Diabetes-induced Functional and Structural Changes in Insulin Receptors from Rat Skeletal Muscle

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Abstract

The effect of diabetes on the structure and function of insulin receptors was studied in rats 7 d after streptozotocin injection, using solubilized, partially purified receptors from rat hindlimb muscles. Diabetes increased the number of insulin receptors per gram of muscle 60–70% without apparent change in insulin binding affinity. Incubation of receptors at 4°C with [γ-32P]ATP and insulin resulted in dose-dependent autophosphorylation of the β-subunit on tyrosine residues; receptors from diabetic rats showed decreased base-line phosphorylation, as well as a decrease in autophosphorylation at maximally stimulating insulin concentrations. These receptors also showed diminished exogenous substrate kinase activity using histone H2b and angiotensin II as phosphoacceptors. The electrophoretic mobility (sodium dodecyl sulfate-polyacrylamide gel electrophoresis) of a subpopulation of β-subunits derived from diabetics was slightly decreased; differences in electrophoretic mobility between control and diabetic-derived β-subunits were enhanced by generating fragments by partial Staphylococcus aureus V8 protease digestion. Endoglycosidase-H or neuraminidase treatment increased the electrophoretic mobility of β-subunits in both groups, but only neuraminidase appeared to decrease or abolish differences in electrophoretic mobility between controls and diabetics, suggesting that excess sialidation may account, in part, for the altered mobility of diabetic derived β-subunits. All structural and functional alterations in insulin receptors were prevented by treating diabetic rats with insulin for 60 h. Peripheral insulin resistance associated with insulinopenic diabetes may be related to modifications in insulin receptor structure, resulting in impaired signal transmission.

Introduction

Insulinopenic diabetes in humans is associated with peripheral insulin resistance (1–4), which appears to be related to prior glycemic control (5). Using a closed-loop subcutaneous insulin infusion system, Mayfield et al. (5) reported a direct correlation between elevated hemoglobin (Hb)A1c levels and increased insulin requirements in type I diabetic patients. Strict control of blood glucose levels increased the subjects’ sensitivity to insulin (5–7). Insulin resistance is also seen in animals with experimentally induced insulinopenia. Reaven et al. (8) observed insulin resistance in alloxan diabetic dogs when fasting plasma glucose levels exceeded 150 mg/dl whereas dogs with lower glycemia responded to insulin as the controls. In rodents, streptozotocin-induced diabetes results in a diminished response to insulin in isolated adipocytes (9–11), liver (12, 13), as well as skeletal (14) and heart muscle (15), whereas insulin binding (e.g., the apparent number of insulin receptors) is increased (9, 10, 13). This has led to the suggestion that the insulin resistance associated with poorly-controlled type I diabetes is primarily a postreceptor defect. Alternatively, it may represent a defect in signal transmission by the occupied insulin receptor.

The initial event in insulin action on target tissues is its binding to specific cell surface receptors. The insulin receptor is a heterotetramer glycoprotein, consisting of two α- and two β-subunits. The α-subunit with an apparent molecular weight of MWα ~ 135,000 appears to contain the insulin-binding domain, whereas the β-subunit MWβ ~ 95,000 may be primarily involved in signal transmission. Insulin binding induces the autophosphorylation of the β-subunit in both intact cells and in cell-free systems (16–23). After this autophosphorylation, the insulin receptor acquires the ability to act as a kinase toward exogenous substrates (18, 21, 23). These phosphorylation events occur on tyrosine residues and have been suggested to trigger the phosphorylation–dephosphorylation cascade seen in cells upon exposure to insulin (16, 20, 22).

Because skeletal muscle is an important site of insulin action, we investigated the effect of streptozotocin-induced diabetes on the functional and structural properties of the solubilized rat skeletal muscle insulin receptor. We found that the insulin-stimulated autophosphorylation and exogenous substrate kinase activity of diabetic-derived receptors was impaired. This alteration was reversed by in vivo insulin treatment. The functional changes were paralleled by the appearance of a subpopulation of insulin receptor β-subunits in diabetic skeletal muscle with a slower electrophoretic migration; the latter may reflect altered glycosylation of the β-subunit.

Methods

Materials: Streptozotocin was a gift from Dr. William Dulin, Upjohn Research Laboratories (Kalamazoo, MI). Polyclonal antis insulin receptor antibody (A410) was a gift from Dr. Steven Jacobs, Wellcome Research Laboratories (Research Triangle Park, NC) and from Dr. C. Ronald Kahn (B9), Joslin Research Laboratories (Boston, MA). Monocompo-

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1. Abbreviations used in this paper: DMSO, dimethyl sulfoxide; MWα β , apparent molecular weight; PAGE, polyacrylamide gel electrophoresis; PIC, protease inhibitor cocktail; PMSF, phenylmethylsulfonyl fluoride; TLC, thin-layer chromatography; TPCK, L-1-tosylamide-2-phenylethyl chloromethyl ketone; WGA, wheat germ agglutinin.
crystalline pork insulin was a gift from Dr. Ronald Chance, Lilly Research Laboratories (Indianapolis, IN). Monodiodinated A14 125I-insulin used in some studies was kindly provided by Dr. B. H. Frank, Lilly Research Laboratories. Wheat germ agglutinin (WGA) was from Vector Laboratories (Burlingame, CA); Aprotinin from FBA Pharmaceuticals (New York); Bactрин from Pfizer (New York). Electrophoresis reagents were from Serva (Heidelberg, Federal Republic of Germany) and the apparatus was from LKB Instruments, Inc. (Houston, TX). Protein A-bearing Staphylococcus aureus (Pansorbin) and neuraminidase (Vibrio cholerae) were from Calbiochem-Behring Corp. (San Diego, CA). S. aureus V8 protease and endoglycosidase H were from Miles Laboratories, Inc. (Elkart, IN). 125I-iode was from New England Nuclear (Boston, MA). \(\gamma\)-P32ATP was prepared from \(\gamma\)P32orthophosphate (ICN, Cambridge, MA) in Gammaprep vials (Promega Biotec, Madison, WI) as described by the manufacturer. Cyanogen bromide (CNBr)-Sepharose 4B beads were from Pharmacia Fine Chemicals (Piscataway, NJ). Thin-layer chromatography (TLC) plates, Polygram CEL300, were from Brinkmann Instrument Co. (Westbury, NY). Autoradiography film, X-Omat AR was from Eastman Kodak Co. (Rochester, NY). Lightening + intensifying screens were from Dupont Co. (Wilmington, DE). All other reagents and chemicals were from Sigma Chemical Co. (St. Louis, MO) unless otherwise specified.

**Animals.** Male Wistar rats (Charles River Breeding Co., Wilmington, MA) 100–140 g of initial body weight were used in all experiments. They were allowed food and water ad libitum. Diabetes was induced by i.p. injection of streptozotocin in citrate buffer, pH 4.5, 130 mg/kg of body weight, to overnight fasted animals. In insulin-reversal experiments, diabetic rats were initially treated with 2 U of regular and 2 U of protamine zinc insulin s.c. at 1700 h on the 5th day after streptozotocin and subsequently with 2 U of protamine zinc insulin at 0900 h and 1700 h the next 2 d for a total of five treatments. Glycerosuria was monitored by Test-Tape (Eli Lilly Co.) and was + in untreated diabetic rats, and trace, or negative in insulin-treated rats by the 2nd day of treatment. Plasma glucose levels were determined with a Beckman Glucose Analyzer 2 (Beckman Instrument Co., Palo Alto, CA) on blood samples obtained when the animals were killed. Diabetic rats with or without insulin treatment were studied on the 7th day after streptozotocin, and compared to control animals. Rats were killed by decapitation, and the hindlimb muscles were rapidly removed and frozen in liquid nitrogen. Insulin receptors from control and experimental animal groups were always processed on the same day and analyzed in the same experiments.

Solubilization and partial insulin receptor purification were performed as previously described (23). Briefly, the frozen muscles, ~5 g, were powdered and membrane components were solubilized by homogenization as a frozen slurry in 4 vol of 25 mM Hepes, pH 7.4, 5 mM EDTA, 1 TIU/ml aprotinin, 2 mM phenylmethylsulfonyl fluoride (PMSF), and 1% Triton X-100. The homogenate was centrifuged at 10,000 g for 10 min and the supernatant was stirred at room temperature for 1 h. The mixture was then centrifuged at 150,000 g for 90 min. The receptors were further purified by adsorption to and elution from agarose-bound WGA. Receptors were eluted from the WGA column with 25 mM Hepes, pH 7.4, 0.1% Triton X-100 containing 0.3 M N-acetylglucosamine (~1 mg/ml of original tissue weight) and used directly or further purified by precipitation with polyclonal antinsiulin receptor antibody. In some preparations 125I-insulin (~1.5 µCi) was added after homogenization of muscles with or without 50 µg/ml unlabeled insulin. Aliquots were taken at various times during the isolation procedure, and 125I-insulin specifically bound was determined by polyethylene glycol precipitation. Recovery and trichloroacetic acid (TCA) precipitability of the 125I-insulin were determined by counting aliquots in a γ-counter as described below.

**Insulin binding.** WGA eluate (50 µl) was incubated with 10,000 cpm 125I-insulin (sp act 125–175 µCi/µg, prepared by the lactoperoxidase method as described in Reference 24) at increasing concentrations of unlabeled insulin for 16 h at 4°C in a buffer containing 25 mM Hepes, 0.1% Triton X-100, 150 mM NaCl, 0.1% bovine serum albumin, and 100 U/ml Bactrin as described by Taylor et al. (25). The receptor and bound insulin were precipitated by the addition of 100 µl of 0.3% bovine γ-globulin and 300 µl of 25% polyethylene glycol and collected by centrifugation. The resulting pellet was washed with 300 µl of 12.5% polyethylene glycol and counted in a Beckman Instruments Gamma 4000 γ counter. Nonspecific binding (10–15%) was estimated as that occurring in the presence of 10 µg/ml unlabeled insulin. Binding was related to the protein concentration in the WGA eluate estimated by the Bradford reaction (26).

**Equalization of receptor number.** In all experiments that assessed functional and structural properties of solubilized insulin receptors derived from experimental and control animals, approximately equal insulin-binding activity was used in comparing groups. Because no change in insulin-binding affinity was apparent in receptors from diabetic animals (see Results), equal numbers of insulin receptors were estimated as follows: 125I-insulin binding per 50 µl of WGA eluate was determined in the presence of 1, 5, 25, 100, 1,000, and 10,000 ng/ml unlabeled insulin (the latter used as estimate of nonspecific binding after overnight incubation at 4°C or 45 min at 25°C). Ratios of the specifically bound counts in the experimental group vs. the control were determined at each insulin concentration, these ratios were averaged, and the mean value used as an estimate of relative number of insulin receptors. Individual values were within 10–15% of this mean. Appropriate volumes of the WGA eluate were then used in subsequent experiments. Identical ratios were obtained after overnight or 45-min incubation with 125I-insulin; the latter resulted in ~60–70% of the equilibrium binding obtained in the former. All dilutions of WGA eluate were made with 25 mM Hepes, pH 7.4, 0.1% Triton X-100. The presence of 0.3 M N-acetylglucosamine had no effect on the functional properties of the insulin receptor (Reference 18, and our data). Experimental groups were always compared to control insulin receptors prepared the same day from untreated animals. 

Insulin specifically bound to receptors in the wheat germ eluate was linear between 12.5 and 75 µl of the eluate assayed in a constant volume (r = 0.97 and 0.96 for control and diabetic receptor preparations, respectively). The 32P incorporated into the β-subunit of the insulin receptor, after exposure to maximally stimulating insulin concentrations (10−6) was also linear between 10 and 50 µl of the wheat germ eluate, (r = 0.98 and 0.97 for control and diabetic receptor preparations, respectively). These values were derived from three different pairs of receptor preparations (diabetic and control), with each point assayed in duplicate or triplicate.

**Receptor autophosphorylation.** 50 µl of WGA eluate from control animals and appropriately diluted receptors from diabetic or insulin-treated diabetic animals were incubated in 25 mM Hepes, pH 7.4, 0.1% Triton X-100, 5 mM MnCl2, 0.05% bovine serum albumin (buffer A) with increasing insulin concentrations in 75 µl of final volume for 45 min at 25°C. The reaction mixtures were then cooled to 4°C, and [γ-32P]ATP (10 µl ~ 15 µCi, 100 µM final concentration) was then added, and after 30 min at 4°C, the phosphorylation reaction was terminated by the addition of 28 µl four-times concentrated Laemmli’s sample buffer, heated at 100°C for 3 min and the proteins were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (see below). The β-subunit of the insulin receptor was localized by autoradiography, excised, and digested in 1 ml of 30% H2O2 at 60°C for 10 h, and the 32P content was determined by counting in 10 ml of Scintiverse II (Fisher Scientific Co., Norcross, GA) in a Beckman LS7000 scintillation counter. Insulin degradation by the receptor preparation during incubation at 25°C in buffer A was estimated by TCA precipitability of added 125I-insulin tracer; ~2% of added insulin became TCA-soluble after 2 h of incubation.

Exogenous substrate kinase activity was determined using angiotensin II as the phosphoacceptor, as described by Rosen et al. (18). Approximately 10–20 fmol insulin-binding activity was incubated with or without insulin at 25°C for 1 h in 225 µl of buffer A. Autophosphorylation was initiated by the addition of [γ-32P]ATP (50 µg, 100,000 cpm final concentration) and allowed to proceed for 30 min at 4°C. Aliquots (35 µl) were then added to 5 µl of buffer B or buffer A containing 2 mM angiotensin II and the phosphorylation was carried out at 4°C for 5 min, which was in the linear part of the velocity curve as determined in preliminary experiments. The reaction was stopped by the addition of 5 µl of 1.5% bovine serum albumin (BSA) and 50 µl 5% TCA. After centrifugation,
duplicate 40-μl aliquots were spotted onto phosphocellulose paper, washed, and counted as described (18).

Kinase activity of insulin receptors bound to insulin-agarose. Control receptors (120 μl) and equal binding activity of diabetic-derived receptors were diluted to a final volume of 320 μl with buffer consisting of 50 mM Hepes, pH 7.4, 0.1% Triton X-100, 1 μg/ml leupeptin, and 2 mM PMSF (buffer B); 50 μl of insulin-agarose beads or agarose-beads inactivated with hot dithiothreitol (27) were added and the mixture incubated for 1 h at 25°C. After centrifugation the pellet was washed twice with 1 ml of buffer B supplemented with 0.5 M NaCl and then with buffer B alone. The pellet was then resuspended in 75 μl of 50 mM Hepes, pH 7.4, 0.1% Triton X-100, 0.025% BSA, and 5 mM MnCl2 (phosphorylation buffer) supplemented with 750 μM ATP and incubated for 15 min at 25°C to induce maximal receptor autophosphorylation (27). The mixture was then diluted with 1 ml of salt-supplemented buffer B, centrifuged, and the pellet washed with 1 ml of buffer B, and was then resuspended in 75 μl of phosphorylation buffer, supplemented with 1 mg/ml histone H2b and 50 μM γ[32P]ATP (2.5 μCi/vial) and incubated for 4 min at 25°C. The reaction was stopped by the addition of hot four-times concentrated Laemmli's sample buffer and the sample was separated on SDS-PAGE and autoradiographed as described below. The bands corresponding to histone H2b were excised and the 32P was quantitated as described above for the insulin receptor β-subunit.

ATPase activity. Equal binding activities of control and diabetic derived wheat germ eluates were incubated as described for the autophosphorylation assays except that the specific activity of the ATP was decreased (∼6 μCi/ml, ATP final concentration 10–100 μM). At various time intervals (30 s–120 min) the reaction was stopped by the addition of 65 μl of 3 M perchloric acid, and after 5-min equilibration at 4°C, 200 μl of D2O were added and the protein was removed by centrifugation. The supernatant (300 μl) was neutralized with 60 μl 2 M KHCO3, clarified by centrifugation, supplemented with 2 μl of a 2 M solution of unlabeled carrier ATP, and 20 μl was analyzed on a Perkin-Elmer high performance liquid chromatograph (Perkin-Elmer Corp., Norwalk, CT) using a 5-cm C18 reverse-phase column. ATP was eluted isocratically in a buffer composed of 20 mM phosphoric acid (pH 6.5 with triethylammonium, 98%) and methanol (2%) at a flow rate of 2 ml/min (28). The first 1.5 min of the run (containing inorganic phosphate) and the ATP peak were collected separately and 32P counts per minute quantitated by liquid scintillation counting. These assays were performed in parallel with assays of insulin-receptor kinase activities. All kinase assays reported here were carried out under conditions where ATP degradation was <10%.

Cross-linkage of solubilized insulin receptors with 125I-insulin. Wheat germ eluates from control and diabetic-derived muscle extracts were dialyzed against 25 mM Hepes, pH 7.6, and concentrated approximately fourfold with Aquacide (Calbiochem-Behring Corp.). Equal binding activities were incubated for 15 h at 4°C with A14125I-insulin (0.5 μCi) with or without 10 μg/ml unlabeled insulin in 250 μl of final volume, then disuccinimidyl suberate was added in dimethyl sulfoxide (DMSO) to a final concentration of 2 mM at 4°C, for 15 min (29). After addition of four-times concentrated Laemmli's sample buffer, the mixture was boiled for 3 min. Electrophoresis and autoradiography were performed as described below.

Immunoprecipitation and partial proteolytic digestion of phosphor ylated insulin receptors: WGA eluate was incubated in the presence of 10⁻⁴ M insulin for 45 min at 25°C in buffer A (100 μl of final volume) and 10⁻²⁴P32P3ATP (10 μM, 10 μCi) added for an additional 10 min. The reaction was stopped by addition of 20 μl of 40 mM ATP, and 10 mM NaPO4, 2 mM PMSF, in buffer A. Antiinsulin receptor antibody (1:400 dilution) was added and incubated overnight at 4°C. Protein A bearing formalin-fixed S. aureus was added and incubated for an additional 2 h and the precipitated complex was washed three times with 25 mM Hepes, 0.1% Triton X-100. Partial proteolytic digestion of the immune complex was achieved by resuspending it in 50 mM sodium acetate buffer, pH 5 (10 μl) containing 75 μg/ml V8 protease and incubating for 5 min at 37°C. The reaction was stopped either by the addition of an equal volume twice-concentrated Laemmli's sample buffer or by the addition of 10 μl of protease inhibitor cocktail (PIC) I and 5 μl of PIC II (PIC I was composed of 1.5 mg/ml benzimidazole, 1.5 mg/ml leupeptin, 1.5 mg/ml benzyl arginine ethyl ester in 10 TIU/ml aprotinin; PIC II was composed of 1 mg/ml PMSF and 1 mg/ml antipain in DMSO). The protease-inhibited mixtures were then treated with glycosidases as described below.

Phosphoamino acid analysis. The phosphorylated, immunoprecipitated β-subunit was identified by autoradiography of the wet, unfixed gel after SDS-PAGE and solubilized by electrophoresis as follows: a 2×2-cm gel slice was placed in a dialysis bag containing electrophoresis buffer (1.5 M Tris, 58 mM glycine, pH 8.8), and thus immobilized between the glass plates of the vertical gel electrophoresis apparatus, was immersed in this buffer. One end of the sandwich was plugged with 15% acrylamide and a potential of 500 V was applied for 1 h. The contents of the bag were removed, 30 μg of bovine γ-globulin was added and the protein was precipitated by making the solution 20% in TCA. The precipitate was dissolved in 250 μl of 6N HCl and hydrolyzed at 110°C for 60 min under N2 atmosphere. The hydrolysate was adsorbed to 0.5 ml of Dowex 1×8, washed with 2 ml of dH2O and eluted with 1 ml 0.1N HCl (30). The lyophilized sample was redissolved in 5 μl of a mixed phosphoamino acid standard containing 0.33 mg/ml each of phosphoserine, phosphothreonine, and phosphotyrosine and spotted on thin cellulose TLC plates (0.1 mm thick). The plates were electrophoresed at 1,000 V for 40 min in pyridine/acetic acid/H2O (5:15:945) towards the anode (30). The phosphoamino acids were localized by ninhydrin staining and by autoradiography at −80°C with intensifying screen.

Tryptic phosphopeptide mapping. Electroleotuted receptors were precipitated at 4°C in 10% TCA with 30 μg of carrier bovine γ-globulin and the precipitate was extracted three times with 750 μl of 1:1 ether/ethanol (31). The washed pellet was oxidized at 25°C for 2 h by the addition of 200 μl of 9:1 formic acid/H2O2, diluted to 2 ml with water, and lyophilized. The lyophilized was resuspended in 0.3 ml of 50 mM (NH4)2CO3, pH 8.0, containing 75 μg/ml l-τosylamide-phenylthlely chloromethyl ketone (TPCK)-treated trypsin and incubated at 25°C for 12 h when an additional 40 μg of tryspin was added and incubation was continued for another 12 h. The mixture was brought to 3 ml with water and lyophilized three times, resuspending in 3 ml of water each time (31). Phosphopeptides were separated in two dimensions on 20 cm × 20 cm × 0.1 mm TLC plates by electrophoresis in 1% ammonium bicarbonate, pH 8.9, for 40 min towards the anode and then by ascending chromatography in pyridine/butanol/acetic acid/water (70:50:15:60) in the second dimension (32). Phosphopeptides were localized by autoradiography at −80°C for 4 h with intensifying screens.

Treatment of immunoprecipitated receptors with endoglycosidase H. Washed, immunoprecipitated receptors with or without prior partial proteolytic digestion were made 0.1% in SDS and heated for 1 min in boiling water. After cooling, 10 μl of PIC I and 5 μl of PIC II were added, followed by 50 μl of 30 μU/ml endoglycosidase H in 50 mM sodium acetate, pH 5.5, 10 mM CaCl2, 0.1% bovine serum albumin and incubated for 4 h at 37°C. The samples were boiled in Laemmli's sample buffer (four-times concentrated) and separated on SDS-PAGE.

Neuraminidase. Immunoprecipitated receptors with or without prior partial V8 digestion were resuspended in 10 μl of PIC I and 5 μl of PIC II, followed by the addition of 40 nU/ml neuraminidase in 50 mM sodium acetate, pH 5.0, 10 mM CaCl2, and 150 mM NaCl and were incubated 4 h at 37°C. As above the reaction was stopped by the addition of four-times concentrated Laemmli's sample buffer and proteins separated on SDS-PAGE. Methods for treatment of insulin receptors with glycosidases were modified from those described by Ronnett et al. (33).

Electrophoresis. SDS-PAGE. Immunoprecipitated proteins were separated on discontinuous buffer system of Laemmli (34) with a 7.5% polyacrylamide resolving gel for nonprotease-digested samples, a 9% polyacrylamide resolving gel for samples previously treated with V8 protease, and a linear 6–18% polyacrylamide resolving gel for reactions containing histone H2b. A 4.5% polyacrylamide stacking gel was used for all samples. Gels were electrophoresed at 150–170 V, constant voltage. Gels were stained, unless indicated, in 50% methanol, 10% acetic acid, and 0.125% Coomassie blue R and destained in 30% methanol and 10%
acetic acid. The gels were dried in vacuo and autoradiography was performed with Kodak AR autoradiography film with or without Dupont Lightening + intensifying screens at −80°C.

Results

Animal condition. Rats rendered diabetic by i.p. injection of streptozotocin gained little or no weight during the subsequent 7 d. Final weights of the diabetic rats were 111±4 g (n = 23) vs. 164±4 g (n = 15) for the controls. Insulin treatment (60 h) of diabetic rats resulted in rapid weight gain (final weight 145±5 g; n = 10). At the time of sacrifice the plasma glucose concentration of control rats was 130±4 mg/dl (n = 15), that of untreated diabetic rats was 510±14 mg/dl (n = 17), and that of insulin-treated diabetic rats was 94±6 mg/dl (n = 12).

Insulin binding. Equilibrium binding of 125I-insulin by the partially purified insulin receptors extracted from hindlimb muscles of diabetic rats increased approximately twofold per milligram of protein in the wheat germ eluate as compared to controls (Fig. 1). When calculated per gram of muscle that increase in receptor number was 60–70% (~100–125 pmol insulin bound/g in control vs. 180–250 pmol bound/g in the diabetic). Administration of insulin for 60 h caused a reduction in insulin receptor number per milligram of protein and per gram of tissue weight to nearly that of controls. The yield of milligrams of glycoprotein/gram of muscle from diabetic rats was ~25% less than that obtained from controls, 0.55±0.06 vs. 0.72±0.07, respectively (n = 10/group, P < 0.01, paired analysis). The increase in receptor number solubilized from muscles of diabetic rats occurred without an apparent change in the affinity of the insulin receptor. These values were obtained after incubation at 4°C for 16 h. Binding of 125I-insulin to control and diabetic-derived insulin receptors for 45 min at 25°C showed that 65–75% of equilibrium binding occurred in this time frame in both groups and the ratio of insulin bound by diabetic vs. control preparations was identical to that obtained with incubations for 15 h at 4°C. Because the kinetics of insulin binding were identical in diabetic and control animals, the tyrosine kinase activity of the solubilized insulin receptors was routinely studied after 45 min to 1 h of incubation with insulin at 25°C. Experiments performed after incubation with insulin for 16 h at 4°C yielded identical results to those shown below.

Insulin-stimulated autophosphorylation. Insulin stimulates the autophosphorylation of the insulin receptor through an intramolecular event (20, 35). As previously reported (23), when the WGA-purified insulin receptors from rat skeletal muscles were incubated with [γ-32P]ATP, 5 mM MnCl2, and increasing concentrations of insulin, there was an insulin dose-dependent increase in the amount of 32P incorporated into a ~97,000-mol wt peptide when the reaction mixture was analyzed by SDS-PAGE and autoradiography. This peptide was identified as the β-subunit of the insulin receptor by its MWapp, its ability to increase labeling after incubation with insulin, and its specific immunoprecipitability with antiinsulin receptor antibodies (see Reference 23).

When equal binding activities of control and diabetic-derived receptors were allowed to autophosphorylate at 4°C for 30 min in the presence of 100 μM ATP, there was a significant decrease (P < 0.05) in the amount of 32P incorporated into the β-subunit of the insulin receptor derived from diabetic animals incubated without insulin and at all insulin concentrations tested (Fig. 2). At maximally stimulating insulin concentrations, the mean autophosphorylation of control receptors was 40% greater than that of the diabetics; however, the insulin-stimulated autophosphorylation over base line (a insulin) was only 20% greater in controls than in diabetics. Although the latter difference was not significant, as will be shown below, it resulted in marked impairment of insulin stimulated exogenous kinase activity. Half-maximal activation of autophosphorylation occurred at ~2 × 10−8 M insulin in controls and diabetics. When diabetic rats were treated with insulin to normalize their blood glucose levels, the insulin receptors derived from these animals showed nearly identical autophosphorylation kinetics as controls (Fig. 2).

Exogenous substrate kinase activity: The insulin receptor displays exogenous substrate tyrosyl kinase activity that is activated by the autophosphorylation event (18, 21). To investigate this property insulin receptors from control, diabetic, or diabetic-treated animals were incubated for 1 h at 25°C with the indicated concentrations of insulin and then allowed to autophosphorylate in the presence of 100 μM [γ-32P]ATP at 4°C for 30 min; exogenous kinase activity was then assessed for 5 min at 4°C using angiotensin II as the phosphoacceptor. Little exogenous kinase activity was observed in any of the groups in the absence of added insulin. At 10−6 and 10−7 M insulin, a significant decrease in the exogenous substrate kinase activity was seen in the diabetic-derived receptors (P < 0.01, Fig. 3). The ability of insulin to stimulate the exogenous kinase activity over basal was significantly less in diabetic-derived receptors than in controls (P < 0.01). Treatment of diabetic rats with insulin restored the

Figure 1. Scatchard analysis of 125I-insulin binding to solubilized, WGA-purified insulin receptors. WGA eluate (50 μl) was incubated for 16 h at 4°C in a solution containing 25 mM Hepes, 0.1% Triton X-100, 150 mM NaCl, 0.1 mg/ml bovine serum albumin, and 100 U/ml bacitracin with 125I-insulin (10,000 cpn; 125–175 μCl/μg) and increasing concentrations of unlabeled insulin in a final volume of 200 μl. The receptor and bound insulin were precipitated by sequential addition of 100 μl of 0.3% bovine γ-globulin and 300 μl of 25% polyethylen glycol. Nonspecific binding was estimated as the radioactivity precipitated in the presence of 10,000 ng/ml unlabeled insulin. Specific binding was normalized to protein concentration in the wheat germ eluate estimated by the Bradford reaction (26). Control (c), diabetic (d), and diabetic-treated (a) muscle-derived receptor population. Points are means of three to five determinations from two separate experiments. Standard error of the mean of picomoles bound per milligram of protein ranged from 7% to 17% of the mean.
exogenous substrate kinase activity of the receptors toward normal (Fig. 3).

The decreased ability of diabetic-derived receptors to phosphorylate exogenous substrates was not temperature-dependent nor substrate-specific, in that similar decreases in the diabetic-derived receptor kinase activity were observed at 25°C using histone H2B as the phosphoacceptor. In these experiments, the wheat germ eluates were preincubated with insulin coupled to agarose, the agarose-bound receptors were extensively washed, and then allowed to fully autophosphorylate in the presence of 750 μM ATP. The exogenous kinase activity was then assessed using 50 μM [γ-32P]ATP. When equal insulin-binding activities derived from control and diabetic muscles were incubated with the beads, there was again a significant decrease in the amount of label incorporated into the histone by the diabetic-derived receptors as compared to the controls (Fig. 4). When receptors were bound to insulin-agarose beads that had been inactivated with hot dithiothreitol (27), histone-H2B phosphorylation was ~10% of that observed with fully activated agarose-bound receptors (data not shown).

A number of mechanisms may explain the decreased kinase activity of the diabetic-derived receptor population, including errors in the estimation of insulin receptor number or contaminants in the wheat germ eluate altering insulin receptor phosphorylation. Insulin that was "carried over" in the preparation of the insulin receptor could lead to an underestimate of the true insulin receptor number, especially in the control-derived preparations. This possibility was examined by the addition of 125I-insulin to the muscle homogenate and by observing its recovery during different stages of purification. We found that ≤0.1% of the added radioactivity was recovered in the final wheat germ eluate and that this was entirely nonspecific binding in that an identical homogenate which was supplemented with 50 μg/ml unlabeled insulin showed identical recovery of 125I-insulin. A second approach was to dialyze the insulin receptors exhaustively in the wheat germ elute to remove insulin. Upon re-binding, identical ratios of estimated insulin receptor numbers from control and diabetic animals were obtained as those before dialysis. Furthermore, no change in basal or insulin-stimulated autophosphorylation was observed in the dialyzed receptors.

The alterations in insulin receptor kinase activities did not appear to be due to contaminating insulinases in the diabetic-derived receptors nor to contaminating phosphatases because <2% of added 125I-insulin was degraded by the WGA eluates over a 2-h period and no change in the amount of 32P associated with the insulin receptor β-subunit or with angiotensin II or histone H2B was seen for 30 min after the addition of unlabeled ATP to previously phosphorylated insulin receptors and kinase substrates in either the control or diabetic derived preparations (data not shown). Although significant ATPase activity was de-
Figure 4. Autoradiogram of histone H2b phosphorylated by skeletal muscle-derived insulin receptors. Wheat germ eluates of 120 μl from controls and equal binding activity from diabetics were incubated with insulin-agarose conjugates for 60 min at 25°C as described in Methods. After washing, the insulin-agarose bound receptors were allowed to autophosphorylate in the presence of 750 μM ATP, 5 mM MnCl2, and 0.025% BSA for 10 min at 25°C, and then washed and incubated in phosphorylation buffer with 1 mg/ml of histone H2b and 50 μM [γ-32P]ATP (2.5 μCi/vial) at 25°C for 4 min. The reaction was stopped by the addition of four-times concentrated Laemmli's sample buffer and the samples were separated by SDS-PAGE and autoradiographed. Histone bands were excised, digested, and counted as described for the β-subunit in Fig. 2. Two separate sets of control and diabetic-derived receptors are shown. Lanes A and C, H2b phosphorylated by control receptors; lanes B and D, H2b phosphorylated by diabetic-derived receptors. Arrow indicates the position of H2b. (Inset) Mean±SEM of counts per minute of 32P incorporated into histone H2b from four different control and four diabetic-derived insulin receptor preparations (all with equal binding activity) each assayed in triplicate.

The results shown here were derived from assays, where <10% of [γ-32P]ATP was degraded.

Fig. 5 shows the phosphoamino acid analysis of control and diabetic-derived insulin receptor β-subunits that were maximally stimulated by insulin and then exposed to 10 μM [γ-32P]ATP for 10 min at 25°C. As with all other insulin receptors phosphorylated in vitro (18, 19), labeling occurred predominantly on tyrosine residues in both groups. Although a small amount of serine phosphorylation was observed, it contributed little to the total amount of phosphoamino acid present in either group.

Two-dimensional phosphopeptide maps generated from trypsin digestion of the β-subunit of the insulin receptors derived from control and diabetic animals and autophosphorylated at 25°C are shown in Fig. 6. Although slight variations occur in the relative labeling of the various phosphopeptides generated by this procedure, no consistent changes occurred in a number of maps analyzed, thus we found no indication that a site or sites are eliminated from the diabetic-derived population that would account for the decrease in overall labeling.

Structure of the insulin receptor subunits. Incubation of wheat germ eluates from control and diabetic animals with A14 125I-insulin with subsequent addition of disuccinimidyl suberate specifically labeled a protein of MWapp 131,000 (Fig. 7), which corresponds to the approximate molecular weight of the α-subunit of the insulin receptor. No differences were observed in the electrophoretic migration of the insulin cross-linked α-subunit between control and diabetic-derived insulin receptor populations. Furthermore, the amount of label bound per receptor was identical when diabetic- and control-derived wheat germ eluates were cross-linked with 125I-insulin.

In exploiting the property of autophosphorylation to localize the β-subunit of the insulin receptor, we investigated the possibility that the functional alterations observed in the insulin receptor from diabetic animals could stem from an identifiable structural alteration in this subunit. Insulin receptors were phosphorylated, immunoprecipitated, and separated under reducing conditions by SDS-PAGE. Autoradiography revealed that the β-subunit of the insulin receptor derived from diabetic animals included a component that showed a small but reproducible slowing in electrophoretic migration as compared to controls (Fig. 8, lanes A and B). This change in electrophoretic migration became more prominent after short proteolytic digestion of the phosphorylated, immunoprecipitated receptor with V8 protease (Fig. 8, lanes C and D). V8 protease treatment reproducibly generated two to four major phosphorylated species each of which showed this altered migration when derived from diabetic muscle.

The major phosphorylated peptides seen after V-8 digestion were shown to be derived from the β-subunit (as opposed to the variably phosphorylated α-subunit) in the following manner. Phosphorylated, immunoprecipitated insulin receptors were electrophoresed under reducing conditions and the unstained, unfixed gel autoradiographed at 4°C. The β-subunit was localized and excised, the gel slice placed in a dialysis bag and the phosphoprotein eluted as described in “Methods.” The dialyzed, ly-
ophilized protein was then subjected to partial V8 digestion, equal numbers of counts from each treatment group (~4,000 cpm) were loaded in each lane and reelectrophoresed. The autoradiograph, (Fig. 8, lanes F and G) shows the same major phosphorylated species as were generated by partial digestion of the immunoprecipitated intact receptor (lanes C and D) with the fragments generated from diabetic-derived β-subunits again showing a component with retarded migration when compared to controls.

As shown in Fig. 8, lanes E and H, the diabetes-associated alteration in electrophoretic mobility was reversed by 60 h of in vivo insulin treatment that normalized blood sugar. Lanes I and J show autoradiograms of control- and diabetic-derived insulin receptors that were immunoprecipitated and partially digested with V8 protease. Brief autoradiography revealed the diabetic derived β-subunit (lane J) to consist of two distinct populations, one that comigrates with the control receptor (lane I) and one with retarded migration.

We next investigated the possibility that sugar side chains associated with this glycopeptide could be altered in the diabetic state. Considering the difficulty in metabolically labeling the

Figure 6. Two-dimensional phosphopeptide analysis and autoradiography of insulin receptor β-subunit. Insulin receptors from control (A) and diabetic (B) animals were phosphorylated, immunoprecipitated, eluted, and TCA-precipitated as described in Fig. 5. The precipitated protein was washed with ether/ethanol (1:1), oxidized with formic acid/H2O2 (9:1), and digested with TPCK-treated trypsin in 50 mM (NH4)2CO3, pH 8, for 24 h. After removal of (NH4)2CO3 by lyophilization the hydrolysate was spotted on 20 × 20-cm TLC plates, electrophoresed for 40 min in the first dimension in 1% (NH4)2CO3, pH 8.9, and ascended chromatographically in the second dimension with N-butanol/pyridine/acetic acid/H2O (30:15:7:37.5). The dried plate was then autoradiographed for 4 d at −80°C with intensifying screens.

Figure 7. Structural analysis of the insulin receptor α-subunit. Control and diabetic-derived insulin receptors were dialyzed against 50 mM Hepes, pH 7.8, and concentrated with "Aquate," and equal binding activities were incubated for 16 h with 0.5 μCi A14125I-insulin with or without 10 μg of unlabeled insulin at 4°C. Disuccinimidyl suberate was added in DMSO to a final concentration of 2 mM and incubated for an additional 15 min when four-times concentrated Laemmli's sample buffer with 5 mM 2-mercaptoethanol were added. The samples were separated on 6% SDS-PAGE and the dried gel was autoradiographed. Lane A, control-derived receptors; lane B, diabetic-derived receptors; lanes C and D, control and diabetic-derived receptors cross-linked in the presence of 10 μg/ml unlabeled insulin.

Figure 8. Structural analysis of insulin receptor β-subunit. Wheat germ eluates with approximately equal insulin binding activities, prepared from muscles of control, diabetic, and diabetic-treated rats were incubated with 10−7 M insulin for 45 min and autophosphorylated for 10 min in the presence of 10 μM [γ-32P]ATP, (~15 μCi/100 μl). Receptors were immunoprecipitated with antiinsulin receptor antibody and Pansorbin. The precipitates were either boiled in Laemmli's sample buffer and analyzed by 7.5% SDS-PAGE under reducing conditions or first digested for 5 min at 37°C in 50 mM sodium acetate buffer, pH 5, with 75 μg/ml S. aureus V8 protease and then analyzed by 9% SDS-PAGE under reducing conditions. Dried, fixed gels were autoradiographed with intensifying screens at −80°C for 12–24 h. Lane A, control; lane B, diabetic-derived receptors without protease digestion. Lanes C, D, and E, receptors derived from control, diabetic, and insulin-treated diabetic rats, respectively, with prior V8 protease digestion. Lanes F, G, and H, β-subunits from control, diabetic, and diabetic-treated, respectively, that were isolated from a 7.5% gel by electrophoresis and subsequently digested with V8 as described in the text. Lane I, control; lane J, diabetic-derived receptors after V8 protease digestion, and short autoradiographic exposure.
sugar side chains and the minute amounts of insulin receptors available for chemical modification, we used enzymatic digestion of the insulin receptor with glycosidases in an attempt to characterize the structure of the glycosyl residues. Incubation of the immunoprecipitated insulin receptor with endoglycosidase H caused an \( \sim 5,000 \) decrease in \( MW_{app} \) in control- and diabetic-derived insulin receptors (Fig. 9a, lanes A, B and E, F) but did not correct (and actually accentuated) the difference between the control and diabetic. Endoglycosidase H digestion of the immunoprecipitated receptor after partial V8 proteolysis was accompanied by adding an excess of protease inhibitors before the addition of the endoglycosidase. Again a decrease in \( MW_{app} \) of \( \sim 4,000-5,000 \) occurred in the major phosphopeptides generated from control- and diabetic-derived \( \beta \)-subunits and allowed the separation of the diabetic derived receptors into two distinct populations (Fig. 9b, lanes E and F), one that comigrates with the control, and one with retarded electrophoretic migration. The decrease in \( MW_{app} \) of the \( \beta \)-subunit after endoglycosidase H treatment suggested that this glycoprotein contains "high mannose" sugar side chains.

Neuraminidase is an exoglycosidase that removes terminal sialic acid residues from the sugar side chains of glycoproteins. When the immunoprecipitated insulin receptor was digested with this enzyme, a decrease in \( MW_{app} \) of \( \sim 2,000 \) occurred in control lanes (Fig. 9a, lanes A and C). A similar decrease occurred in the diabetic-derived insulin receptor (Fig. 9a, lanes B and D); in addition, there was an apparent normalization in the electrophoretic migration of this population as compared to control (Fig. 9a, lanes C and D). Neuraminidase treatment caused a slight decrease in \( MW_{app} \) of the major (\( \sim 54,000 \)) phosphorylated peptide in the diabetic-derived population (Fig. 9b, lanes B and D), and nearly normalized the migration of the diabetic-derived phosphopeptides as compared to control (Fig. 9b, lanes C and D).

However, double glycosidase digestion, i.e., neuraminidase followed by endoglycosidase H, showed that the second, slower migrating band in the diabetic-derived preparation increased its electrophoretic migration but was not completely eliminated after neuraminidase treatment (Fig. 10, demonstrating the main \( \sim 54,000 \) \( MW_{app} \) phosphopeptide generated by V8 protease digestion). This result raises the possibility that an additional component, possibly "core" sugars, may contribute to the heterogeneity in electrophoretic migration of the diabetic-derived \( \beta \)-subunit.

**Discussion**

An increase in the number of insulin receptors associated with insulinopenia has been observed in adipocytes (9, 10), and hepatocytes (13, 36). In the present study the number of insulin receptors solubilized per gram of hindlimb muscle was increased 60-70% after 7 d of diabetes, without an apparent change in affinity. Insulinopenic diabetes in rats has been reported to cause no change (10, 36) or to increase the affinity of insulin receptors (10, 37) in adipocytes and liver. Studies of insulin binding to intact, incubated soleus muscles from streptozotocin diabetic mice reported findings of increased binding affinity without a change in receptor number (38). These discrepancies may reflect the difficulty in interpreting changes in insulin binding kinetics, especially when intact tissues are used, as well as possible differences between solubilized, total receptors assessed in the present study versus surface receptors on intact muscle cells studied in Reference 38. It should be noted that the increased number of receptors observed in muscles after 7 d of diabetes in this report reflects in part differences in growth rate and muscle weight between control and diabetic rats and the \( \sim 25\% \) decrease in the

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**Figure 9. Glycosidase digestion of phosphorylated, immunoprecipitated insulin receptors.** Autophosphorylated insulin receptors were immunoprecipitated and incubated for 5 min without (a) or with (b) V8 protease as described in Fig. 8. After addition of protease inhibitors, samples were incubated with buffer alone (lanes A and B) or with neuraminidase (40 mU/ml, lanes C and D) or with endoglycosidase H (30 mU/ml, lanes E and F) at 37°C for 4 h as detailed in Methods after which four-time concentrated Laemmli's sample buffer was added, the samples analyzed by SDS-PAGE and autoradiographed. Lanes A, C, and E, control-derived receptors; lanes B, D, and F, diabetic-derived receptors. The numbers on the left indicate the positions of molecular weight markers in thousands.

_Muscle Insulin Receptor Changes in Diabetes_ 267
amount of glycoprotein eluted from the wheat germ affinity column in the isolation of insulin receptors from diabetic rats as compared to controls.

The decrease in the ability of the diabetic-derived insulin receptors to incorporate $^{32}$P from [γ-$^{32}$P]ATP into its β-subunit and into exogenous substrates (angiotensin II or histone H$_3$B) was a consistent finding and does not appear to reflect biased estimates of insulin receptor number nor contaminating insulinase, phosphatase, or ATPase activities. Note that cross-linking experiments revealed identical labeling of α-subunits in diabetic and control receptor preparations (see Fig. 7). The decrease in autophosphorylation was presumably not due to a decrease in the number of β-subunits in the diabetic-derived wheat germ eluate in that similar binding kinetics were observed and removal of a significant proportion of the insulin receptor β-subunits would likely result in a noticeable change in receptor affinity. The impairment of insulin-stimulated exogenous kinase activity of diabetic-derived receptors (Fig. 3) was more evident than that of the insulin activated autophosphorylation (Fig. 2). Preliminary data (39) suggest that phosphorylation of different tyrosyl sites on the β-subunit may not be equally important for the activation of the exogenous kinase.

Our results are in essential agreement with those reported by Kadowaki et al. (36) who showed impaired autophosphorylation of the insulin receptors derived from livers of streptozotocin-diabetic rats and by the recent report of decreased autophosphorylation in skeletal muscle from rats rendered diabetic by gold thioglucose treatment (40). The latter report suggests that the defect in the insulin receptor kinase activity is secondary to the diabetic state and not due to insulinopenia per se because the rats in Reference 40 were hyperinsulinemic. However, Ama-truda et al. (41) isolated hepatic plasma membranes from diabetic rats and found no defect in insulin-stimulated receptor auto-phosphorylation. Whether or not the insulin receptor population that manifests diabetes related structural and functional alterations is actually present on the plasmalemma or represents an intracellular species requires further investigation.

The phosphopeptides generated by treating the isolated β-subunit with TPCK-treated trypsin appeared identical in the diabetic and control muscle-derived receptors. We consistently generated 10 or more phosphopeptides from immunoprecipitated, electroeluted β-subunits even after digesting for 72 h. Previous studies utilizing high performance liquid chromatography to separate the products of trypsin digestion reported the isolation of three to five phosphopeptides (27, 35). Whether the larger number observed in our system is due to incomplete digestion or to receptor heterogeneity or to the higher sensitivity of autoradiography after two-dimensional peptide separation is not clear. Visual analysis of these maps failed to reveal salient differences in the relative phosphorylation of these fragments between controls and diabetics. The receptors used in generating these maps were phosphorylated for 10 min at 25°C in order to maximize $^{32}$P incorporation for visualization of phosphopeptides. However, a more detailed, quantitative examination of the phosphopeptides separated from receptors autophosphorylated at lower temperatures or for shorter time periods at 25°C may reveal differences between the two populations.

The presence of a subpopulation of insulin receptor β-subunits in diabetic rat skeletal muscle with retarded migration on SDS-PAGE suggests that diabetes affects the processing of the insulin receptor in muscle. We were able to detect this subpopulation of receptors in some rats 24 h after streptozotocin injection and in most animals by 3 d. In some animals this slower migrating species comprised up to 50% of the phosphorylated β-subunit band detected by autoradiography. It was difficult to discern this altered population consistently on a 7.5% polyacrylamide gel; however, it became distinct after limited proteolytic digestion of the insulin receptor with V8 protease. This is likely due to the inverse logarithmic relationship between molecular weight and relative mobility on isocatic polyacrylamide gels resulting in improved resolution of smaller proteins, as well as the increased relative preponderance of the component with decreased mobility in the fragments. The changes in the insulin receptor β-subunit structure appear to be in response to insulopenia and/or resulting changes in the metabolic milieu rather than to toxic effects of streptozotocin in that the slower migrating component was reduced or eliminated by 60 h of insulin treatment which normalized blood sugar. It is interesting to note that ~10% of control animals studied demonstrated a small amount (most <10%) of a higher molecular weight subpopulation.

The glycoprotein nature of the insulin receptor has been demonstrated by direct labeling of the sugar moiety (33, 42) and indirectly by its ability to bind to lectins (43). Ronnet et al. (33) observed that the insulin receptor is synthesized as a single polypeptide chain, which is cleaved and subsequently glycosylated on both the α- and β-subunits. These intermediates are sensitive to endoglycosidase H and neuraminidase resistant. Salzman et al. (44) and Herzberg et al. (45) reported that the mature α- and β-subunits from 3T3-L1 adipocytes and IM-9 lymphocytes contain both endoglycosidase H-sensitive and -insensitive sugar side chains. The latter report also showed that endoglycosidase F could not remove all the sugar side chains from the β-subunit but treatment with a chemical deglycosylating agent resulted in an additional decrease in molecular weight suggesting the presence of O-linked side chains. The insulin receptor β-subunit isolated here from skeletal muscle of control or diabetic rats appeared to be sensitive to digestion by endoglycosidase H; on autoradiography the 95,000-band migrated at ~89,000 after endoglycosidase H treatment, suggesting that the β-subunit isolated from skeletal muscle also contains “high mannose” oli-
posaccharide side chains. The peptides generated after V8 digestion, especially the prominent 54,000 band was also endoglycosidase H-sensitive. The diabetic-derived β-subunit population separated into two distinct bands after V8 protease and endoglycosidase H treatment, one that comigrated with the control and one with ~1,500 higher MWapp (Fig. 8b). The further improvement in the resolution of the diabetes induced β-subunit populations after endoglycosidase H treatment likely reflects the elimination of oligosaccharides which are known to affect protein migration on SDS-PAGE (46). Treatment of the immunoprecipitated, insulin receptor populations with neuraminidase increased the migration of the diabetic-derived β-subunit more than that of the control, especially that of the prominent 54,000 peptide fragment (Fig. 9). This suggested that the heterogeneity of the diabetic-derived β-subunit was due in part to an increase in its sialic acid content. When V8-digested insulin receptors were treated with neuraminidase followed by endoglycosidase H, the more slowly migrating component of the ~54,000 fragment (seen in diabetic derived β-subunits) appeared to decrease in molecular weight whereas the “control band” did not change its migratory behavior (Fig. 10). This suggests that there may be an additional sugar side chain on some diabetic-derived β-subunits or an alteration in the structure of one or more normally occurring oligosaccharide side chains. We encountered difficulties similar to those of Herzberg et al. (45) in attempting to completely remove the oligosaccharide side chains from the β-subunit with endoglycosidase F or glucanase (broad spectrum N-linked glycosidases [Burant and Buse, unpublished observations] suggesting that the β-subunit may have O-linked sugar side chains. Streptozotocin diabetes results in decreased sialic acid and other carbohydrate content of major glycoproteins on hepatocyte membranes (47) as well as decreased activity of certain glycosyltransferase enzymes, without a detectable change in sialyltransferase activity (48). The insulin receptor, however, is spared from the diabetes-associated decline in hepatic membrane glycoproteins, because hepatic insulin binding and receptor number appear to increase in insulinopenic diabetes (13, 36). The mechanism for the selectivity in hepatic glycoprotein synthesis or degradation is not understood; to our knowledge there is no information concerning the effect of diabetes on glycoprotein turnover or sialyltransferase activity in muscle. We did, however, see in diabetic rats a decline in the amount of glycoprotein per gram of muscle eluted from the wheat germ columns in the insulin receptor purification protocol.

In our studies, there appeared to be a relationship between the appearance of the β-subunit population with decreased electrophoretic mobility and changes in tyrosyl kinase activity, whenever one phenomenon was detected, so was the other. It is tempting to speculate that the metabolic alterations associated with insulin deficiency result in conformational changes in the receptor that renders the coupling between insulin binding and autophosphorylation less efficient. The precise relationship, if any, between the observed alteration in the kinase kinetics in the diabetic-derived insulin receptor and the structural heterogeneity of the β-subunit remains to be defined. Yamaguchi et al. (49) recently reported that neuraminidase treatment of highly purified placental insulin receptors increased insulin binding and tyrosine kinase activity. This observation suggests that alterations in sugar side chains may indeed modulate insulin receptor function.

The insulin resistance which accompanies poorly controlled insulinopenic diabetes involves both the liver and peripheral tissues; skeletal muscle has been implicated as the primary site (4). The work of Kadowaki et al. (36) in liver and the present work in skeletal muscle suggest that diabetes-induced changes in the β-subunit of the insulin receptor may play a role in this insulin resistance. In recent years there has been increasing awareness that insulin resistance associated with different pathologic conditions is caused to variable degrees by a defect in the binding of insulin to its targets, or to defects which occur after insulin binding. The latter abnormalities are generally referred to as postreceptor defects, with the tacit implication that they are distal to the receptor, involve the intracellular metabolic milieu, second messengers or their intracellular targets. Our data as well as those of others (36, 39), including a rare genetic defect of the insulin receptor (50, 51) suggest that postbinding defects may include “intrareceptor” defects, resulting in defective signaling by the receptor in response to hormone binding.

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