Vacuum evaporation and nitrogen-assisted deodorization affects the antioxidant capacity in the olein fraction of red palm oil and its emulsion products [version 1; peer review: 2 approved with reservations]

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Abstract

**Background:** Deodorization of the olein fraction of red palm oil (OFRP) determines not only the taste of a multivitamin emulsion but also its antioxidant capacity. The emulsion product was formulated from OFRP, pumpkin juice (PJ), and dragon fruit juice (DFJ). This study aimed to optimize vacuum evaporation and nitrogen-assisted deodorizations of OFRP, observing levels of β-carotene, α-tocopherol, inhibition percentage of ABTS reduction, and ferric reducing antioxidant power (FRAP) activity.

**Methods:** The deodorizations observed were vacuum evaporation in four conditions: (1) 90°C, 80±5 mmHg, (2) 100°C, 80±5 mmHg, (3) 90°C, 100±5 mmHg, (4) 100°C, 100±5 mmHg, and nitrogen-assisted in two flow durations: (1) 15 min and (2) 30 min. β-carotene, α-tocopherol, and butylated hydroxytoluene (BHT) were employed as standards.

**Results:** The deodorized OFRP had fewer than 2% free fatty acids (FFA), lower than 3% peroxide value (PV), and lower than 4% acidic value (AV). Fluctuations of the β-carotene and α-tocopherol concentrations were observed in the deodorized OFRP. The final emulsion product had β-carotene of 259.9±1.4 to 271.7±2.4 ppm and α-tocopherol of 36.36±0.20 to 39.12±0.20 ppm. The total betacyanin of the emulsions were ±25% than DFJ. The emulsions had 22.93 to 32.11% of ABTS reduction inhibitory activity of the BHT activity and FRAP activity of 16.54±0.19 to 17.69±0.67 mM FeSO₄•7H₂O.

**Conclusions:** The best vacuum evaporation optimized at 90 °C, 100±5 mmHg, 60 RPM for 1 hour. The best nitrogen-assisted deodorization was at 85±3°C and 1 l/minute of nitrogen for 15 minutes. The deodorization process affected the antioxidant activity of OFRP and emulsions.
Keywords
α-tocopherol, antioxidant capacity, β-carotene, deodorization, olein fraction of red palm oil

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Introduction

The attempt to produce multivitamins locally sourced from the emulsion of the olein fraction of red palm oil (OFRP), pumpkin juice (PJ), and dragon fruit juice (DFJ) was hampered by the strong odor and aftertaste of OFRP. The stripping of vapor components from OFRP is theoretically achievable through heating at low pressure or flowing inert gas as a carrier of the evaporated odor compounds. However, application of heat to reduce the off-taste compounds in OFRP may result in the formation of trans-fatty acids and the occurrence of lipid oxidation. Oxidation is a major concern to edible oil quality, deteriorating its chemical, sensory and nutritional properties. The application of heat is also limited the desire to conserve the minor antioxidant components (e.g. tocopherols compounds).

Maintaining healthy contents of natural antioxidants and polyunsaturated fatty acids (PUFA) are key indicators of the quality of refined and deodorized edible oils. In this regard, natural antioxidants protect oils from hydrolysis, oxidation, polymerization, isomerization, and cyclization. Previous deodorization efforts to reduce the off odor of palm oil resulted in α-tocopherol being a better marker for maintaining overall antioxidant contents. An important finding was the need to maintain the critical time of the deodorization, which the destruction of tocot groups of the OFRP was in linear correlation to the deodorization time.

Vacuum evaporation and steam-assisted deodorization are common practices in oil refinery. Applying vacuum pressures of $10^{-5}$ to 1 mbar enables heat application to reach 300°C in edible oil processing. Steam-assisted deodorization requires good contact between the edible oils and the steam as stripping medium. A nitrogen medium is preferable in comparison to steam medium, due to the inert property of the gas and the effectiveness of thermal breakdown of off-taste precursors in the presence of the nitrogen carrier inside the deodorization chamber. Based on various application reports, both methods are capable of deodorizing off-odor compounds in the production of better-tasting and rich-in-antioxidant OFRP.

Concerning the challenges of OFRP deodorization, this stage should be revisited and further developed to produce a higher antioxidant capacity in the final OFRP, PJ, and DFJ emulsion products. This study aims to optimize the vacuum evaporation and nitrogen-assisted deodorization method of OFRP, observed using the levels of β-carotene, α-tocopherol, percentage inhibition of ABTS reduction, and ferric reducing antioxidant power (FRAP) activity.

Methods

Olein fraction of red palm oil

The crude palm oil (CPO) was obtained from PT Rea Kaltim, Indonesia. The olein fraction of red palm oil (OFRP) was prepared from CPO which had been tested for free fatty acid (FFA) levels and then carried out neutralization and gum removal by adding 100 ml warm water (80–90°C). Shuffling of the sample was carried out for 1 minute in a separating funnel before removal of residual water. About 400 ml of 10% NaOH (Emsure® Merck Cat no. 106498, USA) was added. Rinsing with 50 ml of warm water was carried out repeatedly until two phases of mixture were formed. The OFRP was produced with yields ranging from 60 to 70% of CPO (Rahmadi et al., 2015).

Deodorization of olein fraction of red palm oil

Deodorization of the OFRP was prepared by two methods, namely vacuum and nitrogen-assisted evaporation. The liquid fraction was deodorized using a rotary evaporator (Büchi R-200, Switzerland) at a combination of temperature and pressure: (1) olein fraction processed with vacuum evaporation 1 (OFV1): 90°C, 80±5 mmHg, (2) OFV2: 100°C, 80±5 mmHg, (3) OFV3: 90°C, 100±5 mmHg, (4) OFV4: 100°C, 100±5 mmHg, and speed of 60 RPM for 1 hour. The OFVs that has been obtained was then stored in a closed container in the refrigerator (4±2°C) for further processing. Deodorization of the olein fraction by the nitrogen-assisted method was prepared by addition of 100 ml OFRP in a closed container. The sample had added to it 2% (w/v) pharmaceutical-grade activated carbon (Norit, Japan), then pure-grade nitrogen gas (Samator Gas, Indonesia) was flowed using a valve for 15 minutes (OFN1) and 30 minutes (OFN2) at a temperature of 85±3°C, and a flow rate of 1 l/minute.

Preparation of pumpkin and red dragon fruit juices

Preparation of pumpkin and red dragon fruit juice was carried out according to previously described method. A total of 1 kg of each peeled fruit was cut into pieces of approximately 3–5 cm³ and washed using clean water. Then, the pieces of fruit were mashed using a commercial juicer. After that, the juice was put separately in glass bottles and pasteurized at 80°C for 10 minutes, then filtered using a clean filter cloth. The filtered fruit juices were then stored at 4±2°C before being utilized.

Preparation of OFRP, PJ, and DFJ emulsions

A total of 30 ml of the deodorized OFRP was added to 70 ml pumpkin juice, giving a total volume of 100 ml. Food grade carboxymethylcellulose, xanthan gum, and cinnamon powder were added at concentrations of 2% (w/v), 2% (w/v), and 0.5% (w/v), respectively. After homogenization, the product samples had then pure-grade nitrogen gas (Samator Gas, Indonesia) was flowed using a valve for 15 minutes (OFN1) and 30 minutes (OFN2) at a temperature of 85±3°C, and a flow rate of 1 l/minute.

Free fatty acid, peroxide value, and acidic values

FFA levels, peroxide values (PV), and acidic values (AV) were determined as previously described.

Determination of total carotenoid

Carotenoids were measured via measurement of β-carotene by modification of the Palm Oil Research Institute (PORIM) method, with changes in wavelength 446 to 443 nm (Rayleigh model UV 2601, China) and solvent changed from n-hexane to absolute methanol (Fulltime cat. 6501-04, China). Quantitative determination of β-carotene levels in samples was obtained based on calibration curves with β-carotene standards (Sigma-Aldrich cat. no. C9750-10G, UK).
Determination of α-tocopherol
The determination of α-tocopherol was accomplished by interpolating the absorbance of the sample with a standard calibration curve of α-tocopherol at a wavelength of 291 nm (Rayleigh model UV 2601, China). The α-tocopherol standard (Sigma-Aldrich cat. no. T3251-25G, UK) was prepared in absolute ethanol (Smartlab cat. no. A1035, Indonesia) with various concentrations of 50, 75, 100, 125 and 150 mg/L in 10 mL of absolute ethanol. Blank sample was prepared without containing the active substance of α-tocopherol.

Determination of betacyanins
Total betacyanin quantification was carried out as described and calculated using established equation. A total of 1 g of sample was macerated with 5% HCl solution (Merck cat. no. 109063, USA) (1:10) in a dark bottle, then stored at 4±2°C for 24 hours. The mixture was vacuum-filtered with Whattman’s filter paper no. 4. Next, 5 ml filtrate was diluted with 95% ethanol solution: 1.5 N HCl (85:15) to 10 ml. The absorbance was measured at a wavelength of 535 nm (Rayleigh model UV2601, China).

Antioxidant activity
ABTS. For the preparation of antioxidant measurement by the ABTS method, solution A was prepared by dissolving 7.1015 mg of 2,2’-azino-bis(3-ethylbenzothiazoline-6-sulphonic acid) (ABTS) (Sigma-Aldrich cat. no. A1888-1G, UK) in 5 ml distilled H2O, and solution B was prepared by dissolving 3.500 mg K3S2O8 (Sigma-Aldrich cat. no. 216224-100G, UK) in 5 ml of distilled H2O. Solution A and B were separately incubated in dark bottles for 12 hours. After incubation, solution A and B were mixed in a dark room and added with absolute ethanol to produce 25 ml of the final solution. Measurement of absorbance of ABTS blank solution was carried out at a wavelength of 750 nm (Eppendorf Single Beam BioSpectrometer®, Germany). Measurement of free radical-binding activity was carried out by mixing the emulsion sample with ABTS solution at a concentration of 100 ppm. The sample was measured at maximum absorbance wavelength. butylated hydroxytoluene (BHT) was used as a standard (Sigma-Aldrich cat. no. W218405, UK). The percentage of antioxidant activity was determined based on the calculation of the difference between the blank and the sample absorbance divided by the blank absorbance.

FRAP. FRAP reagent consists of 0.1 M acetate buffer (pH 3.6) (Merck cat. no. 107827, USA), 10 mM 2,4,6-tripyridyl-s-triazine (TPTZ) (Sigma-Aldrich cat. no. 93285-5G, UK), and 20 mM FeCl₃•6H₂O (Sigma-Aldrich cat. no. 157740-100G, UK) in a ratio of 10:1:1 in 40 mM of HCl. Standard solutions of FeSO₄•7H₂O (Sigma-Aldrich cat. no. 215422, UK) were prepared at concentrations of 10, 20, 30, 40 and 50 μmol/L. The maximum absorbance of FeSO₄•7H₂O was determined, which was at wavelength of 594-597 nm (Eppendorf Single Beam BioSpectrometer®, Germany). A total of 0.1 ml emulsion sample was prepared and added to 3 ml FRAP reagent in a test tube. The absorbance of the mixture was then read at 574 nm. Quantitative determination of total antioxidants in the sample was obtained based on calibration curve of FeSO₄•7H₂O.

Results and discussion
FFA, PV, and AV of OFRP, deodorized OFRP, and the emulsion products
FFA indicates the palatability acceptance of the OFRP and its emulsion-derived products; a higher presence of FFA contents results in a more pronounced sharp taste. FFA standard for fresh palm oil is less than 2%, while maximum FFA content for processed palm oil is set to 5%. The vacuum evaporation of OFRP produced a higher concentration of FFA in comparison to nitrogen-assisted deodorization. However, the final products had less than 2% of FFA (Figure 1).

PV indicates the presence of hydrogen peroxide in edible oils. The standards for the maximum PV in edible oils are determined based on the product and the processing technology used. For example, the standard PV for virgin olive oil is not more than 20 mEq O2/Kg, while for processed olive oil is not more than 10 mEq O2/Kg. Neither vacuum-evaporated nor nitrogen-assisted deodorized OFRP had PV of more than 3 mEq O2/Kg. The final emulsion products had PV of less than 2 mEq O2/Kg (Figure 1).

OFRP and the derived emulsion products had an AV of less than 4 mEq KOH/g (Figure 1). Pure palm oil had AV on average of 10 mEq KOH/g, in acidic environment. Crude Palm Oil (CPO) produced from small scale palm oil mills had an average AV of 18 mEq KOH/g. From the two deodorization approach, in terms of values of FFA, PV, and AV, nitrogen-assisted deodorized OFRP emerged as the better deodorization process.

Peak wavelengths and standard curves of β-carotene and α-tocopherol
The PORIM method allows for quick determination of the β-carotene content in CPO. Based on the procedure and to ensure high repeatability, it is necessary to recheck the peak wavelength of β-carotene standard, which was at 443 nm, and to generate a β-carotene standard curve (Figure 2a). The contents of β-carotene in OFRP, PJ, DFJ and derived products are given in Table 1. The relative contents of β-carotene to OFRP were highlighted in Figure 2b. As expected, DFJ had the lowest percentage of β-carotene in comparison to OFRP, while the content of β-carotene in PJ was 60% of the content in OFRP. The β-carotene contents for the emulsion made using vacuum-deodorized OFRP (EV) was 271.7±2.4 ppm, while 259.2±1.4 ppm of β-carotene was found in the emulsion made of nitrogen-assisted deodorized OFRP (EN).

The use of higher temperatures in vacuum evaporation (OFV2 and OFV4) slightly reduced β-carotene content in OFRP. A temperature increase without an increase in head pressure resulted in a lower accumulation of β-carotene, as observed in OFV4 vs OFV2 and OFV3 vs OFV1. This phenomenon is in line with that previously reported, stating that a temperature increase while reducing head pressure resulted in lower efficiency of soybean oil neutralization and distillation. Nitrogen-assisted deodorization helped to concentrate β-carotene content in OFRP. The emulsion products contained around 50% of β-carotene in OFRP.
Nitrogen was used as a stripping medium substitute for steam, owing to its inert and easy-to-remove properties of non-reacting. This highlights the possibility of neutralizing distillation by stripping with nitrogen in physical refining to obtain refined oils within specified end product’s acidity and flavour.

To provide a repeatable measurement of the content of α-tocopherol, peak wavelength of α-tocopherol standard was rechecked and occurred at 291 nm and α-tocopherol standard curve was produced as in Figure 3a. Table 1 displays the α-tocopherol content in OFRP, PJ, DFJ and derived products, while Figure 3b highlights the relative contents of α-tocopherol to OFRP. All three sources (OFRP, DFJ and PJ) contained a reasonable concentration of α-tocopherol, (70.61±0.59, 37.02±0.33 and 32.95±0.04 ppm, respectively). The final emulsion products contained between 36.36±0.20 and 39.12±0.20 ppm of α-tocopherol.

Previously, we reported that processing time was critical to maintain α-tocopherol levels in OFRP deodorization. While the deodorization time was fixed at 1 hour, a combination of temperature and vacuum pressure (OFV1 to OFV4) had slight effect in reducing the content of α-tocopherol in OFRP. The phenomenon was similar to that previously reported, which found that a 10°C temperature increase resulted in slightly reduced α-tocopherol content in soybean oil neutralization and distillation. The best treatment for vacuum deodorization was at 100 °C and 100±5 mmHg (OFV3).
Betacyanin

The presence of betacyanin in dragon fruit has led to the use of the color for dye in various applications, from food to solar cells\textsuperscript{16,25}. DFJ juice contains 2.18 ppm of betacyanin, while the contents of betacyanin in the emulsion products were 22–26% of the content in DFJ (Figure 4). This indicated that the function of DFJ was to provide pleasant color while masking the after taste of OFRP in the final emulsion product\textsuperscript{1}.

Antioxidant activity

The antioxidant activity of OFRP, PJ, DFJ, and the derived emulsion products were estimated with ABTS and FRAP assays (Figure 5). The two methods were suitable to measure antioxidant activity in oil based products\textsuperscript{26}. Products containing high concentrations of carotenes, xanthophylls and tocopherols are expected to have moderate-to-strong antioxidant potency, comparable to BHT. In comparison to BHT, the emulsion products had
22.93±0.10 to 32.11±0.04% of free-radical-scavenging activity, as measured by ABTS method. There was no significant difference of percentages of antioxidant activity of OFRP, PJ, DFJ, and the derived emulsion products against the BHT standard. Based on the ABTS and FRAP assays, it was concluded that OFRP, PJ, DFJ, and the derived emulsion products exhibited moderate-to-strong antioxidant activity.

While acting as a putative chain-breaking antioxidant, β-carotene was capable of scavenging peroxyl radicals\(^5\). Tocopherols are strong antioxidants, but their activity depends on the isomers, reactivity of the tocopheryl radical, surrounding temperature, and the type and viscosity of emulsion\(^6\). Oil-based products may contain secondary antioxidants, due to their greater ability to reduce lipid oxidation than acting as free radical scavengers\(^7\).

**Figure 4.** Total betacyanin contents of dragon fruit juice and the emulsion products.

**Figure 5.** Percentage of ABTS inhibition and total antioxidant by emulsion products and components in comparison to standard.

**Dataset 1.** All data on the properties of the emulsions produced in the current study

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Included in labelled files are all raw data for each variable measured in this study, alongside the processed data.

**Conclusion**

The vacuum evaporation of OFRP produced slightly higher FFA and AV in comparison to nitrogen-assisted deodorization, while the final products had less than 2% of FFA, less than 3 mEq O\(_2\)/Kg of PV, and less than 4 mEq KOH/g of AV. The β-carotene contents for the emulsion containing vacuum-deodorized OFRP was at 259.9±1.4 and the nitrogen-assisted deodorized OFRP was 271.7±2.4 ppm; these values were 51 and 53% of the...
References

1. Rahmadi A, Setionoegroh IB, Yuliani Y, et al.: Dragon Fruit Juice Addition in Palm Oil-Pumpkin Emulsion: Panelist Acceptance and Antioxidant Capacity. Jurnal Teknologi dan Industri Pangan. 2017; 28(2): 122–128. Publisher Full Text

2. Dijkstra AJ: Vacuum stripping of oils and fats. In: Gunstone, F.D., Harwood, J.L., & Dijkstra, A.J. (eds.). The Lipid Handbook, 3rd Edition. Taylor & Francis Group LLC, Boca Raton, FL, 2007; 235–253.

3. De Greyt WFJ: Edible oil refining: Current and future technologies. In: W, Hamm, RJ, Hamilton and GH. Callaux (eds.). Edible Oil Processing. John Wiley & Sons, Chichester, 2013; 127–151. Publisher Full Text

4. Kellens M, De Suryay D: Method for vacuum stripping of oils and fats. U.S. Patent No. 7,870,634. Washington, DC: U.S. Patent and Trademark Office. 2010. Reference Source

5. Alireza S, Tan CP, Hamed M, et al.: Effect of frying process on fatty acid composition and iodine value of selected vegetable oils and their blends. International Food Research Journal. 2010; 17(2): 295–302. Reference Source

6. Rahmadi A, Yanti I, Jannah SM, et al.: Quantitation and Optimization of β-Carotene and α-Tocopherol in Emulsion Prototype with Reversed Phase Chromatography. UAESET (submitted), 2018. Reference Source

7. San Ho DS: Recovery of phytonutrients from oils. U.S. Patent No. 7,544,822. Washington, DC: U.S. Patent and Trademark Office. 2009. Reference Source

8. Ham JS, Kim HY, Lim ST: Antioxidant and deodorizing activities of phenolic components in chestnut inner shell extracts. Ind Crops Prod. 2015; 73: 99–105. Publisher Full Text

9. Konsoula Z, Liakopoulou-Kyriakides M: Effect of endogenous antioxidants of sesame seeds and sesame oil to the thermal stability of edible vegetable oils. LWT-Food Sci Technol. 2010; 43(9): 1379–1386. Publisher Full Text

10. Cuelar-Bermúdez SP, Barbá-Davila B, Sema-Saldivar SO, et al.: Deodorization of Arthrosira platensis biomass for further scale-up food applications. J Sci Food Agric. 2017; 97(15): 5123–5130. Published Abstract | Publisher Full Text

11. Chen D, Chen X, Chen H, et al.: Identification of odor volatile compounds and deodorization of Paphia undulata enzymatic hydrolysate. J Ocean U China. 2016; 15(6): 1101–1110. Published Abstract | Publisher Full Text

12. Aliaf T, Tomas V, Ruiz K, et al.: Deodorization by instant controlled pressure drop autodispersion of rosemary leaves prior to solvent extraction of antioxidants. LWT-Food Sci Technol. 2013; 51(1): 111–119. Publisher Full Text

13. Rahmadi A, Puspita Y, Agustin S, et al.: Peneriman Panelis dan Sifat Kimiawi Emulsi Labu Kuning dan Fraksi Olein Sawit. Jurnal Teknologi dan Industri Pangan. 2015; 26(2): 201–212. Publisher Full Text

14. Andrawelid N, Kuswandar F, Herawati D: Analisis Pangan. Dian Rakyat. Jakarta. 2011; 1–41.

15. [PQRI] Palm Oil Research Institute of Malaysia: Test Methods Carotene Content. Palm Oil Research Institute of Malaysia, Kuala Lumpur. Malaysia. 1995.

16. Woo KK, Ngwu Fh, Ngo LS, et al.: Stability of betalain pigment from red dragon fruit (Hylocereus polymorbus). Am J Food Technol. 2011; 8(2): 140–148. Publisher Full Text

17. Inicio MR, de Lima KM, Lopes VG, et al.: Total anthocyanin content determination in intact acai (Euterpe oleracea Mart.) and palmitero-juçara (Euterpe edulis Mart.) fruit using near infrared spectroscopy (NIR) and multivariate calibration. Food Chem. 2013; 136(3–4): 1160–1164. Published Abstract | Publisher Full Text

18. Tagoe SMA, Dickinson MJ, Apetorgbor MM: Factors influencing quality of palm oil produced at the cottage industry level in Ghana. International Food Research Journal. 2012; 19(1): 271–278. Reference Source

19. Eskin AK: Methods to determine the extent of lipid oxidation in foods. In: Decker EA, Elias RJ, & McClements DJ. (eds). Oxidation in Foods and Beverages and Antioxidant Applications: Understanding Mechanisms of Oxidation and Antioxidant Activity. Woodhead Publishing. 2010; 181–195. Publisher Full Text

20. Matthaus B: Oxidation of edible oils. In: Decker EA, Elias RJ, & McClements DJ. (eds). Oxidation in Foods and Beverages and Antioxidant Applications: Understanding Mechanisms of Oxidation and Antioxidant Activity. Woodhead Publishing. 2010; 183–238. Reference Source

21. Hasyan A, Miyaji F, Minhanti M, et al.: Treatment of acidified palm oil for fatty acid methyl esters production. Chem Pap. 2012; 66(1): 39–46. Publisher Full Text

22. Ekop SA, Etuk BA, Eddy NO: Effect of some local additives on the chemical constituent of palm oil. Journal of Applied Sciences and Environmental Management. 2007; 11(1): 85–89. Publisher Full Text

23. Chang AS, Sherazi STH, Kandhoo AA, et al.: Characterization of Palm Fatty Acid Distillate of Different Oil Processing Industries of Pakistan. Journal of Oleo Science. 2016; 65(11): 897–901. Published Abstract | Publisher Full Text

24. Gracianny-Constante E, Rodriguez-Berbel F, Paredes-Torronteras A, et al.: Decadification by distillation using nitrogen as stripper: Possible application to the refining of edible fats. Grassas y Aceites. 1991; 42(4): 286–292. Reference Source

25. Ali RAM, Nayan N: Fabrication and analysis of dye-sensitized solar cell using natural dye extracted from dragon fruit. International Journal of Integrated Engineering. 2010; 2(3): 55–60. Reference Source

26. Christodoulides DC, Fotakis C, Nikokavoura A, et al.: Modified DPPH and ABTS assays to assess the antioxidant profile of untreated oils. Food Anal Methods. 2015; 8(5): 1294–1302. Publisher Full Text

27. Kiekasz S, Varzakas T, Opreoupolov V: In vitro activity of vitamins, flavonoids, and natural phenolic antioxidants against the oxidative deterioration of oil-based systems. Critical reviews in food science and nutrition. 2008; 48(1): 78–93. Published Abstract | Publisher Full Text

28. Kamal-Eldin A, Appleiest LA: The chemistry and antioxidant properties of tocopherols and tocotrienols. Ljndis. 1996; 31(7): 671–678. Published Abstract | Publisher Full Text

29. Bohari B, Muhadir M, Rahmadi A: Dataset 1 in: Vacuum evaporation and nitrogen-assisted deodorization affects the antioxidant capacity in the olein fraction of red palm oil and its emulsion products. F1000Research. 2018. doi:10.2556/f1000research.16545.d221738

Data availability
Dataset 1. All data on the properties of the emulsions produced in the current study. Included in labelled files are all raw data for each variable measured in this study, alongside the processed data. DOI: https://doi.org/10.5256/f1000research.16545.d221738.

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The article reports on optimisation of deodorization methods for the olein fraction of red palm oil by using vacuum evaporation and vacuum evaporation nitrogen-assisted. It is a piece of interesting work, the authors though are focused on the application of the above methods without giving the state of the art in deodorization methods applied so far and what are the advantages of the ones they are reporting. They should add something in the introduction. The conclusions must be rewritten and commented based on their results and efficiency of the methods. It seems that Nitrogen-assisted vacuum evaporation at 85 °C is slightly better in terms of β-carotene, tocopherol and all the other analytical data compared to solely vacuum evaporation at higher temperature, which is expected. Antioxidant activity of the final emulsion and content in β-carotene etc are not very encouraging in case of application. They should discuss it. The paper can be accepted after minor revision.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and is the work technically sound? Yes

Are sufficient details of methods and analysis provided to allow replication by others? Partly

If applicable, is the statistical analysis and its interpretation appropriate? Partly

Are all the source data underlying the results available to ensure full reproducibility? Yes
Are the conclusions drawn adequately supported by the results?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** medicinal chemistry, Natural products chemistry and engineering, food and environmental biotechnology, microbial biotechnology

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Reviewer Report 17 January 2020

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Azis Boing Sitanggang
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1. The statistical analysis and its appropriate interpretation must be performed on the results to see significant differences between treatments (Fig 1-Fig 5)

2. The standard deviations especially in Fig 1 is really high (approx. 50% of the avg. value). By this, the data reproducibility might be questionable.

3. The authors should be consistent in presenting the figures. Sometimes the bars are accompanied with standard deviations, and without labels (numbers) on top of them, but sometimes without.

4. The drawn conclusion seems to be correct and stems for the discussion being presented. However, if the discussion accompanied with statistical analysis, this would make the study more scientifically sound.

Is the work clearly and accurately presented and does it cite the current literature?  
Yes

Is the study design appropriate and is the work technically sound?  
Yes

Are sufficient details of methods and analysis provided to allow replication by others?  
Yes
If applicable, is the statistical analysis and its interpretation appropriate?
Not applicable

Are all the source data underlying the results available to ensure full reproducibility?
Partly

Are the conclusions drawn adequately supported by the results?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: food process engineering, enzymatic membrane reactor, bioprocess engineering

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.