Glyceride structure and sterol composition of SOS-7 halophyte oil.

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1. INTRODUCTION

Egypt suffers from the shortage of edible oils and large tonnages of oils have been imported (about 500,000 tons) from the foreign market to meet the increasing local consumption amounting to 620,000 tons per year. Since local cottonseed oil production does not cover the requirements of local consumption of oils, several trials have been made to increase the cultivated area with oil seed crops, namely, soybean and sunflower with an attempt to fulfill the needs of local requirements. However, desert cultivation with new oilseed crops, as salt and climatic tolerant, have been successfully tried in small desert area in Egypt.

The Environmental Research Lab., ERL, Arizona (1), initiated the work in the field of halophyte plants and in particular, halophyte SOS-7. In continuation to the previous study in this Lab. (2) on characterizing halophyte, SOS-7, oil produced from locally cultivated halophyte plant, it was of interest to carry out further studies dealing with quantitative determination of the glyceride pattern of such oil. In addition, sterol components of the oil was also investigated using high efficient separation column of GLC as well as mass-coupling for confirming the structure of the sterol components.

2. GLYCERIDE STRUCTURE

2.1. Experimental

A sample from 1988 harvest of SOS-7 halophyte planted in Ghardaka, on the border of the Red Sea, Egypt, was used in this study. The oil seeds were ground and extracted for their oil using commercial hexane (b.r. 63 - 68°C) in Soxhlet extractor. The solvent was evaporated under reduced pressure and the resulting oil (25.3% of the weight of the seeds) was kept in well stoppered bottles in refrigerador until use.
Oil Analysis and Lipolysis

The oil characteristics were determined in a previous work (2). Part of the oil was converted into methyl esters using HCl gas as a catalyst (3) and the fatty acid composition of the oil was determined by gas liquid chromatographic analysis. Perkin Elmer Fractometer - FID7 with flame ionization detector and a column DEGS on chromosorb A was used at 190°C. The nitrogen, hydrogen and oxygen flow rates were 70, 25 and 300 ml/min, respectively.

Lipolysis of one gram sample was carried out in duplicate using 100 mg pork lipase according to Coleman and Fulton (4). After 30 minutes from the beginning of hydrolysis, about 60-70% of the original oil was converted into 2-monoglycerides, and the reaction was stopped by acidification. The hydrolyzed lipids were extracted with diethyl ether and washed till neutral, dried over anhydrous sodium sulphate and filtered. The solvent was removed under reduced pressure to obtain the residual lipids. The latter were dissolved in chloroform and separated by TLC using 0.5 mm layer silica gel G preparative plates and petroleum ether –diethyl ether– acetic acid (60 : 40 : 1 V/V/V) as the developing solvent. The located monoglyceride band was scraped off the TLC plate and eluted with chloroform. The solvent was removed and the monoglycerides were converted to their methyl esters (3) and the fatty acid composition of the 2-monoglycerides (2-MGS) was determined by GLC. The procedure and conditions of the analysis were the same as those used for the analysis of the fatty acid composition of the original oil.

From the fatty acid composition of the 2-MGS and the calculated fatty acids in 1- and 3-position, the distribution of fatty acyls in the original triglycerides was calculated according to Coleman (5). The relative proportion (r.p.) representing the percentages of fatty acid in question in the 2-MG, was also calculated (6) (7).

2.2. Results and discussion

The fatty acid composition of the halophyte oil seed is recorded in table I. This oil contains high content of unsaturated fatty acids (84.7%) which is characteristic of the vegetable oils, among them linoleic acid is the major constituent and forms 69% of the total fatty acids. The other two unsaturated fatty acids, namely oleic and linolenic constitute 14.3 and 1.4% of the mixed fatty acids. Saturated fatty acids represented by palmitic and stearic acids are detected in the halophyte oil forming 10.5 and 4.8%, respectively.

The component fatty acids of the 2-MGS resulting from lipolysis in comparison with those present in the whole triglyceride are shown in table I. r.p. values as a measure of the preference of each fatty acid in question for 2- and 1,3- positions are recorded also in table I. From the results, it can be seen that palmitic acid is preferentially esterified at 1- and 3-positions are recorded also in table I. The results, it can be seen that palmitic acid is preferentially esterified at 1- and 3-positions in the whole triglycerides since the r.p. value is lower than the random value of 33.3% (8). This is in agreement with the general distribution pattern of the saturated fatty acids reported for vegetable oils (6) (7) (8) (9) (10). Oleic and linolenic acids show high r.p. values corresponding to 51.3 and 50, respectively, indicating that oleic and linolenic acids compete nearly equally for the 2-position. Whereas linoleic having r.p. value of 35.7 shows comparatively low preference for the 2-position than oleic and linolenic acids. Gene-

Table I
Fatty Acid Composition of the Halophyte Oil Seed and the Corresponding 2-monoglycerides (2-MG) (Mole %)

<table>
<thead>
<tr>
<th>Fatty Acid</th>
<th>Whole TG (Fatty Acid compn.)</th>
<th>2-MG</th>
<th>r.p.*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Palmitic</td>
<td>10.5</td>
<td>1.8</td>
<td>5.7</td>
</tr>
<tr>
<td>Stearic</td>
<td>4.8</td>
<td>zero</td>
<td>zero</td>
</tr>
<tr>
<td>Oleic</td>
<td>14.3</td>
<td>22.0</td>
<td>51.3</td>
</tr>
<tr>
<td>Linoleic</td>
<td>69.0</td>
<td>74.1</td>
<td>35.7</td>
</tr>
<tr>
<td>Linolenic</td>
<td>1.4</td>
<td>2.1</td>
<td>50.0</td>
</tr>
</tbody>
</table>

*r.p. (Relative proportion) = F.A.% in 2-MG x 100/F.A.% in whole TG x 3; r.p. values exceeding 33.3% show more preference to 2-position.
Table II
Component glycerides (Mole %)

<table>
<thead>
<tr>
<th>Glyceride Type</th>
<th>Mole %</th>
<th>Glyceride Type</th>
<th>Mole %</th>
<th>Glyceride Type</th>
<th>Mole %</th>
<th>Glyceride Type</th>
<th>Mole %</th>
<th>Glyceride Type</th>
<th>Mole %</th>
</tr>
</thead>
<tbody>
<tr>
<td>P_3</td>
<td>0.05</td>
<td>P_2 O</td>
<td>0.62</td>
<td>P_2 O</td>
<td>0.72</td>
<td>0.3</td>
<td>0.23</td>
<td></td>
<td></td>
</tr>
<tr>
<td>St_2P</td>
<td>0.04</td>
<td>St_2 O</td>
<td>0.11</td>
<td>PL_2</td>
<td>16.37</td>
<td>L_2 O</td>
<td>19.39</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P_2 St</td>
<td>0.01</td>
<td>PStO</td>
<td>0.52</td>
<td>Pln_2</td>
<td>0.01</td>
<td>L_2 O</td>
<td>3.71</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>P_2 Ln</td>
<td>2.30</td>
<td>POl</td>
<td>7.25</td>
<td>L_2 O</td>
<td>0.07</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>St_2 Ln</td>
<td>0.40</td>
<td>Poln</td>
<td>0.14</td>
<td>LnO_2</td>
<td>0.75</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>PStLn</td>
<td>1.88</td>
<td>PLln</td>
<td>0.72</td>
<td>LnO_2</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>P_2 Ln</td>
<td>0.06</td>
<td>StO_2</td>
<td>0.32</td>
<td>Ln_2 O</td>
<td>1.93</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>St_2 Ln</td>
<td>0.01</td>
<td>StL_2</td>
<td>6.86</td>
<td>Ln_2 O</td>
<td>0.04</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>PStln</td>
<td>0.05</td>
<td>StLn_2</td>
<td>0.00</td>
<td>L_3</td>
<td>0.00</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>StOL</td>
<td>3.10</td>
<td>L_3</td>
<td>31.98</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>StOLn</td>
<td>0.06</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>StLLn</td>
<td>0.31</td>
<td>-</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>0.10</td>
<td>Total</td>
<td>5.95</td>
<td>Total</td>
<td>35.86</td>
<td>Total</td>
<td>58.11</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

rally, the proportionally high concentration of oleic, linolenic and linoleic acids in the 2-position has been ascribed to the fact that specific distribution of saturated acids (in this case palmitic and stearic acids) in 1- and 3-position forces a higher proportion of oleic, linoleic and linolenic in the 2-position of the triglyceride molecules (6) (11). Concerning the type of glycerides calculated, namely, trisaturated (GS3), disaturated (GS2U), monosaturated (GSU3) and triunsaturated (GU3), the results are recorded in table II.

The component triglycerides of GS3 type, contain tripalmitin P_3 and distearo-palmitin (St_2P) as major components constituting 0.05 and 0.04% respectively, whereas dipalmito-stearin (P_2St) is found in lower concentration of 0.01%.

Concerning the GS2U, it can be noticed that dipalmito-olein (P_2L) and palmito-stearo-olein (PStL) are present as major components in halophyte oil. Dipalmito-olein (P_2O), palmito-stearo-olein (PStO) and distearo-olein (St_2L) are found in comparatively low concentrations (0.62, 0.52 and 0.4% respectively). Minor triglyceride components of the GS2U type, namely, St_2O, P_2 Ln, St_2 Ln and PStL are also detected.

With reference to component glycerides of the GSU3 type (table II), palmito-dilinolein (PL_2) is the major component which constitutes 16.37% of the total GSU3 type. Palmito-oleo-linolein (POL), stearo-dilinolein (StL_2) and stearo-oleo-linolein (StOL) are present in moderate concentrations amounting 7.25, 6.86 and 3.10% respectively, whereas POl, PLln, SlO_2 and SlLn are found in low concentrations (0.72, 0.72, 0.32 and 0.31% respectively). Other minor triglycerides of the type GSU_3, namely, PLn_2, POLn, StLn_2 and StOLn are also detected.

From the results representing glyceride components of the GU3 type, it can be seen that trilinolein (L_3) constitutes 31.98 which corresponding to more than 50% of the GU3 type. Oleodilinolein (L_2O) comes after L_3 (19.39 mole%) which corresponding to nearly 33% of the total GU3 type. LO_2 and LnL_2 are present in comparatively low concentrations (3.71 and 1.93 respectively). O_3, LnO_2, LnO_3, LnOL, LnOLn, LLn_2 and Ln_3 are detected in minor amounts.

From the above results, it can be concluded that the 2-position in the halophyte triglycerides is mainly occupied by the unsaturated C_18 which agrees with other investigators (4) (6) (7) (9) and this also confirms the non-random character of the acyl group.
distribution in vegetable oil glycerides and has led to the wide acceptance of the theory of positional distribution (5) (12) (13) (14) in place of other theories.

3. STEROL COMPOSITION

3.1. Experimental

Identification and quantitative determination of sterols were carried out using reference standards in ethanol (campsterol, stigmasterol, B-sitosterol and 7-stigmasterol) as well as cholesterol as internal standard (IS). The procedure of Slover et al (15) was applied for the identification and determination of the sterol components.

Preparation of Pure Sterols

A mixture of oil sample (0.5 g) and 5 ml of IS was saponified with alcoholic 1N KOH for one hour on a water bath. The sterol fraction was separated from the unsaponifiable matter by preparative TLC on silica gel G plates (0.5 mm thick) using chloroform / ether / acetic acid (94 / 5 / 1, V/V/V). Sterols mixture was subjected to silylating reaction using two parts of hexamethyl-disilazane and one part of trimethyl-chlorosilane with ten parts of dry pyridine. After standing for at least 15 minutes at room temperature, the mixture was injected directly into gas chromatograph (15).

Gas Liquid Chromatography GLC*

Hewlett Packard - HP 5890 A, gas chromatograph equipped with flame ionization detector and 2m x 4mm glass column packed with 3% OV-1 on Gas Chrome Q (80-100 mesh) was used. The operating temperatures were: Column 270-280°C, injection and detector 280°C. Nitrogen gas was used as carrier gas at 40 ml/min.

Gas Liquid Chromatography - Mass Spectrometry GC - MS*

An instrument Hitachi M-80 B was used for this purpose under the following conditions: Column Diasolid ZS 2m x 4mm; carrier gas, helium at 35 ml/min; oven temperature, 270°C; interface temperature, 280°C; ionization voltage, 20 ev; ionization mode, electric (EI) and ion source temperature 200°C. The effluent arriving the mass spectrometer was detected by total ionization monitor.

<table>
<thead>
<tr>
<th>Campsterol</th>
<th>Stigmasterol</th>
<th>B-Sitosterol</th>
<th>7-Stigmasterol</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \Delta^5, C_{28} )</td>
<td>( \Delta^5,22, C_{29} )</td>
<td>( \Delta^5, C_{29} )</td>
<td>( \Delta^7,22, C_{29} )</td>
</tr>
<tr>
<td>M/e</td>
<td>M/e</td>
<td>M/e</td>
<td>M/e</td>
</tr>
<tr>
<td>473 ( M^+ )</td>
<td>465 ( M^+ )</td>
<td>487 ( M^+ )</td>
<td>487 ( M^+ )</td>
</tr>
<tr>
<td>458 ( M-CH_3 )</td>
<td>470 ( M-CH_3 )</td>
<td>472 ( M-CH_3 )</td>
<td>472 ( M-CH_3 )</td>
</tr>
<tr>
<td>382 ( M-OTMS )</td>
<td>394 ( M-(OTMS+2H) )</td>
<td>396 ( M-(OTMS+2H) )</td>
<td>396 ( M-(OTMS+2H) )</td>
</tr>
<tr>
<td>367 ( M-(OTMS+CH_3+2H) )</td>
<td>379 ( M-(OTMS+CH_3+2H) )</td>
<td>381 ( M-(OTMS+CH_3+2H) )</td>
<td>381 ( M-(OTMS+CH_3+2H) )</td>
</tr>
<tr>
<td>343 ( M-(side chain+H_2O) )</td>
<td>344 ( M-(side chain+2H) )</td>
<td>345 ( M-(side chain +1) )</td>
<td>345 ( M-(side chain +1) )</td>
</tr>
<tr>
<td>255 ( M-(side chain +CH_3+2H) )</td>
<td>255 ( M-(side chain +OTMS+2H) )</td>
<td>255 ( M-(side chain +OTMS+2H) )</td>
<td>255 ( M-(side chain +OTMS+2H) )</td>
</tr>
</tbody>
</table>

3.2. Results and discussion

The GLC analysis of mixed sterols of halophyte oil reveal the presence of four sterols which stand for campsterol, stigmasterol, B-sitosterol and 7-stigmasterol at ratios of 1.3, 7.3, 39.0 and 52.4% respectively. Since the oil appears to be unique in its sterol composition, it was advisable that the sterol components should be subjected to GC - MS for elucidating their structures. The effluent passing to mass spectrometer from the gas chromatograph is detected by Total Ion Monitor (TIM) and the mass spectra of the components are obtained. Mass spectra of 7-stigmasterol, as a representative sterol, is shown in the figure. It was found that the sterols detected by GLC are confirmed by GC - MS to be campsterol, stigmasterol, B-sitosterol and 7-stigmasterol as shown from the m/e fragments (table III). The fragments from 7-stigmasterol of halophyte, major sterol, found to be similar to those reported by Itoh et al (16). The presence of 7-
stigmasterol as major constituent of halophyte sterols is peculiar and is unlike other plants which has β-sitosterol as a major component.

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REFERENCES


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