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Timothy J. Lyons
Karen E. Bailie
Daniel G. Dyer
John A. Dunn
John W. Baynes

University of South Carolina - Columbia, john.baynes@sc.edu

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Decrease in Skin Collagen Glycation with Improved Glycemic Control in Patients with Insulin-dependent Diabetes Mellitus

Timothy J. Lyons,*† Karen E. Baille,* Daniel G. Dyer,§ John A. Dunn,§ and John W. Baynes**

*Department of Medicine, Altnagelvin Hospital, Londonderry, Northern Ireland, United Kingdom; †Sir George E. Clark Metabolic Unit, Royal Victoria Hospital, Belfast, Northern Ireland, United Kingdom; and **Department of Chemistry, and
†School of Medicine, University of South Carolina, Columbia, South Carolina 29208

Abstract

Glycation, oxidation, and nonenzymatic browning of protein have all been implicated in the development of diabetic complications. The initial product of glycation of protein, fructosyllysine (FL), undergoes further reactions, yielding a complex mixture of browning products, including the fluorescent lysine-arginine cross-link, pentosidine. Alternatively, FL may be cleaved oxidatively to form N'- (carboxymethyl) lysine (CML), while glycated hydroxylsine, an amino-acid unique to collagen, may yield N'- (carboxymethyl) hydroxylsine (CMHl). We have measured FL, pentosidine, fluorescence (excitation = 328 nm, emission = 378 nm), CML, and CMHl in insoluble skin collagen from 14 insulin-dependent diabetic patients before and after a 4-mo period of intensive therapy to improve glycemic control. Mean home blood glucose fell from 8.7±2.5 (mean±1 SD) to 6.8±1.4 mM (P < 0.005), and mean glycated hemoglobin (HbA1c) from 11.6±2.3% to 8.3±1.1% (P < 0.001). These changes were accompanied by a significant decrease in glycation of skin collagen, from 13.2±4.3 to 10.6±2.3 mmol FL/mol lysine (P < 0.002). However, levels of browning and oxidation products (pentosidine, CML, and CMHl) and fluorescence were unchanged. These results show that the glycation of long-lived proteins can be decreased by improved glycemic control, but suggest that once cumulative damage to collagen by browning and oxidation reactions has occurred, it may not be readily reversed. Thus, in diabetic patients, institution and maintenance of good glycemic control at any time could potentially limit the extent of subsequent long-term damage to proteins by glycation and oxidation reactions. (J. Clin. Invest. 1991. 87:1910–1915.) Key words: glycation • nonenzymatic browning • Maillard reaction • diabetes • oxidation

Introduction

The mechanisms underlying the development of the complications of diabetes are not fully understood. Even the relationship between glycemic control and the risk of developing complications remains unclear, although there is now a consensus that hyperglycemia does, in itself, play an important role in the development of retinopathy, nephropathy, and neuropathy (1). The processes of glycation and nonenzymatic browning of proteins provide an attractive hypothesis to link hyperglycemia with the development of complications (2–4). The first step in this reaction pathway, glycation, involves the nonenzymatic condensation of glucose with free amino groups in the protein, primarily the α-amino groups of lysine residues, forming the Amadori adduct, fructosyllysine (FL) (Fig. 1). FL may react further to initiate a complex series of reactions, which lead to the accumulation of covalently attached brown and fluorescent products, cross-links, and other chemical modifications in proteins. These reactions are known collectively as Maillard or nonenzymatic browning reactions (5, 6). This reaction is readily illustrated by the browning and cross-linking of proteins on incubation with glucose in physiological buffers in vitro.

A number of chemical and physical changes occur in human skin collagen with age. Fluorescence, cross-linking, and resistance to enzymatic degradation increase with age, while solubility and elasticity decrease (reviewed in 7). In diabetes, in concert with increased glycation (8–10) and nonenzymatic browning (11–13) of collagen, these age-related changes in the physical and chemical properties of collagen appear to be accelerated (11–15). This suggests a role for glucose and nonenzymatic browning reactions in the development of age-like chemical and functional alterations of collagen in diabetes. Although the extent of glycation of skin collagen does not appear to correlate directly with the presence of complications in diabetes (9, 10), the long-term effect of increased glycation, i.e., the browning reaction, may be more relevant. Thus, in groups of insulin-dependent diabetic patients, matched for age and duration of diabetes, there is a significant correlation between skin collagen fluorescence and the severity of retinopathy, nephropathy, arterial stiffness, and joint stiffness (16). Increased free-radical–mediated oxidative damage to biomolecules, including both lipids and proteins, has also been proposed as a mechanism contributing to the development of diabetic complications (reviewed in 17, 18). Indeed, free radical reactions may in themselves generate fluorescent products and cross-links in proteins (19), and thus the changes in collagen with age and in diabetes may be the combined result of increases in glycation, browning, and oxidation reactions.

Several distinct chemical products of glycation and browning reactions of protein have now been measured in human proteins. These compounds (Fig. 1) include the Amadori adduct, FL, and three products of later stages of the Maillard

1. Abbreviations used in this paper: CMHl, N'- (carboxymethyl) hydroxylsine; CML, N'- (carboxymethyl)lysine; FL, N'- (1-deoxy-fructose-1-yl) -lysine; HbA1c, hemoglobin A1c; MHBG, mean home blood glucose.
Figure 1. Maillard reaction pathways for formation of fructoselysine, pentosidine, and N\(^\alpha\)-(carboxymethyl)lysine. FL is the Amadori compound, the first stable intermediate in the Maillard reaction. CML and the analogous compound CMH-L (not shown), are formed by oxidative cleavage of Amadori adducts to lysine and hydroxylysine, respectively. The mechanism of formation of the fluorescent crosslink, pentosidine, is unknown.

Reaction: pentosidine\(^2\) (20–23), N\(^\alpha\)-(carboxymethyl)lysine (CML) (24–27), and N\(^\alpha\)-(carboxymethyl)hydroxylysine (CMH-L) (27). The concentration of the initial product, FL, in long-lived proteins, such as lens proteins (26, 28) and skin collagen (27), increases in response to hyperglycemia in diabetes. Among the later products of the Maillard reaction, pentosidine is a fluorescent cross-link formed between lysine and arginine residues during the browning process (20–23), while CML and CMH-L, which are colorless, are formed by oxidative cleavage of carbohydrate adducts to lysine and hydroxylysine residues in protein, respectively (24–27). All three of these late-stage products of the Maillard reaction require oxygen for their formation (20, 24, 25), i.e., they are either direct products of oxygen radical reactions or are formed by further reaction of oxidation products. In addition, they all accumulate gradually with age in skin collagen (27), and at an accelerated rate in diabetes (21, 23).

In this study, we have measured FL, pentosidine, CML, and CMH-L and Maillard-type fluorescence (excitation (Ex) = 328 nm, emission (Em) = 378 nm) in insoluble skin collagen from patients with insulin-dependent diabetes, both before and after a period of improved glycemic control. Our aim was to discover if improvements in control would lead to a decrease in the level of any of these Maillard reaction products in diabetic skin collagen, and thus, perhaps, to a reversal of the potentially damaging effects of glycation, browning, and oxidation to long-lived proteins in diabetes.

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2. The compound Maillard Fluorescent Product #1 (MFP-1) described in references 22 and 23 was originally identified as a fluorescent cross-link formed during browning and cross-linking of proteins by glucose. MFP-1 was isolated and characterized recently in our laboratory and shown by nuclear magnetic resonance spectroscopy and mass spectrometry to be identical to the compound, pentosidine, previously characterized by Sell and Monnier (20, 21). There is some uncertainty about whether pentosidine is formed in vivo from ribose (20, 21), glucose (22, 23), or other sugars (23a).

**Methods**

**Patient selection.** Patients (eight male, six female) with insulin-dependent diabetes mellitus, who were in relatively poor glycemic control, but motivated to improve, were recruited from the diabetes clinics of Altnagelvin Hospital, Londonderry, and the Royal Victoria Hospital, Belfast, Northern Ireland. Their mean age was 31.9±10.5 (19–51) yr (mean±SD, range), and mean duration of diabetes 12.5±10.6 (0–38) yr. Three patients had newly diagnosed diabetes. Of the remainder, 10 were receiving twice daily injections of regular and NPH (Isophane) insulin, and one a single daily injection. Mean daily insulin dose was 74±11 (52–88) U. The study was approved by the Ethical and Human Subjects Committees of the participating institutions, and informed consent was obtained from all volunteers.

**Study design.** After an initial assessment, each patient was taught to perform home blood glucose monitoring using a Memory Glucometer (Ames Div., Miles Laboratories Inc., Elkhart, IN) and was given a Memory Glucometer on loan for the duration of the study. Patients performed blood glucose measurements four times daily (before meals and at bedtime) throughout the study. For a 2-wk “run-in” period before the first skin biopsy, no effort was made to alter glycemic control. During this period, home blood glucose monitoring results were recorded, and hemoglobin A\(_1\) (HbA\(_1\)) was measured on two occasions. At the end of the run-in period, a full-thickness, elliptical (1.0 x 0.5 cm) skin biopsy was obtained, under local anesthesia, from the upper inner aspect of the buttocck. The biopsy samples were washed in saline and stored at -70°C. Patients then entered a program of intensive management to improve glycemic control, and were seen on an individual basis by one of us (T. Lyons or K. Bailie) at least once every two weeks. At each visit, the patients received appropriate individual education, including advice on the adjustment of insulin dosage. Between visits, one of us was available at all times to give advice by telephone. Each patient was also assessed initially, and subsequently reviewed, as necessary, by a diettitian. Throughout the study, HbA\(_1\) measurements were performed every two weeks. The results of home blood glucose monitoring were analyzed using the Ames "Glucofacts" program, and were expressed as "mean home blood glucose" (MHBG) on a weekly basis. At the conclusion of the study, a second skin biopsy was obtained from the corresponding site on the opposite buttocck. The mean interval between first and second skin biopsies was 120±19 (92–157) d.

**Analytical procedures.** HbA\(_1\) was measured by agar gel electrophoresis (29); the normal range in our laboratory is 3.6–7.2%. Insoluble collagen was isolated from skin biopsy samples by mechanical scraping and solvent extraction as previously described (9). The preparation of N\(^\alpha\)-formyl-N\(^\alpha\)-(fructoselysine) as the standard (for the measurement of FL), CML, and CMH-L, have been described previously (24, 25, 27). A radioactive pentosidine standard was prepared from glucose, N\(^\alpha\)-acetylarginine, and N\(^\alpha\)-acetyl-[4,5-\(^3\)H]lysine of known specific radioactivity, and purified by reversed phase HPLC (23a). Levels of FL, CML, CMH-L, and lysine in insoluble collagen were measured by gas chromatography/mass spectroscopy with selected ion monitoring (SIM-GC/MS), as previously described in detail (26, 27). Briefly, for measurement of FL, collagen samples were hydrolyzed in 7.8 N HCl (24 h, 110°C, under N\(_2\)), yielding 40% conversion of FL to the analyte, furosine (26, 27, 30). Because of partial conversion of FL to CML during the hydrolysis reaction (and possible conversion of glycated hydroxylysine to CML), FL, CML, and CMH-L were measured separately in NaBH\(_4\)-reduced samples (27). Furosine (formed during acid hydrolysis of FL), CML, and CMH-L were measured as their N\(_2\)-N-trifluoroacetyl methyl ester derivatives by SIM-GC/MS. The assays were standardized using standard curves prepared by standard addition, and the concentrations were normalized to the lysine (for FL and CML) or hydroxylysine (for CMH-L) content of the collagen. The pentosidine content of skin collagen was measured, following NaBH\(_4\)-reduction and hydrolysis in 6 N HCl (24 h at 110°C), by reversed phase HPLC using fluorometric detection (Ex = 328nm, Em = 378nm) (22, 23a). Quantitation was based on fluorescence area units, using a standard curve prepared with known amounts of authentic pentosidine, and normalized to the lysine content of the protein. For

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measurement of fluorescence (Ex = 328 nm, Em = 378 nm), collagen was digested with pepsin (5%, wt/wt) in 0.5 M acetic acid (adjusted to pH 2 with 6 M HCl), for 24 h at 37°C. The samples were clarified by centrifugation for measurement of fluorescence. More than 98% of the hydroxyproline content of the skin sample, 95% of which was digested with pepsin (5%, wt/wt) in 0.5 M acetic acid (adjusted to pH 2 with 6 M HCl), for 24 h at 37°C. The samples were clarified by centrifugation for measurement of fluorescence. More than 98% of the hydroxyproline content of the skin sample, 95% of which was described (9), was solubilized by this method. The fluorescence readings were normalized to the hydroxyproline content of the solution. For each analysis all samples were analyzed together in a single batch to avoid inter assay variations. All laboratory work on the samples was also done “blind”, i.e., with no knowledge of sample identity.

Statistics. Results are expressed throughout as mean ± 1 SD. Differences between groups before and after improved glycemic control were compared using a paired t test.

Results

Measurements of both MHBG and HbA1 (Fig. 2) show that there were significant improvements in glycemic control in the patient group during the course of this study. The overall differences and statistical analyses for these and other measurements discussed below are summarized in Table I. Fig. 3 shows the results of all the analyses for individual patients. It is apparent that, in addition to MHBG (Figs. 2 A and 3 A) and HbA1 (Figs. 2 B and 3 B), the extent of glycation of skin collagen also decreased significantly between the beginning and end of the study (Fig. 3 C and Table I). Glycation of collagen correlated significantly with MHBG (r = 0.65, P < 0.02) and HbA1 (r = 0.89, P < 0.0001) at the beginning, but not at the end, of the study. Similarly, MHBG during the 2-wk run-in period correlated significantly with initial HbA1 (r = 0.56, P < 0.05), but, as might be expected, this correlation was lost with improved glycemic control. Over the course of the study, the relative decrease in individual patient HbA1 (initial−final)/initial) correlated significantly with the relative decreases in MHBG (r = 0.68, P < 0.01) and FL in collagen (r = 0.72, P < 0.005). Thus, changes in mean blood glucose concentration were mirrored by changes in glycation of both hemoglobin and collagen. This is apparent in Fig. 3, A–C, which shows visually that the greatest reductions in collagen FL occurred in those patients with highest initial FL values and who achieved the greatest decreases in MHBG and HbA1.

As shown in Fig. 3, D–G and summarized in Table I, levels of CML, CMhL, pentosidine, and total fluorescence were unaffected by the 4-mo period of improved glycemic control. All of these parameters showed wide variations among individuals, reflecting differences in age (Fig. 3), as well as duration and severity of diabetes. There were no significant correlations of MHBG, HbA1, or collagen FL with CML, CMhL, pentosidine, or total fluorescence.

One other important observation made in this study is summarized in Fig. 4 and its legend. In diabetic patients, either at the beginning or end of the study, there was a strong correlation between any two of the measures of long-term chemical modification of collagen, i.e., nonenzymatic browning products (pentosidine), oxidation products (CML and CMhL), and total fluorescence. Thus, the four parameters measured in this study provide a consistent assessment of the extent of Maillard reaction damage to collagen.

Discussion

Collagen, in its various forms, is a ubiquitous protein in the body. Collagen abnormalities may therefore have widespread consequences, such as interference with basement membrane function in small vessels and glomeruli, or with the processes of growth and remodelling of tissues. Such problems are well recognized features of long-term diabetes. Increased glycation and nonenzymatic browning of collagen and other structural proteins by glucose is thought to be one of perhaps several mechanisms contributing to pathophysiological changes characteris-
tic of aging (31) and the development of the complications of diabetes (3, 5, 6, 16). The extent of glycation of long-lived proteins is directly related to ambient glucose concentration (26–28) and does not change significantly with age in the nondiabetic population (26–28, 32, 33). While a number of earlier studies have shown that glycation of collagen is increased in response to hyperglycemia in diabetes (8–10, 15) and have found a strong correlation between glycation of hemoglobin and collagen (9, 10), our work demonstrates that glycation of human skin collagen may be significantly reduced within as short a time as a 4-mo period of improved glycemic control. To our knowledge only one other study (34) has addressed the question of reversibility of glycation of collagen. In that case the investigators found no decrease in glycation of tail tendon collagen in diabetic rats treated with insulin for an 8-wk period. The failure to detect differences in collagen glycation in these animal experiments may be explained by the shorter period studied.

Because of the impracticality of maintaining constant glycemia and obtaining multiple skin biopsies from patients, it is difficult to estimate an exact half-time for reversal of glycation of human skin collagen. However, the results of this study suggest that the half-time for response of collagen glycation to improvements in glycemic control may be as short as 4 mo. This conclusion is drawn from analysis of individual percent decreases in HbA1 and collagen FL, normalized to the percent decrease required to achieve the means of the nondiabetic population (5.5% HbA1, and 4.6 mmol FL/mol Lys in skin collagen) (27). For example, the response in collagen glycation was calculated as 100 × (FLF − FL0)/(FLA − FL0), where FL0, FLA refer to the initial and final measurements of mmol FL/mol Lys in collagen, and FL0 = 4.6 mmol FL/mol Lys in collagen, the mean value for the nondiabetic population (27). By this analysis there was an average 45% decrease in HbA1 and 25% decrease in glycation of collagen (r = 0.68, P < 0.01) toward the nondiabetic means. The greater response in HbA1 is consistent with the increased rate of turnover of hemoglobin, compared to skin collagen. The 25% decrease in glycation of collagen within a 4-mo period indicates that the upper limit for the half-time for reversal of collagen glycation is ~ 8 mo. The actual half-time is undoubtedly shorter since none of the patients achieved normoglycemia instantaneously or maintained normoglycemia throughout the course of the study. For the six patients in poorest control at the beginning of the study (initial HbA1 > 12.0%, marked by an asterisk in Fig. 3 B), the average decreases toward the nondiabetic means during the course of the study were 66% for MHBG, 64% for HbA1, and 36% for glycation of collagen (see Fig. 3, A–C). These results suggest that if glycemia were completely and instantaneously normalized at the beginning of the study (i.e., 100% decrease to normal in MHBG), it should have been possible to achieve a 50% decrease towards normal in glycation of collagen within the 4-mo period of our study, suggesting an actual half-time closer to 4 mo for reversal of the excessive glycation of collagen. A more rigorous mathematical analysis cannot be justified because of the lack of information about the kinetics of collagen glycation and the mechanisms by which collagen FL content was decreased.

There are a number of possible explanations for the decrease in glycation of collagen during improved glycemic control. All assume a decrease in the rate of glycation in response to the fall in ambient glucose concentration, plus some reaction in which the existing glucose adducts are consumed. Assuming negligible turnover of insoluble skin collagen (35–38), one possible explanation for the decrease in glycation is that the reaction of glucose with lysine may be reversible, the hexose being released as glucose and mannose stereoisomers, regener-
ating the unmodified lysine residue. As reviewed in reference 39, there is firm evidence for reversal of the Amadori rearrangement with model Amadori compounds (24), but little information on the rate or extent of reversal of glycation of proteins in vivo. It is also possible that the Amadori adduct is consumed in a forward reaction, resulting, for example, in the release of the carbohydrate in a modified form, such as 1- or 3-deoxyglucone (40), again regenerating the unmodified lysine residue in collagen. The precision of our analyses do not permit us to determine whether the loss of FL in collagen is accompanied by an increase in the lysine content of the protein. Thus, it is also possible that the FL may be consumed in fragmentation and oxidation reactions, leading to formation of browning and oxidation products, including pentosidine, CML (C(MHL)), and other species. However, the further progress of these reactions should be limited by the decrease in the precursor, FL.

In contrast to FL, levels of CML, C(MHL), pentosidine, and total fluorescence in collagen did not respond to improved glycemic control within the period of this study. This is consistent with the fact that, to our knowledge, these compounds are stable and irreversible chemical modifications of protein; they are known to accumulate with age in long-lived proteins, such as lens crystallins (26) and the insoluble fraction of skin collagen (21, 27). Their constant concentration in collagen during the limited course of this study is also consistent with the metabolic inertness of insoluble human skin collagen (35-38), while the strong correlations between the concentrations of these compounds in collagen (Fig. 4) emphasize that they are all sensing the same chemical environment and stresses. While our results show that glycation of collagen may be reduced by improved glycemic control, they also indicate that browning and oxidation products formed during advanced stages of the Maillard reaction result in relatively permanent, perhaps irreversible, modification of the protein. These considerations add further weight to the argument that the establishment and maintenance of good glycemic control may inhibit the development of diabetic complications.

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