N. ocularata for edible oil application instead of MWE.

![Figure 3](image-url)

Figure 3. Oxidized site of polyunsaturated fatty acids (ex: Methyl cis-5,8,11,14,17-Eicosapentaenoate, EPA, C20:5)

These results contradict those of other researchers who studied the effect of microwave on microalgae for biofuel and biodiesel application. The presence of a shorter carbon chain < 19 carbon guarantees a low viscosity which is more superior. Based on conducted study, the microwave was preferred in producing shorter carbon chains than longer carbon chain > 19 carbons. Moreover, microwave irradiation is also capable of producing reasonable cetane number, CN and iodine value, IV compared to conventional extraction techniques [45]. Iodine value, IV and cetane number, CN are other indicators for biodiesel and biofuel quality evaluation where lower IV values reflect better lubricating properties and higher CN values reflect better ignition property [46]. Thus, microwave appears to be an appropriate extraction technique to be applied on microalgae for biofuel and biodiesel but not for edible oil production.

Based on the overall results, it was proven statistically that reaction time does not cause significant changes towards fatty acids quantity (p > 0.05). However, the type of extraction technique resulted in significant changes to fatty acid profiles qualitatively. Similar results were found by Olabemiwo, Esan, Adepoju, & Omodara, [47] which claimed fatty acids of three Nigerian oils (soya oil, groundnut and palm oils) were slightly quantitatively affected by microwave radiation. Meanwhile, qualitatively, they found that only microwave treatment on groundnut oil led to the disappearance of C14:0. In another research, Shreelalitha & Sridhar [48] discovered additional essential fatty acids are present when hot-extraction was applied in dry seeds compared with other extraction techniques. Thus, their results also proved that different extraction techniques give an effect on fatty acids profiles qualitatively.

Table 2 compares the total unsaturated (TPUFA and TMUFA), TSFA and ω-6/ω-3 ratio. It can be seen from the data in Table 2 that the TPUFA, TMUFA and ω-6 from CE samples increased with longer extraction time. In contrast, only TPUFA and ω-6 from MWE samples increase with longer extraction time. Comparison study between MWE and CE shows that, MWE exhibited higher TPUFA, TMUFA and ω-6 at lower extraction time. Meanwhile, at longer extraction time, yielded higher TPUFA and ω-6. The overall findings showed that at a longer extraction time, both CE and MWE techniques developed less TSFA and more TPUFA as well as ω-6. Similar findings from previous studies showed that microwave irradiation of the three Nigerian vegetable oils for 30 minutes resulted in a marginal decrease in total SFA, resulting in a marginal increase in total unsaturated fatty acids (TUSFA) [45]. The fatty acid compositions of microalgae compared well with those of vegetable oils from conventional oil-producing crops. Palm oil's saturated and unsaturated fatty acids are about 50% each. Soybean and peanut oils are unsaturated oils with around 80% unsaturated fatty acids [49]. Thus, the percentage of TSFA and unsaturated fatty acids (TMUFA and TPUFA) in N. ocularata is in accordance with palm oil fatty acids profile which reflects its suitability for edible oil.

Table 2. GC-FID – Percentage and ratio of TPUFA, TMUFA, ω-6 and ω-3 of Nanochloropsis Oculata, N. ocularata after conversion to fatty acid methyl ester with different transesterification treatment

| Extraction Technique | CE, 2 min | CE, 10 min | MWE, 2 min | MWE, 10 min |
|----------------------|-----------|------------|-------------|--------------|
| TPUFA, %             | 9.16      | 11.04      | 13.72       | 15.40        |
| TMUFA, %             | 41.69     | 42.76      | 72.73       | 40.94        |
| TSFA, %              | 49.15     | 46.20      | 44.17       | 43.66        |
| ω-6, %               | 2.98      | 5.14       | 13.72       | 15.40        |
| ω-3, %               | 6.18      | 5.90       | ND³         | ND³          |
| TPUFA/TSFA           | 0.19      | 0.24       | 0.31        | 0.35         |
| ω-6/ω-3              | 0.49      | 0.87       | NA²         | NA²          |

ND³: not detected; NA²: not applicable - unable to calculate because ω-3 was not detected
The ω-3 was not detected in MWE but was present in CE. Hence, ω-6/ω-3 ratio for MWE cannot be determined. Apparently CE for 10 minutes achieved ω-6/ω-3 ratio of 0.87 which reflects a value close to 1 indicating healthier edible oil [50]. However, the ω-6/ω-3 ratio attained by CE for 10 minutes is still far from the recommended ratio set by the World Health Organization, WHO (5 - 10). Very high ratio of ω-6/ω-3 can cause many diseases such as cancer, cardiovascular disease, CVD, an inflammatory and autoimmune disease. Meanwhile, low ω-6/ω-3 ratio exerts suppressive effects. Moreover, increased of ω-6 fatty acids and ω-6/ω-3 ratio in red blood cell (RBC) membrane phospholipids caused the increment of obesity risk [51]. Furthermore, MWE, 10 minutes obtained TPUFA/TSFA ratio which nearly complies with WHO recommendation, 0.8 – 1.

Table 3. The comparison of TPUFA/TSFA and ω-6/ω-3 ratios for various types of edible oil sources

| Ratio | TPUFA/TSFA | ω-6/ω-3 |
|-------|------------|---------|
| N. oculata, µg/ml (CE) | 0.19-0.24 | 0.48-0.87 |
| N. oceanica, µg/ml (MWE) | 0.31-0.35 | NA † |
| Chlorella vulgaris, µg/ml [25] | 0.69 | 0.50 |
| Scenedesmus obliquus, µg/ml [25] | 1.92 | 1.26 |
| Palm Oil, µg/ml [41],[54] | 0.44 | 1.03 |
| Coconut oil, µg/ml [52] | 0.21 | NA † |

NA †: not applicable

Table 3 displays some important ratio for healthy cooking oil. TPUFA/TSFA as well as ω-6/ω-3 ratio are some of the important indicators for determining a healthy diet for edible oil [52]. The values obtained from CE and MWE samples were compared with other microalgae genus and Nannochloropsis species as well as two types of well-known cooking oil sources in the Malaysia current market (palm oil and coconut oil). Based on comparison results of TPUFA/TSFA with previous findings, N. oceanica almost meet the recommended value set by the WHO (0.8 – 1). However, ω-6/ω-3 for N. oceanica and other types of microalgae still do not meet the WHO requirement (5 – 10). Nevertheless, N. oculata managed to achieve ω-6/ω-3 ratio close to 1 which can be classified as a healthier edible oil source. This is because it fits the target ratio of ω-6/ω-3 fatty acids for appropriate human nutrition and health. Simopoulos [53] stated that the ratio of ω-6/ω-3 which is approximately 1 is determined as a healthier diet intake for human. Based on these comparison results, it can also be summarized that N. oculata are significantly different from those of N. oceanica in several key features although both of these microalgae species are in the same genus. The significant differences may be due to other aspects apart from different species such as microalgae source and cultivation techniques. Furthermore, palm [54] and coconut oil from the mentioned references did not provide the ω-6/ω-3 ratio as both edible oil sources were unable to produce ω-6 and ω-3 fatty acids. In accordance to that, the TPUFA/TSFA and ω-6/ω-3 of N. oculata, as well as its trans-fat, were being compared with current cooking oil in the market to benchmark its potential for edible oil application as illustrated in Table 4.

Table 4. Comparison TPUFA/TSFA between Nannochloropsis Oculata, N. oceanica and numbers of current market cooking oil in Malaysia.

| Type of cooking oil | TPUFA/TSFA | ω-6/ω-3 | Trans fat |
|---------------------|------------|---------|-----------|
| N. oculata          | 0.19-0.35  | 0.46-0.87 | 0         |
| Refined Palm Oil, A | 0.28       | NA †    | 0         |
| Refined Palm Olein, B | 0.27     | NA †    | 0         |
| Double Fractionated | 0.27       | NA †    | 0         |
| Super Palm Olein, C | 0.28       | NA †    | 0         |
| Sunflower Oil (with ω 3, 6 & 9), E | 5.9 | 564 | 0.3g/100g serving |
| Refined and Virgin Olive Oil, F | 0.5 | NA † | 0 |
| Refined Palm Olein, G | 0.25       | NA †    | NA †     |
| Corn Oil, H         | 3.41       | NA †    | 0.3 g/100 ml serving |
| Rice Bran Oil, I     | 1.42       | NA †    | 0.1 g/100 ml serving |
| Canola Oil, J        | 2.56       | NA †    | 0.1 g/100 ml serving |
| Canola Oil, K        | 4          | 2       | 0.4 g/100 g serving |
| Sunflower Oil, L     | 1          | 0       | NA †     |

NA †: not applicable

It is apparent from Table 4 that all palm-based cooking oil attained a similar ratio of TPUFA/TSFA which is 0.25-0.28. Among all types of cooking oil, sunflower oil, L complied with WHO standard for TPUFA/TSFA ratio. However, sunflower oil, L did not detect ω-6 and ω-3. Thus, the ratio of ω-6/ω-3 is not applicable. Sunflower oil, L also did not contain trans-fatty acid in its oil. Besides N. oculata, only sunflower oil, E and Canola oil, K provide the ω-6 and ω-3 information. Hence, the ratio of ω-6/ω-3 can be determined as listed in Table 4. The ω-6/ω-3 for N. oculata, sunflower oil, E and Canola oil, K are 0.46-0.87, 564 and 2 respectively. Based on the Table 4, N. oculata obtained the nearest value (0.87) that complied with healthier diet indicator, ~ 1 for ω-6/ω-3 ratio compared to other cooking oil [53]. Moreover, most of the cooking oil did not contain trans-fatty acid except sunflower oil E, corn oil H, rice bran oil I, canola oil J and canola oil K. Each of mentioned cooking oil content is 0.3 – 0.4 g per 100 ml or 100 g serving. The findings suggested that N. oculata is potentially a suitable alternative source for edible oil application due to the positive results from ratio of TPUFA/TSFA (0.19 – 0.35) which aligned with current commonly used cooking oil from palm-based source (0.25 – 0.28). Interestingly both techniques, CE and MWE on N. oculata did not produce trans-9- elaic acid methyl ester, trans-fatty acids. It is believed that trans-fatty acids intake was associated with the raised risk.
of coronary heart disease [55]. In fact, many cooking oil from vegetable and seed oils as depicted in Table 4 and previous findings contributed to more than 1% of trans-fatty acids. Trans-fatty acids can also be found in sunflower oil E, corn oil H, rice bran oil I, canola oil J and canola oil K. The effect of microwave radiation towards morphology of *N. oculata* can be observed via FESEM as shown in Figure 4.

### 3.4. Morphological Change

Figure 4 illustrates *N. oculata* samples before and after undergoing CE and MWE for 10 minutes. The images of Figure 4 (a) and (b) show *N. oculata* with the intact cell wall before extraction process at different magnifications (100 and 1.00kX). Meanwhile, Figure 4 (c) and (d) show FESEM micrograph of *N. oculata* after CE. FESEM images Figure 4 (e) and (f) show *N. oculata* after exposed to MWE. It was very obvious that cell walls of *N. oculata* were ruptured after being extracted via MWE. The smooth surface of round shape *N. oculata* cell, was totally ruptured into smaller fragments depicted in Figure 4 (c – f) releasing the lipids stored in the biological matrix [56]. However, *N. oculata* cell of CE sample obtained severe cracks compared to MWE after extraction. Thus, this probably explained CE attained higher lipid yield than MWE at 10 minutes.

![Figure 4](image_url)

Figure 4. Micrograph of *N. oculata* before extraction (a) at 100 X and (b) 1.00kX, After CE for 10 minutes (c) at 100 X and (d) 1.00kX, After MWE for 10 minutes (e) at 100 X and (f) 1.00kX.
3.5. Recommended Best Practices

A set of guidelines are provided to encourage best practices in the production of lipids and fatty acids so that they can be applied to other related systems. All recommendations are based on the study’s observation.

(1). To obtain higher lipid yield, lower extraction time (2 minutes) should be applied for both CE and MWE.

(2). To produce a shorter carbon chain, C4:0, lower extraction time should be applied for CE. It only can be detected at 2 minutes of CE. MWE is unable to produce this type of fatty acids.

(3). CE can only produce fatty acids that have a carbon chain length of up to C20 at longer extraction time, i.e., 10 minutes.

(4). MWE should be applied to solvent extraction to produce longer carbon chain, C22:2 which is good for anti-cancer applications. Both CE at 2 and 10 minutes unable to produce this type of beneficial and important fatty acids.

(5). CE and MWE techniques produced less TSFA and more TPUFA as well as ω-6 at longer extraction time.

(6). CE for 10 minutes achieved ω-6/ ω-3 ratio of 0.87 (with a value close to 1) indicating healthier edible oil.

(7). N. oculata is potentially a suitable alternative source for edible oil application due to the positive results from the ratio of TPUFA/ TSFA (0.19 – 0.35) which aligned with current commonly used cooking oil from palm-based source (0.25 – 0.28).

(8). CE and MWE on N. oculata did not produce trans-9-elaidic acid methyl ester (trans-fatty acids). Trans-fatty acid intake was associated with the raised risk of coronary heart disease.

4. Conclusions

The results showed that the CE technique is more appropriate to be used when extracting lipids from N. oculata for edible oil production. The time frame in CE influences both the lipid yield and quantity of FAMEs. Although MWE when used for extraction of lipids from N. oculata produces higher lipid yield but it does not contain ω-3. Finding shows MWE tend to oxidize EPA present in N. oculata. However, both techniques are successful in extracting the two most important types of fatty acids (C16 and C18) which are beneficial for edible oil application. Moreover, other beneficial fatty acids which have anti-cancer effect were also detected from N. oculata. Lipids from N. oculata has ω-6/ ω-3 ratio close to 1 and does not produce trans-fatty acid which makes it a suitable candidate for healthier edible oil. N. oculata is suitable to be applied as a functional food (healthier edible oil). The presence of beneficial functional lipids such as ω-3 and ω-6 from N. oculata offers a balance of fatty acids for existing cooking oil such as palm-based cooking oil, canola oil and other vegetable oils. Thus, N. oculata has the potential to be blended with other oils and formulated as super oil or blended oil (a mixture of two or more oils) which is supposed to be healthier than individual oils. Further research regards blending oil or more comprehensive testing on edible oil quality such as peroxide value (PV) and acid value (AV) should be conducted in the future.

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