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# Alternative solvents for lipid extraction and their effect on protein quality in black soldier fly larvae

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## Highlights

- Comparison of alternative solvents for lipid extraction from black soldier fly.
- Lipid extraction was based on theoretical and experimental data.
- 2-methyloxolane (2-MeO) was found to be the ideal green solvent.
- Defatted BSF flour with 2-MeO had relatively better protein quality parameters.

# Alternative solvents for lipid extraction and their effect on protein quality in black soldier fly larvae

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#### 1 Abstract

Scrutiny of alternative solvents for the extraction of lipid constituents from black soldier fly 2 3 (BSF) was the main theme of this research. The present investigation compared a wide array of solvents for the extraction of desired components theoretically using tools like Hansen 4 5 solubility parameters (HSP), conductor-like screening model for real solvents (COSMO-RS) 6 and technical data (ACD labs) with the application of hurdle technology for solvent 7 screening. The ideal solvent selected was 2-methyloxolane (2-MeO) which was employed for conventional and multistage cross-current lipid extraction and the experimental data obtained 8 9 was compared with that of n-hexane extract. Fatty acid profile, lipid class, bioactivity of the oils were analysed and were in good correlation with the theoretical prediction. The kinetics 10 and diffusion modelling for the extraction system was proposed. The effect of solvent on 11 protein quality parameters like protein dispersibility index, solubility in alkaline solution, 12 urease index in the defatted flour was elucidated. The defatting step had no deleterious effect 13 on the molecular weight distribution of soluble proteins. Overall the study manifested that 2-14 MeO had better lipid recovery, enhanced bioactivity in the BSF oil, and relatively better 15 protein quality in the defatted flour. 16

17 **Keywords:** Alternative solvents; Black soldier fly; 2-methyloxolane; Lipids; Protein quality.

18

#### 19 **1. Introduction**

In order to feed the growing population which is roughly estimated to touch 9.6 20 billion by 2050 (FAO, 2017) and alleviate its negative impact on the food supply chain many 21 research units, start-ups, think-tanks and other entities are coming up with innovative, greener 22 alternative solutions to fix the anticipated demand-supply gap. Among the multifarious 23 24 options being considered to handle the predicaments, utilisation of insects for food and feed applications are gaining significant traction. In particular, Hermetia illucens or the Black 25 26 Soldier Fly larvae (BSFL) belonging to the Diptera order is being contemplated as a promising substitute to replace the conventional protein sources to an extent. Industrial 27 rearing, processing, and valorization of BSFL comes with its own challenges, yet their nature 28 to aggregate the micro, macro-nutrients (Barroso et al., 2017) present in the feeding medium 29 thereby giving proteins, lipids, chitin derivatives, bioactive peptides, and organic manure 30 makes them an attractive candidate for numerous applications. 31

Hermetia illucens or BSFL for bio-conversion of discarded industrial by-products or 32 33 side streams, municipal wastes from urban activities have been extensively studied and the economic value it imparts makes it a valuable contender among other solutions 34 recommended. A comprehensive and concise review by Gold et al., 2018 on the 35 decomposition of biowaste types such as human, animal manures, fruits, vegetable wastes, 36 etc. was reviewed recently. Manure management system for laying hens treated with BSFL 37 diminished the manure accumulation by 50% and yielded 42% protein, 35% fat feedstuff is 38 one such example (Sheppard et al., 1994). 39

Due to the higher lipid concentration (St-Hilaire et al., 2007; Liland et al., 2017), 40 41 defatting BSFL should be the primary processing step in the downstream valorization of the insect biomass. For the production of protein, fat/oil and chitin, to be used in animal feed, the 42 43 raw insect materials must undergo a heat treatment process as described in the legislation on animal-by-products (Regulation (EC) No 1069/2009) (EFSA scientific committee., 2015). 44 There is a sharp increase in the number of insect companies foraying into the feed market 45 across continental Europe and North America capitalizing on the new regulation on Novel 46 47 Foods (EU 2015/2283) passed by the European Parliament. The fortification or replacement of conventional feed with BSFL to augment the protein content has been a time tested idea. 48 The efficacy of BSFL as a feed additive for poultry and a sustainable aquafeed ingredient has 49 been widely advocated (Bondari & Sheppard., 1981; Kroeckel et al., 2012; Vargas-Abundez 50

et al., 2019). The benefits of BSFL as a feed component is not limited to its protein content
and quality, its oil as a potential replacement for soybean oil in Jian Carp diets without any
negative effect on growth, feed efficiency in fish fillets was suggested (Li et al., 2016).

54 In the case of oilseeds, the solvent extraction of oil with n-hexane is preferred as it promotes easier oil recovery and has a narrow boiling point (69 °C). But, n-hexane reacts 55 with free pollutants to form ozone, photo chemicals and is said to affect the neural system 56 when inhaled by humans (Kumar et al., 2017). Though the idea of complete replacement of 57 petrochemical solvents for oil extraction is far-fetched, it is imperative that efforts to reduce 58 the dependency on them and find suitable, economically feasible alternatives for the same 59 must be given due consideration. Several research articles have articulated the effectiveness 60 of green solvents for oil extraction from various biomass. Bio-based solvents for the 61 extraction of oil from rapeseed (Sicaire et al., 2015), the green extraction of lipids from 62 oleaginous yeast biomass (Breil et al., 2016), the green extraction of Litsea Cubeba kernel 63 oil using alternative solvents (Zhuang et al., 2018) are few examples where a green and eco-64 friendly approach for the extraction of oil was proposed. 65

The objective of this work was to probe and identify an optimal green solvent from a 66 wide array of solvents including alcohols, ethers, esters and terpenes for defatting the BSFL 67 matrix by combining theoretical and experimental methods. Theoretical data for determining 68 the interactions between the solute and solvent were realized with tools such as Hansen 69 solubility parameters, consequently supported with relatively precise data obtained from 70 conductor-like screening model for real solvent (COSMO-RS) and finally with the 71 72 application of hurdle technology for decisive selection of suitable solvent based on the absolute theoretical data retrieved. It was within the purview of this study to establish an 73 74 industrial scale simulation for oil extraction and to develop kinetic modelling for a better 75 understanding of lipid diffusivity in the solvent medium. Furthermore, to elucidate the lipid 76 composition data by identifying the fatty acid profile, lipid class constituents, the bioactivity of BSF oil and compare the effect of solvent on the protein quality parameters in the defatted 77 BSF flour. 78

79

80 2. Materials and methods

#### 81 2.1. Larvae harvesting conditions

Black Soldier Fly Larvae (BSFL) was provided by a local insect rearing plant based in
Avignon region. The larvae were freeze-dried, milled (<1mm), and stored at -18 °C until</li>
further analysis. The proximate values of the freeze-dried BSFL were crude nitrogen:
40.27±0.62; crude lipid: 36.41±1.29; ash content: 9.01±0.13; moisture: < 3% (AOAC, 1990)</li>

86 2.2. Solvents, Standards and Reagents

87 All solvents for extraction and chromatographic analysis were of analytical grade and purchased from VWR international (Darmstadt, Germany). The solvent 2-methyloxolane also 88 known as 2-methyl tetrahydrofuran (CAS: 96-47-9) was sourced from Honeywell, Sigma-89 Aldrich Co, St. Louis (MO, USA). Standards: Supelco 37 FAME mix, DL-a-palmitin, 90 glyceryl 1,3-dipalmitate, glyceryl tripalmitate, palmitic acid, phospholipid mixture were 91 purchased from Sigma-Aldrich (USA). Ergosterol, 98% was procured from Acros organics 92 (Germany). Milli-Q water was used for electrophoresis and the protein ladder was sourced 93 from Bio-Rad. 94

#### 95 **2.3. Extraction**

#### 96 2.3.1. Conventional Soxhlet

97 The freeze-dried BSFL powder 25 g was taken in a cellulose thimble and subjected to exhaustive Soxhlet extraction for a period of 6 h with 250 mL of n-hexane and 2-MeO 98 99 solvents respectively. To achieve complete lipid removal the reflux was temporarily stopped every two hours and the sample inside the thimble were mixed thoroughly to facilitate 100 101 percolation and reduce agglomeration in the matrix. The solvents were collected after 6 - 7 h to establish approximately similar extraction cycles and then evaporated under reduced 102 103 pressure in a rotavapor. The yield was calculated gravimetrically and extractions were carried out in triplicates. 104

105 2.3.2. Multistage cross-current

To mimic the industrial scale oil extraction a multistage cross-current extraction system was set up, wherein three stages of sequential conventional extraction (Fig. 3) was used for maximum oil solubilization in the solvents. The powdered matrix 10 g was mixed with 100 mL solvent for 1 h under constant agitation (150 rpm). Later the solvent was 110 collected and the residue matrix was subjected to another stage of extraction by infusing 100 111 mL of fresh solvents with both n-hexane and 2-MeO respectively. This above-mentioned step 112 was repeated again, where the solvent was collected and the residual matrix was extracted 113 again with fresh solvents. The extractions were performed at 55 °C and temperature was 114 maintained using a Huber Pilot system. Therefore in total three stages of conventional 115 extraction was executed and the yield at every stage was noted and the cumulative yield at the 116 end of three stages was compared to conventional Soxhlet extraction results.

117 2.3.3. Kinetics

The kinetics study comparison to find the solubilization efficacy of both the solvents (n-hexane and 2-MeO) was established by placing 10 g of freeze-dried powder in 100 mL solvent at 55 °C. To understand the lipid extraction with respect to time in both solvents approximately 1 mL of clear solvent was collected from the extraction flask within specific intervals (1,3,5,10,15,20,30,60,90,120,150, and 180 min) and evaporated at 40 °C under nitrogen using a block heater. The yield was calculated and extrapolated to get results for 100g of dry powder.

125 2.3.4. Diffusion model

During extraction the solvent system was perfectly agitated, therefore, the major mass resistance during extraction was the internal diffusion of lipid within larvae powders (Baümler et al., 2017). In this case, the extraction process can be theorized by the diffusion model based on Fick's second law. Several hypotheses were given prior to simulation (Tao et al., 2017):

a) Larvae powders were regarded as spherical geometry and lipids were initially
homogeneously distributed within larvae powders. The radius of the larvae powders was 0.5
mm.

b) The diffusion coefficient of lipid did not change throughout the extraction process. c)Lipid content in larvae particles changed with time and position.

136 d) No external mass resistance was taken into consideration due to external agitation.

137 The diffusion equation for spherical geometry is written as (Tao et al., 2017):

138 
$$\frac{\partial C_{\rm S}}{\partial t} = D_{\rm e} \left( \frac{1}{x^2} \frac{\partial}{\partial x} \left( x^2 \frac{\partial C_{\rm S}}{\partial x} \right) \right) \tag{1}$$

7

where  $C_s$  is lipid concentration within larvae particle (g/m<sup>3</sup>), *De* is effective diffusion coefficient for lipid (m<sup>2</sup>/s), *x* is the radial distance in the diffusion direction (m), and *t* is time (s).

142 The initial conditions are  $C_S = C_{S,0}$  for the solid phase and  $C_L = 0$  for the liquid phase. 143  $C_{S,0}$  is the initial concentration of lipid in larvae particles (g/m<sup>3</sup>) and  $C_L$  is the concentration of 144 lipid in solvent (g/m<sup>3</sup>).

145 The boundary conditions for Eq. 1 are:

146 
$$\left(\frac{\partial C_S}{\partial x}\right)_{x=0} = 0$$
 (2)

147 
$$-D_e A \left[ \frac{\partial C_s(x,t)}{\partial x} \right]_{x=r} = V \frac{d C_L(t)}{dt}$$
(3)

where A is the surface area of larvae particles  $(m^2)$ , V is the volume of solvent used for extraction  $(m^3)$ , r is the radius of larvae particles (m).

The "*pdepe*" function in Matlab, R2010a (The MathWorks, Inc., MA, USA) was used to solve the aforementioned parabolic partial differential equation (Tao et al., 2019). To be exact, the original partial differential equation (Eq. 1) was first discretized spatially. After that, the resulting ordinary differential equations were integrated in time and solved by the *pdepe* solver. The  $D_e$  value was adjusted iteratively to fit the experimental data, thus minimizing the root mean square error (*RMSE*) between experimental and predicted the content of lipid in larvae particles:

157 
$$RMSE(g/m^3) = \sqrt{\frac{1}{n} \sum_{i=1}^{n} [C_{S,p}(t) - C_{S,e}(t)]^2}$$
 (4)

where  $C_{S,e}$  and  $C_{S,p}$  is the experimental and predicted values of lipid content in larvae particles (g/m<sup>3</sup>), respectively. *n* is the number of experimental points.

160 Once the optimized  $D_e$  value was obtained, three statistical indicators, including  $R^2$ 161 (coefficient of determination), *RMSE* and absolute average deviation (*AAD*) were calculated 162 using the data about lipid extraction yield to test the predictive accuracy of the diffusion:

163 
$$R^{2} = 1 - \frac{\sum_{i=1}^{n} (Y_{e} - Y_{p})^{2}}{\sum_{i=1}^{n} (Y_{e} - Y_{m})^{2}}$$
(5)

164 
$$RMSE(mg/100g DM) = \sqrt{\frac{1}{n} \sum_{i=1}^{n} [Y_p(t) - Y_e(t)]^2}$$
 (6)

8

165 
$$AAD(\%) = \left[\frac{\sum_{i=1}^{n} (|Y_e - Y_p|)/Y_{S,e}}{n}\right] \times 100$$
 (7)

where  $Y_e$ ,  $Y_p$  and  $Y_m$  are the experimental, predicted values of lipid extraction yield, respectively (mg/100g DM).  $Y_m$  is the average lipid extraction yield (mg/g DM).

Following the numerical simulation results, the distributions of lipid content withinlarvae powders at different stages of extraction were visualized by programming in Matlab.

- 170 **2.4. BSFL oil analyses**
- 171 2.4.1. Fatty acid profile

172 Fatty acid methyl esters (FAMEs) were prepared from BSF oil samples by acidcatalyzed transmethylation (Breil et al., 2016). Ten to fifteen milligram of oil sample was 173 174 taken to which 1 mL of methanolic sulfuric acid was added, the mixture was heated in a heating block to facilitate transmethylation and 1.5 mL of 0.9% NaCl solution, and 1 mL of 175 176 GC-FID grade hexane was added after it reached room temperature. The organic layer was collected and subjected to further analysis. Triheptadecanoin (C17:0; TAG) was used as 177 internal standard and FAME mix was used to calibrate the system. An Agilent (Kyoto, Japan) 178 gas chromatography coupled with a flame ionization detector (GC-FID) system was used to 179 determine the fatty acid profiles of BSF oils. The system was equipped with a BD-EN14103 180 capillary column with dimensions 30 m  $\times$  320  $\mu$ m  $\times$  0.25  $\mu$ m. The carrier gas (He) velocity 181 was set at 33 cm.s<sup>-1</sup>. The sample injection volume was 2 µL and two washes with n-hexane 182 183 were executed to avoid carryovers after every sample run. The detection was enabled in split mode (split ratio 1:20), the injection temperature was set at 250 °C. The oven temperature 184 gradient was initially 50 °C for 1 min and then increased at a constant rate of 20 °C/min from 185 50 °C to 180 °C and then raised from 180 °C to 220 °C at a rate of 2 °C/min. Once it reached 186 230 °C the temperature was maintained for a period of 10 min and the fatty acids were 187 188 identified based on retention time and standards used for calibration.

189 2.4.2. Polar and neutral lipids quantitation

High-performance thin-layer chromatography (ATS 5 automatic TLC sampler, ADC
2 automatic developing chamber, CAMAG 3 TLC scanning densitometer) was used for
separation, identification and relative quantitation of the lipid classes of BSF oil extracted
with n-hexane and 2-MeO. All stock solutions were prepared in chloroform and 10 mg of
sample lipid fraction was used for quantitation. A known volume of lipid extract was loaded

on  $20 \times 10$  cm Silica gel 60 F254 HPTLC plates. Polar lipids were separated with eluent A, a 195 mixture of methyl acetate/isopropanol/chloroform/methanol/KCl (0.25%) in a ratio of 196 25:25:25:10:9 (v/v/v/v). Neutral lipids were eluted with eluent B, a mixture of n-197 hexane/diethyl ether/glacial acetic acid in a ratio of 70:30:2 (v/v/v). Both plates were allowed 198 199 to reach a height of 7 cm from the origin, then dried and was dipped in a primuline dye reagent (10 mg of primuline, 160 mL of acetone, 40 mL of distilled water) for better 200 201 visualization of the lipid classes. Lipid standards were used to identify and quantify the lipid classes in BSF oil. 202

203 2.4.3. Total polyphenol content (TPC) and radical scavenging capacity (RSC)

Twenty microliters of appropriate dilutions of the lipid samples or gallic acid (standard) in methanol were taken in a 96-well microplate and 80  $\mu$ L of 7.5 % Na<sub>2</sub>CO<sub>3</sub> solution was added and allowed to equilibrate at room temperature for 10 mins. Rapid addition of 100  $\mu$ L of 1N Folin-Ciocalteu reagent was completed and absorbance was read at 750 nm for every 5 min over a period of 60 min. Distilled water was used as blank and results were calculated as gallic acid equivalents (GAE).

Similarly, for RSC, 50  $\mu$ L of samples in the methanolic phase was allowed to react with 0.5 mM methanolic DPPH<sup>•</sup> radical for 60 mins and the absorbance was measured at 520 nm. Methanol was used as blank, trolox was used for generating the standard curve and results were expressed in trolox equivalent (TE).

#### 214 **2.5. Defatted BSFL flour analyses**

#### 215 2.5.1. Protein molecular weight distribution

216 Sodium Dodecyl Sulfate-Polyacrylamide Gel Electrophoresis (SDS-PAGE) was used to determine the molecular weight distribution of soluble proteins from BSFL. The proteins 217 were extracted according to Janssen et al., 2017 with few modifications. The dry BSFL 218 powder was placed in citric acid – disodium phosphate buffer (1:10; w/v) and was gently 219 mixed with a magnetic stirrer. After 60 min, the mixture was centrifuged at 9000 rpm for 20 220 min and the supernatant collected was dialyzed at 18 °C with a dialysis tubing having a cut-221 off value of 12- 14 kDa and then the dialyzed fraction was subjected to lyophilization to 222 obtain the soluble proteins. The soluble protein 2 mg/ml in milli-Q water was denatured by 223 addition of equal volume of Laemmli sample buffer (65.8mM Tris-HCl at pH 6.8, 26.3% 224 225 (w/v) glycerol, 2.1% SDS, 0.01% bromophenol blue, 5% 2-mercaptoethanol) at 90 °C for 5

mins and the electrophoresis was run in a Bio-Rad mini-PROTEAN system using TGX pre-cast gels (4-15%).

228 2.5.2. Protein quantification, dispersibility index, solubility

The nitrogen content in different BSFL solid fractions were analysed by Kjeldahl (Buchi speed digester K-425 system) and the protein content in liquid fractions was calculated by Lowry method with BSA as standard. The defatted BSFL flour protein quality was evaluated according to the quality analyses manual for soybean products in the feed industry (Van Eys et al., 2004) where the protein dispersibility index (PDI), protein solubility in 0.2% potassium hydroxide solution, urease index, and the absorbance at 420 nm were identified.

#### 236 **2.6.** Theoretical predictions

For an effective understanding of theoretical data that are generated by tools such as 237 Hansen, COSMO-RS and ACD labs it is essential to establish a local solute solvent database 238 which is specific to the sample chosen for analysis (BSF) and chemical constituents of 239 interest present in them. In this case solvents of different polarity (n-hexane, ethanol, iso-240 propanol, methyl acetate, ethyl acetate, ethyl lactate, dimethyl carbonate, 2-methyl 241 242 tetrahydrofuran, cyclopentyl methyl ether, a-pinene, d-limonene, p-cymene) and solutes generally present in the lipid fraction of BSF were chosen after careful literature review. 243 244 Preliminary parameters was set up with solutes belonging to various classes of the lipid fraction such as free fatty acids (FFA- Lauric Acid), monoglycerides (MAG- Glyceryl 1-245 246 laurate), diglycerides (DAG- Glyceryl 1,2-dipalmitate), triglycerides (TAG- Lauric triglyceride), Vitamin E (VE1–  $\alpha$ -tocopherol; VE2–  $\gamma$ -tocotrienol), sterols (ST1-  $\beta$ -sitosterol; 247 248 ST2- Cholesterol), and pigments (CA1- β-carotene). The solutes were selected based on previously reported data (Liland et al., 2017; Ushakova et al., 2016; Caligiani et al., 2018) on 249 250 the lipid fraction of BSF.

251

#### 252 2.6.1. Hansen Solubility Parameters (HSP)

HSP helps in understanding the solubility of two compounds, more precisely the 253 miscibility of two components in a medium, it is based on a simple, yet classical principle 254 "like dissolves like" phenomenon. This principle serves as the rule of thumb in characterizing 255 256 the solute-solvent interactions, theoretically the total cohesive energy density ( $\delta_{total}$ ) is equal to the square root of sum of the energy densities required to overcome atomic dispersion 257 forces ( $\delta d^2$ ), molecular polar forces due to dipole moments ( $\delta p^2$ ) and hydrogen bonds 258 (loss/gain of proton and exchange of electrons) between molecules ( $\delta h^2$ ) and mathematically 259 represented by the following equation: 260

261 
$$\delta_{\text{total}} = \sqrt{(\delta d^2 + \delta p^2 + \delta h^2)}$$
(8)

The magnitude of affinity between solute and solvent is directly proportional to the  $\delta_{total}$  value, higher the  $\delta_{total}$  greater the affinity. The relative energy difference (RED) is another parameter that indicates the miscibility of solute in solvents and is calculated:

$$RED = \frac{R_a}{R_b}$$
(9)

where R<sub>a</sub> is the distance of a solvent located inside the Hansen solubility sphere and R<sub>b</sub> is the
radius of the Hansen solubility sphere. The chemical structures and simplified molecular
input line entry syntax (SMILES) notations were fabricated using ACD/ChemSketch
(Toronto, Canada).

### 270 2.6.2. Conductor-Like Screening Model for Real Solvents (COSMO-RS)

COSMO-RS is used for the calculation of the thermodynamic properties for solvation, 271 without any experimental data. It is the best tool for molecular description and solvent 272 screening based on quantum-chemical approach. The usability of COSMO-RS for 273 determining the relative solubility index log<sub>10</sub>(x-solub) for sample-specific solutes and 274 solvents has been well documented in our previous work (Ravi et al., 2018). The sheer 275 amount of data generated based on the parameters like  $\sigma$ -surface,  $\sigma$ -profile,  $\sigma$ -potential can be 276 used to predict the compatibility of the solvent for the solubilization of solutes (Fig. 2.). The 277 calculations were executed in a COSMOthermX'17 program (version C30 release 13.01). 278 The 279 standard

280 quantum chemical method triple-valence polarized basis set (TZVP) was used in this study

and 55 °C was the temperature used for the solubility prediction to draw parallels with
 industrial processing conditions

283

284 
$$\log_{10}(x_j) = \log_{10}\left[\frac{\exp\left(\mu_j^{pure} - \mu_j^{solvent} - \Delta G_{j,fusion}\right)}{RT}\right]$$
(10)

285  $\mu_{j}^{\text{pure}}$ : chemical potential of pure compound j (Joule/mol)

286  $\mu_i^{solvent}$ : the chemical potential of j at infinite dilution (Joule/mol)

287  $\Delta$ Gj, fusion: free energy of fusion of j (Joule/mol)

288  $x_i$ : solubility of j (g/g solvent).

 $\alpha$ -tocopherol ( $\sigma$ -surface) was the solute used in the representative image and for its solubilization in respective solvents ( $\sigma$  -profile,  $\sigma$ -potential) were chosen for theoretical calculations. This process is repeated for all potential solutes, thereby generating the log<sub>10</sub>(xsolub) values which eventually was compared and tabulated (Table 2.)

### 293 2.6.3. Hurdle technology for solvent screening

Application of the hurdle concept has been predominantly used for food preservation, 294 food quality and safety assessment (Khan et al., 2016). It is an excellent decision-making tool 295 that can assist in numerous applications once the factual hurdles are established. For instance, 296 297 Fig. 1. clearly depicts the impediments that are associated with the solvent selection and how the technical parameters of the solvents are used for solvent screening purposes. A list of 298 299 candidate solvents are considered and the technical properties such as Log P, boiling point, toxicity index and solvent origin were chosen as appropriate hurdles in this work. These 300 301 parameters are consolidated (Table 3.) and paves way for taking an informed decision based on the theoretical data available. 302

303

#### **304 3. Results and discussion**

#### 305 3.1. Solvent selection based on theoretical studies

306 Relative energy difference (RED) is the empirical value that denotes the ability of a 307 solvent to dissolve the solutes of interest present in the sample matrix. The major constituents of BSFL lipid fraction were selected based on literature review, a generalized approach was 308 309 taken for the solute scrutiny, this way a broader class of compounds can be analysed for their solubility based on Hansen solubility parameters. A representative compound for each class 310 of solutes: lauric acid for fatty acids, glyceryl 1-laurate for monoglycerides, glyceryl 1,2-311 dipalmitate for diglycerides, lauric triglyceride for triglycerides,  $\alpha$ -tocopherol and  $\gamma$ -312 313 tocotrienol for vitamin E, cholesterol and β-sitosterol for sterols and finally β-carotene for pigments were selected. It is important to understand that the dietary components of the 314 315 BSFL heavily influence its chemical composition and is subject to vary drastically based on rearing conditions. 316

The solvent n-hexane was chosen as a reference and table 1 summarizes the RED 317 scores that were colour coded for easy identification of solvents better than reference. Among 318 the overall class of solvents chosen, ethers and terpenes outperformed the rest. Ultimately, 2-319 methyloxolane and cyclopentyl methyl ether had the best RED scores for all the solutes 320 considered indicating their theoretical ability to solubilize a wide range of chemical 321 322 constituents. In case of 2-MeO the RED score ranged between 0.73 for TAG and 1.75 for 323 MAG, similarly, CPME had 0.56 for TAG and 1.76 for MAG as its boundary values. These 324 values symbolise the solvents relative capacity to solubilize the solutes and n-hexane had better theoretical solvation than alcohols and esters collectively. If we look at the data to 325 326 identify solvents for better solubilization of solutes individually no trend can be found for example a-tocopherol had the best solvation in d-limonene and so did cholesterol which in 327 328 some cases may not be desirable, hence the model works perfectly for a collectively similar 329 class of compounds (polar or non-polar).

The relative solubility of the lipid contents (Table 2) in different solvents was given by COSMO-RS software which uses quantum calculations from sequential, iterative integration of data generated in the conductor like environment where initially the  $\sigma$ -surface was generated, then  $\sigma$ -profile and  $\sigma$ -potential were used to calculate log<sub>10</sub>(x\_solub) value. The solubilization of solute  $\alpha$ -tocopherol as indicated in the example (Fig. 2) was compared with the solvent list wherein the above-mentioned calculations are executed iteratively to generate the relative value. The ideal solvents had a value of 0 meaning theoretically they were the best solvents for better solubilization of their respective solutes. Again, 2-MeO and CPME proved to be relatively better than other solvents considered. Thus, the collective theoretical result with regard to the solvation power of each solvent for select solutes using Hansen and COSMO-RS determined that 2-MeO and CPME were the best solvents for extraction of lipid-based chemical constituents from BSFL.

Hurdle technology was employed to screen the solvents according to their technical 342 properties (Table 3) such as Log P, boiling point, toxicity index, enthalpy of vaporization, the 343 energy required to evaporate 1 metric ton of solvent, and the nature of solvent (petroleum 344 based, bio-based etc.,). These parameters guide in assessing the ecological footprint 345 associated with usage of solvent for extraction purposes. In the scope of food preservation, 346 347 hurdle technologies provide a framework for combining a number of preservation techniques for achieving an enhanced level of product safety and stability (Gupta et al., 2012). Similarly, 348 the same concept can be retrieved and applied for solvent screening, the parameters 349 considered were clustered together as hurdles (Fig. 1) and the solvents were assorted based 350 on their classes to observe if there was any trend exhibited by any particular class of solvent. 351 Interestingly, ethers had the upper hand in this scrutiny as well, they were able to surpass all 352 353 the hurdles showing the versatility and advantages of employing them for extraction. With all parameters and theoretical data considered, CPME and 2-MeO were deduced to be the best 354 suitable solvents and out of the two only 2-MeO is truly a bio-based solvent and produced 355 from lignocellulosic biomass. CPME manufacturing involves the methylation of 356 cyclopentanol or the addition of methanol to readily available cyclopentane (Wanatabe et al., 357 2007). The mission was to find a green solvent that could potentially replace n-hexane and 358 can be used industrially for oil extraction and 2-methyloxolane met all the criteria put forth 359 360 and therefore was used for further experimental analyses.

361 3.2. Solvent performance comparison: yield, fatty acid profile, lipid class

Conventional Soxhlet extraction with n-hexane, 2-MeO recovered  $32.51 \pm 0.39\%$  and 35.83  $\pm 1.12\%$  of lipids respectively. The lipid fraction of BSFL comprises of a complex set of substances, the main reason behind such lipid accumulation is that the adult larvae don't feed after the pupal instar ends. This is because of the lack of development of functional mouthparts, rendering them to rely only on the reserves accumulated during larval stages (Gobbi et al., 2013).

The fatty acid profile of oils extracted using n-hexane and 2-MeO were relatively 368 similar (Table 4), lauric acid (C12) was the major fatty acid with 42.29 %, followed by 369 370 linoleic acid (C18:2n6) with 13.91%, palmitic acid (C16) with 13.83%, oleic acid (C18:1n9) with 11.43%, myristic acid (C14) with 9.36% in decreasing order. These five acids combined 371 372 made up almost 90% of the fatty acid profile of BSF oils in both cases. The saturated fatty acids were the largest class of fatty acids accounting for 69.13%, followed by 373 374 polyunsaturated fatty acids responsible for 15.44 % and closely succeeded by monounsaturated fatty acids taking up 14.34%. The data were consistent with previously 375 published results (Liland et al., 2017) with a similar trend of fatty acids being reported. 376 Across all previous research articles that probed the fatty acid profile of BSF, lauric acid was 377 found to be the principal fatty acid with almost 35-40% present in BSF fed average diets, 378 379 without any extreme feed formulations (Caligiani et al., 2018; Ushakova et al., 2016).

The neutral and polar lipid classes of BSF oils were identified and quantified using 380 381 HPTLC. Monoglycerides, diglycerides, triglycerides, ergosterol and fatty acid classes were 382 relatively quantified. Free fatty acids were the largest neutral lipid class among the two oils 383 compared, accounting for almost 62% in the 2-MeO lipid fraction and 48% in the n-hexane lipid fraction. In BSF oil extracted with 2-MeO, the sequence was as follows FFA > DAG > 384 385 ERGO > TAG > MAG, whereas in n-hexane it was FFA > ERGO > DAG > TAG > MAG. The relative quantity of each class is detailed (Fig. 5) with the HPTLC plate of neutral lipids 386 387 that shows the migration distance of each lipid class. Polar lipids in particular phospholipids were quantified in the 2-MeO BSF oil and were not present in the oil extracted by n-hexane, 388 the solvent polarity plays a major part in extracting polar lipids, sterols, pigments and waxes. 389 This explains the increased oil yield when using 2-MeO as a solvent for defatting as it 390 391 enhances the overall oil profile by eluting other non-polar constituents present in the matrix. 392 Phosphatidylethanolamine was the primary polar lipid (42%), followed by phosphatidylinositol and phosphatidylcholine present in the 2-MeO BSF oil. This was the 393 first time polar lipids were identified in the lipid fraction of BSF. The absence of polar lipids 394 in n-hexane BSF oil was anticipated. Authors, Ushakova et al., 2016 presented the 395 composition of glycerides in BSFL and attempted to identify other components of the lipid 396 fraction. Dodecanoic ethyl ester, stigmasterol and cholesterol are few of the compounds they 397 quantified, reiterating the importance of understanding lipid prospects of larval biomass and 398 its importance in black soldier fly artificial breeding. Likewise, authors (Liland et al., 2017) 399 vividly elucidated the dietary modulation of BSF with seaweed-enriched media to alter the 400

401 lipid profile of BSF, where the total vitamin-E content increased by four folds in the 100% 402 seaweed feed based diet fed to BSF when compared to that of the control. Thus making BSF 403 a unique one-of-a-kind bioreactor like system which aggregates the desired components 404 present in the feed. This particular aspect of BSF as an enriching system is underexplored and 405 can be properly exploited in all field of life sciences and more.

406 3.3. Oil diffusion kinetics and industrial modelling

While comparing the solvents efficacy to elute lipids from biomass its kinetics aid in 407 408 understanding the diffusion mechanism with respect to time under predetermined conditions like temperature (55 °C), pressure, agitation (200 rpm). In solid-liquid extraction, the solute 409 410 of interest in this case oil is readily available and at the very instant that it comes in contact with the solvent it is freely dispersed in the solvent medium. The readily solubilized oil at t =411 412 0-5 min is termed starting accessibility ( $\delta X_s$ ; g of extract / g of dry matter) signifying the amount of solute solubilized in a limited time frame via the convection of solvent interacting 413 with the surface of the biomass. The oil yield in solvents n-hexane and 2-MeO were plotted 414 against time (Fig. 4a) and the data was extrapolated to 100 g of dry material. The individual 415 oil yield using the solvents n-hexane and 2-MeO indicate the effective diffusivity achieved, 416 exhaustive oil recovery was not accomplished with this setup as it was only meant to give 417 data of oil solubilized as a function of time. 418

In mass transfer terms, solvent extraction occurs in two stages a) solvent-surface
interaction transpires for a short duration, followed by the main mass transfer mechanisms
mediated by various b) penetration processes in the solvent-solute system (capillary forces,
molecular diffusivity, etc.).

423 A multistage cross-current system was incorporated to witness the oil recovery 424 efficacy of both the solvents in an industrially scalable system. Three stages of extraction 425 with each stage lasting 1 hour and the solvent replenished at the beginning of every new stage was executed. Incidentally, the cumulative oil yield extracted with this system was precisely 426 the same as the results obtained by conventional Soxhlet system which acknowledges the 427 efficiency and robustness of such system for enhanced oil recovery in a short period of time. 428 429 After each stage the oil yield was calculated (Fig. 4b) and 2-MeO had relatively higher yield gravimetrically in every stage but when the percentage oil recovery was considered n-hexane 430 (70.2%) had better oil solubilization in stage 1 when compared to that of 2-MeO (66.7%), in 431 432 subsequent stages 2-MeO exhibited better recovery than n-hexane. The number of stages for exhaustive lipid recovery can be effectively reduced by supplementing the system with other
intensification techniques like ultrasonication, microwave. Ultrasonication in synergy with
this system amplified the oil recovery from rapeseed cake (Sicaire A.G. et al., 2016).

436 3.4. Diffusion modelling and numerical simulation for lipid extraction

The experimental and simulated results of lipid extraction kinetics are illustrated in 437 438 Fig. 6a. Evidently, the diffusion model provided a satisfactory description of the evolution of lipid yield during extraction, although there were certain divergences in some areas, especial 439 in the early stage of extraction, the overall model was justified for the extraction system. 440 Meanwhile,  $R^2$  values for both cases exceeded 0.96, and AAD values were also quite low 441 442 (Table 5). Accordingly, the diffusion model was qualified to model the extraction process and investigate the mass mechanism using different solvents. The extraction yield at t=0 min was 443 444 set at 0, so as to enhance the predictive accuracy.

The distributions of lipid content during extraction at 5 and 20 min were visualized in Fig. 6b. There was a relatively high concentration gradient of lipid within larvae powders at the early stage of extraction using n-hexane as a solvent. Along with the increase of extraction time, the lipid concentration gradient decreased. When 2-MeO was used as extraction solvent, the distribution of lipid within larvae powder was more homogenous than that using n-hexane as solvent at the beginning of extraction, probably due to the high extraction rate and fast movement of lipid within the powders.

452 3.5. Bioactivity of BSF oil

The total polyphenol content (TPC) and radical scavenging capacity (RSC) of the oils 453 were examined (Table 6). BSF oil extracted with 2-MeO had a polyphenol content of 454 19.03±1.11 mg GAE which was 2.5 times higher than what was present in BSF oil extracted 455 456 using n-hexane. Comparable results were found in the RSC activity as well, BSF oil (2-MeO) had higher DPPH radical scavenging capacity. The enhanced bioactivity in BSF oil obtained 457 using 2-MeO might facilitate lowered lipid peroxidation at ambient and elevated 458 459 temperatures and could be used as a functional oil for various dietary supplements in animal feed applications. The vitamin E content contributes to the bioactivity in the lipid fraction, 460 pigments like carotenoids can boost the overall crude oil profile of BSF. The results obtained 461 were compared with refined sunflower oil to establish a reference. Recently, a skincare 462 product with sterilized BSF oil in their formulation was patented (US 2018/0256483 A1). 463 Using alternative solvents to extract oil with enhanced functional properties from insect 464

biomasses can lead to more innovations and its incorporation in cosmetic and therapeuticformulations.

467 3.6. Effect of solvent on protein quality in defatted flour

Protein dispersibility index (PDI), is a parameter mostly used to ascertain soybean meal protein quality. It measures the amount of protein dispersed in water after high-speed blending of the biomass. Measuring the PDI for BSF meal (defatted flour) gives a rough idea of protein solubility in water after lipid removal. In this study, the PDI value suggested that 31.6% of the protein in 2-MeO defatted flour and 29.1% of the protein in n-hexane defatted flour was soluble in water indicating a reduced availability of protein in the n-hexane defatted meal.

The presence of non-protein nitrogen in insects specifically in BSFL results in the overestimation of true protein available, therefore a nitrogen-to-protein conversion factor (Kp) value of  $4.76 \pm 0.09$  and  $5.6 \pm 0.39$  was proposed for larvae as such and after protein extraction respectively (Janssen et al., 2017). This might explain the low PDI values in the defatted flours as the crude protein quantitation includes the chitin, glucosamine nitrogen as well, and a Kp value of 6.25 was used for protein assessment. This hypothesis was not probed further as it was out of the scope of this study.

Similarly, protein solubility in 0.2% potassium hydroxide (KOH), urease index (UI), 482 483 and the absorbance at 420 nm were quantified (Table 7) as these are the generic parameters considered for soybean products in the feed industry. The protein solubility in alkaline 484 485 conditions for 2-MeO BSF meal was higher than its n-hexane counterpart. As the extraction and analytical conditions were similar the variation in values can be attributed to the nature of 486 487 the solvent. The crude protein content in 2-MeO BSF and n-hexane meal was 62.16±0.62% 488 and 59.09±0.62% respectively, these results are concurrent, as the lipid recovery was higher 489 in case of 2-MeO thereby concentrating protein content in its corresponding defatted meal. These values combined help determine the protein quality, the state of anti-nutritional factors, 490 and further processing techniques to be employed in soybean meal. The presence of anti-491 nutritional factors in BSF might depend on the feed substrate, for now very little is known 492 about its presence in BSF reared for feed purposes. Effectively, establishing these values for 493 insect biomass will help potential feed manufacturers relate to the properties of the defatted 494 flour of their respective insect products and compare them to existing industry standards. 495 496 Insects particularly BSF as a dietary feed for livestock, artic salmon, turbot, crustaceans were

497 adequately reviewed (Barroso et al., 2013; Makkar et al., 2014; Wang & Shelomi., 2017)
498 projecting the inclusion and infiltration of BSF in feed formulations.

The molecular weight distribution of soluble proteins extracted from 2-MeO defatted 499 500 BSF flour, n-hexane defatted BSF flour were compared to that of freeze-dried BSF flour (Fig. 501 7). Almost all proteins belonging to the soluble fractions were within the 25 and 75 kDa range. The most abundant protein band was close to 75 kDa followed by a sharp band next to 502 50 kDa, these proteins can be enzymes, muscle proteins, exoskeleton proteins or other 503 proteins like melanisation-inhibiting proteins as reported by Yi et al., 2013 for Tenebrio 504 *molitor* larvae. Ideally, since there was no visible change in the protein bands observed by 505 electrophoretic separation it can be assumed that no deleterious effects were imparted during 506 the defatting step at least in the soluble protein fraction. Further protein extraction, 507 fractionation, purification might help throw some light on the elusive protein profile of BSF. 508 The techno-functionality properties of flours and proteins from mealworm and BSFL was 509 510 compared by Bubler S. et al., 2016 who also found two soluble proteins from BSFL of which the most abundant was identified to have a molecular weight of 80.5 kDa and the other 14.3 511 kDa. Only one protein was designated in the reviewed section of the UniProt database, it was 512 cecropin-like peptide 1 with accession number: L7VIN3 and molecular mass of 4.84 kDa, 513 514 supposedly an antimicrobial peptide active against gram-negative bacteria.

The solvent 2-methyloxolane was efficient in lipid recovery and had improved the 515 protein quality in defatted BSF flour while considering it to that of n-hexane defatted flour, 516 making it a better solvent than n-hexane in all technical aspects. Addition of intensification 517 518 techniques, pre-treatment of the biomass will enhance the extractability and should also be considered to maximise efficiency. Microwave, ultrasound, thermal treatments were 519 effectively used for lipid extraction from yeast (Meullemiestre et al., 2016). A novel 520 521 Simultaneous Distillation and Extraction Process (SDEP) with alternative terpene solvents for lipid extraction was successfully demonstrated (Tanzi et al., 2013) proving the growing need 522 to find alternative green bio-based solvents to replace toxic petrochemical solvents. 523

#### 524 **4.** Conclusion

525 Numerous solvents were studied for the extraction of lipid constituents from BSFL 526 biomass. The solvent 2-methyloxolane (2-MeO) was found to be the ideal solvent based on 527 theoretical prediction and experimental results obtained. Extraction kinetics, diffusivity 528 modelling, and industrial simulation for lipid recovery were established. The yield was significantly higher in 2-MeO BSF oil in conventional as well as the industrial system.
Lauric, linoleic, palmitic, oleic, and myristic acids were the major fatty acids. Among the
lipid classes, free fatty acids were the major component, phospholipids were identified and
quantified in BSF oil (2-MeO) and were absent in BSF oil extracted with n-hexane.

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### Figure caption.

Figure 1. Application of hurdle technology for solvent screening.

Figure 2. Representative image for theoretical solubility prediction using COSMO-RS.

Figure 3. Graphical representation of the design of experiment.

Figure 4. a) BSF oil extraction kinetic curve b) Oil yield in multistage cross-current extraction system.

**Figure 5. a**) Relative content of neutral lipids in BSF oil **b**) Relative content of polar lipids in BSF oil **c**) HPTLC plate of neutral lipids.

**Figure 6. a)** Experimental versus predicted values of lipid extraction yields. •: n-hexane; •: 2-MeO; The solid lines represent the diffusion model. **b**) Visuals of lipid content distribution within larvae powders during extraction using the numerical simulation results.

**Figure 7.** Molecular weight distribution (kDa) of soluble proteins of freeze dried and defatted flour, FDBSF – Freeze Dried Black Soldier Fly; HR – n-hexane residue; 2-MeO – 2-methyloxolane residue.

Solute	FFA	MAG	DAG	TAG	VE1	VE2	ST1	ST2	CA1
Solvent	Relative Energy Difference : RED score								
n-hexane	2.24	3.21	2	1.24	1.4	2.03	1.5	1.6	1.49
Ethanol	3.22	2.23	3.42	4.39	4.39	4.25	4.42	4.28	4.9
Iso-propanol	2.3	1.34	2.47	3.48	3.44	3.33	3.49	3.35	3.97
Methyl acetate	0.89	1.09	1.15	1.69	1.88	2.07	1.91	1.79	2.41
Ethyl acetate	0.42	1.05	0.64	1.28	1.42	1.62	1.47	1.35	1.98
Ethyl lactate	1.55	0.65	1.81	2.69	2.73	2.69	2.76	2.62	3.26
DMC	1.34	0.97	1.62	2.29	2.44	2.54	2.47	2.34	2.97
2-MeO	0.89	1.75	0.98	0.73	0.89	1.09	0.84	0.75	1.27
CPME	0.83	1.76	0.85	0.56	0.73	1.02	0.71	0.61	1.17
α-pinene	1.56	2.55	1.35	0.46	0.49	1.11	0.57	0.68	0.67
d-limonene	1.07	2	0.92	0.55	0.24	0.48	0.23	0.13	0.7
p-cymene	1.45	2.39	1.34	0.53	0.43	0.72	0.29	0.38	0.42

**Table 1.** Solvent selection based on RED scores of Hansen Solubility parameters

Gray - Reference solvent; Green - Equivalent or better than reference; Red - Worse than reference \*FFA- Lauric acid; MAG- Glyceryl 1-laurate; DAG- Glyceryl 1,2-dipalmitate; TAG- Lauric triglyceride; VE1-  $\alpha$ -tocopherol; VE2-  $\gamma$ -tocotrienol; ST1-  $\beta$ -sitosterol; ST2- cholesterol; CA1-  $\beta$ -carotene

\*\*DMC- Dimethyl carbonate; 2-MeO- 2-methyloxolane; CPME- Cyclo pentyl methyl ether

Solute*	FFA	MAG	DAG	TAG	VE1	VE2	ST1	ST2	CA1
Solvent**	Solubility index : log <sub>10</sub> (x-solub)								
n-hexane	-1.0307	-2.1617	-0.6419	-0.1963	0	-0.345	-0.4479	-0.1823	0
Ethanol	0	0	-0.5865	-1.0797	-0.8543	-0.0836	-0.5312	-0.7166	-1.7267
Iso-propanol	0	0	-0.1874	-0.7162	-0.5073	0	-0.2727	-0.4068	-1.3477
Methyl acetate	0	0	0	-0.0394	0	0	-0.2694	-0.255	0
Ethyl acetate	0	0	0	0	0	0	-0.0307	0	0
Ethyl lactate	0	0	0	0	0	0	-0.0798	-0.0743	-0.1278
DMC	-0.2469	-0.3211	-0.7839	-0.6539	-0.5882	0	-0.8265	-0.8554	-0.5794
2-MO	0	0	0	0	0	0	0	0	0
CPME	0	0	0	0	0	0	0	0	0
α-pinene	-0.9093	-1.9464	-0.5993	-0.1173	-0.0119	-0.2729	-0.4623	-0.232	0
d-limonene	-0.7317	-1.6106	-0.3792	0	0	-0.0869	-0.3972	-0.1979	0
p-cymene	-0.7266	-1.5291	-0.4172	0	0	-0.0739	-0.4638	-0.2939	0

**Table 2.** Solvent selection based on the solubility index log10(x\_solub) of COSMO-RS

Gray - Reference solvent; Green - Ideal solvent; Yellow - Equivalent or better than reference; Red - Worse than reference \*FFA- Lauric acid; MAG- Glyceryl 1-laurate; DAG- Glyceryl 1,2-dipalmitate; TAG- Lauric triglyceride; VE1- α-tocopherol; VE2 + togetrignel: ST1 - β situatenel: ST2 - shelesterel: CA1 - β seretere

VE2-  $\gamma$ -tocotrienol; ST1-  $\beta$ -sitosterol; ST2- cholesterol; CA1-  $\beta$ -carotene

\*\*DMC- Dimethyl carbonate; 2-MO- 2-methyloxolane; CPME- Cyclopentyl methyl ether

Solvents / Parameters	Log P	Boiling point	Toxicity index	Energy required to evaporate 1 metric ton of solvent (kW.h)	Enthalpy of vaporization $[\Delta_{vap}H(T_{bp})]; kJ/mol$
n-hexane	3.9	68	6	120.1	28.85
Ethanol	-0.2	78.37	5	268.6	38.56
Iso-propanol	0.2	82.5	5	225.8	39.85
Methyl acetate	0.2	57.1	5	130.8	30.32
Ethyl acetate	0.7	77.1	5	128.7	31.94
Ethyl lactate	-0.2	154	5	171.0	45.57
DMC	0.2	90	5	134.8	33.05
2-MeO	0.8	80.2	4	126.1	30.74
CPME	1.4	106	4	132.4	33.00
α-pinene	4.4	155	4	142.5	37.83
d-limonene	4.4	176	5	153.8	37.83
p-cymene	4.0	177	5	155.8	39.34

## Table 3. Technical parameters for solvent screening

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Green – Good score; Yellow – Average score; Red – Poor score (relative comparison). DMC- Dimethyl carbonate; 2-MO- 2-methyloxolane; CPME- Cyclopentyl methyl ether.

Fatty acids	n-hexane (%)	2-MeO (%)
C10	$1.02 \pm 0.01$	$1.02 \pm 0.00$
C12	$42.27 \pm 0.13$	$42.29 \pm 0.18$
C14	$9.41 \pm 0.01$	$9.36 \pm 0.01$
C14:1	$0.19 \pm 0.01$	$0.18\pm0.00$
C15	$0.18 \pm 0.01$	$0.18\pm0.00$
C15:1	$0.06 \pm 0.01$	$0.06\pm0.00$
C16	$13.91 \pm 0.05$	$13.83 \pm 0.00$
C16:1	$2.73 \pm 0.02$	$2.67\pm0.00$
C18	$2.28\pm0.01$	$2.23 \pm 0.01$
C18:1n9	$11.84 \pm 0.05$	$11.43 \pm 0.12$
C18:2 n6 trans	$14.29\pm0.09$	$13.91 \pm 0.03$
C18:3n3	$1.41 \pm 0.01$	$1.37 \pm 0.01$
C20	$0.16 \pm 0.01$	$0.16 \pm 0.00$
C22	$0.06 \pm 0.00$	$0.06\pm0.00$
C22:2 n6	$0.17 \pm 0.01$	$0.16 \pm 0.00$
Σ SFAs	69.29	69.13
$\Sigma$ MUFAs	14.82	14.34
Σ PUFAs	15.87	15.44
Others	0.02	1.09

Table 4. Relative percentage of fatty acid profiles of BSF oil

SFA - Saturated Fatty acids

MUFA - Mono Unsaturated Fatty Acids

PUFA - Poly Unsaturated Fatty Acids

**Table 5.** Effective diffusion coefficients of lipids at different extraction conditions and accuracy of the diffusion model

Solvent	De $(m^2/s)$	$\mathbb{R}^2$	RMSE (mg/100g DM)	AAD (%)
n-hexane	$2.17 \times 10^{-9}$	0.99	0.438	3.27
2-MeO	$6.67 \times 10^{-10}$	0.97	1.013	4.75

De – Effective diffusion coefficient for lipid; R<sup>2</sup> – Coefficient of determination; RMSE – Root Mean Square Error; AAD – Absolute Average Deviation

 Table 6. Total Polyphenol Content and Radical Scavenging Capacity of BSF oil

	6 6	1 2
Description	TPC (mg GAE/g of oil)	RSC (mg TE/g of oil)
Refined sunflower oil (reference)	$3.60 \pm 0.09$	$0.79 \pm 0.17$
BSF oil (n-hexane)	$7.42 \pm 0.51$	$0.41 \pm 0.03$
BSF oil (2-MeO)	$19.03 \pm 1.11$	$5.42 \pm 0.76$

GAE - Gallic Acid Equivalent; TE - Trolox Equivalent.

 Table 7. Protein quality evaluation of defatted BSF flour

Protein quality parameters	Defatted BSF (n-hexane)	Defatted BSF (2-MeO)
Crude protein; %	$59.09 \pm 0.62$	$62.16 \pm 0.62$
PDI; %	29.09	31.55
PS - KOH; %	78.29	80.6
UI; $\Delta$ pH units	0.13	0.15
Abs @ 420nm	0.063	0.071

PDI -Protein Dispersibility Index; PS - Protein solubility in 0.2 % KOH.

UI - Urease Index; Abs - Absorbance at 420 nm.

## Figure 1.







Figure 3.





# Figure 4.









Figure 7.

