THE BIOSYNTHESIS OF THE FATTY ACIDS OF THE PLASMA OF MAN.
I. THE FORMATION OF CERTAIN CHROMATOGRAPHICALLY
SEPARATED HIGHER FATTY ACIDS OF THE MAJOR
LIPIDE COMPLEXES FROM ACETATE–1–C\textsuperscript{14}

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One of the outstanding difficulties precluding the use of radioactive tracers to study the dynamics of lipide metabolism in man has been the lack of adequate micromethods for the isolation and identification of the major lipide complexes present in the plasma and the individual fatty acids associated with these groups. In recent years, with the advent of new chromatographic procedures, two significant achievements have occurred in this sphere. By the use of silicic acid columns, Borgström (1, 2) in 1952 and Fillerup and Mead (3) in 1953 separated lipide extracts into sterol ester, triglyceride, free fatty acid and phospholipide fractions. Crombie, Comber, and Boatman (4) extended the reverse phase partition chromatographic technique of Howard and Martin (5), and while encountering considerable overlapping of zones, demonstrated the separation of milligram quantities of certain common saturated and unsaturated fatty acids contained in natural mixtures.

The present investigation was undertaken in an effort to combine and modify these methods in order to determine: (a) the nature and quantity of the specific higher saturated and unsaturated fatty acids that form ester linkages with the lipide complexes of the plasma of man and (b) the rates of synthesis of these acids following the administration of acetate–1–C\textsuperscript{14}.

METHODS AND MATERIALS

Part I. The separation of the total lipide extract of the plasma into the total fatty acids of the sterol esters, the triglycerides and the phospholipides

a) Extraction. Four patients with limited life expectancies due to carcinoma of the lung without evidence of metastases were selected for study. All were considered to be in a good nutritional state and clinically free from gross metabolic disturbances. The subjects were maintained on normal hospital diets during the course of the investigation. Two hundred microcuries of acetate–1–C\textsuperscript{14} (specific activity 1.0 mc. per mM) were dissolved in a convenient volume of tap water and administered orally to all patients in the postabsorptive state on the morning of the experiment. Serial blood samples of approximately 60 ml. were taken in heparinized syringes beginning one hour after the administration of acetate and continued at intervals for 96 hours. Thirty-five ml. of plasma was then removed after centrifugation and rapidly blown into a one-liter Erlenmeyer flask containing 525 ml. of a 4:1 mixture of dimethoxymethane: methanol (6). At this point in some preliminary experiments, individual carbon-14 or tritium labelled lipides of known specific activity were dissolved in one ml. of petroleum ether and added to the flask. The mixture was allowed to boil momentarily by cautiously rotating the flask under a hot water tap. After cooling for fifteen minutes, with frequent swirling, the mixture was filtered through an 18.5 cm. Whatman No. 1 filter paper into a one-liter round bottom flask with a side arm. The precipitate was washed twice with an additional 30 ml. of the dimethoxymethane:methanol mixture and the washings added to the lipide extract contained in the flask. The flask was placed on a constant temperature water bath maintained at 50 °C and the clear yellow solution was concentrated to a small volume under reduced pressure. Throughout this procedure oxygen-free nitrogen was allowed to flow onto the surface of the liquid via the side arm. The capillary tip by which the nitrogen was delivered was not placed below the surface of the liquid since this tended to induce excessive foaming as evaporation proceeded. Forty ml. of warmed

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\textsuperscript{2}The labelled radiochemicals were obtained from the following sources: a) Triolein—courtesy of Dr. David Krichevsky, American Cyanamid Co., Pearl River, N. Y. (7). b) Tripalmitin, tristearin, palmitic acid, and stearic acid—Isotopes Specialties Co., Burbank, Calif. c) Sterol Ester—prepared according to the procedure of Borgström (8). d) Phospholipides—prepared biosynthetically following the administration of acetate–1–C\textsuperscript{14} to a subject. The phospholipides of the lipide extract of the plasma were obtained by precipitation with cold acetone and ethanolic MgCl\textsubscript{2} followed by repeated washings.
petroleum ether (b.p. 30° to 60° C) was then added to the flask, the flask was stoppered with a ground glass stopper and shaken vigorously for two minutes. The supernatant petroleum ether layer was transferred to a 300-ml. round bottom flask with a side arm by means of a simple thin glass tube siphon device attached to water suction. The petroleum ether extraction was then repeated four more times. It was found that for quantitative recoveries, this technique of transfer was superior in all respects to that employing the use of separatory funnels.

The petroleum ether extracts were combined and the solvent was evaporated off under nitrogen at reduced pressure as described. The total lipid extract was then stored under nitrogen at -4° C until it was to be chromatographed.

b) **Chromatographic separation on silicic acid columns.**

A snugly fitting piece of coarse filter paper was placed on top of the sintered disc at the bottom of the chromatography tube depicted in Figure 1. Twenty grams of silicic acid (100 mesh, Mallinckrodt) was thoroughly mixed with 4 grams of "Hyflo-Supercel" (Johns-Manville) and added to the tube. This ratio of filter aid to absorbent was found to provide excellent resolution with an adequate flow rate. The mixture was then packed by tapping the tube gently on a firm surface. Another filter paper disc was placed on top of the column and the column was washed with 70-ml portions of absolute methanol, acetone, anhydrous peroxide-free ether and petroleum ether (3). The lipid extract to be chromatographed was dissolved in four to eight ml. of warmed 1 per cent ether in petroleum ether and quantitatively added dropwise to the top of the column. The sides of the column were then washed down with an additional five to ten ml. of solvent. In preliminary experiments (Table 1), elution was carried out with the solvent systems recommended by Fillerup and Mead (3, 9). Since it was noted that some of the phospholipid fraction was eluted prematurely by solvent systems containing moderate concentrations of diethyl ether, the following technique was adopted to separate the sterol ester, triglyceride and phospholipid components of the plasma. Following the concentration of the total lipid extract prior to chromatographic separation, this material was taken up in a small volume of petroleum ether and quantitatively transferred to a heavy walled 40-ml centrifuge tube. The final volume was reduced to about two to four ml. by evaporating off the excess of petroleum ether under a stream of nitrogen. Thirty ml. of cold acetone and ten drops of saturated MgCl₂ in ethanol were then added to the tube to precipitate the phospholipides. The tube was agitated and then placed in a refrigerator and allowed to stand overnight. Following centrifugation, the supernatant which contained the entire lipid extract with the exception of the phospholipides was siphoned off into a 200-ml. round bottom flask with a side arm. The precipitated phospholipides were washed two to three times with 30-ml. portions of cold acetone. After repeat centrifugation the acetone supernatants were then combined. Recoveries for phospholipides by this technique ranged from 97 to 102 per cent. Control cross contamination experiments performed by the addition of carbon-14 labelled sterol, fatty acid, triglyceride, and biosynthetically prepared radioactive phospholipide usually demonstrated less than 1 per cent contamination of the phospholipide precipitate by non-phospholipide constituents. Similar results were obtained when the acetone supernatants were assayed for contamination by the phospholipides. These findings are in accord with the observations of Borgström (1) employing the same technique.

The combined acetone supernatants were evaporated to dryness under nitrogen at reduced pressure. The residue was taken up in two to five ml. of warmed 1 per cent ether in petroleum ether and applied to the top of the silicic acid column as described. The sterol esters were eluted with 300 ml. of 1 per cent ether in petroleum ether; the triglycerides with 350 ml. of 3 per cent ether in petroleum ether. An elution rate of approximately 150 ml. per hour was maintained in all experiments by means of a "Manostat" attached to a vacuum pump. After separation the eluates were evaporated down to dryness as previously described, taken up in petroleum ether and then quantitatively transferred to a 100-ml. round bottom flask with a side arm. The phospholipide precipitate was also dissolved in a small volume of warm petroleum ether and transferred in a similar fashion.

c) **Saponification.** To each flask were added 20 ml.
of 95 per cent ethanol and 0.7 ml. of 90 per cent aqueous KOH. A small Soxhlet water condenser was attached to the top of each flask, a stream of nitrogen introduced via the side arm, and the material refluxed for 90 minutes in a boiling water bath. During the procedure, evaporation of approximately 50 to 75 per cent of the ethanol occurred. The samples were then removed from the bath and allowed to cool. Five ml. of H₂O and twenty ml. of 95 per cent ethanol were added to each flask. The mixture was extracted three times with petroleum ether. The petroleum ether washings were combined and washed once with alcoholic KOH. This ethanol wash was added to the original aqueous material remaining in the flask and the petroleum ether extracts were then discarded. The mixture was made acid to phenolphthalein with 6 N H₂SO₄. Ten to twenty ml. of H₂O were added and the mixture was then extracted three times with 20-ml. aliquots of petroleum ether. The combined petroleum ether extracts were washed once with 5 per cent sodium bicarbonate, three times with distilled water, and then dried over anhydrous sodium sulfate. The solvent was evaporated off under nitrogen at reduced pressure and the fatty acid residue was transferred to a 25-ml. volumetric flask and brought up to volume with petroleum ether.

d) **Assay.** Portions of these samples were then taken in duplicate, placed in small test tubes and the solvent blown off with a stream of nitrogen. The residue was taken up in 2 ml. of 65 per cent acetone in water (v/v) and two drops of bromothymol blue added. The total fatty acids of each class of lipides was then determined by titrating the sample under nitrogen against 0.005 N KOH in a microburette calibrated in 0.001 ml. In experiments where radioactive tracers were employed, another sample was removed from the volumetric flask, transferred to a 30-ml. counting vial (Kimble K-10 "Opticlear") and evaporated to dryness. To this vial was added 15 ml. of toluene containing 0.4 per cent of the phosphor, 2,5 diphenylloxazole and 0.005 per cent of the wave length shifter 1, 4-di-[2-(5-phenyloxazolyl)]-benzene (10). The radioactivity contained in the vial was determined in a dual channel liquid scintillation counter 8 which had an efficiency of 75 per cent for carbon-14 (with an 11:1 energy acceptance window and a background of 30 cpm.) and 22 per cent for tritium. All counts were corrected for background and counted to a standard error of 2 per cent.

**Part II. The isolation of individual higher saturated and unsaturated fatty acids commonly found in ester linkage with the major lipide complexes**

a) **Preparation of kieselguhr columns.** Five pounds of "Hyflo Supercel" (Johns-Manville) were thoroughly

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**TABLE I**

Distribution of added radioactivity appearing in the various chromatographed fractions of plasma lipides in order of elution

<table>
<thead>
<tr>
<th>Solvent system</th>
<th>Radioactive tracer*</th>
<th>Sterol esters</th>
<th>Triglycerides</th>
<th>Free fatty acids</th>
<th>Sterol</th>
<th>Combined fraction</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>Triolein-H²</td>
<td>3,120</td>
<td>36</td>
<td>36</td>
<td>30</td>
<td>19</td>
</tr>
<tr>
<td></td>
<td>Stearic acid-1-C₁⁴</td>
<td>15,140</td>
<td>30</td>
<td>37</td>
<td>24</td>
<td>29</td>
</tr>
<tr>
<td></td>
<td>Palmitic acid-1-C₁⁴</td>
<td>11,017</td>
<td>37</td>
<td>43</td>
<td>97</td>
<td>24</td>
</tr>
<tr>
<td></td>
<td>Cholesterol-4-C₁⁴</td>
<td>20,980</td>
<td>24</td>
<td>97</td>
<td>14,070</td>
<td>228</td>
</tr>
<tr>
<td></td>
<td>Cholesterol-4-C₁⁴</td>
<td>20,797</td>
<td>36</td>
<td>106</td>
<td>13,895</td>
<td>185</td>
</tr>
<tr>
<td>B</td>
<td>Sterol Ester-C₁⁴†</td>
<td>8,508</td>
<td>52</td>
<td>83</td>
<td>42</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>Triolein-1-C₁⁴</td>
<td>13,987</td>
<td>26</td>
<td>83</td>
<td>42</td>
<td>44</td>
</tr>
<tr>
<td></td>
<td>Stearic acid-1-C₁⁴</td>
<td>11,737</td>
<td>26</td>
<td>83</td>
<td>11,521</td>
<td>228</td>
</tr>
<tr>
<td></td>
<td>Cholesterol-4-C₁⁴</td>
<td>16,335</td>
<td>56</td>
<td>145</td>
<td>15,636</td>
<td>238</td>
</tr>
<tr>
<td></td>
<td>Phospholipide-C₁⁴†</td>
<td>1,991</td>
<td>28</td>
<td>83</td>
<td>170</td>
<td>1,715</td>
</tr>
</tbody>
</table>

* Counts per minute added to the column.
† Typical examples of the separation of certain lipide fractions by chromatography with silicic acid using various solvent systems.
mixed with several gallons of distilled water in a high wide-mouth glass container. The mixture was allowed to settle for ten minutes, after which time the fine particles that float were removed by aspiration. The washing process was repeated three to four more times. The Celite was then transferred to large glass trays, excess water decanted, and the tray placed in a drying oven maintained at 110° for 24 hours. The material should be thoroughly dry before further processing. The Celite was then transferred to large evaporating dishes and placed in a large vacuum desiccator at the bottom of which was 20 ml. of dichlorodimethylsilane in a wide-mouth container. The desiccator was partially evacuated and the silane vapors allowed to remain in contact with the Celite for four hours. The Celite was then removed and placed in a 3-liter beaker filled with water. The mixture was vigorously stirred and the particles that did not float were discarded. The coated Celite was transferred to a large Buchner funnel and washed several times with water and then methanol until the washings appeared neutral when tested with pH paper. The material was then thoroughly dried in an oven maintained at 80° C.

The most satisfactory separation of certain mixtures of the higher saturated and unsaturated fatty acids was obtained by using a jacketed column 1 cm. in diameter and 130 cm. in height equipped with a small reservoir at the top and an "Ultramax" valve at the bottom. For a column of this size, 48 grams of coated Celite and 34.8 grams of Standard White Oil No. 15 USP 4 were placed in separate beakers. One hundred ml. of anhydrous peroxide-free ether was added with stirring to the mineral oil. The Celite was then added. The beaker was placed in a warm bath and the mixture gently stirred to hasten the evaporation of the solvent. The coarse powder was then dried in a vacuum oven at 60° C for two hours. A convenient volume of 70 per cent acetone in water (v/v) was added and the mixture agitated for one minute in a Waring Blender to dissolve any lumps. To expel the air bubbles, the Celite was transferred to a beaker and warmed gently on a steam bath until bumping occurred. The slurry was then added to a chromatographic column filled with 70 per cent acetone by means of a large-bore dropping pipette. The Celite was uniformly packed by gravity by opening the column and allowing it to flow at a maximum rate. Air-free columns have been obtained by observing two precautions: a) the tip of the pipette by which the slurry is delivered should be placed below the meniscus of fluid at the top of the column before discharging the contents; b) the temperature of the water flowing through the jacket of the column should be the same as that of the slurry and the solvent systems (room temperature). A filter paper disc was placed on top of the Celite column and the column was then washed with 150 ml. of 55 per cent acetone before use.

b) The loading of fatty acids on the column (II). Ten to twenty-ml. samples of the total fatty acids of various lipide classes that were dissolved in petroleum ether and placed in volumetric flasks were removed and placed in a 50-ml. beaker. The solvent was evaporated off on a warm water bath under a stream of nitrogen. The residual fatty acids were then dissolved in 0.4 ml. of Standard White Oil and 15 ml. of anhydrous peroxide-free ether. Then 0.52 gram of coated Celite was added. The beaker was placed in a tray of warm water, and with constant stirring under a stream of nitrogen, the ether was blown off. The thoroughly dried powdery mull was then added to the top of the chromatographic column with a spatula. The beaker was rinsed several times with several ml. of warmed 55 per cent acetone and the washings were also added to the top of the column. The Mull containing the acids was gently agitated with a glass rod and allowed to settle by gravity. Another filter paper disc was then placed on top of the Mull.

* Standard Oil Company of California.
The synthesis of specific fatty acids from acetate

FIG. 4. The Chromatographic Separation of the Total Fatty Acids of Sterol Esters of the Plasma of Subject J

The solvent systems employed in sequence in the separation of known mixtures or plasma extracts containing the common higher fatty acids were as follows:

- 55 per cent acetone elutes arachidonic acid
- Linolenic acid
- Lauric acid
- 60 per cent acetone elutes linoleic acid
- Myristic acid
- 65 per cent acetone elutes palmitic acid
- Oleic acid
- 70 per cent acetone elutes stearic acid

The systems were changed at intervals noted in Figures 2 to 6. The capacity of the column for total fatty acids was approximately 60 milligrams. The columns may be used repeatedly (5 to 10 times) provided they are kept at a constant temperature. Under these conditions, elution was quantitative and the removal of individual fatty acids with predetermined volumes of the various solvent systems could be anticipated.

c) Collection and assay of samples. The columns were mounted on an automatic fraction collector adjusted to collect 5-ml. samples every three minutes. The samples were titrated under nitrogen with 0.005 N KOH to a blue-green endpoint using bromthymol blue as an indicator. The contents of the tubes representing individual fatty acids were pooled, made acid with 6 N H₂SO₄ and concentrated under nitrogen at reduced pressure. The fatty acids were then extracted with petroleum ether, washed, dried and assayed for content and radioactivity as previously described.

§ For weight measurements this fraction was subjected to the Bertram oxidation procedure (4) and then rechromatographed for the quantitative estimation of palmitic acid. The amount of oleic acid present in the original mixture was then obtained by subtracting the weight (by titration) of the recovered palmitic acid from that of the acids eluted by 65 per cent acetone during initial chromatography.

RESULTS

Nature and quantity of the fatty acids found in ester linkage with the various lipide complexes of the plasma

Typical examples of the resolving power of the silicic acid columns for the separation of the sterol esters and the triglycerides of the plasma using the solvent system recommended by Fillerup and Mead (3) are shown in Table 1. In a series of quantitative experiments, recoveries of 95 per cent or more were constantly achieved for these fractions. Attempts at further fractionation of the lipide extract by this technique, however, were generally unsuccessful. Considerable cross contamination between "free fatty acids" and sterols was noted. The gross overlapping of zones that occurred could not be obviated by either increasing the column length or altering the concentrations of the solvent systems employed in this investigation. Although the phospholipide fractions were generally eluted in a pure state by 25 per cent methanol in ether, the yields were frequently 10 to 20 per cent lower than those obtained by precipitation of the phospholipides in the cold by acetone and MgCl₂. Losses appeared to be due primarily to the premature separation of small quantities of

FIG. 5. The Chromatographic Separation of the Total Fatty Acids of the Triglycerides of the Plasma of Subject J

§ Added radioactive fatty acids.
phospholipide when higher concentrations of di-
ethyl ether were used in the elution of "free fatty
acids" and sterols from the column.

The quantities of total fatty acids found in ester
linkage with each of the major lipide complexes of
the plasma are noted in Table II. These values
compare favorably with those obtained by indirect
measurements (12). The fatty acids of the trigly-
cerides in the post-prandial plasma were usually
present in the highest concentration, followed
by the fatty acids of the phospholipides and then
those of the sterol esters.

The results of the qualitative and quantitative
resolution of known mixtures of some of the com-
mon straight-chain higher saturated and unsatu-
rated fatty acids are depicted in Figures 2 and 3.
Highly satisfactory separation of linolenic, linoleic,
palmitic or oleic and stearic acids was accom-
plished by using kieselguhr column 1 cm. in
diameter and 120 cm. in height. Preliminary ex-
periments with shorter columns led to adequate
resolution of palmitic and stearic acids. However,
considerable overlapping was noted in the zones
distinguishing linolenic from linoleic acid and lin-
oleic from palmitic or oleic acid. As noted by pre-
ioius investigators (4), the presence of each double
bond some distance from the carboxyl group
causcd an acid to act like that of a saturated acid
having a chain length of two less carbons. Thus,
oleic, linoleic, and linolenic acids behave as and
are indistinguishable from C_{18}, C_{14}, and C_{12} straight-
chain saturated acids, respectively. Similarly the
C_{20} tetraene, arachidonic acid (an acid not avail-
able in pure form at this time) is theoretically
eluted in a manner akin to that of the C_{18} triene,
linolenic acid (Figures 2 and 3) or the saturated
C_{12} lauric acid.

The chromatographic separation of the fatty
acids of the sterol esters, the triglycerides, and the
phospholipides of the plasma is depicted in Fig-
ures 4, 5, and 6. It is noted that the predominant
components of all fractions are presumably linoleic,
oleic, palmitic, and stearic acids. The various con-
centrations of the individual fatty acids associated
with the major lipide complexes are listed in
Table III. It should be stated that some titratable
acidity, approximately two to four times that of
the blank, occurred in the fractions eluted by 55
per cent acetone in water (v/v). Although this
may represent very small quantities of either lauric
(13), linolenic and arachidonic acids (4), or the
products of partial oxidation of some of the higher
unsaturated acids, no definite "hill and valley" elu-
tion pattern was noted. The very small band that
did form tended to spread widely. This was prob-
ably due to the high water content of the solvent
system employed under these circumstances (4,
5). Since the small quantities of these acids that
were recovered precluded further accurate chemi-
cal identification at this time, they have been ex-
cluded from the table. It is, therefore, apparent
that the small amounts of the higher unsaturated
acids of the plasma, as determined by the alkali
isomerization technique (14), are not easily dis-
cernible by this method at this time.

From an analysis of Table III, it appears that
oleic and palmitic acids, respectively, represent
the major unsaturated and saturated fatty acids of
the plasma of man. It is also noted that the fatty acids
associated with the sterol esters are predominantly
unsaturated—linoleic and oleic acids constituting
approximately 86 per cent of the titratable acidity

![Fig. 6. The Chromatographic Separation of the Total Fatty Acids of the Phospholipides of the Plasma of Subject J](image)
THE SYNTHESIS OF SPECIFIC FATTY ACIDS FROM ACETATE

TABLE III
The distribution of the individual fatty acids associated with the major lipide complexes in the plasma of subject J

<table>
<thead>
<tr>
<th>Fatty Acids</th>
<th>Linoleic Acid mg. %</th>
<th>Oleic Acid mg. %</th>
<th>Stearic Acid mg. %</th>
<th>Palmitic Acid mg. %</th>
<th>Total Fatty Acids of Major Complex mg. %</th>
<th>Per cent Recovery of Total Fatty Acids Added to the Column</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sterol esters</td>
<td>57.7</td>
<td>31.9</td>
<td>2.2</td>
<td>11.5</td>
<td>103.3</td>
<td></td>
</tr>
<tr>
<td>Triglycerides</td>
<td>18.9</td>
<td>68.9</td>
<td>7.3</td>
<td>45.1</td>
<td>140.2</td>
<td>89</td>
</tr>
<tr>
<td>Phospholipides</td>
<td>43.1</td>
<td>31.2</td>
<td>16.1</td>
<td>23.1</td>
<td>113.5</td>
<td></td>
</tr>
</tbody>
</table>

Per cent distribution of the individual fatty acids recovered from the total fatty acid extract

- Sterol esters: 33.5
- Triglycerides: 37.0
- Phospholipides: 7.2

Per cent distribution of the individual fatty acids present in each major complex

- Sterol esters: 55.8
- Triglycerides: 13.4
- Phospholipides: 37.9

* Obtained by analysis of the chromatographically separated fractions.

The distribution of the individual fatty acids recovered from the kieselguhr column after the addition of this fraction. Similarly, 65 per cent of the titratable acidity associated with the phospholipides and 62 per cent of that found with the triglycerides were attributable to these unsaturated acids. Palmitic and stearic acids, on the other hand, were more widely distributed in the triglyceride and phospholipide complexes, with only small quantities noted in the sterol esters.

It is highly probable that both the composition and concentration of the individual fatty acids associated with each of the major lipide complexes vary considerably with the diet (15-17).

Incorporation of acetate-1-C\(^{14}\) into the total fatty acids of triglycerides, the sterol esters and the phospholipides (Figure 7, Table IV)

The appearance of radioactivity in the total fatty acids of the triglycerides following the administration of the C\(^{14}\)-labelled two-carbon fragment was extremely rapid in all subjects. The specific activity of these substances invariably reached a maximum at two hours, declined sharply within a 24-hour period and then slowly declined during the next 72 hours. A semilogarithmic plot of the data demonstrates that decay does not occur by a single exponential process, and if additional points were taken over several more days the curves describing the decline of radioactivity probably could be resolved into a series of exponential rates. Thus, the calculation of a composite half-life for these fatty acids would not be very meaningful at this time. Undoubtedly a number of metabolic processes contribute to the disappearance of these materials from the blood stream (18).

The curve describing the specific activity of the fatty acids of the phospholipides increased more gradually to reach a plateau between 12 and 24 hours, at which point it intersected the specific activity curve of the fatty acids of the triglycerides and then fell off more slowly within the next 72 hours.

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**Figure 7.** The Incorporation of Acetate-1-C\(^{14}\) into the Total Fatty Acids of the Triglycerides, the Phospholipides, and the Sterol Esters of the Plasma of Subject J
the triglyceride creased very
synthesis
approximately 50
fatty acids of the
height
slowly. At
very
point
which
cay
curve
the fatty acids
ester
linkages
and
more
activity
cific
of sterol
Subject
hours
Specific activity of
V
1
D
Subject
Time in
hours
CPM/
0.005 mM*  
CPM/
0.005 mM  
CPM/
0.005 mM
J
1
2
4
8
12
24
48
96
2,660 29 170
2,600 48 378
1,680 80 628
1,050 96 320
406 120 766
202 126 532
96 81 202

V
1
2
4
8
12
24
48
96
2,840 25 100
5,705 41 251
4,631 74 561
3,589 110 933
1,905 122 960
729 151 1,280
387 178 770
187 103 320

S
1
2
4
8
12
24
48
96
2,025 10 51
8,381 16 130
5,267 25 270
3,761 42 473
2,470 45 627
1,386 63 472
540 69 395
190 40 165

D
1
2
4
8
12
24
48
96
1,300 27 30
3,625 60 80
3,481 90 146
2,733 152 322
1,375 191 397
560 226 332
302 252 260
140 140 110

* Counts per minute per 0.005 millimole.

hours. When plotted semilogarithmically the
decay curve extending from 24 to 96 hours declined
exponentially and the half-life of this fraction was
approximately 50 to 60 hours. On the other hand,
the synthesis of labelled fatty acids entering into
ester linkages with the sterols of the plasma in-
creased very slowly during the first 48 hours, at
which point it reached its maximum and then fell
very slowly. At no time did the specific activity
of the fatty acids of sterol esters exceed that of
the triglyceride or phospholipide fatty acids. At
the height of incorporation of C\(^{14}\) into the total
fatty acids of the major lipide complexes, the
specific activity of the fatty acids of the triglycerides
was two to four times that of the phospholipides
and more than twenty times that of the fatty acids
of sterol esters.

Incorporation of acetate-1-C\(^{14}\) into certain of the
chromatographically separated fatty acids of the
triglycerides, phospholipides and sterol esters (Table V)

The incorporation of radioactivity into certain of
the individual fatty acids of the various lipide com-
plexes followed more or less the same general pat-
tern noted in the formation of the total fatty acids
of each group. Thus, the specific activity of the
palmitic, stearic, and oleic acids found in ester
linkage with the triglycerides was maximum two
to four hours after the administration of the ace-
tate. Despite the difference in concentrations of
these fatty acids in the plasma, palmitic acid
showed the greatest specific activity at its peak,
with less activity in stearic, and still less in oleic
acid.

The presence of radioactivity in the fraction
eluted by 60 per cent aqueous acetone (4) is prob-
dly due to contamination of the linoleic acid by the

<table>
<thead>
<tr>
<th>Time in hours</th>
<th>Palmitic acid</th>
<th>Stearic acid</th>
<th>Oleic acid</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sterol Esters</td>
<td>96</td>
<td>30</td>
<td>NSC‡</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>2,910</td>
<td>1,929</td>
</tr>
<tr>
<td></td>
<td>24</td>
<td>263</td>
<td>252</td>
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<tr>
<td></td>
<td>48</td>
<td>402</td>
<td>243</td>
</tr>
<tr>
<td></td>
<td>96</td>
<td>171</td>
<td>96</td>
</tr>
</tbody>
</table>

| Phospholipides | 183         | 72          | 63         |
|                | 432         | 210         | 159        |
|                | 942         | 465         | 297        |
|                | 1,563       | 723         | 534        |
|                | 1,578       | 672         | 555        |
|                | 1,890       | 1,110       | 825        |
|                | 1,287       | 804         | 414        |
|                | 96          | 351         | 120        |

* Counts per minute per 0.005 millimole.
† Radioactivity in this fraction is presumably due to presence of small quantity of myristic acid (see Results).
‡ No significant counts above background.
very small quantity of myristic acid thought to be present in the plasma of man (13), which is also eluted from the chromatographic column by this solvent system. If this be the case, the myristic acid fraction would undoubtedly be diluted by the presence of presumably unlabelled linoleic acid (19–22). The specific activity of the myristic acid would therefore be greater than that stated, since the total acidity of the combined fraction was used in the calculation. This fraction was not subjected to the Bertram oxidation procedure (4) for the separation of the saturated from unsaturated acids because of difficulties encountered in handling very small quantities of saturated acids by this method, as would be the case with myristic acid in this particular instance.

The curves describing the appearance of C¹⁴ in the palmitic, stearic, and oleic acids attached to the phospholipides closely resemble those of the total fatty acids of this fraction. The maximum specific activity was again attained approximately 24 hours after the introduction of the precursor. The relative distribution of counts within the individual fatty acids was somewhat similar to that noted in the triglyceride fractions. No significant radioactivity was found, however, in that fraction containing linoleic acid and possibly traces of myristic acid.

The relatively small amount of radioactivity noted in the fatty acids of the sterol esters was predominantly due to the presence of labelled palmitic and oleic acids. Although some labelling occurred, the quantities of stearic acid isolated from this complex were too small for accurate analyses for specific activity. Again there was no discernible radioactivity in the relatively large quantity of linoleic acid that was isolated.

**DISCUSSION**

It is known that the fatty acids entering into ester linkage with each of the specific major lipide complexes are mixtures which may vary in quantity and composition. In contrast to the triglycerides, the fatty acids of the cholesterol esters under normal circumstances contain large quantities of linoleic acid. It has been shown that this polyunsaturated acid is not readily synthesized by the mammalian organism (19–22) and is presumably derived in toto from the diet. The specific activity of the saturated and monoethenoid fatty acids of the “non-phospholipide” fraction in this and other studies (16, 18, 23, 24) would tend to be greater if the contribution derived from the large unlabelled linoleic acid fraction of the sterol esters and from the somewhat smaller fraction from the triglycerides was excluded from the calculations. Similarly, other polyunsaturated fatty acids such as linolenic and arachidonic acids thought to reside in the phospholipide and sterol ester complexes (25) would also tend to alter the estimation of the specific activity of these fractions.

The degree to which this occurs, however, would depend upon certain characteristics of some of the polyethenoid acids. Although relatively little is known about the intermediary metabolism of the higher unsaturated fatty acids, it has been shown in animals that some of the processes leading to the formation of one essential fatty acid from another may involve endogenous synthesis. Thus, Widmer and Holman (21) noted that fat-starved rats apparently synthesized arachidonic acid from linoleic, but not from linolenic acid. Further investigation of the problem by Mead, Steinberg, Howton, and Slaton (20, 22) demonstrated the formation of arachidonic acid from an exogenous precursor presumably derived from linoleic acid by the addition of a two-carbon fragment obtained from administered radioactive acetate. It is, therefore, apparent that the variation between the actual and theoretical specific activities of the total fatty acids, particularly of the phospholipide and sterol ester fractions, would also be dependent upon the extent to which labelled acetyl coenzyme A is utilized to form tetaenes and conceivably other highly unsaturated acids that may exist in small quantities in the plasma of man. Because of difficulties inherent in the isolation of these acids in the pure state, the specific activities listed in this study are only relative.

The curves (Figure 7) depicting the synthesis of the labelled fatty acids of the triglycerides (A) and the phospholipides (B) resemble that of a time course curve demonstrating the relationship between a precursor (A) and product (B) when a labelled precursor (acetate) is furnished at zero time (24). Since it has been implied on the basis of experiments in animals (23, 26) as well as man (18) that the triglyceride fraction is apparently the major vehicle for the transport of fatty acids in

THE SYNTHESIS OF SPECIFIC FATTY ACIDS FROM ACETATE 241
the plasma, it seems likely that at least certain of the individual fatty acids of the phospholipid such as palmitic, stearic, and oleic acids are derived in part from or are exchanging with the fatty acids of the triglycerides. The same reasoning may apply to a limited extent to the fatty acids of the sterol esters. However, the lack of adequate studies concerning the relative rates of synthesis of this fraction in the mammalian liver and its presence in other tissues precludes further speculation on this point.

Some information concerning the interconversion of fatty acids may be derived from an analysis of the relative specific activities of the individual acids. It may be noted that despite the different concentrations of these substances in the plasma, the turnover rates of palmitic, stearic, and oleic acids within a major lipide complex are approximately comparable. Furthermore, the greatest degree of radioactivity was found in the palmitic acid fraction of each group, followed then by stearic and oleic acids. It would appear, from the work of Dauben, Hoerger, and Peterson (27), that palmitic acid for the most part is synthesized directly from two-carbon units and the amount derived from the process of elongation of an intermediate fatty acid such as myristic by the addition of a two-carbon fragment is exceedingly small. On the other hand, Stetten and Schoenheimer (28) and Zabin (29) have indicated that a significant quantity of stearic acid is formed by direct elongation of the carbon chain of palmitic acid by two-carbon atoms. Furthermore, the formation of the monounsaturated oleic acid probably occurs in a manner similar to that of the saturated acids. Indeed, Anker (30) studied the relative distribution of the isotope in the various higher fatty acids after feeding myristic acid-1-C^{14} to rats and concluded that the 14 carbon atoms of myristic acid were utilized for carbon atoms 5 to 18 of oleic acid by way of palmitic and stearic acids. A similar investigation employing acetate-1-C^{14} by Dauben, Hoerger, and Peterson (27) gave support to this postulate by showing that the pattern of distribution of the isotope in the degraded unsaturated C_{18} acids was identical with that of the saturated acids. Additional information on the metabolic interrelationships of these acids was provided by Weinman, Chaikoff, Dauben, Gee, and Entenman (31) who found that when palmitic acid was catabolized in vivo, it was primarily converted to small carbon units without any appreciable formation of acids of intermediary carbon length. In contrast, the catabolism of stearic acid (32), while similarly breaking down to short chain units, also gave rise to an appreciable quantity of palmitic acid. The relative concentration of C_{14} in the palmitic, stearic, and oleic acid fractions isolated in this study would tend to support the occurrence of these overall reactions in man. Thus, it can be assumed that palmitic acid is the major higher saturated fatty acid intermediate formed from acetate. At least three pathways seem to be involved in the further metabolism of this acid. First, some palmitic acid is undoubtedly deposited in the fat depots as such. Second, a quantity is utilized as fuel by breaking down to two-carbon fragments which can enter the tricarboxylic acid cycle and provide a source of energy. Lastly, a portion of the palmitic acid pool is converted to stearic acid. Only small quantities of this C_{18} acid are found in the plasma of man under normal circumstances. Presumably much of this acid is either oxidized for energy, desaturated to form oleic acid or reconverted to palmitic acid.

The absence of appreciable radioactivity in the linoleic acid fraction of the major lipide complexes of the plasma lends support to the contention that this diene cannot be detectably synthesized by the mammalian liver (19, 20, 22, 28). However, it has been shown recently by Mead, Slaton, and Decker (33) that at least the carboxyl carbon of linoleic acid can be utilized to a limited extent in the formation of some of the higher saturated fatty acids in the rat. The metabolic pathway involved in this reaction is not clear at this time. It may be mediated via a higher unsaturated acid such as arachidonic which can be formed from a linoleate derivative (20, 22) by the addition of a two-carbon fragment or by the direct conversion of linoleic acid to short chain or more intermediate units which then can be partially utilized in the synthesis of the saturated acids. This latter concept seems to be the more likely possibility.

**Summary**

Certain specific saturated and unsaturated fatty acids associated with the major lipide complexes
of the plasma of man were isolated and measured by chromatographic methods. The predominant fatty acid components of the sterol esters, the phospholipides, and the triglycerides were linoleic, oleic, palmitic, and stearic acids. The largest quantity of unsaturated acids was noted in the sterol ester fraction. The majority of the saturated fatty acids were present in the triglycerides. The rates of synthesis of the total and certain of these individual higher fatty acids of man following the administration of acetate-1-C14 were also studied. Some of the fatty acids of the phospholipides appeared to be derived in part from those of the triglycerides. The triglycerides probably represent the major vehicle for the transport of fatty acids in man as well as animals. The highest concentrations of radioactivity appeared in the palmitic acid fraction of each of the major lipid complexes, followed by those in the stearic and oleic acids. The absence of demonstrable activity in the isolated linoleic fractions signified the lack of endogenous formation of this polyunsaturated fatty acid in man.

REFERENCES


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