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Heterogeneity of Plasma High Density Lipoproteins *

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The plasma high density lipoproteins, which are isolated between the densities 1.063 and 1.21 in the ultracentrifuge and which correspond to alpha lipoproteins on electrophoresis, have been the subject of far fewer studies than their counterparts of lower density. In recent years, however, they have served as useful models for examination of lipoprotein structure and have become of particular interest to this laboratory since the discovery of Tangier disease, in which they are congenitally absent or seriously deficient (1).

When these lipoproteins are distributed along a density gradient, they seem to form a continuum without stepwise compositional changes (2). Subfractions isolated over different portions of the broad density band by which they are defined have been found to contain similar lipid composition and protein of the same amino acid composition (3). It is also usually accepted that the high density lipoproteins (HDL) are antigenically homogeneous (4, 5) and form a single electrophoretic band.

These data, however, have never been completely reconciled with other evidence that the high density lipoproteins may be discontinuously distributed over their common density band. For example, the separation of HDL into two major peaks in the analytical ultracentrifuge, a technique developed by deLalla and Gofman (6), is a reproducible phenomenon; it has been repeatedly demonstrated in different clinical situations that these fractions may independently vary in concentration (7). Preparative ultracentrifuge fractions corresponding to these two HDL peaks obtained in the analytical ultracentrifuge have also been found to contain protein moieties of quite different molecular weights (8). Subsequently, Shore (9) has presented evidence suggesting that HDL sub-

fractions may contain different multiples of a basic peptide, and this has since been partially confirmed by others (10, 11).

Finally, although only one major immunochemical study of high density lipoproteins (12) has concluded that they may be antigenically heterogeneous, there are other instances in which more than one form of this lipoprotein would appear to be present (13-16). Thus, there is both direct and indirect evidence of inhomogeneity among high density lipoproteins. The latter is of uncertain character and has required further definition. This is of particular importance in preparation of pure lipoprotein fractions required in current experimental work.

An intensive study of high density lipoproteins, with particular reference to their immunochemical identity, has therefore been undertaken. By using specific antisera to HDL, the lipoproteins present in plasma and ultracentrifugal fractions have been carefully characterized. It has been demonstrated that two forms of alpha lipoprotein exist (17). Their nature and relationship to certain transformations of the lipoproteins from their native state and the manner in which they help to reconcile discrepancies among older data are subjects of this report.

Methods

Unless otherwise indicated, blood was collected in 0.1% EDTA from normal males and females between ages 20 to 45 fasted overnight. Either individual plasma samples or that pooled from ten subjects was used. Analyses and fractionation were begun on the day of collection, and the remaining plasma was stored at 4° C. In a few studies the low density lipoproteins (density <1.063) of fresh plasma were completely precipitated with 0.05 ml per ml 0.1 M manganese chloride and 2 mg per ml heparin in the cold (18), and the supernate was used as a source of alpha lipoprotein. Preparative ultracentrifugation. Lipoprotein fractions were separated in the preparative ultracentrifuge by the method of Havel, Eder, and Bragdon (19). The density of plasma of lipoprotein isolates was raised by
addition of NaCl and KBr (19) and checked by pyk-
nometry at 20° C. Samples were centrifuged at 10° C
in the Spinco model L preparative ultracentrifuge at
40,000 rpm (ca. 105,000 g). With the no. 40.3 rotor,
centrifugation was carried out for 16 hours at density
1.063 and 22 hours at densities 1.1 and 1.21. With the
no. 40 rotor, the centrifuge times at comparable densities
were 24 and 48 hours, respectively. As desired, one
or more of the four fractions were usually isolated. These
are subsequently designated as LDL (density < 1.063),
HDL (density 1.063 to 1.21), HDL₂ (density 1.063 to
1.1), HDL₃ (density 1.1 to 1.21), and the fraction of
density > 1.21.

For use as antigen some LDL was prepared as the
fraction of density 1.019 to 1.063. The infranate of this
material was not further used to prepare HDL.

The lusteroid tubes were sliced through the inter-
mediate colorless zone with a standard tube slicer, and
the top and bottom fractions were centrifuged once
again at each separation density. This was done be-
cause of frequent immunochemical evidence of trace con-
tamination of HDL by LDL and of HDL fractions by
albumin and other nonlipoprotein proteins. After re-
centrifugation none of these contaminants was present in
amounts detectable by immunoelectrophoresis or double
diffusion studies except for trace amounts of LDL that
could not be removed from the HDL fractions by as
many as three recentrifugations. However, precipitation
of LDL (18) before centrifugation eliminated all LDL
contamination in subsequent HDL fractionation begin-
ning with density 1.1. All fractions were dialyzed at
4° C against 40 vol or more of 0.15 M NaCl containing
0.01 M EDTA at pH 6.5 to 7.3. The dialysis fluid
was changed three times during the 24-hour period.

Analytical ultracentrifugation. Lipoprotein fractions
of densities 1.063 to 1.1 and 1.063 to 1.21 were cen-
trifuged in the Spinco model E analytical ultracentrifuge
within 24 hours of their preparation without dialysis.
A double sector cell was employed, one side containing
the 10 to 30 mg per ml of lipoprotein, the other a solu-
tion of NaCl-KBr of the same solvent density, provid-
ing the reference boundary. The rotor speed was 52,640
or 56,100 rpm and the temperature 20° C. The Schlieren
patterns were photographed at intervals of 4 to 16 min-
utes for 2 to 3 hours at a bar angle of 65 to 70.

In the analytical ultracentrifuge at density 1.21, HDL
(density 1.063 to 1.21) appeared as an asymmetric single
peak at 16 to 32 minutes. By 48 minutes it had re-
solved into at least two components (Figure 1). HDL₃
(density 1.063 to 1.1) centrifuged at either solvent
density of 1.1 or 1.21 appeared as a relatively homogene-
ous peak with a small tail of more slowly floating ma-
terial apparent with time. In the analytical ultracen-
trifuge at density 1.1 HDL (density 1.063 to 1.21)
separated by 32 minutes into two peaks. One was of
an area equal to that previously determined for the
HDL₃ component; the other was a somewhat broader
and sedimenting peak (HDL₄). The Schlieren patterns
were the same as those reported earlier by deLalla and
Gofman (6), who introduced the nomenclature HDL₄
and HDL₅ to describe these major components of HDL
distinguishable in the analytical ultracentrifuge. No
small peak, comparable to their fraction HDL₅ (6, 7)
was seen, probably because this lower density material

![FIG. 1. ANALYTICAL ULTRACENTRIFUGAL ANALYSIS OF HIGH DENSITY LIPOPROTEIN (HDL)
(D 1.063 to 1.21). Medium density 1.21, 52,650 rpm, concentration 10 mg of lipoprotein protein
per ml, 26° C. An asymmetric single peak can be seen at 30 minutes (left frame); a hy-
paphic peak is apparent at 60 minutes (right frame).](image-url)
TABLE I

Lipid and alpha lipoprotein composition of plasma samples studied*

<table>
<thead>
<tr>
<th></th>
<th>Plasma</th>
<th></th>
<th>HDL</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cholesterol</td>
<td>Phospholipid</td>
<td>mg per 100 ml</td>
<td>Cholesterol</td>
</tr>
<tr>
<td>Pool A</td>
<td>176</td>
<td>249</td>
<td>48</td>
<td>104</td>
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<td>Subject 1</td>
<td>195</td>
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<td>Subject 2</td>
<td>121</td>
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<td>Subject 3</td>
<td>163</td>
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<td>96</td>
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<td>Subject 4</td>
<td>198</td>
<td>239</td>
<td>47</td>
<td>100</td>
</tr>
</tbody>
</table>

* All high density lipoprotein (HDL) fractions reported above were obtained with the number 40 rotor.

had been removed by repeated preparative centrifugation at density 1.063.

**Lipid analysis.** Two-ml samples of plasma lipoproteins were extracted in 50 ml of chloroform:methanol (2:1), and the cholesterol (20) and phospholipid (21) contents determined. Protein was estimated by the method of Lowry, Rosebrough, Farr, and Randall (22). Representative examples of the total plasma lipid and HDL (density 1.063 to 1.21) lipid concentrations in the samples studied are given in Table I. HDL₄ contained from 25 to 32% of the total HDL cholesterol and phospholipid, and 20 to 25% of the total HDL protein with the resulting average lipid to protein ratio of about 1.1 for HDL₄ and 0.85 for HDL₅. The fraction of density > 1.21 contained less than 2 mg per 100 ml cholesterol and 10 to 20 mg per 100 ml of phospholipid, or 6 to 10% of the total plasma phospholipid.

**Partial delipidation.** Preparations of HDL₄ and HDL₅ were partially delipidated by injecting them through a no. 25 needle into 50 vol of pure ethanol:ether (3:1) at 4°C for 24 hours. The resulting protein precipitate, containing 10 to 20% of the cholesterol and up to 50% of the phospholipid in the starting material, was then washed twice with 25 vol of cold redistilled diethyl ether. The washed precipitate readily redissolved in 0.15 M NaCl.

**Immunological methods. Preparation of antisera.** Antisera to LDL or HDL were prepared in both sheep and rabbits. Lipoprotein fractions used as antigens were mixed in an equal volume of complete Freund's adjuvant and injected intramuscularly. Some fractions were first concentrated 5- to 25-fold with Carbowax (15). The final protein concentration of the antigens was 20 to 50 mg per ml.

The total dose, administered in three divided doses at intervals of 3 weeks, was approximately 80 mg of antigen protein to Cheviot sheep (wt. 40 to 50 kg) and approximately 30 mg to rabbits (3 to 5 kg).

Antigen was stored at 4°C and prepared in adjuvant just before injection. Blood was collected 1 week after injection.

**TABLE II**

Characterization of antisera used for immunochemical study

<table>
<thead>
<tr>
<th>Animal</th>
<th>Sensitizing antigen</th>
<th>Reactivity</th>
<th>LDL</th>
<th>HDL</th>
<th>Albumin</th>
<th>Other proteins</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sheep</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
</tr>
<tr>
<td>S₁</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td></td>
</tr>
<tr>
<td>S₂</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>S₃</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rabbits</td>
<td></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₁</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₂</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₃</td>
<td>HDL (D 1.063-1.21)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
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</tr>
<tr>
<td>R₄</td>
<td>HDL (D 1.063-1.21)</td>
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<td>+</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₅</td>
<td>HDL (D 1.063-1.21)</td>
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<td>+</td>
<td>+</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₆</td>
<td>LDL (D 1.019-1.063)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
<td></td>
</tr>
<tr>
<td>R₇</td>
<td>LDL (D 1.019-1.063)</td>
<td>+</td>
<td>+</td>
<td>-</td>
<td></td>
<td></td>
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<tr>
<td>Horses</td>
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<tr>
<td>H₁</td>
<td>Whole serum</td>
<td>+</td>
<td>+</td>
<td>+</td>
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<td></td>
</tr>
<tr>
<td>H₂</td>
<td>Whole serum</td>
<td>+</td>
<td>+</td>
<td>±</td>
<td></td>
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</tr>
<tr>
<td>H₃</td>
<td>Whole serum</td>
<td>+</td>
<td>+</td>
<td>±</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Absorbed with excess albumin (A) or LDL (low density lipoproteins) (B).
† H₁ = Pasteur Institute (223-224) antiwhole serum; H₂ = Walter Reed Army Hospital (Liddy) antiwhole serum; H₃ = Hyland Laboratories antiwhole serum.
HETEROGENEITY OF PLASMA HIGH DENSITY LIPOPROTEINS

FIG. 2. IMMUNOELECTROPHORESIS OF FRESH PLASMA BEFORE AND AFTER PRECIPITATION OF THE LOW DENSITY LIPOPROTEINS (LDL). In the left hand slide the trough contains antiserum $S_a$ (Table II); on each of the middle and right hand slides, one trough was filled with antiserum $R_s$ and the other with antiserum $R_t$ (anti-LDL). All precipitation lines on this and succeeding figures are unstained. All the lines shown here were subsequently shown to stain for lipids.

the final dose and the serum frozen after the addition of 0.01% merthiolate. Each antigen was carefully characterized by immunoelectrophoresis before use. The lipoprotein fractions used as antigen for $S_a$, $S_b$, and $R_s$ through $R_t$ all were derived from pool A (Table I). With each antiserum the precipitating antibodies were shown by immunoelectrophoresis to be almost entirely G immunoglobulins (7 S globulin) (23). Certain of the antisera were partially purified by sodium sulfate precipitation by the method of Kekwick (24).

Horse antiwhole human serum 1 and commercial horse antiwhole serum, rabbit anti-beta lipoprotein, and anti-albumin 2 were used.

Characterization of antisera. HDL (density 1.063 to 1.21) used as antigen produced adequate anti-HDL titers in both of two sheep and four of six rabbits. All of the eight antisera are characterized in Table II. Seven cross-reacted with LDL. Several also formed precipitation lines with albumin, and two reacted with other unidentified proteins. Four rabbit and one sheep antisera could be absorbed with albumin or LDL leaving sufficient anti-HDL titer for use in the experiments to be described. LDL produced antibodies of relatively higher titer that did not appear to cross-react with HDL or other proteins. Pure HDL antisera were thus very difficult to produce, presumably because the antigen could not be freed of undetected LDL, which is so much stronger in provoking antibody response. Antisera, such as $R_s$ (Table II), however, could be freed of all anti-LDL activity by absorption without appreciable decrease in anti-HDL titer.

Hyland antialbumin and anti-LDL were relatively pure. They did not react with HDL fractions and gave sharp precipitation bands with albumin and LDL, respectively. All the antihuman sera contained antibodies to both HDL and LDL. The Pasteur antiserum ($H_p$, Table II) gave the sharpest precipitation line with alpha lipoprotein at serum concentration and was used for most experiments with antihuman serum. The other sera were used only to check ultracentrifugal fractions for nonlipoprotein impurities.

Absorption of antibodies was carried out at equivalence as determined by serial dilution using the double diffusion tube method of Preer (25). The precipitates were allowed to stand for 5 to 10 days before absorbed antisera were harvested.

1 Obtained from Walter Reed Army Hospital (Leddy) and the Pasteur Institute (223-224).

2 Purchased from Hyland Laboratories, Los Angeles, Calif.
Agar diffusion. Ouchterlony plates or Preer tubes of 1 mm i.d. were made with 1% Difco special noble agar in saline. Plates with wells requiring 5 to 10 μl or 40 to 50 μl of antigen or antibody were used. They were allowed to develop at 4°C and observed for at least 2 weeks. The Preer tubes were sealed after preparation with molten paraffin and allowed to develop at room temperature. The appearance of precipitin bands and their location in relationship to the antigen-agar boundary were plotted daily for 10 days.

Immunoelectrophoresis. The technique of Grabar and Williams (26) as modified for microscope slides by Scheidegger (27) was used for immunoelectrophoresis. Standard microscope slides were coated with a thin film of 2% Difco special noble agar in 0.05 Veronal buffer, pH 8.2, and allowed to dry. Two ml of the agar in buffer was then applied in a coat of uniform thickness. To assure uniform water content, plates were always used within 2 hours of preparation. The wells were filled with 2 to 3 μl of antigen, and electrophoresis was carried out in 0.05 M Veronal buffer at pH 8.2 using constant current (58 ma) for exactly 40 minutes. Proteins were either precipitated immediately after electrophoresis with 2% acetic acid or allowed to diffuse against 50 to 60 μl of antiserum for 24 to 72 hours at room temperature in a high humidity chamber. Slides were then washed overnight with saline, followed by distilled water, dried, and exposed to a freshly prepared saturated solution of oil red O in 60% ethanol at 40°C for at least 12 hours and then counterstained for protein with SF light green for 2 hours. Photographs of the immunoprecipitation lines were made before washing, after oil red O staining, and again after counterstaining.

Protein analyses. Ultracentrifugal fractions of LDL, HDL, HDLα, and HDLβ were extracted with ethanol:ether (3:1) for 18 to 24 hours at 24°C (3). The precipitate was washed twice with ethanol and once with ether. Portions of the precipitate were sealed in an oxygen-free atmosphere and hydrolyzed at 110°C for 24 hours in 6 N HCl. Some of the delipidated proteins were oxidized with performic acid for 1 hour and were then lyophilized before acid hydrolysis (28). Hydrolysates representing 0.5 to 5 mg of protein were analyzed for amino acid content with the Spinco Moore-Stein analyzer. Quantification was standardized using known references (29). Two of the plasma fractions were analyzed four times each, and the amounts of individual amino acids obtained on each run were in agreement within 3%. Amino terminal acids were determined by the dinitrophenol method of Levy (30).

Results

Heterogeneity of alpha lipoprotein. Immunoelectrophoretic studies. Fresh, whole plasma or plasma from which all low density lipoproteins had been removed by precipitation with polyhans usually yielded on immunoelectrophoresis a single
precipitation arc attributable to alpha lipoprotein with any of the eight anti-HDL antisera described in Table II (Figure 2). However, under appropriate conditions and especially in certain ultracentrifugal fractions, heterogeneity of the alpha lipoproteins became apparent with all of the antisera. As shown in Figures 3 and 4 a single broad arc with anti-HDL serum was present in fresh whole plasma and in the ultracentrifugal fraction containing HDL2 (density 1.063 to 1.1). There was in addition a second band of slower migration in ultracentrifugal fractions of HDL (density 1.063 to 1.21) and HDL3 (density 1.1 to 1.21). Only the second band was present in the fraction density > 1.21.

Both of these precipitation bands were lipid staining, although the second band was occasionally only faintly so. The positions of these bands were relatively constant, the first migrating with and just behind the albumin, the second in the
alpha₁-alpha₂ region. With the antisera, only the first band could be detected in HDL₂, but both bands were always detected in HDL₃. Only the second band was ever detected in the fraction density > 1.21. As shown in Figures 3 and 4 and subsequently in Figures 5 and 6, these arcs blended completely, crossed, or appeared as several interrupted lines depending upon the antiserum used. The precipitation lines obtained with HDL (density 1.1 to 1.21) in Figures 3 and 4 represented the extreme in complexity obtained, and these were observed only with two antisera and with concentrations of HDL₂ considerably in excess of that in plasma. The small precipitation lines close to the junction of the major bands were interpreted as also representing reactions between antisera and the HDL antigens and not unrelated contaminants. They were not seen when only one or the other major band was present alone as in concentrated preparations of HDL₂ or the fraction of density > 1.21. Furthermore, all the lines stained for lipid, and they were removed by absorption along with the major bands (see below). The multiplicity of lines is considered to reflect the fact that the HDL was composed of very similar but not identical antigens. All of the antigenic preparations used for antibody production contained HDL over the density spectrum from 1.063 to 1.21. It appears that a family of antibodies was produced, its members differing in reactivity to certain antigenic sites, only some of which were common to HDL throughout its density spectrum. It was concluded that there were only two major antigens in HDL, these being detected with all the antisera.

Recombination and absorption studies. Recombination of the lipoprotein fractions with each other or with plasma (Figure 5) confirmed that

![Figure 5](image-url)
the alpha lipoprotein in whole plasma and that in HDL\textsubscript{2} was identical, and different from the second more slowly migrating band. By similar combination experiments (Figure 6), it could easily be demonstrated that neither of these bands was related to either albumin or lipoprotein of density 1.019 to 1.063 (LDL).

With Preer tubes (25), the titer of antiserum S\textsubscript{1AB} to each of the alpha lipoprotein bands were determined separately with a sample of HDL\textsubscript{2}, containing only the faster migrating band, and a sample of fraction > 1.21 that contained only the slower migrating band. This antiserum easily discriminated both bands in HDL\textsubscript{2}. An amount of HDL\textsubscript{2} slightly in excess of equivalence was then added to S\textsubscript{1A}. This absorbed antibody was then tested against whole plasma and several HDL fractions by immunoprecipitation, double-diffusion, and immunoelectrophoresis. No reactions were now obtained with either plasma or HDL\textsubscript{2}. Precipitation was obtained with HDL\textsubscript{3} and the density > 1.21 fraction, which proved only to represent the slower migrating band on immunoelectrophoresis.

Similar absorption of antiserum S\textsubscript{1AB} with the density > 1.21 fraction in amounts slightly in excess of equivalence left reactivity to plasma, HDL\textsubscript{2}, and HDL\textsubscript{3}. This was shown to represent only the fast migrating band on immunoelectrophoresis. No reaction was obtained with the density > 1.21 fraction. Quantitatively about two-thirds of the reactivity of antiserum S\textsubscript{1AB} could be removed with either HDL\textsubscript{2} or the > 1.21 fraction. Results of similar absorption
experiments using other antisera varied depending on their ability to discriminate between the two forms of alpha lipoprotein.

**Nomenclature.** On the basis of the above experiments, the two forms of alpha lipoprotein distinguishable by immunoelectrophoresis have been designated as alpha LP\(_A\), the band of greater migration seen in plasma, HDL\(_1\), HDL\(_2\), or HDL\(_3\); and alpha LP\(_B\), the second band present only in HDL\(_3\) and the density \(> 1.21\) fraction (Figure 7). This nomenclature incorporates the designation “LP” to differentiate the lipoprotein from the delipidated protein, alpha P.\(^3\)

**Diffusion and precipitation studies.** Small amounts of alpha LP\(_B\) could be detected in fresh plasma by immunoelectrophoresis if the plasma was concentrated 5- to 10-fold by polyethylene glycol (15) or triple-loading of the antigen wells.

\(^{3}\) In a preliminary communication (17), alpha LP\(_B\) was referred to as alphaD, a simpler term, but offering less flexibility for possible new forms of alpha lipoprotein that conceivably may be demonstrated in the future.

![Figure 7. Immunoelectrophoretic patterns obtained with antiserum R\(_1\) and lipoprotein fractions HDL\(_d\) (D 1.063 to 1.1) and density greater than 1.21 undergoing electrophoresis separately (left hand slide) and in combination (right hand slide). HDL\(_d\) contains only the form designated alpha LP\(_A\), and the density greater than 1.21 fraction contains only the form designated alpha LP\(_B\). All the lines shown here stain for lipid.](image-url)
FIG. 8. IMMUNOELECTROPHORETIC PATTERN OF HDL$_2$ (ALPHA LP$_A$) AND ITS PARTIALLY DELIPIPETED PRODUCT. Antiserum $S_{14}$ was used. Both lines shown here stain for lipid.
Usually very small amounts of this second alpha band could be detected in unconcentrated plasma by Ouchterlony plates or especially by the method of Preer. Even with these more sensitive techniques, only one line was seen with HDL₂ using any of the antisera, emphasizing the immunologic homogeneity of this lipoprotein fraction.

With either the Preer or Ouchterlony techniques, repetition of the absorption studies with antiserum S₁A, described above, using immunoelectrophoresis, again demonstrated that alpha LPₐ and alpha LPₜ differed in some antigenic sites for this antiserum.

As could be predicted from immunoelectrophoresis, the two forms of alpha lipoprotein could also easily be distinguished by differing mobility on simple electrophoresis on agar.

Conversion of alpha LPₐ to alpha LPₜ. The absence of significant amounts of alpha LPₜ from fresh serum and its obvious appearance in certain of the ultracentrifugal fractions implied that the latter process was capable of producing alpha LPₜ, quite probably through the conversion of some of alpha LPₐ.

The appearance of alpha LPₜ during ultracentrifugation proved to be dependent upon the exposure of lipoprotein to a centrifugal field and not to the other manipulations attending the routine preparative procedure. Serum brought to a density of 1.063 by addition of NaCl-KBr, allowed to stand at 15°C for 1 week, and then dialyzed against saline containing 0.01 M EDTA contained no alpha LPₜ detectable by immunoelectrophoresis. In contrast, alpha LPₜ became readily apparent at serum concentration in the infranate after a single 16- to 24-hour centrifugation at density 1.063. Furthermore, repeated centrifugation at density 1.21 of either HDL or HDL₂ yielded additional alpha LPₜ in the infranate of each successive run indicating continuous production of this form of the lipoprotein during ultracentrifugation.

Storage, as well as ultracentrifugation, was found to lead to the production of alpha LPₜ. Although fresh serum or plasma obtained with EDTA, sodium oxalate, or heparin as anticoagulant contained little or no detectable alpha LPₜ, it became increasingly demonstrable during storage of the unconcentrated samples over a period of 2 to 10 days. Storage of plasma in EDTA minimized the generation of alpha LPₜ.

Alpha LPₜ was also produced by repeated freezing and thawing between 4°C and –20°C or by addition of urea to plasma in a final concentration of 8 M. As alpha LPₜ became apparent, the titer of alpha LPₐ decreased. As noted earlier, the supernate remaining after removal of LDL by precipitation (18) contained only alpha LPₐ (Figure 2), but the B form was then easily produced by either simple storage of the supernate without EDTA, or by freezing and thawing. This experiment provided further proof that alpha LPₜ was derived from alpha LPₐ and not from other lipoprotein precursors.

Final proof of the conversion of the A form to the B was provided in the following experiment, the results of which are shown in Figure 8. An HDL₄ fraction, containing no alpha LPₜ, was partially delipidated with ethanol:ether (3:1) and washed with ether. The resulting material was completely soluble in 0.15 M NaCl. It was subjected again to immunoelectrophoresis and consisted almost exclusively of alpha LPₜ. Partial delipidation of the fraction 1.1 to 1.21, in the same way, again eliminated most of the alpha LPₐ with a concomitant gross increase in the amount of alpha LPₜ. The slower migrating material produced by delipidation of alpha LPₐ was recombined with plasma, HDL₂, HDL₃, and the fraction density > 1.21. Both immunoelectrophoresis and double diffusion of these mixtures

<table>
<thead>
<tr>
<th>TABLE III</th>
<th>Amino acid compositions of the proteins of serum LDL, HDL₂, and HDL₃ subfractions*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LDL</td>
</tr>
<tr>
<td>Aspartic</td>
<td>100</td>
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<tr>
<td>Threonine</td>
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<td>Serine</td>
<td>74</td>
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<td>Glutamic</td>
<td>118</td>
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<td>Proline</td>
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<td>Glycine</td>
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<tr>
<td>Cystine</td>
<td>7</td>
</tr>
<tr>
<td>Valine</td>
<td>52</td>
</tr>
<tr>
<td>Methionine</td>
<td>13</td>
</tr>
<tr>
<td>Isoleucine</td>
<td>53</td>
</tr>
<tr>
<td>Leucine</td>
<td>108</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>33</td>
</tr>
<tr>
<td>Phenylalanine</td>
<td>49</td>
</tr>
</tbody>
</table>

* All values are expressed as moles relative to aspartic acid taken as 100. LDL = D 1.019 to 1.063.
† Cystine determined as cysteic acid after pretreatment of hydrolysate with performic acid.
on Ouchterlony plates indicated that the delipidated product was immunochemically identical to alpha LP\textsubscript{B}.

**Protein analyses.** In Table III are presented the relative amino acid compositions of the lipoprotein fractions LDL\textsubscript{2}, HDL\textsubscript{3}, HDL\textsubscript{4}, and HDL\textsubscript{5}. The values given for each fraction represent the means obtained on 3 to 5 separate preparations of each fraction. These, as well as separate determinations on the same fractions, all agreed within 3\%. The amino acid compositions of these HDL fractions were indistinguishable and different from those previously published for LDL and albumin (3, 31). The basic amino acids, arginine, histidine, and lysine are not shown in Table III. They were determined once on a single sample of each fraction. The contents of these amino acids were indistinguishable in HDL and its fractions. No attempt was made to correct for possible losses of amino acids during hydrolysis, nor was there an attempt to quantify tryptophan or amide ammonia. As previously noted by others (3, 9, 11, 32), the HDL fractions all contained amino terminal aspartic acid with trace amounts of serine and threonine. In view of the similarity of the amino acid analyses of the HDL fractions, the peptides contained in alpha LP\textsubscript{A} and LP\textsubscript{B} are very probably identical. The analyses of HDL amino acids (Table III) are comparable to those previously published except for discrepancies in cystine content (3, 9-11, 33). For the latter our results are in agreement with the data of Shore and Shore (33), who used a titration method for determining sulphydryl groups, but in disagreement with the data of Scanu and Hughes (3) and Sanbar and Alau-povic (10), which showed little or no cystine. This is probably explained by our pretreatment of protein hydrolysates with performic acid converting the cystine and cysteine present to the more stable cysteic acid before acid hydrolysis (28).

**Discussion**

Two forms of alpha lipoprotein, designated A and B, have been demonstrated. The native state of this lipoprotein in plasma appears to be almost entirely alpha LP\textsubscript{A}. As plasma ages, undergoes ultracentrifugation, or is otherwise manipulated, alpha LP\textsubscript{B} appears. It is distinguished from alpha LP\textsubscript{A} by differences in electrophoretic migration, immunochemical behavior, and flotation in the ultracentrifuge. Proof has been presented that alpha LP\textsubscript{B} can be derived from alpha LP\textsubscript{A} by partial delipidation, but there is other evidence that this transformation may involve more than simple loss of lipid.

The importance of these findings lies particularly in three areas: First is the practical problem of having to deal with several forms of alpha lipoprotein in any immunochemical studies of plasma lipoproteins, particularly those aimed at defining the purity of lipoprotein isolates. The second is the implication that use of the ultracentrifuge to isolate and quantify high density lipoproteins is associated with molecular rearrangements that give rise to heterogeneity that probably does not exist in normal plasma. Finally, the observed transformation of alpha lipoprotein correlates with other data relevant to the peptide portions of high density lipoprotein in such a way as to offer further insight into the structure of lipoproteins.

The eight different antisera employed in this study varied in their specificity for the A and B forms of alpha lipoprotein. All, however, when used with immunoelectrophoresis, distinguished these two forms. Most of the antisera showed only partial cross-reactivity with each by either immunoelectrophoresis, double diffusion, or differential absorption. In assessing the validity of these findings it has been necessary to reconsider why, if the differences between alpha LP\textsubscript{A} and alpha LP\textsubscript{B} now seem apparent, has heterogeneity of alpha or high density lipoproteins not been more widely recognized in many previous immunochemical studies of these lipoproteins. In reviewing these in light of the present experiments, it is impressive how much evidence of heterogeneity has actually been present, and how closely it seems to corroborate the present demonstration of more than one form of alpha lipoprotein.

Aladjem, Lieberman, and Gofman (12) made the first immunochemical studies of HDL and concluded that these lipoproteins were not (antigenically) homogeneous. Interestingly, they found evidence of antigenic differences between the centrifugal subfractions HDL\textsubscript{2} and HDL\textsubscript{3}. The only other study employing a specific anti-
HDL serum was that of DeLalla, Levine, and Brown (5). Although their experiments are often cited as proof of the homogeneity of HDL, their data are actually more consistent with their single antiserum having been primarily reactive with what is now called alpha LP_B. Its titer of reactivity for alpha lipoprotein was much higher with aged plasma, HDL fractions, and the fraction of density > 1.21 than with comparable amounts of fresh plasma. They also obtained precipitation lines showing only partial identity between plasma and ultracentrifugally isolated HDL fractions in reacting with their antiserum.

Heterogeneity of alpha lipoprotein can also be seen in most published experiments using immuno-electrophoresis with antiwhole human sera. This was recognized by Ayrault-Jarrier, Levy, and Pólonožvski in their work (15), but it also appears to be present in the patterns published by Uriel and Grabar (13) and Grabar and Burtin (14). If one assumes that the absolute migration rates of the three lipid-staining bands obtained by Uriel and Grabar (13) differed from ours due to differences in water content of the agar (34), their "rho-lipoprotein" and "lipalbumin" lines represent alpha LP_A and LP_B, respectively. Neither line was present in their precipitate of LDL obtained by polyanions, the latter yielding only a single lipoprotein band, which they called "alpha." Burstin and Fine, using a technique that precipitates some of the alpha lipoprotein from material first freed of LDL, have also recently found two precipitation bands (16).

Scanu, Lewis, and Page also obtained two immuno-electrophoretic bands with HDL (4), but attributed one of these to albumin contamination. It is interesting how closely their published patterns resemble those we have obtained with the two forms of alpha lipoprotein.

Today the most widely used techniques for the preparation and quantification of lipoproteins depend upon the ultracentrifuge. Column chromatography has been attempted without demonstrated success (35). Precipitation methods with high molecular weight substances, which quite successfully permit separation of pure low density lipoproteins (18, 36), have so far proved capable of isolating only a small fraction of the total alpha lipoprotein in plasma (16). There still is, therefore, no method for isolating large amounts of high density lipoproteins that does not involve high speed centrifugation at least as a purification step.

The present studies are disturbing in their demonstration that ultracentrifugation is associated with alterations in high density lipoproteins that affect both their structure and quantitative recovery. The native form of HDL, alpha LP_A, is progressively converted to a lipoprotein form of higher density (alpha LP_B) with the loss of some to the infranate of the accepted density limit of 1.21. Such losses probably have been responsible in part for the tendency of concentrations of high density lipoproteins to be lower than that of their counterparts isolated by Cohn fractionation and electrophoresis (37). The conversion of alpha LP_A to alpha LP_B also probably accounts for much of the great variation in relative concentrations of HDL subfractions reported in the literature (2, 3, 7, 19, 38).

The two forms of alpha lipoprotein have been shown to bear a constant relationship to the two peaks into which HDL separates in the analytical ultracentrifuge by the technique of deLalla and associates. HDL_2 (density 1.063 to 1.1) represents only the A form, whereas HDL_3, the fraction of greater density (1.1 to 1.21), contains both A and B. Theoretically the relative amounts of HDL_2 and HDL_3 can be expected to vary with the degree of conversion of alpha A to B. It is interesting in this regard to compare analyses of HDL_2 and HDL_3 obtained in the analytical ultracentrifuge. Two such studies, in two similar populations with similar total HDL concentrations, have been reported in the past few years from the Donner Laboratory (7, 38). The ratios of HDL_2 to HDL_3 in the two sets of data are very different. There may be other possible explanations for these differences, but the fact remains that conversion of alpha lipoprotein during ultracentrifugation must be taken into account in assessing the significance of HDL subfractions. It is possible that the conventional subfractions, HDL_2 and HDL_3 (5), may be "artifactual" in the sense of their being more related to the transformability of alpha LP_A than to any specific metabolic influences.

The nature of the transformation of alpha LP_A to alpha LP_B can only be inferred from the pres-
ent data. B certainly contains relatively less lipid than A, as judged from their comparative flotation characteristics and lipid staining, and as confirmed by lipid analyses before and after conversion of A to B by treatment with ethanol: ether. The density range in which A and B are found is broad and overlapping, suggesting that the lipid contents of the A and B molecules are both variable and frequently similar. This suggests that other differences also probably exist between these two forms.

A possible difference in the protein component of the two forms of alpha must be considered. As indicated, the amino acid compositions of the polypeptide forms A and B are indistinguishable. It may be presumed that differences in their electrophoretic mobility and antigenicity therefore can most likely be due to some change in the structure of the protein of the lipoprotein. From previous work it is possible that such a structural change could be the presence of different multiples of an identical peptide subunit.

Shore (9) has shown that the protein contained in HDL subfractions of density 1.093 and 1.149 had the same amino acid analyses and end groups. The protein in density 1.093 molecules had twice the molecular weight of that in the 1.149 fraction and twice the number of amino terminal groups. He postulated that the protein in the lighter fraction was a dimer of that in the heavier fraction. Scanu, Lewis, and Bumpus (11) and Sanbar and Alaupovic (10) have found a single protein peak after delipidation of HDL over the entire 1.063 to 1.21 density range. The molecular weight of this protein was calculated to be about 75,000 or comparable to that of the monomer of Shore (9). Sanbar and Alaupovic found that this delipidated protein separated into two peaks on standing. The second peak appeared to be an aggregate of the first and could readily be converted to the first peak in the presence of 1 to 4 M urea or increased pH (10). More recently Shore and Shore (33) have presented evidence that the delipidated density 1.125 to 1.20 lipoproteins can be chemically divided into identical repeating protein subunits of about 36,000 mol wt.

The report of Scanu and Hughes (3) that their delipidated protein of 75,000 mol wt migrated upon electrophoresis more slowly on agar and more rapidly on starch gel than did alpha lipoprotein is consistent with the relative mobilities of alpha LPB and LPA in these same media.

It is considered that alpha LPB could represent a "monomer" form of alpha lipoprotein and alpha LPA a "dimer" or larger aggregate than alpha LPB. It will be necessary to obtain the molecular weight of the protein in each form before this hypothesis can be proved. How aging and exposure of plasma to a high centrifugal field remove lipid from the lipoprotein and quite possibly transform the polypeptide portion are clearly problems that both need to be solved, and should, in the course of their solution, throw much light on the nature of lipoprotein structure.

The demonstration of partially delipidated alpha lipoprotein (alpha LPB) in the ultracentrifugal fraction of density > 1.21 is quite consistent with its accounting for some of the phospholipid long known to be in this fraction (19, 39-41). Indeed this phospholipid has been shown by Kunkel and Trautman to migrate with the alpha1 globulin on starch block electrophoresis (41). The interesting suggestion has been made that an apoprotein circulates in plasma, at a density of > 1.21 that can be converted to a lipoprotein by the liver (42). A partially delipidated lipoprotein such as alpha LPB could be identical to such a carrier and this possibility will have to be explored.

**Summary**

The homogeneity of alpha1 or high density lipoprotein (density 1.063 to 1.21) (HDL) has been studied by immunochemical techniques in combination with electrophoresis, preparative and analytical ultracentrifugation, and amino acid analysis. Lipoproteins from the following sources have been examined: fresh plasma, plasma aged for a few days at 4°C, or freeze-thawed repeatedly, delipidated, or exposed to 8 M urea, and ultracentrifugal fractions including HDL (density 1.063 to 1.21), HDL2 (density 1.063 to 1.1), HDL3 (density 1.1 to 1.21), and the fraction of density greater than 1.21.

With the exception of fresh plasma and HDL2, alpha lipoprotein from all the other sources was reproducibly shown to exist in two forms. These have been designated alpha LPA and LPB. These two forms have differing mobility on agar gel electrophoresis, and 7 of 8 antisera prepared in
three different animal species have found them to be only partially cross-reactive as antigens.

Only very small amounts of alpha LPB were found in fresh plasma, and presumably alpha LP\textsubscript{A} is the native form in which these lipoproteins circulate. Both forms contain protein of the same amino acid composition and amino terminal groups. Alpha LP\textsubscript{B} contains relatively less lipid and is judged to be of a greater average density by its flotation characteristics. Alpha LP\textsubscript{B} was shown to be formed from alpha LP\textsubscript{A} by a partial delipidation.

It is concluded that repeated ultracentrifugation of alpha LP\textsubscript{A} is always associated with its transformation in part to alpha LP\textsubscript{B}. It was also demonstrated that the standard high density lipoprotein fractions obtained by ultracentrifugation are consistently related to, and their relative concentrations possibly explained by, this transformation. The HDL\textsubscript{2} peak of density 1.063 to 1.1 obtained by Gofman and colleagues in the analytical ultracentrifuge consists solely of LP\textsubscript{A}, whereas the HDL\textsubscript{3} peak (density 1.1 to 1.21) consists of alpha LP\textsubscript{B} plus some alpha LP\textsubscript{A}. Alpha LP\textsubscript{B} is also consistently demonstrated by immunochromal techniques in the fraction of density greater than 1.21, indicating that at least after ultracentrifugation, some alpha lipoprotein is present in a form of greater density than that usually attributed to the lipoproteins.

The relationship of these findings to earlier discrepancies in studies involving the immunochrometry or quantitation of high density lipoproteins is discussed.

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References


