Kir6.2 Variant E23K Increases ATP-Sensitive K⁺ Channel Activity and Is Associated With Impaired Insulin Release and Enhanced Insulin Sensitivity in Adults With Normal Glucose Tolerance

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OBJECTIVE—The E23K variant in the Kir6.2 subunit of the ATP-sensitive K⁺ channel (KATP channel) is associated with increased risk of type 2 diabetes. The present study was undertaken to increase our understanding of the mechanisms responsible. To avoid confounding effects of hyperglycemia, insulin secretion and action were studied in subjects with the variant who had normal glucose tolerance.

RESEARCH DESIGN AND METHODS—Nine subjects with the E23K genotype K/K and nine matched subjects with the E/E genotype underwent OGTTs. Functional studies of the wild-type and E23K variant genotyped for the E23K variant also underwent graded glucose infusion, and hyperinsulinemic-euglycemic clamp with stable-isotope–labeled tracer infusions to assess insulin secretion, action, and clearance. A total of 461 volunteers consecutively genotyped for the E23K variant also underwent OGTTs. Functional studies of the wild-type and E23K variant potassium channels were conducted.

RESULTS—Insulin secretory responses to oral and intravenous glucose were reduced by ~40% in glucose-tolerant subjects homozygous for E23K. Normal glucose tolerance with reduced insulin secretion suggests a change in insulin sensitivity. The hyperinsulinemic-euglycemic clamp revealed that hepatic insulin sensitivity is ~40% greater in subjects with the E23K variant, and these subjects demonstrate increased insulin sensitivity after oral glucose. The reconstituted E23K channels confirm reduced sensitivity to inhibitory ATP and increase in open probability, a direct molecular explanation for reduced insulin secretion.

CONCLUSIONS—The E23K variant leads to overactivity of the KATP channel, resulting in reduced insulin secretion. Initially, insulin sensitivity is enhanced, thereby maintaining normal glucose tolerance. Presumably, over time, as insulin secretion falls further or insulin resistance develops, glucose levels rise resulting in type 2 diabetes. Diabetes 58:1869–1878, 2009
diabetes in subjects with the E23K variant. Since hyperglycemia can cause defects in insulin secretion, we focused on subjects with normal glucose tolerance. This allowed us to define the changes in insulin secretion and action that antedate diabetes onset and are due to the effects of the E23K variant per se. To resolve questions regarding the underlying mechanisms, we performed comprehensive assessments of the effect of E23K on K-ATP channel activity.

RESEARCH DESIGN AND METHODS

Nondiabetic subjects, aged <65 years in good health and with stable weight for 6 months, were recruited using advertisements. The studies were approved by the Human Research Protective Office.

Study 1 (intensive metabolic studies). Nine subjects with the K/K genotype and nine age-, sex-, and BMI-matched subjects with the E/E genotype participated in three separate protocols designed to test insulin secretion, action, and clearance. All subjects were unrelated and had no family history of type 2 diabetes. Insulin secretion was assessed using the 5-h oral glucose tolerance test (OGTT) and graded glucose infusion (GGI). Insulin sensitivity was assessed during the hyperinsulinemic-euglycemic clamp and OGTT. Insulin clearance was assessed during OGTT and GGI.

Study 2 (cross-sectional study). A total of 461 volunteers who responded to advertisements were genotyped for the E23K variant and underwent 5-h OGTTs for assessments of insulin secretion, action, and clearance.

Study protocols. Subjects adhered to their regular diet and refrained from exercise for 3 days before the studies.

5-h OGTT. Participants ingested a 75-g glucose load. Blood samples were collected at 0, 0.5, 1, 1.5, 2, 3, 4, 5, and 60 min after ingestion to determine plasma glucose, insulin, and C-peptide concentrations. GGI involved the intravenous administration of glucose at increasing rates (1, 3, 5, 7.5 g/kg body wt) and was continued for 3.5 h. Insulin was infused at 40 mU/m² per min. Dextrose (20%) with 6,6-²H₂ glucose (40 mmol/l) was added to maintain plasma glucose concentration at ~5.6 mmol/l. The infusion of ²H₂ glucose was decreased by 75% of basal during the OGTT and GGI. The slope of the line relating these two variables provided a measure of the slope of the line relating these two variables provided a measure of the dose-response relationship between ISR and glucose over the physiological range.

Whole-body insulin sensitivity was estimated using the oral glucose minimal model (25,26). Composite insulin sensitivity index was calculated using the 50% of basal during the OGTT and GGI. The slope of the line relating these two variables provided a measure of the dose-response relationship between ISR and glucose over the physiological range.

In vitro studies of the E23K variant. Recombinant K_ATP channel Kir6.2 (mouse) + SUR1 (hamster) was transiently expressed in COS7 cells as described (28). Of the missense variants reported in humans, the Kir6.2 clone contained only variants, while the SUR1 clone contained the Kir6.2 substitution.

Statistical analyses. Group differences were compared using the Student’s t test for unpaired samples for continuous variables and the Fisher’s exact test for categorical variables. Where appropriate, we used ANOVA and Tukey’s test for post hoc analyses for continuous variables and χ² test for categorical variables. ANCOVA were used to adjust for age, race, and BMI. SPSS version 15.0 (SPSS, Chicago, IL) was used for all analyses. A P value <0.05 was considered to be statistically significant. Results are reported as means ± SE.

RESULTS

Insulin secretion in subjects with E23K variant. We initially examined insulin secretion in subjects homozygous for the E23K variant, since they might exhibit the greatest difference if E23K genotype affected insulin secretion. The two groups were well matched for age, sex, and BMI (Table 1). Both groups had normal glucose concentrations in the fasting state and following oral and intravenous glucose administration (Table 1) (Fig. 1). The respective AUCs for glucose between E/E and K/K subjects during the OGTT (17.6 ± 0.1 vs. 18.7 ± 0.1 × 10⁵ min · mol/l) were compared using the Student’s t test. The results showed no significant difference between the groups (P = 0.29).
TABLE 1
Characteristics of study participants (study 1)

|          | E/E   | K/K   | P value |
|----------|-------|-------|---------|
| n        | 9     | 9     | 1.0     |
| Women (%)| 6 (67)| 6 (67)| 1.0     |
| Caucasian (%)| 9 (100)| 9 (100)| 1.0     |
| Age (years) | 45.9 ± 2.8 | 46.6 ± 3.9 | 1.0     |
| Weight (kg) | 75.5 ± 5.7 | 72.9 ± 4.2 | 0.73    |
| Height (cm) | 166.1 ± 2.5 | 164.2 ± 3.9 | 0.69    |
| BMI (kg/m²) | 27.8 ± 1.3 | 26.4 ± 1.6 | 0.53    |
| Fat mass (kg) | 31.1 ± 6.1 | 22.0 ± 3.0 | 0.20    |
| Fat-free mass (kg) | 43.7 ± 3.6 | 49.2 ± 3.7 | 0.30    |
| Truncal fat (kg) | 13.4 ± 1.8 | 10.0 ± 1.7 | 0.20    |
| Fasting glucose (mmol/l) | 5.1 ± 0.2 | 5.2 ± 0.1 | 0.73    |
| 2-h glucose (mmol/l) | 6.9 ± 0.3 | 6.8 ± 0.3 | 0.62    |
| A1C (%)     | 5.6 ± 0.1 | 5.5 ± 0.2 | 0.72    |

Data are means ± SE.

mmol⁻¹·l⁻¹ and GGI (17.6 ± 0.1 vs. 17.6 ± 0.1 × 10⁴ min⁻¹·mmol⁻¹·l⁻¹) were not different (Fig. 1). By contrast, the AUC for insulin in the OGTT (52.6 ± 5.7 vs. 71.0 ± 7.0 × 10³ min⁻¹·pmol⁻¹·l⁻¹) and GGI (24.5 ± 2.1 vs. 36.1 ± 4.8 × 10³ min⁻¹·pmol⁻¹·l⁻¹) was ~40% lower (P < 0.05) in the K/K group. The AUC for C-peptide was also reduced (P < 0.05) in the K/K subjects in both the OGTT (30.2 ± 2.7 vs. 58.9 ± 15.1 × 10⁴ min⁻¹·pmol⁻¹·l⁻¹) and GGI (26.4 ± 2.1 vs. 36.9 ± 3.3 × 10⁴ min⁻¹·pmol⁻¹·l⁻¹). Static, dynamic, and overall β-cell response to glucose were also ~40% lower (P < 0.05) in the K/K group (Fig. 2A) based on the OGTT and GGI. In addition, the dose-response curve relating glucose and ISR during the GGI was shifted downward and to the right in the K/K group (Fig. 2B). Accordingly, the ISR AUC (17.2 ± 1.2 vs. 26.1 ± 0.5 × 10⁴ pmol/l) and mean ISR (70.6 ± 4.9 vs. 106.7 ± 9.9 pmol·l⁻¹·min⁻¹) were lower (P < 0.05) in the K/K group. No significant differences were observed in insulin clearance rates in this study (P > 0.05).

Alterations in ATP sensitivity of reconstituted E23K channels in vitro. To examine the molecular basis for the differences in insulin secretion, we transiently expressed Kir6.2 with residue 23, being either glutamate (E23) or lysine (K23), together with the SUR1 subunit and measured ATP sensitivity in excised membrane patches (Fig. 3A). Homomeric K23 channels (K/K) exhibit a modest, yet significant, decrease in ATP inhibition compared with homomeric E23 (E/E) channels (K₁/₂,ATP = 16 μmol/l [n =

FIG. 1. Plasma glucose insulin and C-peptide concentrations during the oral glucose tolerance test (A) and intravenous GGI (B) in the K/K (○) and E/E (●) groups. P values indicate the significance of the differences between AUC values between groups. Values are means ± SE.
cells expressing E/E or K/K channels (201 ± 59 and 128 ± 50 channels/patch, respectively; n = 11–19 patches). To recapitulate the heterozygous E23K genotype (E/K), cells were transfected with a 1:1 mixture of E/E and K/K cDNAs. Since four subunits generate the channels, five different ratios of subunits will be present in the resultant channels (1 of 16 channels will be homozygous E/E and homozygous K/K). The ensemble of expressed channels display intermediate ATP sensitivity (K_{1/2,ATP} = 10.0 μmol/l [n = 10 patches] compared with homomer K/K and E/E channels) (Fig. 3B). Our data are similar to those reported by Schwanstecher et al. (5) for recombinant K/K channels. For comparison, the dose-response curves are shown in Fig. 3B for two Kir6.2 mutations that impair insulin release and underlie NDM. The I182V mutation underlies a transient NDM, whereas the I296L mutation underlies a syndromic form of NDM (32,33). Importantly, the ATP sensitivities of mutant channels correlate with the severity of the disease (K_{1/2,ATP} = 39 μmol/l for homomer I182V [n = 17 patches] and 771 μmol/l for homomer I296L channels [n = 5 patches]).

**Increased cellular activity of reconstituted E23K channels.** K_{ATP} channel activity in metabolically intact cells was screened by 86Rb^{+}-efflux from transfected COSM6 cells. Efflux from cells transfected with homomeric E/E channels was low under basal conditions and was activated by metabolic inhibition to lower cellular [ATP]/[ADP] (Fig. 4A). Rb^{+}-efflux from cells transfected with recombinant K/K channels was also low under basal conditions and increased with metabolic inhibition. Quantitative estimation of K_{ATP} channel conductance (see research design and methods) indicates that for K/K channels, basal conductance was a higher fraction of fully activated conductance than E/E channels (Fig. 4B). In the β-cell, the increased basal flux is expected to impair glucose sensing and account for the association of K/K genotype with reduced insulin secretion.

**E23K variant decreases ATP sensitivity by stabilizing the open state of the channel.** Kir6.2 mutations can reduce ATP sensitivity by directly reducing ATP binding to the Kir6.2 subunit or indirectly by affecting the intrinsic opening ability (28,29,34). In the latter case, an increase in the open probability (P_o,zero) decreases the frequency with which the channel enters the ATP-accessible closed state, resulting in a decrease in ATP sensitivity. We have modeled this nonlinear relationship between P_o,zero and K_{1/2,ATP} (29), and this kinetic model describes the diabetes-causing effects of Kir6.2 mutations that underlie NDM (e.g., Q52R, I296L) (Fig. 5C). To estimate the open probability of K_{ATP} channels, both nonstationary noise analysis (NA) and phosphatidylinositol biphosphate (PIP2) application were used to independently examine K_{ATP} channel gating in isolated membrane patches. As shown in Fig. 5A, the estimated open probability (P_o,zero) of K/K channels in the absence of ATP (0.68 ± 0.04 [NA method]; 0.67 ± 0.08 [PIP2 method]) is higher than that of E/E channels (0.49 ± 0.05 [NA method]; 0.46 ± 0.06 [PIP2 method], P < 0.05), and this increase can fully account for the shifted ATP sensitivity (K_{1/2,ATP} = 9.4 ± 1.6 μmol/l [E/E] and 21.6 ± 3.3 μmol/l [K/K] from curve fit of individual membrane patches, P < 0.01) (Fig. 5B). A similar increase in open probability was reported for K/K channels by Schwanstecher et al. (5), using analysis of single-channel records. That multiple methodologies reiterate the same findings strengthens the conclusion that the changes associated with the E23K variant are significant and real. As with
more severe NDM, the E23K variant indirectly affects ATP sensitivity by increasing the $P_{o,zero}$ (Fig. 5C). The predicted consequence will be reduced excitability of the $\beta$-cell, with increasingly severe consequences for insulin secretion (E23K < Q52R < I296L).

**Sulfonylurea sensitivity is reduced in E23K channels.**

Mutations in Kir6.2 that allosterically decrease ATP sensitivity by stabilizing the open state of the channel also reduce high-affinity block by sulfonylureas, a feature of NDM-associated mutations (28). We next examined the

![Graph 1](image1.png)

**FIG. 3.** Reduced ATP and sulfonylurea sensitivity of mutant E23K channels. A: Representative currents (at −50 mV) from inside-out membrane patches from COS cells expressing KATP channel (Kir6.2 + SUR1): homomeric E23 channels (E/E), K23 channels (K/K), or heteromeric E23 and K23 channels (E/K). Patches were exposed to differing [ATP], and baseline current was determined by exposure to ATP (5 mmol/l). B: Steady-state dependence of membrane current on [ATP] (relative to current in zero ATP [Irel]) for E23- and K23-containing channels. $K_{1/2\text{ ATP}}$ = 7.5 μmol/l (E/E) and 16 μmol/l (K/K). Data points represent means ± SE (n = 24–28 patches). The fitted lines correspond to least-squares fits of a Hill equation (see RESEARCH DESIGN AND METHODS). **P < 0.01 vs. E/E channels by unpaired Student’s t test (two tailed assuming equal variance). C: Representative currents recorded from inside-out membrane patches containing homomorphic E/E or mutant K/K channels at −50 mV and in response to varying [tolbutamide]. Zero-channel current was determined by application of ATP (5 mmol/l). D: Steady-state dependence of current on [tolbutamide] (relative to current in zero tolbutamide [Irel]) for E/E (○) and K/K (●) variant channels (from records such as those shown in C). Data points represent the means ± SE (n = 6–19 patches). For all channels, the lines are fits of the product of two Hill components, each of the form ($I_{\text{rel}} = I_0/(1 + ([\text{Tolb}] / K_{1/2}))$, with $H$ fixed at 1.3 in each case (see RESEARCH DESIGN AND METHODS). The relative fraction and $K_{1/2}$ values of each component were varied. The high-affinity component was 53 and 44% for wild-type and K/K channels, respectively. *P < 0.05 vs. wild-type KATP channel by unpaired Student’s t test. The shaded box shows the reported range of serum tolbutamide concentrations from a cohort of 37 type 2 diabetic subjects receiving sulfonylurea therapy (49).

![Graph 2](image2.png)

**FIG. 4.** Increased basal activity in intact cells expressing K/K variant channels. A: Representative efflux of $^{86}$Rb$^+$ as a function of time in basal conditions or in the presence of metabolic inhibition for reconstituted E/E and variant K/K channels and untransfected controls. B: Ratio of KATP channel–dependent efflux rate constant (k2) in basal conditions relative to metabolic inhibition (MI) for E/E or homomorphic K/K channels (see RESEARCH DESIGN AND METHODS). Graphs show compiled data (means ± SE) from six experiments in which each transfection was done in triplicates. *P < 0.05 vs. E/E channels by paired one-tailed Student’s t test.
**Insulin sensitivity in subjects with the E23K variant.** The clinical findings described above (of reduced insulin secretion with normal glucose concentrations) suggest a simultaneous change in insulin sensitivity. Hyperinsulinemic-euglycemic clamp experiments revealed that hepatic insulin sensitivity was significantly greater (2.4 ± 0.4 vs. 1.5 ± 0.2 [1,000/µmol · kg fat-free mass⁻¹ · min⁻¹ · pmol⁻¹ · l⁻¹]; *P < 0.05) in subjects with the K/K genotype (Fig. 6A). There was also a strong trend (0.19 ± 0.03 vs. 0.14 ± 0.02 [1,000/µmol · kg fat-free mass⁻¹ · min⁻¹ · pmol⁻¹ · l⁻¹]; *P < 0.10) for an increase in peripheral insulin sensitivity, although the differences were not statistically significant. Similar trends were observed for whole-body insulin sensitivity as assessed from the OGTT (Fig. 6B). Glucose infusion rate, plasma glucose, insulin, glucagon, and tracer-to-tracee ratio are presented in the online appendix (available at http://diabetes.diabetesjournals.org/cgi/content/full/db09-0025/DC1).

**Confirmation of changes of insulin secretion and action.** The human studies described above were conducted in a small group of E/E or K/K subjects. To confirm the findings on insulin secretion and action in a larger cohort, we carried out 5-h OGTTs in 461 additional subjects (Table 2). Subjects were divided into three groups based on E23K genotype. The three groups had similar glucose concentrations during fasting and 2-h post-glucose challenge. The relative frequencies of E/K (40%) and K/K (13%) in our sample are comparable with those reported (7,8,35), conferring a relative diabetes risk of 1.15–1.65 (3). Consistent with the above findings, the K/K subjects had reduced insulin and C-peptide concentrations, reduced insulin secretory responses, reduced β-cell responsiveness to glucose, and enhanced insulin sensitivity and clearance rates. Interestingly, β-cell responsiveness was also lower in the E/K variant than the E/E.

**DISCUSSION**

**Association of Kir6.2 E23K variant with impaired insulin secretion.** Type 2 diabetes is a multifactorial disease in which genetic and environmental factors interact to determine the level of predisposition, and, recently, genome-wide association studies have identified common polymorphisms in various genes that are associated with increased diabetes susceptibility (2,36–39). These studies often involve the comparison of genetic and phenotypic characteristics of diabetic and control subjects. They have made important contributions to identifying novel genetic loci that determine diabetes risk. However, studies of subjects with diabetes do not provide information on the effects of the respective polymorphisms prior to the onset of glucose intolerance and diabetes. Since hyperglycemia, per se, adversely affects insulin secretion, it is difficult to differentiate between the effects of hyperglycemia and the effects of the polymorphism in studies of established diabetes.

The present study was focused on subjects without a prior history of diabetes who had normal glucose tolerance. Our results demonstrate that the Kir6.2 E23K variant is associated with multiple insulin secretory defects, including a reduction in overall β-cell responsiveness to low-affinity (IC₅₀ = 2 mmol/l) site. The fractional block by the high-affinity therapeutically relevant component is ~53%. For the K/K channels, the value for the high-affinity block (IC₅₀ = 1.0 µmol/l) is unaltered; however, the fractional block is significantly decreased from 53 to 44%.

**Effect of the first-generation sulfonylurea, tolbutamide, on channel activity (Fig. 3C).** As shown in Fig. 3D, homeric E/E channels exhibit a typical biphasic block by tolbutamide, with both a high-affinity (IC₅₀ = 1.1 µmol/l) and a

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**FIG. 5.** An increase in maximum open probability underlies the reduced ATP sensitivity of K/K variant channels. A: Open probability in zero ATP (P₀ zero) calculated using the PIP₂ method and noise analysis (NA) (see research design and methods) for membrane patches expressing either homeric E/E or K/K channels. Data points represent means ± SE (n = 21–44 patches for NA method; n = 6–8 patches for PIP₂ method). *P < 0.05 and **P < 0.01 by two-tailed Student’s t test assuming equal variance. B: Relationship between P₀ zero (calculated using the NA method) and ATP sensitivity (K1/2 ATP) for individual membrane patches expressing E/E or K/K channels together with averaged data (triangles). For E/E: K1/2 ATP = 9.4 ± 1.6 µmol/l, P₀ zero = 0.49 ± 0.05 (n = 18 patches). For K/K: K1/2 ATP = 21.6 ± 3.3 µmol/l, P₀ zero = 0.68 ± 0.04 (n = 18 patches), **P < 0.01 for both K1/2 ATP and P₀ zero values of E/E compared with K/K channels (unpaired Student’s t test). Data points represent means ± SE. C: Relationship between P₀ zero and K1/2 ATP (mmol/l). Solid line represents prediction of kinetic model II (inset) of Enkvetchakul et al. (29). Symbols represent data points as in B together with mean values for Q52R and I296L channels. The key feature of the model is that ATP acts by binding to a closed state and in consequence ATP sensitivity is reduced by shifting the equilibrium between the Cᵢ and O states toward the open state (increasing KCO).
glucose and a rightward shift in the dose-response curve between glucose and insulin secretion. Overall insulin secretory responses to glucose were reduced by ~40%, and the extent of the reduction was similar to both oral and intravenous glucose.

**Association of E23K variant with enhanced insulin sensitivity.** We did not anticipate that severe reductions in insulin secretion would be compatible with normal glucose tolerance. The significant increase in insulin sensitivity is presumably the mechanism that makes this possible, and the results of the hyperinsulinemic-euglycemic clamp suggest that the major effects are in the liver. This is the first study to report the coexistence of decreased insulin secretion with increased insulin action in subjects with the E23K polymorphism. Previous studies that evaluated insulin action yielded inconsistent results. One study (11) suggested an increase in insulin sensitivity, while others (8,12,13) found no effect. Moreover, findings from the hyperinsulinemic clamp suggested an increase in peripheral (skeletal muscle) insulin sensitivity, although this did not reach statistical significance ($P < 0.10$) because of the small sample size. This interpretation is supported by the results obtained in the larger cohort study in which statistically significant increases in insulin sensitivity were observed after oral glucose ingestion. Although BMI was different between groups, measurements of total and regional fat (truncal fat) showed no differences. Nevertheless, because of the differences in BMI, we controlled for BMI using ANCOVA. The differences in insulin sensitivity remained significant ($P < 0.001$), despite similar and normal glucose tolerance.

**TABLE 2**
Characteristics of study participants (study 2)

| Kir6.2 genotype | Kir6.2 genotype | Kir6.2 genotype | Kir6.2 genotype | $P$ value |
|-----------------|-----------------|-----------------|-----------------|-----------|
| E/E             | 216             | 184             | 61              | 0.40      |
| Women (%)       | 158 (73)        | 120 (65)        | 45 (74)         | 0.001     |
| Caucasian (%)   | 127 (59)        | 155 (84)        | 55 (90)         | 0.001     |
| Age (years)     | 38.0 ± 0.9      | 38.2 ± 1.1      | 36.6 ± 12.4     | 0.74      |
| BMI (kg/m²)     | 29.5 ± 0.5      | 27.5 ± 0.5*     | 27.1 ± 0.9*     | 0.001     |
| Fat mass (kg)   | 33.5 ± 0.7      | 31.9 ± 0.7      | 31.8 ± 1.3      | 0.23      |
| Truncal fat (kg)| 13.2 ± 0.5      | 12.2 ± 0.4      | 11.4 ± 0.9      | 0.16      |
| AIC (%)         | 5.6 ± 0.0       | 5.5 ± 0.0*      | 5.5 ± 0.1*      | 0.002     |
| OGTT            |                 |                 |                 |           |
| Fasting glucose (mmol/l) | 5.3 ± 0.1 | 5.1 ± 0.0 | 5.2 ± 0.1 | 0.28 |
| 2-hour glucose (mmol/l) | 7.8 ± 0.1 | 7.6 ± 0.1 | 7.4 ± 0.2 | 0.43 |
| Glucose AUC × 10^4 (min · mmol⁻¹ · l⁻¹) | 19.6 ± 0.2 | 19.1 ± 0.3 | 18.8 ± 0.4 | 0.34 |
| Insulin AUC × 10^4 (min · pmol⁻¹ · l⁻¹) | 90.2 ± 4.2 | 73.6 ± 3.5 | 62.5 ± 4.9* | <0.001 |
| C-peptide AUC × 10^4 (min · pmol⁻¹ · l⁻¹) | 51.6 ± 1.4 | 48.9 ± 1.3 | 43.4 ± 2.2* | 0.010 |
| Insulin sensitivity ($S_I$) |             |                 |                 |           |
| $S_{IMM} = 10^{-1}$ (dl · kg⁻¹ · min⁻¹ · pmol⁻¹ · l⁻¹) | 1.7 ± 0.1 | 1.8 ± 0.7 | 2.6 ± 0.2* | <0.001 |
| $S_{COM} = 10^{-3}$ (dl · kg⁻¹ · min⁻¹ · pmol⁻¹ · l⁻¹) | 5.1 ± 0.2 | 5.8 ± 0.3 | 6.7 ± 0.5* | 0.007 |
| Insulin secretion |                 |                 |                 |           |
| ISR AUC × 10³ | 30.0 ± 0.9 | 28.1 ± 0.6 | 25.4 ± 1.2* | 0.01 |
| β-Cell responsivity ($F_o$) (10⁹/min) | 13.7 ± 0.8 | 13.3 ± 0.4* | 11.5 ± 0.7* | 0.02 |
| Insulin clearance ($I_C$) |             |                 |                 |           |
| IC₁ (ISR AUC/insulin AUC) | 2.8 ± 0.1 | 3.1 ± 0.1* | 3.2 ± 0.1* | 0.001 |
| IC₂ (C-peptide AUC/insulin AUC) | 47.8 ± 1.1 | 53.3 ± 1.4* | 55.9 ± 2.3* | 0.001 |

Data are means ± SE or n (%). ANCOVA was used to adjust for age, race, and BMI. *$P < 0.05$ vs. E/E; †$P < 0.05$ vs. E/E and E/K using Tukey’s test for post hoc analyses. ‡$S_{IMM} =$ insulin sensitivity from glucose minimal model (25). §$S_{COM} =$ insulin sensitivity from Matsuda (23).
The mechanistic basis of this change in insulin sensitivity is not clear. Kir6.2 is expressed in multiple tissues, including skeletal muscle, brain, and heart. The increase in insulin sensitivity could be due to direct effects of the altered K\textsubscript{ATP} channel activity in one or more of these tissues, and a link between the K\textsubscript{ATP} channel in the hypothalamus and the liver has recently been established (40). An increase in activation of K\textsubscript{ATP} channels in the hypothalamus decreases hepatic gluconeogenesis, and this central effect could serve to counterbalance peripheral actions to maintain glucose homeostasis (40,41). It is interesting to note that basal and insulin-stimulated muscle glucose transport is increased in Kir6.2-null mice, which suggests a role for this channel in the regulation of muscle glucose metabolism (10).

Whether the enhanced insulin sensitivity results from direct effects of K\textsubscript{ATP} channel activity in muscle or is secondary to effects on insulin secretion is unclear. There is precedent for an increase in insulin sensitivity secondary to reduction in insulin secretion. In the evolution of type 1 diabetes, insulin secretion is reduced before glucose levels rise (42), and increased insulin sensitivity has been demonstrated in normoglycemic carriers of HNF1\textalpha mutations with reduced insulin secretion prior to diabetes onset (43). These observations, coupled with those of the present study, suggest that in the evolution of type 2 diabetes, increased insulin sensitivity may compensate for a reduction in insulin secretion, resulting in normal glucose tolerance. Over time, glucose intolerance develops due to progression in the severity of the secretion defect and/or exposure to factors that reduce insulin sensitivity. However, we also cannot completely exclude the likelihood that these E23K variants have not developed diabetes because of their increased insulin sensitivity. In such cases, it might reflect a feature of nondiabetic E23K variants rather than of subjects with E23K variants, per se. Although Kir6.2 levels are high in pancreatic \alpha-cells, we did not find differences in glucagon secretion. An additional new finding is the association of E23K variant with an increase in insulin clearance, as demonstrated in study 2. An increase in insulin clearance may contribute to the reduction in peripheral insulin concentrations that is due largely to the decrease in insulin secretion. The liver is the major site of insulin clearance under physiologic circumstances, and this change in insulin clearance is likely due to a change in hepatic insulin metabolism (44). A link between hepatic insulin receptor binding and action and degradation by the liver has been reported (45).

**Molecular basis of the E23K phenotype.** In heterologous expression studies, reconstituted E23K channels exhibit a mild, yet significant, decrease in ATP sensitivity and a relative increase in basal activity in the intact cell. In the \beta-cell, decreased ATP inhibition and consequent channel overactivity is predicted to suppress glucose sensing. In contrast to activating mutations in Kir6.2 that underlie NDM (28), the E23K variant has a less radical effect on channel activity and is associated with type 2 diabetes. The molecular consequence of the E23K variant has been controversial. Given the high K\textsubscript{ATP} channel density in \beta-cells, it is predicted that a change in K\textsubscript{ATP} channel activity (<1%) could significantly affect insulin secretion, and, therefore, subtle effects of E23K on channel activity could be physiologically relevant. In support of the conclusion that E23K does not alter channel activity, whole-cell K\textsuperscript{+}-currents were similar in Xenopus oocytes expressing E/E or K/K channels (4). A subsequent study (6) reported an increase in ATP sensitivity of K/K channels, relative to E/E, but a subsequent decrease in ATP inhibition upon application of long-chain acyl-CoAs. The most detailed study was carried out by Schwanstecher and colleagues (5,46). Our data parallel their initial findings of an approximately twofold reduction in ATP sensitivity and increase in open probability and reduced sulfonylurea sensitivity (46) associated with the E23K variant. The similarity of our findings, in a completely independent study, utilizing different methodologies to assess open probability, strongly supports the conclusion that the E23K variant decreases ATP inhibition of the K\textsubscript{ATP} channel.

Mechanistically, the observed increase in open probability can account for both the reduced insulin secretion and decrease in sulfonylurea inhibition of E23K channels. Whether the E23K variant affects sulfonylurea dosing is unknown, but it is notable that the E23K variant is associated with risk for secondary failure to sulfonylureas in type 2 diabetic patients (47,48), and human islets isolated from E23K donors (E/K and K/K) exhibit a decrease in sulfonylurea-induced insulin secretion (47).

**Conclusions.** Subjects with the Kir6.2 E23K variant have multiple insulin secretory defects to glucose and decreased responsiveness of the \beta-cell over a physiologic range of glucose concentrations, and these can be explained by the molecular properties of the E23K channels. These defects in insulin secretion are accompanied by an increase in insulin sensitivity, possibly a compensatory response to reduced insulin secretion. The \beta-cell secretory defects are present prior to diabetes onset and are likely responsible for the increased risk of type 2 diabetes in subjects with the at-risk E23K genotypes, if superimposed on lifestyle factors causing insulin resistance.

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