Receptor-mediated Catabolism of Low Density Lipoprotein in Man

QUANTITATION USING GLUCOSYLATED LOW DENSITY LIPOPROTEIN

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ABSTRACT Low density lipoprotein (LDL) catabolism occurs by LDL receptor-dependent and LDL receptor-independent pathways. We have shown previously that nonenzymatic glucosylation of LDL in the presence of cyanoborohydride irreversibly blocks the lysine residues of LDL. Glucosylated LDL (GLC-LDL) was not degraded by the LDL receptor of fibroblasts, and its degradation by macrophages was similar to that of native LDL. This suggested that GLC-LDL should be a good tracer of LDL receptor-independent catabolism, and if combined with a tracer of total LDL catabolism, should enable one to calculate the extent of LDL receptor-dependent catabolism.

To determine the contribution of each pathway in man, we prepared 125I-GLC-LDL and 131I-control LDL and simultaneously determined the fractional catabolic rate (FCR) of each tracer in four subjects. In preliminary experiments, we showed that the conditions for glucosylation did not affect LDL turnover. In the four subjects, the FCR for total LDL catabolism ranged from 0.345 to 0.724 d⁻¹ with a mean of 0.57±0.16 d⁻¹. The FCR of GLC-LDL varied from 0.071 to 0.141 d⁻¹ with a mean of 0.11±0.03 d⁻¹. The latter is similar to the FCR reported for native LDL in subjects with homozygous familial hypercholesterolemia, supporting the interpretation that GLC-LDL traces only the receptor-independent pathway. Despite the wide range of total LDL catabolism in these subjects, LDL receptor-independent catabolism accounted for only 19.5–20.6% of total catabolism. In turn, LDL receptor-dependent catabolism accounted for 80% of total clearance in each person. Furthermore, while the decay curve of LDL showed the usual biphasic pattern, the decay curve of GLC-LDL was monoexponential in each subject even when followed for as long as 48 d. This suggests that LDL receptor activity is responsible for the biphasic nature of LDL decay.

These studies emphasize the central role of LDL receptor activity in normal LDL metabolism in man.

INTRODUCTION

Because low density lipoproteins (LDL) play a central role in the development of atherosclerosis, a detailed understanding of their catabolism is of particular importance. Clearance of LDL from plasma is mediated via a high affinity LDL receptor-dependent pathway as well as one or more LDL receptor-independent pathways (1–3). The receptor-mediated process involves the recognition of apolipoprotein (apo)1-B, the major protein of the LDL particle, by a high affinity receptor present on cell membranes in various tissues, including liver (1–4). The classic work of Goldstein and Brown (1) has shown that receptor-mediated endocytosis of LDL leads to an orderly series of biochemical events, which serve to maintain cellular cholesterol homeostasis. In the absence of LDL receptors, as in homozygous familial hypercholesterolemia (HFH) (1) and in the Watanabe heritable hyperlipidemic rabbit (3–5), fractional catabolic rates (FCR) of LDL are markedly decreased, plasma LDL levels are grossly

1 Abbreviations used in this paper: apo, apolipoprotein; C, control (LDL); CHD, cyclohexanediene-modified (LDL); GLC, glucosylated (LDL); FCR, fractional catabolic rate; HFH, homozygous familial hypercholesterolemia; U/P, urinary excretion rate of 125I and 131I radioactivity to the 125I and 131I radioactivity in plasma.

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elevated, and LDL apo-B production rates are increased (3–7). The increased synthesis and degradation of LDL apo-B, of necessity, occurs almost entirely via LDL receptor-independent pathways. It is precisely in this setting of a high LDL flux via nonreceptor pathways that severe and premature atherosclerosis is observed. Thus, it would be important to know the relative contribution to total LDL catabolism of the receptor-independent pathway in man.

We have previously demonstrated that incubation of LDL in the presence of glucose and cyanoborohydride results in irreversible glucosylation of lysine groups of LDL and permanently blocks the ability of LDL to interact with the LDL receptor of fibroblasts (8). We have also shown that the metabolism of glucosylated (GLC) LDL by mouse peritoneal macrophages that express very few high affinity receptors for native human LDL, is similar to that of native LDL (8, 9). This suggested that GLC-LDL might be useful for quantitating receptor-independent LDL catabolism in vivo. By combining such a measurement with simultaneous measurement of native LDL catabolism, receptor-dependent and receptor-independent, it is possible to calculate that portion of LDL catabolism accounted for by the LDL receptor pathway (10, 11). This report describes our initial results using this technique in normoglycemic human subjects.

METHODS

Subjects. All studies reported here were performed on subjects admitted to the Special Diagnostic and Treatment Unit (Metabolic Unit) of the Veterans Administration Medical Center, San Diego, CA. The clinical characteristics of the six individuals studied are presented in Table I. Two of the subjects had suffered myocardial infarction (No. 4 and 5) and one (No. 6) had undergone endarterectomy for carotid vascular disease. Routine clinical laboratory tests indicated that at time of study all were free of thyroid, hepatic, or renal dysfunction, and had normal fasting and 2-h post-prandial glucose values. Table I also shows the plasma lipid and lipoprotein values for each of the subjects, based on multiple measurements while on a defined diet as described below. Subjects 1, 2, and 3 had normal plasma lipids, while the other three had mild hypertriglyceridemia consistent with phenotype IV. None of the subjects had followed a therapeutic diet or received hypolipidemic drug therapy for the 3 mo before the study period. Subjects 1 and 2 participated in the control turnover studies and subjects 3–6 participated in the GLC-LDL turnover studies. All patients gave informed consent for the investigations, which were approved by the Human Studies Committee of the University of California, San Diego.

The subjects were fed an isocaloric, weight-maintenance diet of mixed solid food and liquid formula (Hospital Diet Products, Organon Inc., Buena Park, CA). The diet contained 45% of calories as carbohydrate, 15% as protein, and 40% as fat with a polyunsaturated/saturated fat ratio of 0.2. Cholesterol intake averaged 100–150 mg/d. A detailed description of this diet has been previously published (12–13). Three liquid meals and one solid meal were given per day with appropriate vitamins and mineral supplementation. Each subject was weighed daily, and caloric intake was adjusted to maintain constant body weight throughout the study period.

Preparation and iodination of LDL. After the subjects had been on the mixed solid food and liquid formula diet for 1–2 wk, plasmapheresis was carried out by collecting blood (500 ml) in sterile, pyrogen-free plastic bags containing 500 mg of the disodium salt of EDTA. Plasma was separated by centrifugation (4°C) at 5,500 g for 5 min, and cells were reinfused. Isolation of LDL (d = 1.025–1.060 g/ml) was carried out according to the method of Lindgren et al. (14), by using sterile techniques as we previously described (13). Plasma was adjusted to d = 1.025 g/ml with a NaCl-NaBr solution and centrifuged in a 60 Ti rotor in an L 2-65B preparative ultracentrifuge (Beckman Instruments, Inc., Palo Alto, CA) at 59,000 rpm (15°C) for 18 h. The infranate was adjusted to d = 1.060 g/ml and subjected to further ultracentrifugation for 14 h in a 65 Ti rotor at 64,000 rpm (15°C). A final ultracentrifugation was done in a 40.3 Ti rotor at 39,000 rpm (15°C). After addition of an equal volume of a NaCl-NaBr solution of d = 1.070 g/ml the concentrated LDL was then dialyzed against 0.15 M NaCl containing 0.01% EDTA, pH 7.4. (EDTA-saline). The LDL was divided into two equal portions and one was labeled with 125I and the other with 123I by the iodine monochloride

| TABLE I

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<th>Clinical Data and Plasma Lipids</th>
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*Averages of eight determinations (mean±SEM).

CHD, coronary heart disease; CVD, carotid vascular disease.

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method as modified by Bilheimer et al (15). Reactants were added so that the molar ratio of iodine to protein was <1. Free radioiodide was removed by extensive dialysis of the iodinated lipoprotein against EDTA-saline. The amount of free iodide after dialysis was determined by thin-layer chromatography of the iodinated LDL. Specific activities of the labeled LDL preparations ranged between 7.0 and 50 cp/mg. The degree of lipid labeling was determined in each iodinated lipoprotein as previously described (13) and averaged 2.5±1.3% for C-LDL and 2.9±1.8% for GLC-LDL (mean±SD).

Glycosylation of LDL. Glucose reacts with epsilon amino groups of lysine to form a labile Schiff base intermediate, which can spontaneously dissociate, or rearrange to give a stable ketoamine adduct (Amadori reaction). Because the latter reaction is slow, addition of a reducing agent is required to produce a LDL in which greater than one-third of lysines are glucosylated (8). Addition of a reducing agent traps the Schiff base form, yielding a glucolysine adduct that is stable even when labeled LDL is incubated under sterile conditions for 5 d at 37°C in phosphate-buffered saline (PBS), pH 7.4, containing 80 mM glucose and 200 mM cyanoborohydride. (The cyanoborohydride and glucose were added as fresh sterile solutions for each incubation). The C-LDL was incubated under identical conditions except for omission of glucose. At the end of incubation, both GLC-LDL and C-LDL were exhaustively dialyzed for 18 h against EDTA-saline. Before incubation, a typical concentrated native LDL preparation contained 22 ng/mL of cytidine, and after incubation with glucose and cyanoborohydride and extensive dialysis, 0.8 µg of cytidine was detectable per milliliter sample; this resulted in a total cytidine dose injection of 0.96 µg. After equilibration in the average person, this would result in whole blood levels <1 ng/mL, which is below the range detected in whole blood in normal individuals (0–150 ng/mL; SmithKline Corp., SmithKline Clinical Laboratories, Burbank, CA).

Special control preparations. To exclude an effect on LDL metabolism due to the conditions needed for glucosylation, we prepared three control preparations: LDL incubated in PBS for 5 d at 37°C in the presence of cyanoborohydride (C-LDL); LDL incubated in PBS for 5 d at 37°C in the absence of cyanoborohydride (LDL-37°C, no borohydride); and to exclude an effect due to incubation at 37°C, LDL incubated in PBS for 5 d at 4°C in the absence of cyanoborohydride (LDL-4°C, no borohydride). The metabolism of these three controls was compared in cell culture experiments and in vivo turnover studies described below.

Preparation of labeled LDL for injection. After removal of glucose and cyanoborohydride by dialysis, the labeled LDL preparations were subjected to centrifugation at 20,000 g for 30 min (4°C), the supernatant was then removed and passed through a 0.22-µm Millipore filter. Sterile human serum albumin in a 5% solution was added to the labeled LDL to dilute the original preparation 10-fold (13). This solution was then passed again through a 0.22-µm Millipore filter and aliquots set aside for bacterial and fungal testing and for pyrogen testing.

Study protocol and sample analysis. 12 d after plasma-
calculated from the mean of four measured values of the ratio of LDL cholesterol to LDL protein and the multiple measurement of LDL cholesterol (17). LDL protein was determined by the method of Lowry et al. (20).

**Tissue culture studies.** In the present studies, we tested each pair of labeled GLC-LDL and C-LDL preparations in tissue culture to demonstrate that the degree of glucosylation of the GLC-LDL was sufficient to completely block degradation via the LDL receptor. The cell culture techniques used have been described in detail previously (8, 21). The special control preparations were also tested in fibroblasts to show that incubation at 37°C and/or treatment with cyanoborohydride did not affect degradation via the LDL receptor. In addition, to exclude an effect of these treatments on degradation by other pathways, LDL from patient No. 1 were also tested in J774 cells, a macrophage-like cell line (obtained from J. Unkeless, The Rockefeller University, New York). These cells were grown in minimal essential medium with 10% fetal calf serum, and were incubated with labeled LDL preparations for 5 h; otherwise the techniques were the same as those used with the fibroblasts (8).

**RESULTS**

**Cell culture studies.** We have previously shown that with conditions used in these incubations, 45–60% of lysine residues of LDL are irreversibly glucosylated (8, 9). We, (8) and others (22) have shown that when more than one-third of lysine residues of LDL are blocked, LDL is no longer recognized by the LDL receptor. To document the effectiveness of glucosylation, the degradation of each GLC-LDL preparation was tested on normal human fibroblasts and compared with that of the corresponding C-LDL; in every case, saturable, high affinity degradation was completely blocked for GLC-LDL, but was normal for C-LDL.

Fig. 1 (left panel) shows the degradation of C-LDL and GLC-LDL vs. LDL concentration for the labeled preparations of subject 6; virtually identical results were found for each of the other subjects.

The results with the "special control" preparations indicated that in fibroblasts the degradation of LDL was unaffected by cyanoborohydride treatment or incubation at 37°C in the absence of glucose. Fig. 1 (right panel) shows the degradation curve in fibroblasts comparing C-LDL with untreated LDL kept at 4°C. Similar results were seen comparing C-LDL with LDL incubated at 37°C in PBS alone. In addition, control and native LDL were all metabolized equally by J774 macrophage cells (data not shown).

**Lack of effect of cyanoborohydride and incubation on C-LDL catabolism.** To exclude an independent effect of cyanoborohydride treatment on the catabolism of LDL, we compared the turnover of 125I-LDL incubated for 5 d at 37°C in the presence of cyanoborohydride (C-LDL) with 125I-LDL incubated for 5 d at 37°C without cyanoborohydride (LDL-37°C, no borohydride). When these tracers were simultaneously injected into subject 1, the FCR for the control LDL was 0.42 d⁻¹ and for the LDL incubated in the absence of cyanoborohydride, 0.46 d⁻¹ (Fig. 2A). To rule out any effect of the 5-d incubation at 37°C on the catabolism of C-LDL, an iodinated LDL preparation kept at 4°C (LDL-4°C, no borohydride) and C-LDL were injected simultaneously into subject 2. The FCR for the LDL-4°C was 0.57 d⁻¹ and for the C-LDL, 0.52 d⁻¹ (Fig. 2B).

**Figure 1** Degradation of various LDL preparations by human fibroblasts. 24 h before cells were exposed to labeled LDL preparations, medium was replaced with Dulbecco's modified Eagle's medium containing 5% lipoprotein-deficient serum. After addition of indicated concentrations of labeled lipoprotein, cells were incubated overnight and then content of TCA-soluble, noniodide radioactivity was determined. Results are given as micrograms of LDL protein degraded per dish vs. micrograms of LDL added per milliliter of medium. Left panel; comparison of rates of degradation of C-LDL (○) and GLC-LDL (▲) obtained from subject 6. Right panel; comparison of rates of degradation of C-LDL (○) and LDL-4°C (▲) obtained from subject 2. C-LDL is LDL that was incubated for 5 d at 37°C in PBS in presence of cyanoborohydride, and LDL-4°C is LDL that was simply kept at 4°C for the equivalent period of time.

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Catabolism of GLC-LDL and C-LDL. The catabolism of GLC-LDL was remarkably slower than that of C-LDL in each of the four subjects studied as shown in Fig. 3. The kinetic parameters for GLC-LDL and C-LDL turnover are shown in Table II. The FCR of C-LDL ranged from 0.345 to 0.724 d⁻¹, while that for GLC-LDL ranged from 0.071 to 0.141 d⁻¹ (mean 0.11±0.03). However, in all four subjects, the FCR of GLC-LDL as a percentage of that for C-LDL was remarkably constant, 19.5-20.6%. When the U/P ratio data were used to calculate the FCR of each tracer, GLC-LDL again accounted for only 20% or less of total LDL catabolism (Table II). The FCR for C-LDL derived from the U/P ratios, however, must be qualified by the observation that the U/P ratios were not constant over time, but showed an initial rise and subsequent slow decline with time (Fig. 4). Others (23, 24) have described a similar finding for the turnover of native LDL. In contrast, after 2 d, the U/P ratio for GLC-LDL was remarkably constant. Assuming GLC-LDL traces only the clearance of the receptor-independent pathway, it follows that the LDL receptor pathway accounted for 80% of total LDL clearance in each subject. In terms of absolute catabolic rates, receptor-mediated catabolism of LDL protein ranged from 10.5 to 20.5 mg/d per kg among these four subjects, while receptor-independent catabolism ranged from 2.7 to 4.9 mg/d per kg. Assuming that all LDL cholesterol is catabolized as a unit with LDL protein, it can be estimated that receptor-mediated clearance of LDL cholesterol ranged from 14.9 to 20.2 mg/d per kg (Table III).

In contrast with the two exponentials found in each decay curve of the control LDL, the die-away curve of GLC-LDL was monoeponential in each of the patients even when followed for 48 d after injection. The monoexponential decay was observed even during the first few hours after injection during which numerous separate measurements were made. Fig. 5 shows the decay of GLC-LDL during the first 6 h for subject 3 whose complete decay curve can be seen in Fig. 3C. The slow monoexponential decay was also reflected in the low and constant U/P ratio noted in all subjects. The monoexponential decay of GLC-LDL cannot be explained by a different initial volume of distribution because the calculated plasma volumes were the same for both C-LDL and GLC-LDL tracers (Table II).

Accelerated catabolism of GLC-LDL in some subjects. Subsequent to the studies above, we had the opportunity to study an individual with hypothyroid-
isolation. Because hypothyroidism leads to a decreased expression of LDL receptors in vivo (25), we sought to measure the extent of the receptor-dependent pathway in this individual. The plasma decay curves for the control and GLC-LDL tracers in this subject are shown in Fig. 6. The control LDL showed the usual

**Figure 3** Turnover of C-LDL (□) and GLC-LDL (△) in human subjects. For each subject, an equal amount of $^{125}$I-GLC-LDL and $^{131}$I-C-LDL was injected simultaneously and plasma decay curves subsequently determined over the following 14–18 d (note different time scales for each subject). In subject 6 (D), the isotope assignment was reversed to rule out an isotope effect as etiology of the markedly different rates of decay. Kinetic parameters for each pair of turnovers can be found in Table II. A is for subject 4; B, 5; C, 3; and D, 6.

**Table II**

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<th>Patient</th>
<th>C-LDL</th>
<th>GLC-LDL</th>
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* Fraction of intravascular apo-LDL pool metabolized each day was calculated from the plasma die-away curve and from U/P ratio.
† Plasma volume calculated separately for C-LDL and GLC-LDL by isotope dilution technique as described in Methods.

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biphasic decay pattern. However, after 4 d of monoeponential decay, there was the abrupt onset of a phase of rapid decay of the GLC-LDL tracer. We have subsequently seen a similar phenomenon in three of four diabetic subjects and in one other normoglycemic control. In each case, after 4–8 d of monoeponential decay, we noted the abrupt onset of a phase of rapid clearance of the GLC-LDL tracer, similar to that noted in Fig. 6.

DISCUSSION

LDL catabolism occurs by LDL receptor-dependent and LDL receptor-independent pathway(s). To estimate the extent of each pathway in vivo, Mahley et al. (10) and Shepherd et al. (11) introduced the concept of performing a double-labeled turnover study using native LDL as the tracer of total LDL catabolism and a chemically modified LDL as the tracer of receptor-independent catabolism. The difference in FCR between the two tracers is taken to reflect LDL catabolism occurring by the receptor-dependent pathway. For this technique to be valid, the chemically modified LDL must be irreversibly modified so that it is not recognized by the LDL receptor throughout the period of study. In addition, the modification of LDL must not interfere with clearance by any receptor-independent pathway(s).

Nonenzymatic glucosylation of LDL specifically blocks lysine groups of LDL (8) and presumably it is this modification that inhibits LDL recognition by the LDL receptor. We previously demonstrated that the conditions needed for glucosylation, as opposed to the covalent addition of glucose, did not affect LDL metabolism in cell culture or LDL turnover in guinea pigs (8). In the present study, we extend the prior obser-

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<td><strong>Receptor-mediated and Receptor-independent Catabolism of LDL Cholesterol</strong></td>
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vations to show that in man as well, the conditions for glucosylation do not have any significant effect on LDL turnover. Therefore, any difference in metabolic behavior between C-LDL and GLC-LDL is due to the derivatization of lysine by glucosylation.

Several lines of evidence suggest that GLC-LDL is an adequate tracer for the receptor-independent pathway. The modification is irreversible and total blockade of receptor-mediated degradation of GLC-LDL in fibroblast culture was verified for each injected GLC-LDL of the present study (Fig. 1). The observations that all plasma radioactivity originally used to label GLC-LDL retained enhanced electrophoretic mobility, the monoexponential decay of GLC-LDL for up to 48 d, and a rather flat U/P ratio all attest to the in vivo stability and persistent homogeneity of the GLC-LDL tracer.

Does glucosylation of LDL cause LDL to be uniquely recognized by any LDL receptor-independent pathway such as those that exist on macrophages? We have previously shown that degradation of GLC-LDL by macrophages is no different than that of native or C-LDL (8) and the current observation that it has a slow monoexponential decay from plasma strengthens the suggestion that GLC-LDL has no enhanced clearance by any reticuloendothelial system. Finally, does glucosylation of LDL inhibit its in vivo clearance by any receptor-independent pathway(s)? Two lines of evidence suggest it does not. First, in the guinea pig, the clearance of GLC-LDL is identical to the clearance of methyl LDL (9). Second, and of particular relevance, we (9) have also shown that the turnover of GLC-LDL is similar to that of native LDL in the Watanabe heritable hyperlipidemic rabbit, an animal model for HFH, which has <5% of normal LDL receptors (3–5). In sum, our data suggest that GLC-LDL is an appropriate tracer for the receptor-independent pathway.

The FCR values for total LDL catabolism in our subjects ranged from 0.345 to 0.724 d⁻¹. In a recent review (26) of 15 separate published reports of LDL turnover in normal subjects, we calculated a mean FCR of 0.371 d⁻¹, which is similar to the value of 0.345 found for normolipidemic subject 3. The FCR for subjects 4–6 were much higher, possibly reflecting their obesity and/or associated hypertriglyceridemia. We (27) and others (28) have reported that such subjects have LDL FCR values that range between 0.504 and 0.768 d⁻¹. Despite the wide range of total LDL catabolism noted in the subjects of this report, it is remarkable that LDL receptor-dependent catabolism accounted for ~80% of total LDL catabolism in each subject. This was true whether the receptor-dependent contribution was determined from the difference in the plasma die-away curves or as calculated from the U/P ratio data. These data are in agreement with the predictions of Goldstein and Brown (1) who estimated that at least two-thirds of LDL catabolism should occur by an LDL receptor-dependent process. Using GLC-LDL (or methyl LDL) as the tracer of receptor-independent pathway, we found that 66–78% of LDL catabolism occurs by an LDL receptor-mediated pathway in the guinea pig (8, 9). Using GLC-LDL in the rabbit, we found that 75% of catabolism was receptor-mediated (9), which agrees with the value of 67% found by Bilheimer et al. (29) using methyl LDL, and 67% found by Pittman et al. (3) using still a third technique. These data in man and other species serve to emphasize the central importance of the LDL receptor for normal LDL metabolism.

The FCR for GLC-LDL ranged from 0.071 to 0.141 d⁻¹ with a mean value of 0.112 d⁻¹. These values are exactly in the range previously reported for LDL turnover of native LDL in patients with HFH, (range of 0.06–0.178 d⁻¹) (30–32). The similarity in values of FCR for GLC-LDL in our subjects and for native LDL in HFH subjects strongly supports the view that GLC-LDL traces all of the receptor-independent pathways of LDL catabolism and only those pathways. In contrast, the FCR in normal subjects of cyclohexanedione (CHD)-modified LDL, which has also been used as a tracer of the receptor-independent pathway, has ranged from 0.163 to 0.261 d⁻¹ (11, 25–33), values higher than those for native LDL in HFH subjects. This suggests that CHD-LDL overestimates the receptor-independent pathway. In fact, using CHD-LDL, Shepherd et al. (11) and Thompson et al. (32) estimated that in man LDL receptor-dependent catabolism accounted for only 33–40% of total LDL catabolism. We found in the guinea pig that turnover of CHD-LDL is consistently greater than that of either GLC-LDL or methyl LDL (which are equal) (9) and Slater et al. (34) have reported that CHD-LDL is cleared more rapidly than methyl LDL in the rabbit (34). The reasons for the more rapid clearance of CHD-LDL are not clear but could be due to enhanced clearance of CHD-LDL by a nonreceptor mechanism (34), or to slow spontaneous reversal of the CHD modification, as first demonstrated by Mahley et al. (10) and confirmed by Slater et al. (34) and by us (3).

The decay curve for native LDL has an initial rapid phase of decay that lasts for 2–5 d, followed by a second, slower phase. The U/P data also suggest heterogeneity in the metabolism of LDL. In sharp contrast, the decay of GLC-LDL was monoexponential even when followed for as long as 48 d. The exact explanation for this is unknown, but to our knowledge the only physiologic difference between the two tracers is the inability of GLC-LDL to interact with the LDL receptor. This suggests that the biphasic curve for na-

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tive LDL clearance is caused by reversible binding to readily accessible LDL receptors (as in liver, spleen, or endothelial cells) or that reversible exchange with an extravascular compartment occurs chiefly by an LDL receptor-dependent process.

Originally, we planned to study a larger number of subjects with this technique. However, in three of four diabetic patients, one hypothyroid subject, and one normal control after several days of monoeponential decay, we noticed the abrupt onset of a phase of rapid clearance of GLC-LDL that began 4–10 d after injection. We are currently investigating this phenomenon and preliminary evidence suggests that it may be immunological in origin. In guinea pigs, nonenzymatic glucosylation of homologous LDL renders it immunogenic (35). Furthermore, in two of the human subjects, in whom the rapid decay was seen, (including the subject of Fig. 6) preliminary evidence suggests that antibodies against GLC-LDL, but not C-LDL, were present in plasma obtained before injection of the tracers. However, in none of the subjects of the present report have such antibodies been found. Although we have noticed no adverse effects in any of our patients, we would urge caution in further in vivo studies with GLC-LDL until a better understanding of this phenomenon has been achieved.

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