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Research Article

Genetics

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Correlation of rare coding variants in the gene encoding human glucokinase regulatory protein with phenotypic, cellular, and kinetic outcomes

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Defining the genetic contribution of rare variants to common diseases is a major basic and clinical science challenge that could offer new insights into disease etiology and provide potential for directed gene- and pathway-based prevention and treatment. Common and rare nonsynonymous variants in the *GCKR* gene are associated with alterations in metabolic traits, most notably serum triglyceride levels. *GCKR* encodes glucokinase regulatory protein (GKRP), a predominantly nuclear protein that inhibits hepatic glucokinase (GCK) and plays a critical role in glucose homeostasis. The mode of action of rare *GCKR* variants remains unexplored. We identified 19 nonsynonymous *GCKR* variants among 800 individuals from the ClinSeq medical sequencing project. Excluding the previously described common missense variant p.Pro446Leu, all variants were rare in the cohort. Accordingly, we functionally characterized all variants to evaluate their potential phenotypic effects. Defects were observed for the majority of the rare variants after assessment of cellular localization, ability to interact with GCK, and kinetic activity of the encoded proteins. Comparing the individuals with functional rare variants to those without such variants showed associations with lipid phenotypes. Our findings suggest that, while nonsynonymous *GCKR* variants, excluding p.Pro446Leu, are rare in individuals of mixed European descent, the majority do affect protein function. In sum, this study utilizes computational, cell biological, and biochemical methods to present a model for interpreting the clinical significance of rare genetic variants in common disease.

Introduction

Common human diseases result from the combined effects of genetic susceptibility and environmental factors. Understanding the genetic contribution to disease may offer new insights into disease etiology and provide potential for directed gene- and pathway-based prevention and treatment. Genome-wide association studies (GWAS) have been successful in identifying common genetic variants associated with complex disease heritability by their individual statistical associations (1, 2). Individual rare variants conferring low to moderate risk are not as tractable by this approach (2–4). However, emerging evidence suggests rare variants are also important contributors to complex disease susceptibility (5–10). Rapid advances in exome and whole-genome sequencing, as well as collective efforts such as the 1000 Genomes Project, will uncover many novel variants. However, the fundamental challenges in using sequencing for individual diagnostics lie in developing methodologies for distinguishing variants that have demonstrable functional effect from those that are neutral and in relating this information back to clinical phenotypes. The *GCKR* gene, encoding glucokinase regulatory protein (protein name, GKRP), is a logical candidate for exploration

of these issues, as both common and rare variation in *GCKR* have been suggested as being clinically important and the biology of *GCKR* has been extensively studied (11–15).

GKRP posttranslationally regulates hepatic glucokinase (GCK) (16). GCK is the predominant hexokinase in pancreatic β cells and hepatocytes, but *GCKR* is not appreciably expressed in β cells (13, 17–20). In the liver, GKRP binding inhibits GCK competitively with respect to glucose (15, 21). This inhibition is also associated with nuclear sequestration of GCK at low glucose concentrations (14). Glucose-mediated dissociation of GCK from GKRP activates GCK and exposes its nuclear export signal (22). Additionally, GKRP is regulated by binding of fructose 6-phosphate (F6P) or fructose 1-phosphate (F1P). F6P binding to GKRP promotes GKRP-GCK association, while F1P disrupts this interaction (23).

Studies in model systems have suggested GKRP not only serves as a GCK inhibitor, but also plays a somewhat paradoxical role in enhancing GCK protein stability. *Gckr*-knockout mice exhibit normal fasting glucose levels, but suffer from postprandial hyperglycemia due to lower hepatic Gck protein levels and activity (24, 25). Similarly, cats lack endogenous hepatic GKRP expression and display low hepatic GCK activity (26). Additionally, adenoviral overexpression of *GCK* and *GCKR* in HepG2 cells results in elevated GCK protein levels and activity compared with overexpression of *GCK* alone (27). GKRP-bound

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GCK may therefore serve as a functional nuclear reserve that can be rapidly activated and mobilized to the cytoplasm following a glucose challenge.

Given its central role in hepatic glucose metabolism, it is probably not surprising that *GCKR* has emerged as an important locus for susceptibility to diabetes and related traits (12, 28–40). GWAS initially identified a 400-kb region on chromosome 2 significantly associated with serum triglycerides (28). Comprehensive fine mapping identified rs1260326 (c.1337 C>T, p.Pro446Leu) as the likely causative variant associated with an inverse modulation of fasting glucose and triglyceride levels (12). This association has been widely replicated in additional studies and populations (40–42). Further genome-wide analyses have documented associations of this region with risk of type 2 diabetes (T2D), total cholesterol, fasting insulin, C peptide, homeostasis model assessment of insulin resistance (HOMA-IR), and concentrations of a number of other metabolites, including C-reactive protein (CRP), mannose, the liver enzyme γ -glutamyl transferase, and urate (29–40). Functionally, p.Pro446Leu-GKRP has been shown to decrease F6P-mediated inhibition of GCK and reduce nuclear sequestration of GCK (13, 43). Decreased inhibition and sequestration of GCK may lead to increased GCK activity in the liver, resulting in increased de novo triglyceride and cholesterol synthesis and export, as suggested by recently observed associations of *GCKR* with VLDL particle concentrations (44). Increased hepatic glucose disposal could act to decrease plasma glucose concentrations, consistent with the observed inverse effects on glucose and lipid levels.

A recent study highlighted the clinical relevance of the collective burden of rare alleles in *GCKR*, reporting that nonsynonymous variants of minor allele frequency (MAF) of less than 0.01 in *GCKR* are enriched in cases of extreme hypertriglyceridemia (11). This suggests that many nonsynonymous *GCKR* variants will have significant effects on protein structure, function, and/or expression and that these changes will lead to demonstrable metabolic effects. However, this study was not able to explore the effects of individual alleles with respect to biochemical function or to assess the impact of *GCKR* variants on less extreme phenotypes in subjects with trait values more representative of the general population. Understanding such relationships on an individual level will be essential for interpretation of medical sequencing data, especially in the context of genetically complex traits in which rare alleles may be contributory rather than completely penetrant. Such challenges will only become more pronounced due to increasing availability of whole-exome and whole-genome sequence information for large numbers of individuals. Therefore, we aimed to identify and clinically evaluate subjects with nonsynonymous *GCKR* variants and to couple this work with comprehensive functional analysis.

Results

Identification and clinical characteristics of individuals with *GCKR* variants. We sequenced the exons of the *GCKR* gene in 800 members of the ClinSeq cohort ascertained by April 2010. The majority (88.5%) of ClinSeq participants are non-Hispanic individuals of mixed European descent, reflecting the ethnic and sociodemographic characteristics of Bethesda, Maryland, where the NIH Clinical Center is located (45, 46). Cohort selection was implemented to enrich for coronary atherosclerosis and has been described previously (46). Individuals were between the ages of 45 and 65 years, with a mean age of 56 years. Nineteen (10 novel) nonsynonymous

(16 missense, 1 nonsense, and 2 frameshift) *GCKR* variants were identified by exonic Sanger sequencing in ClinSeq participants (Table 1). All variants apart from p.Pro446Leu had a MAF of less than 0.02 in the cohort.

p.Pro446Leu had significant effects on fasting triglyceride levels in the expected direction (Supplemental Figure 1 and Supplemental Table 1; supplemental material available online with this article; doi:10.1172/JCI46425DS1) (28). The effect of p.Pro446Leu on fasting glucose did not reach significance (Supplemental Figure 1), most likely due to limited statistical power, as the previously reported effect of this allele on fasting glucose levels is small (0.5 mg/dl per allele in ref. 34) and unlikely to be detected in a cohort of this size. Individuals with either T2D ($n = 51$) or potential familial hyperlipidemia (family history of dyslipidemia and triglyceride levels > 500 mg/dl; $n = 6$) were excluded from phenotype comparisons because of direct effects on glucose and lipid phenotypes. Exclusion of individuals on statins or niacin did not alter significance of triglyceride or glucose levels. p.Pro446Leu was in Hardy-Weinberg equilibrium (HWE) in the whole sample and the subset of non-Hispanic individuals of mixed European descent. Allele frequencies were in accordance with data from the International HapMap project (<http://www.hapmap.org>).

There were 42 ClinSeq individuals with *GCKR* variants other than p.Pro446Leu. Consistent with the findings of Johansen et al. (11), collective analysis of the group of individuals with rare *GCKR* variants revealed a significant association with plasma triglycerides compared with the WT reference group (individuals homozygous for Pro at position 446 but with no rare *GCKR* variants; Figure 1; $P = 0.03$). Of these 42 individuals, 28 were also heterozygous and 4 were homozygous for the Leu allele at position 446 (Supplemental Table 2). We were able to determine the likely phase for 24 of 28 of the compound heterozygotes through family studies and/or inference from other individuals harboring the same rare allele (Supplemental Table 2). Six individuals carried multiple rare alleles: 1 individual carried p.[Gln234Pro;Arg540X] in *cis*, 1 individual carried p.[Tyr307Asp(;Arg540Gln] (phase unknown), and 4 individuals carried p.[Ser183CysfsX34;Ala519Thr] in *cis*, suggesting p.Ala519Thr may not be relevant at the protein level in these individuals because it occurs downstream of a frameshift variant. One of these 4 individuals also carried p.Arg540Gln in *trans*. No rare variant produced extreme outlier phenotypes greater than 4 SD from the mean for the ClinSeq cohort (data not shown). Efforts to expand pedigrees did not provide sufficient data to establish segregation.

While preliminary comparisons of the collective group of individuals with rare variants with the WT reference group suggested a relationship to triglycerides, we were concerned that the merging of data from certain nonsynonymous variants of potentially no consequence with that of those that cause significant loss or gain of function could considerably diminish the power of the analysis. Finding large numbers of individuals with specific *GCKR* variants for detailed phenotyping was impractical, given the rarity of these particular variants in 1000 Genomes (47) and previously published studies (11, 48). It thus became imperative to explore evolutionary, cell biological, and biochemical characterization of each variant to evaluate their potential contribution to phenotypes.

Evolutionary characteristics and bioinformatic predictions for *GCKR* variants. To aid in the interpretation of genetic and clinical data, we aimed to characterize nonsynonymous variants at the protein level. There is no crystal structure available for GKRP. How-



Table 1
GCKR variants discovered by exonic sequencing in the ClinSeq cohort

Amino acid change	Nucleotide change	dbSNP ID	Number of individuals	Ethnicity	Previous reference
p.Arg51Gln	c.152G>A		4	Mixed European	11
p.Glu77Gly	c.230A>G	rs8179206	1	Asian	11
p.Val103Met	c.307G>A		2	Mixed European/Hispanic	Novel
p.Ser183CysfsX34	c.548_549del		4	Ashkenazi	Novel
p.Ile219Val	c.655A>G		1	Mixed European	Novel
p.Gln234Pro	c.701A>C		4	Mixed European	11
p.Tyr307Asp	c.919T>G	rs34498189	1	Mixed European	dbSNP
p.Thr379AsnfsX36	c.1135dup		2	Mixed European	11
p.Pro383Thr	c.1147C>A		1	Mixed European	Novel
p.Ile396Asn	c.1187T>A		1	Mixed European	Novel
p.Pro446Leu	c.1337C>T	rs12360326	368 C/T 157 T/T	Various	Numerous
p.Arg478His	c.1433G>A		1	Mixed European	Novel
p.Ile500Ser	c.1499T>G		1	Mixed European	Novel
p.Ala519Thr	c.1555G>A		4	Ashkenazi	Novel
p.Arg540X	c.1618C>T		1	Mixed European	11
p.Arg540Gln	c.1619G>A	rs8179249	17	Mixed European	dbSNP
p.His590Tyr	c.1768C>T	rs34792470	2	Mixed European	48
p.Gly607Glu	c.1820G>A		1	Hispanic	Novel
p.Arg612Cys	c.1834C>T		1	Mixed European	Novel

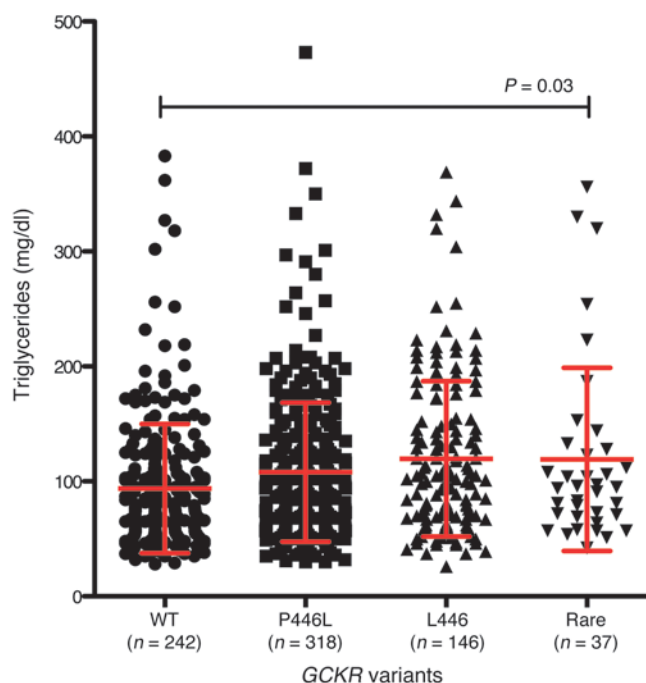
ever, GGRP bears homology to bacterial proteins of the sugar isomerase (SIS) family and contains 2 separate SIS domains that combine to form a single site capable of binding F1P or F6P (23, 49). The location of critical sugar-binding residues appears highly conserved (23). Mapping of variants on these predicted GGRP subdomains is presented in Supplemental Figure 2A. Variant p.Val103Met is located directly within a predicted sugar-binding motif, while p.Ile500Ser immediately precedes a binding motif (Supplemental Figure 2B). p.His590Tyr, p.Gly607Glu, and p.Arg612Cys are C-terminal of SIS domains, as predicted by NCBI Conserved Domains search (<http://www.ncbi.nlm.nih.gov/Structure/cdd/wrpsb.cgi>). p.Ile219Val and p.Gln234Pro are predicted to localize to a eukaryote-specific α -helix (Supplemental Figure 2, B and C). All variants are in highly conserved residues apart from p.Arg51Gln, p.Arg478His, p.Arg540Gln, and p.His590Tyr (Supplemental Table 3). The effect of amino acid substitution was predicted by SIFT (<http://sift.jcvi.org/>) and PolyPhen (<http://genetics.bwh.harvard.edu/pph/>) (Supplemental Table 4). These algorithms predict mutation severity based on a number of parameters, including sequence conservation and known functional motifs (50, 51), and their use has been proposed as a method for classifying variants identified by resequencing (52). Together, SIFT and PolyPhen predicted 7 of 15 missense variants as damaging and 5 of 15 as benign, disagreeing on 3 variants. However, the reliability of these prediction algorithms has not been extensively tested. For example, Johansen et al. (11) found a significant excess in the proportion

of rare variants predicted by PolyPhen as benign in hypertriglyceridemia cases compared with controls, and SIFT predicted all 8 *GCKR* missense variants identified by Johansen et al. as benign (data not shown).

The majority of *GCKR* variants alter cellular localization of the regulatory protein. As we had concerns about the reliability of prediction algorithms and wanted to compare evolutionary features to protein function, we biochemically characterized *GCKR* variants to determine their molecular and cellular effects. As GGRP function is intimately linked with nuclear localization, we generated N-ter-

Figure 1

Triglyceride levels of ClinSeq participants separated by *GCKR* genotype. Red bars indicate unadjusted means \pm SD. Two-tailed *P* values are compared with WT (see also Table 3). WT, individuals homozygous for Pro at position 446; P446L, heterozygous individuals at position 446; L446, individuals homozygous for Leu at position 446; rare, individuals heterozygous for 1 or more rare *GCKR* nonsynonymous variants.



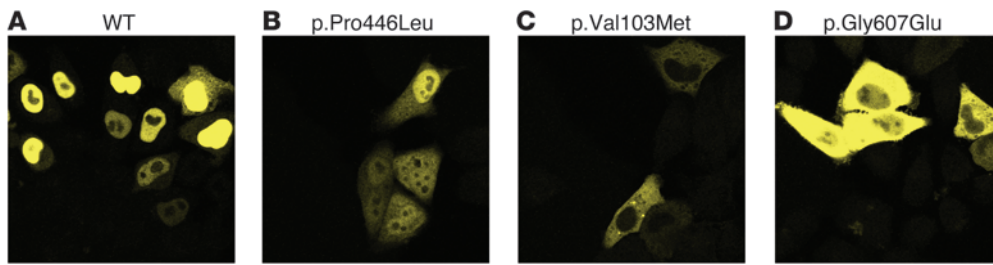


Figure 2

Cellular localization of representative variant YFP-GKRPs on transient transfection into HeLa cells. (A) WT GKRPs localized primarily to the nucleus, as did p.Arg51Gln, p.Glu77Gly, p.Pro383Thr, p.Arg478His, p.Arg540Gln, and p.His590Tyr. (B) p.Ile396Asn, p.Pro446Leu, and p.Ile500Ser localized to both the cytoplasm and nucleus. (C) p.Val103Met, p.Ile219Val, p.Gln234Pro, and p.Tyr307Asp localized primarily to the cytoplasm and showed comparatively lower YFP fluorescence intensity. (D) p.Gly607Glu and p.Arg612Cys were also cytoplasmic, but showed higher levels of YFP expression. Original magnification, $\times 63$. Images use the same laser settings and intensity and are representative of at least 2 transfections of 2 independent plasmid preparations for each variant. A representative image of each localization pattern is shown.

minimal yellow fluorescent protein (YFP) fusions to WT or variant GCKR-coding sequences and determined their subcellular location in HeLa cells. The interaction of GCK and GKRPs has been previously extensively characterized in this cell type (22, 43). WT YFP-GKRPs localized primarily to the nucleus 24 hours after transient transfection into HeLa cells (Figure 2A). Predominantly nuclear localization of WT YFP-GKRPs was also observed in COS-1 and HepG2 cells (data not shown).

Eighteen nonsynonymous human variants from the ClinSeq project were introduced into the YFP-GKRPs plasmid and transiently transfected into HeLa cells (p.Arg51Gln, p.Glu77Gly, p.Val103Met, p.Ser183CysfsX34, p.Ile219Val, p.Gln234Pro, p.Tyr307Asp, p.Thr379AsnfsX36, p.Pro383Thr, p.Ile396Asn, p.Pro446Leu, p.Arg478His, p.Ile500Ser, p.Arg540X, p.Arg540Gln, p.His590Tyr, p.Gly607Glu, and p.Arg612Cys). Only truncating variants p.Ser183CysfsX34, p.Thr379AsnfsX36, and p.Arg540X failed to generate YFP fluorescence, except in a minority of cells where punctate fluorescent aggregates were observed. Both of these phenomena are associated with protein misfolding (53). These results suggest that the p.Gln234Pro variant will not have any additional effect on p.Arg540X *in cis* for the individual in which these variants were discovered.

The remaining nonsynonymous YFP-GKRPs variants showed 4 distinct patterns of localization. First, variants p.Arg51Gln,

p.Glu77Gly, p.Pro383Thr, p.Arg478His, p.Arg540Gln, and p.His590Tyr localized primarily to the nucleus like WT GKRPs (see Supplemental Figure 3 for all variants). Second, consistent with our previous findings, common variant p.Pro446Leu localized to both the cytoplasm and nucleus (Figure 2B and ref. 43). Rare variants p.Ile396Asn and p.Ile500Ser also displayed this behavior. Third, p.Val103Met, p.Ile219Val, p.Gln234Pro, and p.Tyr307Asp showed almost exclusive cytoplasmic localization and comparatively low YFP fluorescence (Figure 2C). Finally, variants p.Gly607Glu and p.Arg612Cys localized to the cytoplasm and exhibited greater fluorescence intensity than other cytoplasmic variants (Figure 2D). Western blot analysis showed that expression levels of p.Gly607Glu and p.Arg612Cys were similar to those of WT YFP-GKRPs, while other cytoplasmic variants showed reduced expression (data not shown).

For 3 rare variants (p.Glu77Gly, p.Pro383Thr, and p.Arg540Gln), the individuals harboring them were confirmed as also having the Leu allele at position 446 *in cis*. We therefore assessed the combined effect of these variants. While p.[Pro446Leu;Arg540Gln] showed diffuse localization similar to that of p.Pro446Leu, p.[Glu77Gly;Pro446Leu] and p.[Pro383Thr;Pro446Leu] showed almost exclusive cytoplasmic localization, suggesting the p.Pro446Leu variant may amplify otherwise modest cellular effects for these 2 variants (Supplemental Figure 4).

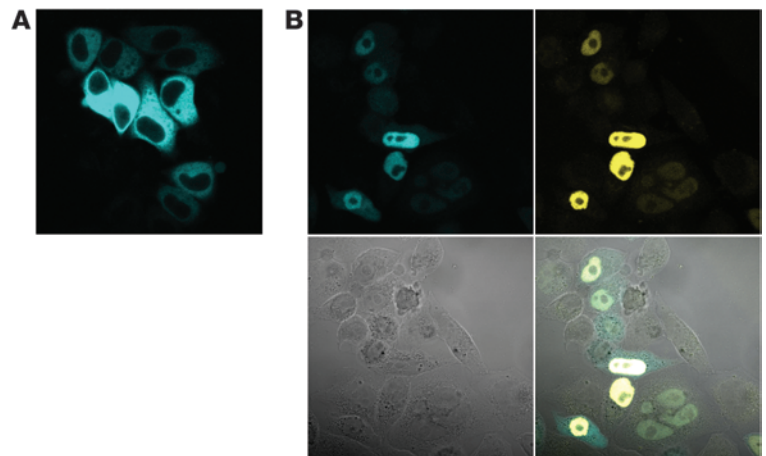
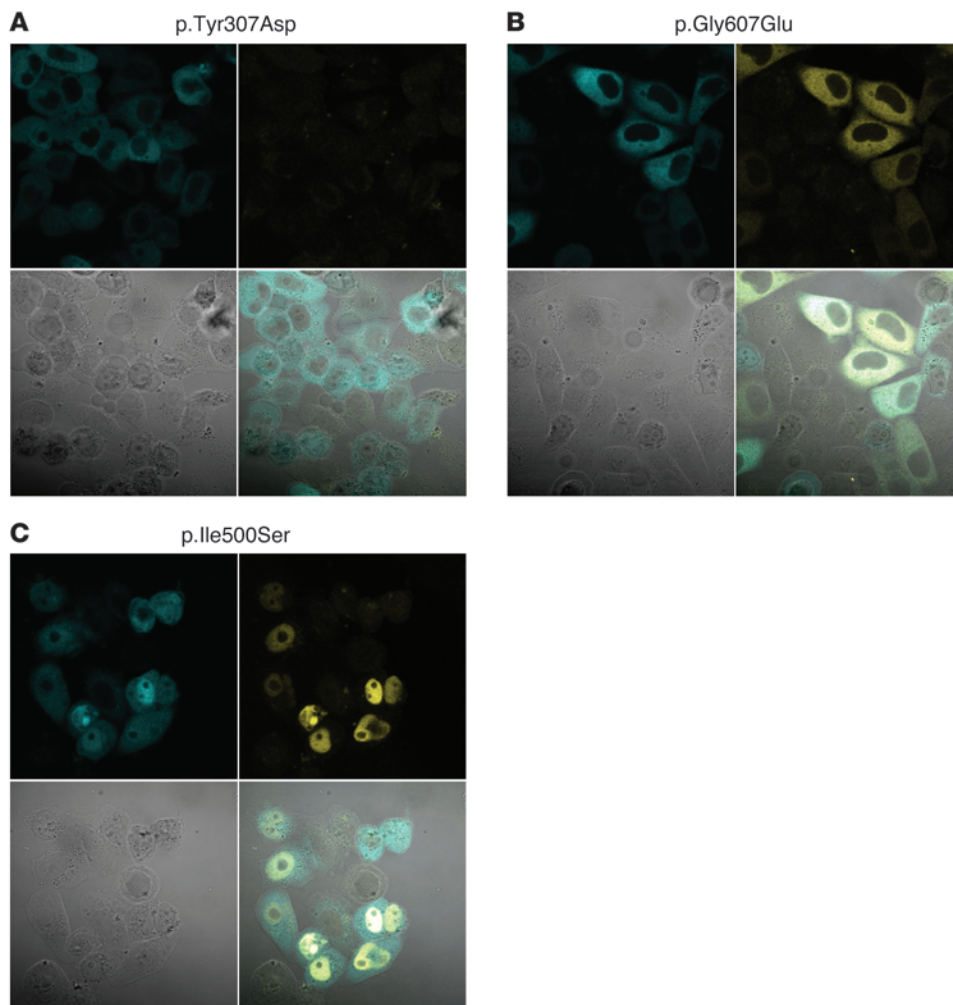


Figure 3

WT YFP-GKRPs is necessary and sufficient to sequester CFP-GCK to the HeLa cell nucleus. (A) CFP-GCK localized to the cytoplasm upon transient transfection into HeLa cells. (B) Cotransfection of YFP-GKRPs expressed from the same plasmid sequestered CFP-GCK to the nucleus in HeLa cells. All YFP-GKRPs variants that localized primarily to the nucleus were capable of sequestering CFP-GCK. Top left, CFP channel; top right, YFP channel; bottom left, phase channel; bottom right, combined channels. Original magnification, $\times 63$. Images were taken using the same laser settings and intensity and are representative of at least 2 transfections of 2 independent plasmid preparations for each variant.

**Figure 4**

Diffuse and cytoplasmic YFP-GKRP variants show distinct classifications of behavior in the presence of CFP-GCK on transient transfection into HeLa cells. **(A)** Variants p.Val103Met, p.Ser183CysfsX34, p.Tyr307Asp, p.Thr379AsnfsX36, and p.Arg540X did not sequester CFP-GCK to the nucleus and did not show appreciable YFP fluorescence when intensity matched to WT YFP-GKRP. **(B)** Variants p.Gly607Glu and p.Arg612Cys did not sequester CFP-GCK, but showed appreciable YFP fluorescence intensity. **(C)** Variants p.Ile219Val, p.Gln234Pro, p.Ile396Asn, p.Pro446Leu, and p.Ile500Ser interacted with GCK and were relocated to the nucleus. Original magnification, $\times 63$. Images were made using the same laser settings and intensity and are representative of at least 2 transfections of 2 independent plasmid preparations for each variant. Top left, CFP channel; top right, YFP channel; bottom left, phase channel; bottom right, combined channels. A representative image of each localization pattern is shown.

Cellular interaction of GKRP variant proteins with GCK emphasizes distinct mislocalization subtypes. To understand the consequences of GKRP mislocalization further, we assessed the effect of WT and variant GKRP on the cellular localization of GCK. Human liver GCK was cloned as an N-terminal cyan fluorescent protein (CFP) fusion protein. Without cotransfected GKRP, CFP-GCK localized exclusively to the cytoplasm in HeLa cells (Figure 3A). The same result was seen in HepG2 and COS-1 cells (data not shown). To assess interaction between GCK and GKRP, a single plasmid containing separate promoters, coding sequences, and polyadenylation signals for CFP-GCK and YFP-GKRP was utilized to ensure consistent coexpression of GKRP and GCK in individual cells across experiments. This was particularly important because a number of variant GKRP showed very low levels of expression.

Transient transfection of the CFP-GCK and YFP-GKRP dual-expression plasmid resulted in nuclear localization of WT GKRP and nuclear sequestration of CFP-GCK (Figure 3B). In accordance with previous observations, GCK exhibited similar subcellular localization patterns for multiple glucose concentrations tested (data not shown) (43). Therefore, images from 1 glucose concentration (25 mM) are presented and representative of other concentrations. The 6 variants exhibiting WT-like localization as YFP fusion proteins (p.Arg51Gln, p.Glu77Gly, p.Pro383Thr,

p.Arg478His, p.Arg540Gln, and p.His590Tyr) remained localized to the nucleus and sequestered GCK. Evolutionary analysis of these variants revealed that 4 of these variants (p.Arg51Gln, p.Arg478His, p.Arg540Gln, and p.His590Tyr) were in amino acid residues showing the lowest conservation in mammals among the *GCKR* variants tested. The observed variant amino acid at residues 51, 478, and 540 is the reference amino acid in 3 or more nonhuman mammalian species (Supplemental Table 3).

Diffuse and cytoplasmic GKRP variants showed distinct responses to coexpression of GCK. Variants p.Gly607Glu and p.Arg612Cys did not sequester GCK to the nucleus, but showed significant cytoplasmic YFP fluorescence (Figure 4A). Five variants (p.Val103Met, p.Ser183CysfsX34, p.Tyr307Asp, p.Thr379AsnfsX36, and p.Arg540X) did not traffic GCK to the nucleus and showed low or undetectable intensity of fluorescent GKRP (Figure 4B). In the presence of GCK, GKRP variant proteins p.Ile219Val, p.Gln234Pro, p.Ile396Asn, p.Pro446Leu, and p.Ile500Ser relocated to the nucleus and sequestered GCK (Figure 4C). Previous studies have suggested GCK promotes nuclear localization of GKRP (22, 54), which may in part explain this observation. However, we have shown by quantification of nuclear and whole-cell fluorescence intensity that the degree of GKRP nuclear relocalization and GCK sequestration is significantly reduced for p.Pro446Leu compared with WT

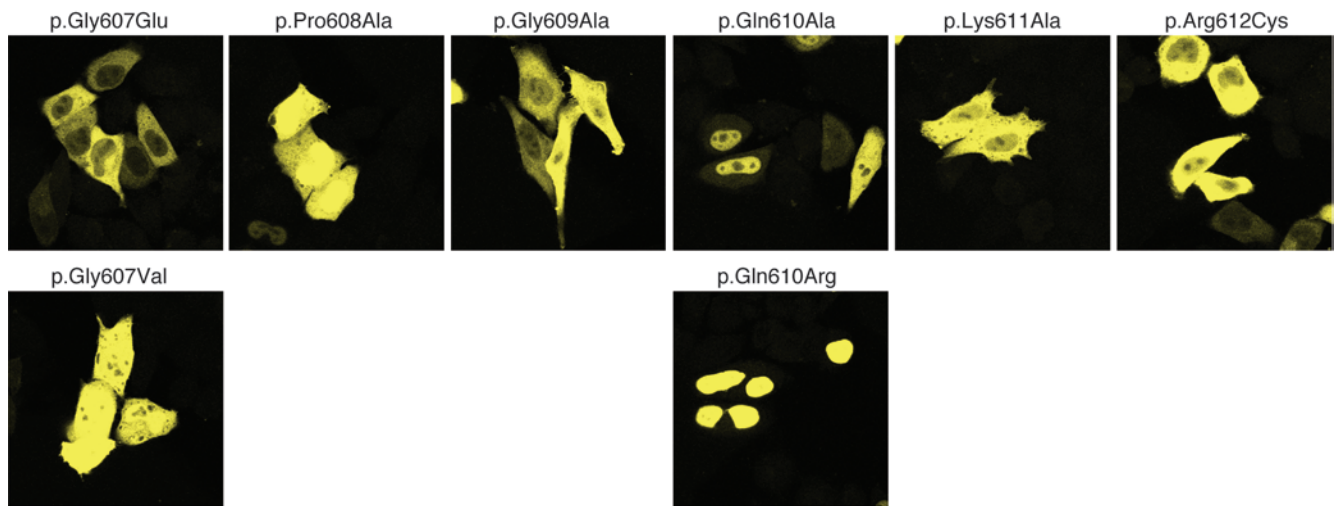


Figure 5

YFP-GKRP C-terminal variants have an important role in nuclear localization. Mutations of YFP-GKRP conserved residues to ClinSeq variants p.Gly607Glu and p.Arg612Cys, residues present in other GKRP orthologs (p.Gly607Val, p.Gln610Arg) or Ala (p.Pro608Ala, p.Gly609Ala, p.Gln610Ala, p.Lys611Ala). Original magnification, $\times 63$. Images were taken the same laser settings and intensity and are representative of at least 2 transfections of 2 independent plasmid preparations for each variant.

GKRP (43). p.[Glu77Gly;Pro446Leu], p.[Pro383Thr;Pro446Leu], and p.[Pro446Leu;Arg540Gln] also relocated to the nucleus and sequestered GCK.

Residues near the C terminus of the regulatory protein affect nuclear localization and GCK sequestration. Two variants near the C terminus of the protein, p.Gly607Glu and p.Arg612Cys, were exclusively cytoplasmic, demonstrated high fluorescence intensity, and did not sequester GCK. Alignment of C-terminal GKRP residues 595–625 with 3 other GKRP orthologs known to localize to the nucleus showed decreased amino acid conservation, particularly in the case of *Xenopus laevis* GKRP (Supplemental Table 5 and Supplemental Figure 5). The region containing ClinSeq variants p.Gly607Glu and p.Arg612Cys was more highly conserved. To determine whether mutation of intervening residues had similar effects, residues 607–612 were mutated to Ala or to residues present in *X. laevis* GKRP. Mutation to *X. laevis* residues p.Gly607Val or p.Gln610Arg maintained partial and complete nuclear local-

ization, respectively. Variants p.Gly609Ala and p.Lys611Ala localized to the cytoplasm and did not sequester GCK, while variants p.Pro608Ala and p.Gln610Ala showed a mixture of nuclear and cytoplasmic localization. All variants showed fluorescence intensity similar to that of p.Gly607Glu and p.Arg612Cys (Figure 5).

Kinetic analysis of selected variant regulatory proteins supports distinct functional variant classes. The combination of these cellular results assessing human *GCKR* nonsynonymous variants suggested GKRP variant proteins have a spectrum of effects on localization and interaction with GCK (Table 2). To further explore these observations, we selected a subset of variants for recombinant expression in *E. coli*, purification, and kinetic comparison with WT GKRP. We selected 4 variants that each showed reduced protein expression, reduced nuclear localization, and potential reduction in GCK sequestration (p.Val103Met, p.Ile219Val, p.Gln234Pro, and p.Ile500Ser). We also selected p.Arg612Cys to represent C-terminal variants displaying high cytoplasmic YFP

Table 2

Subdivision of *GCKR* variants into classes according to cellular localization, cellular interaction with GCK, and kinetic effects

	WT like	Putative loss of function	Putative gain of function
Human variants	WT, p.Arg51Gln, p.Arg478His, p.Arg540Gln, p.His590Tyr	p.[Gly77Glu;Pro446Leu], p.Val103Met, p.[Ser183CysfsX34;Ala519Thr], p.Ile219Val, p.Gln234Pro, p.Tyr307Asp, p.Thr379AsnfsX36, p.[Pro383Thr;Pro446Leu], p.Ile396Asn, p.Pro446Leu, p.Ile500Ser, p.Arg540X	p.Gly607Glu, p.Arg612Cys
GKRP localization	Predominantly nuclear	Decreased nuclear localization	Fully cytoplasmic
GKRP expression	WT like	Reduced to none	WT like
Observed or predicted GCK sequestration	WT like	Reduced (43) to none	None
Observed or predicted kinetic effect	WT like	F1P/F6P response reduced	WT like

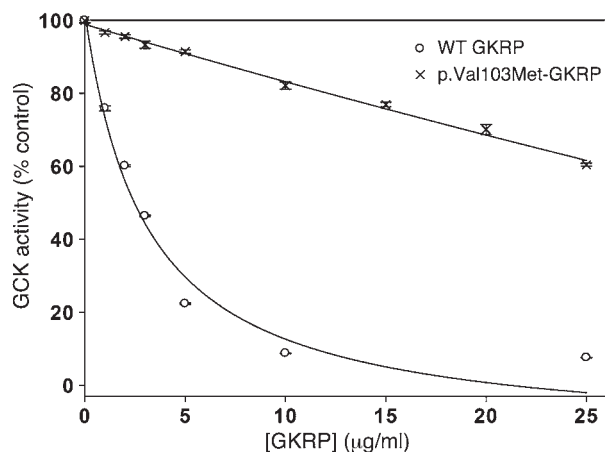


Figure 6

p.Val103Met-GKRP shows significantly reduced inhibition of GCK. Inhibition of 10 mU/ml recombinant GCK by increasing concentrations of recombinant WT GKRP (circles) and p.Val103Met-GKRP (X's). GCK activity (mean \pm SEM; $n = 8$; $P < 0.001$ for all GKRP concentrations tested) is plotted as a percentage of that obtained in the absence of regulatory protein at 5 mM glucose.

expression. Finally, we selected variant p.Pro383Thr, as this variant was strongly sensitive to p.Pro446Leu genotype in *cis*, was in a highly conserved residue, was predicted to be deleterious by both SIFT and PolyPhen, was observed only once in the ClinSeq cohort, and had not been previously reported.

Variant regulatory proteins appreciably inhibited recombinant human GCK, with the exception of p.Val103Met-GKRP, a variant that showed very low expression in HeLa cells. p.Val103Met-GKRP had significantly reduced ability to inhibit GCK over a 0 to 25 μ g/ml concentration range, with $60.5 \pm 0.4\%$ GCK activity remaining at 25 μ g/ml (mean \pm SEM; $P < 0.001$ for all GKRP concentrations). In contrast, GCK activity was only $7.6 \pm 0.1\%$ that of uninhibited GCK in the presence of 25 μ g/ml WT GKRP (Figure 6). As protein misfolding is dependent on temperature, the assay temperature was reduced to 30°C. This led to only a slight improvement in p.Val103Met-GKRP function, with 50% inhibition of GCK predicted at 22.5 μ g/ml protein (data not shown).

We calculated 50% inhibition of 10 mU/ml GCK at 5 mM glucose and 37°C (defined as one GKRP unit) for WT GKRP and all variants excluding p.Val103Met (Supplemental Table 6). p.Ile219Val-GKRP and p.Gln234Pro-GKRP showed a reduction in activity, while p.Pro383Thr-GKRP, p.Ile500Ser-GKRP, and p.Arg612Cys-GKRP were nearly indistinguishable from WT GKRP. GCK inhibition by 1 unit of WT GKRP was compared with inhibition by 1 unit p.Ile219Val-GKRP, p.Gln234Pro-GKRP, p.Pro383Thr-GKRP, p.Ile500Ser-GKRP, or p.Arg612Cys-GKRP over a 0 to 100 mM glucose concentration range. No significant differences were observed between WT and variant proteins (Supplemental Figure 6). One unit of each of these variant proteins was then tested for ability to interact with the phosphate esters F6P and F1P. C-terminal variant p.Arg612Cys showed no difference in response to either phosphate ester (Figure 7A). Accordingly, this variant appears to result in a GKRP protein that is a fully active, exclusively cytoplasmic inhibitor of GCK and hence may be considered as a potential gain-of-function mutation. p.Pro383Thr, a

variant that localizes to the nucleus, showed decreased response to F1P (Figure 7B), but showed no difference in response to F6P. p.Ile219Val, a variant relocalized to the nucleus in the presence of GCK, also showed no difference in response to F6P and had a decreased response to F1P (Figure 7C). Variants p.Gln234Pro and p.Ile500Ser, which were also relocalized to the nucleus by GCK, had significantly diminished response to both F1P and F6P (Figure 7, D and E). The relative amplitudes of response to F1P and F6P are listed in Supplemental Table 6.

To assess the kinetic effect of Leu at position 446 in *cis*, p.[Pro383Thr;Pro446Leu]-GKRP and p.Pro446Leu-GKRP were generated. Consistent with previous observations, p.Pro446Leu-GKRP showed reduction in activity compared with WT GKRP (13). p.[Pro383Thr;Pro446Leu]-GKRP showed reduced activity compared with WT GKRP, p.Pro446Leu-GKRP, and p.Pro383Thr-GKRP (Supplemental Figure 7A). However, comparison of activity-matched p.Pro446Leu-GKRP with p.[Pro383Thr;Pro446Leu]-GKRP showed no additional defects in response to F1P or F6P beyond those observed when comparing p.Pro383Thr-GKRP and WT GKRP (Supplemental Figure 7B). This suggests there is no combined kinetic effect of these 2 variants on F6P and F1P affinity. As p.Pro446Leu-GKRP showed no significant difference in response to F1P and a very modest decrease in response to F6P (Supplemental Table 6), it is probable that Leu446 in *cis* would contribute little to the F1P- and F6P-binding characteristics of rare variants. However, our results suggest *cis* effects on protein activity and cellular localization may be more prominent.

The p.Pro383Thr-GKRP and p.[Pro383Thr;Pro446Leu]-GKRP proteins were also used to model the potential kinetic effect of the p.Pro446Leu variant in *trans*. Protein activities were calculated for WT GKRP, p.Pro383Thr-GKRP, p.Pro446Leu-GKRP, and p.[Pro383Thr;Pro446Leu]-GKRP in conjunction with 1:1 mixtures (by concentration) of these proteins (Supplemental Table 7). The maximal responses of these proteins and mixtures of proteins to F1P and F6P were also calculated. All mixtures resulted in intermediate activity values and responses to F1P and F6P when compared with variant proteins assessed in isolation, suggesting none of these proteins had dominant kinetic effects.

Classification of phenotypes based on variant functional effects. The combination of cellular and kinetic assays suggested 3 broad classifications of variant GKRP (Table 2). Based on our biochemical analysis, we subdivided *GCKR* rare variant heterozygotes ($n = 42$) into subgroups with variants that were functionally similar to WT (*Rare, WT-like*; $n = 22$), C-terminal potential gain-of-function variants (*Rare, GOF*; $n = 2$), and putative loss-of-function variants that were functionally similar to p.Pro446Leu in showing reduced to complete loss of protein expression, reduced nuclear localization, and reduced interaction with F1P and/or F6P (*Rare, LOF*; $n = 18$) (Table 3). Four individuals were excluded because of T2D (1 individual with the p.Thr379AsnfsX36 variant, p.Ile396Asn, 1 individual with the p.Arg540Gln variant, and 1 individual with the p.His590Tyr variant). A second individual with p.Arg540Gln was excluded on the basis of potential familial hyperlipidemia, leaving 37 individuals for analysis. Compared with the WT reference group, the *Rare, WT-like* group showed no significant differences for all phenotypes tested ($P > 0.1$). However, the *Rare, LOF* group had significantly higher levels of total cholesterol ($P = 0.005$), LDL cholesterol ($P = 0.03$), and triglycerides ($P = 0.01$) (Table 3 and Supplemental Figure 8). The difference in total cholesterol ($P = 0.002$), LDL cholesterol ($P = 0.02$), and triglycerides ($P = 0.001$) for the

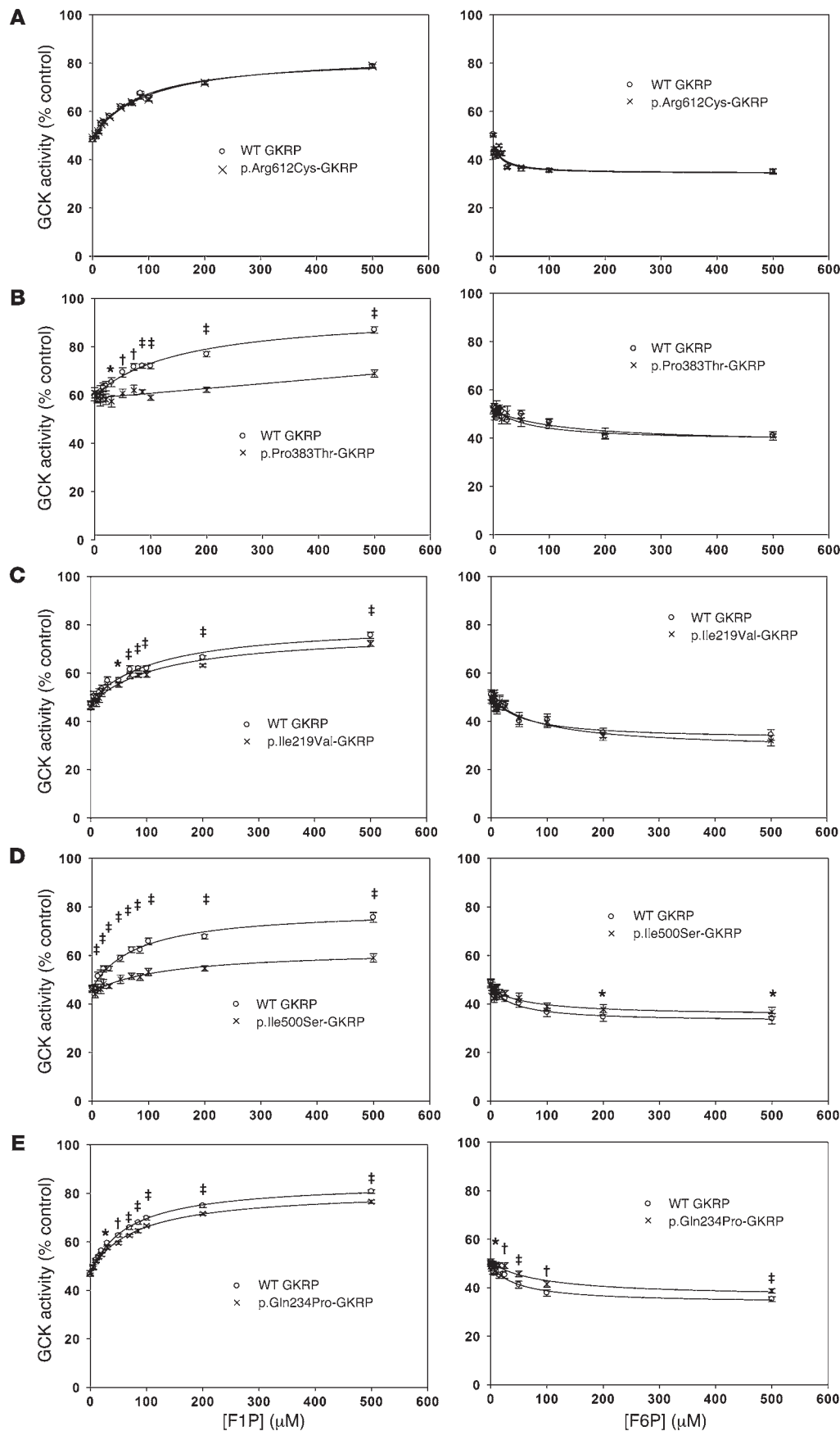


Figure 7
Effect of 0–500 μM F1P or F6P on inhibition of 10 mU/ml recombinant GCK by 1 unit of selected GKRPs. Comparisons of recombinant proteins with WT GKRPs. (A) p.Arg612Cys showed no significant difference ($P > 0.1$) in response to F1P ($n = 12$) or F6P ($n = 12$). (B and C) p.Pro383Thr ($n = 12$; $P < 0.02$ for 30–500 μM F1P < 0.04 for 5–500 μM F1P) showed a significantly reduced response to F1P but not F6P ($P > 0.1$). (D and E) p.Ile500Ser and p.Gln234Pro showed a significantly reduced response to F1P ($n = 12$ and $P < 0.03$ for 15–500 μM for p.Ile500Ser; $n = 38$ and $P < 0.02$ for 20–500 μM for p.Gln234Pro) and F6P ($n = 19$ and $P < 0.05$ for 200–500 μM for p.Ile500Ser; $n = 41$ and $P < 0.02$ for 10–500 μM for p.Gln234Pro). GCK activity is plotted as a percentage of that obtained in the absence of regulatory protein at 5 mM glucose. Data points are mean \pm SEM. * $P < 0.05$; † $P < 0.01$; ‡ $P < 0.001$.

**Table 3**Comparison of baseline phenotypic characteristics of *GCKR* WT and individuals with rare *GCKR* variants

Clinical features	<i>GCKR</i> rare subgroups							
	<i>GCKR</i> WT	<i>GCKR</i> rare ^A	Rare vs. WT	WT like	WT like vs. WT	Loss of function	Loss of function vs. WT	Gain of function
	Mean ± SD <i>n</i> = 242	Mean ± SD <i>n</i> = 37	<i>P</i> (2-tailed)	Mean ± SD <i>n</i> = 19	<i>P</i> (2-tailed)	Mean ± SD <i>n</i> = 16	<i>P</i> (2-tailed)	Mean ± SD <i>n</i> = 2
Age (years)	56.6 ± 5.5	51.7 ± 4.7	0.71	57.5 ± 5.1	0.56	57.0 ± 4.1	0.86	54.0 ± 7.1
Sex (% female)	55.8%	40.5%	0.11	36.8%	0.15	37.5%	0.20	100%
T2D exclusion (%)	8.3%	11.9%	0.39	9.5%	0.69	11.1%	0.66	0%
BMI (kg/m ²)	27.0 ± 4.5	26.8 ± 5.2	0.82	26.5 ± 4.4	0.80	28.1 ± 5.7	0.47	19.4 ± 1.7
Fasting glucose (mg/dl)	97.4 ± 9.0	97.7 ± 9.6	0.77	99.5 ± 9.5	0.52	97.0 ± 9.7	0.60	86.5 ± 0.7
Fasting insulin (mcU/ml)	8.9 ± 6.3	9.6 ± 9.1	0.43	7.8 ± 5.5	0.40	12.6 ± 11.9	0.70	1.5 ± 0.7
C peptide (ng/ml)	2.2 ± 1.1	2.1 ± 1.2	0.21	2.1 ± 1.3	0.30	2.2 ± 1.0	0.91	1.1 ± 0.1
CRP (mg/dl)	0.21 ± 0.31	0.20 ± 0.20	0.77	0.23 ± 0.21	0.26	0.17 ± 0.20	0.79	0.04 ± 0.02
Total cholesterol (mg/dl)	184.6 ± 37.5	193.9 ± 44.5	0.35	175.4 ± 30.3	0.26	216.7 ± 51.1	0.005	180.5 ± 21.9
HDL cholesterol (mg/dl)	60.0 ± 16.3	56.9 ± 15.9	0.25	56.3 ± 15.9	0.27	56.9 ± 17.1	0.43	62.0 ± 7.1
LDL cholesterol (mg/dl)	106.7 ± 31.6	113.6 ± 39.1	0.52	99.1 ± 24.7	0.28	131.8 ± 47.8	0.03	105.5 ± 20.5
Triglycerides (mg/dl)	93.8 ± 56.2	119.1 ± 79.7	0.03	97.3 ± 39.0	0.19	153.7 ± 104.4	0.01	50.0 ± 11.3

GCKR subgroups were defined as *GCKR* WT (individuals who are homozygous for Pro at position 446) or *GCKR* rare (individuals with *GCKR* variants of MAF < 0.02). *GCKR* rare was further broken down into WT like (individuals with rare *GCKR* variants that show WT-like cellular localization), loss of function (individuals with rare *GCKR* variants showing predicted or observed cellular localization defect, reduced protein expression, and a kinetic defect), and gain of function (individuals with rare C-terminal *GCKR* variants showing cytoplasmic localization and no kinetic defects). Fisher's exact test was used to compare differences in frequency distribution of sex and prevalence of T2D. The Mann-Whitney *U* test was used for pair-wise comparison of unadjusted means for *GCKR* WT versus *GCKR* rare groups (excluding the gain-of-function subgroup; *n* = 2) for all other clinical variables. ^AThe 5 variants excluded from the *GCKR* rare group for clinical analysis were p.Thr379AsnfsX36 (*n* = 1), p.Ile396Asn (*n* = 1), p.His590Tyr (*n* = 1), and p.Arg540Gln (*n* = 2), as these individuals were affected with T2D or potential familial dyslipidemia.

Rare, LOF group remained significant after adjustment for covariates including genotype at p.Pro44Leu (Supplemental Table 8 and Supplemental Methods). An effect estimate for the Rare, LOF group (using *P* trend, see Supplemental Table 9) was 50.1 ± 14.7 mg/dl (mean ± SEM; *P* < 0.001) for triglycerides. This compares with an estimate of 11.3 ± 3.0 mg/dl (mean ± SEM; *P* < 0.001) for the Leu allele at position 446 alone. The p.Pro44Leu variant was not overrepresented in either the overall *GCKR* rare group (MAF = 0.37) or the Rare, LOF group (MAF = 0.37) compared with the entire cohort (MAF = 0.42).

We also sought to determine whether rare *GCKR* variants with more severe effects on protein function such as null alleles would have an even more significant effect on phenotype. Loss-of-function variant GKRPs show distinctive kinetic parameters, cellular localization, and interactions with GCK. However, data from *Gckr*-knockout mice suggest more severe LOF variants may predispose to development of diabetes-associated phenotypes, particularly in the context of additional factors such as high-fat and high-sugar diets (24). The variants with most severe cellular and kinetic defects (severe LOF, *n* = 11) appeared to be associated with even greater elevations of triglycerides, cholesterol, BMI, and fasting insulin in the ClinSeq cohort compared with the WT reference group (all *P* < 0.05; Supplemental Figure 9), although numbers of individuals were too small to make definitive conclusions.

Replication genotyping efforts. Four variants shown to affect protein function were observed multiple times in the ClinSeq cohort (p.Val103Met, p.[Ser183CysfsX34;Ala519Thr], p.Gln234Pro, and p.Thr379AsnfsX36). For these variants, the prospects for identifying additional carriers through population-based genotyping were greatest, and we genotyped them in T2D case-control cohorts. Variants p.Gln234Pro and p.Thr379AsnfsX36 were present in ClinSeq samples of non-Hispanic mixed European ancestry and were therefore genotyped in well-characterized sample sets of Finnish and German origin (*n* = 1800–11000; Supplemental Table 10 and refs. 31, 55–57). Both variants were more common in samples of German origin compared with samples of Finnish origin (p.Gln234Pro, MAF = 0.001 and 0.0001, respectively; p.Thr379AsnfsX36, MAF = 0.002 and 0.0003, respectively). Variant p.[Ser183CysfsX34;Ala519Thr] was observed in 4 ClinSeq samples of Ashkenazi descent but was not detected in 1000 Genomes or Finnish or German samples. This variant had a MAF of 0.006 in 1206 individuals of Ashkenazi descent (Supplemental Methods). p.Val103Met was detected in 2 ClinSeq individuals and in 2 1000 Genomes samples of reported Mexican-American descent. This variant had a MAF of 0.009 in 1,528 samples of Mexican-American ancestry (Supplemental Methods and ref. 58). Accordingly, all of these variants were rare (MAF < 0.01) even when screened in populations matched for ethnicity. Consistent with our phe-



notype analyses, *severe* LOF variants p.Val103Met, p.[Ser183Cys fsX34;Ala519Thr], and p.Thr379AsnfsX36 appeared enriched in individuals with impaired glycemia (Supplemental Table 11).

Discussion

Many observers are predicting that whole-genome sequencing will become part of standard medical care within the next decade (59). That potential has heightened interest in interpretation of genetic variants that might provide insight into assessing future risks of illness, refining present diagnoses, or predicting drug response. Hundreds of common variants shown to be associated with disease risk or quantitative traits have emerged in the last 5 years as a result of the GWAS approach. However, most of these have been shown to have a relatively modest effect on risk, and there appears to be “missing heritability” for many diseases and traits that may at least in part be attributable to rare variants, some of which may have larger effects (2).

While successful discovery of such rare variants has been achieved in some studies by focusing on individuals at the extremes of a quantitative trait (6–8, 10), a major challenge for the future is how to interpret such variants when they occur in a less selected population. For that purpose, we studied a clinical population modestly enriched for cardiovascular disease risk and looked for rare variants in *GCKR*.

Variants in *GCKR* have been previously shown to affect a range of metabolic processes. For common variant p.Pro446Leu, it has been relatively easy to detect such associations because of the availability of phenotype information on large numbers of individuals. As an alternative approach to finding rare alleles, sequencing *GCKR* in individuals with extremely high triglycerides has shown utility in relating such variants collectively to phenotypes (11). However, for *GCKR* or indeed any gene, determining whether a specific mutation is functionally important in an individualized clinical setting remains a major challenge. Bioinformatic methods such as mutation prediction algorithms and assessment of evolutionary conservation are useful preliminary tools in analysis of whether a particular variant might have functional consequences, but these approaches showed limitations in accuracy and consistency between prediction programs both in this study and in the study by Johansen et al. (11).

Accordingly, we utilized existing information about GKR protein function and undertook individual molecular characterization of all 18 *GCKR* variants from the ClinSeq project. We observed defects in cellular localization for the majority of these variants (12/18) as YFP-tagged GKR constructs in HeLa cells and refined our analysis by GCK coexpression and kinetic characterization. Functional rare variants could be broadly subdivided into putative LOF and GOF subtypes (Table 2).

Potential GOF mutations in conserved C-terminal residues of GKR abolished GKR nuclear localization and GCK sequestration, while p.Arg612Cys-GKR showed no significant differences in kinetic properties compared with WT GKR (Figure 2D and Figure 5). This suggests that the region surrounding residues 607–612 could be part of the unknown mechanism by which GKR is localized to the nucleus. As only 2 individuals carried potential GOF mutations (Table 3 and Supplemental Figure 9), further studies will be needed to determine the phenotypic effects of such variants. However, physiologically, these mutations increase cytoplasmic GKR and thus may serve to decrease GCK activity by decreasing both the pool of sequestered nuclear GCK and of active, cytoplasmic GCK. This would be predicted to decrease hepatic glycogen, triglyceride, and cholesterol synthesis.

Analysis of the subset of ClinSeq individuals heterozygous for rare *GCKR* LOF variants collectively showed a significant increase in total cholesterol, LDL cholesterol, and triglyceride levels (Table 3). These variants were both functionally and phenotypically similar to p.Pro446Leu, showing reduced expression, reduced nuclear localization, potential reduction in GCK sequestration, and reduced interaction with F6P and/or F1P. As has been proposed for p.Pro446Leu (13, 43), reduced nuclear localization and GCK sequestration would likely increase fasting hepatic glucose uptake and disposal through synthetic pathways including *de novo* lipogenesis. For p.Pro446Leu, the phenotypic effect was much stronger on triglycerides than on fasting glucose; presumably the same phenomenon is present in our collection of rare LOF variants, as even with small numbers, we were able to detect the effect on lipids, but not on fasting glucose. However, while variants within this group are qualitatively similar, they displayed a range in the magnitude of cellular and kinetic effects.

The most severe loss-of-function variants, such as p.Val103Met, appear to form very little, if any, functional protein, characterized by low cellular fluorescence and dramatically reduced ability to inhibit GCK. Physiologically, these variants may be indistinguishable from null mutations and might therefore be compared with heterozygous *Gckr*-knockout mice. *Gckr*^{+/−} mice show reduced liver GCK levels and activity (24, 25) as well as trends toward lower hepatic glycogen content and higher blood glucose levels 30 and 60 minutes after an oral glucose tolerance test (24). The reduction in glycemic control is likely attributable to a decrease in the Gkr-bound Gck nuclear pool that is normally mobilized in response to a glucose challenge. This loss of nuclear stabilization and/or glucose-dependent translocation has been shown to be associated with impaired glucose tolerance in rodent models (24, 60). Phenotype comparisons and results of genotyping for the severe LOF subgroup were consistent with these findings.

As the common p.Pro446Leu variant has been shown to have both cellular and kinetic effects (13, 43), it is useful to consider the effect of this variant in *cis* or in *trans* with rare variants. Experiments assessing Leu446 in *cis* suggested this variant may amplify defects in protein expression, localization, and activity, but will have fairly mild effects on F1P and F6P interaction. Kinetic results modeling the effect of Leu446 in *trans* suggested intermediate effects on activity and phosphate ester response (Supplemental Table 7). Using our cellular model system, it is difficult to assess the effect of Leu446 in *trans* on GCK sequestration. However, as loss-of-function variants are similar to p.Pro446Leu (Table 2), cellular effects of Leu446 in *trans* with a LOF rare variant are likely to include reduced GCK sequestration (43), GKR activity, and inhibition from both chromosomes. Phenotypic effects may range from those of Leu446 homozygotes (as suggested by GWAS) to those approaching *Gckr*-knockout mice for more severe rare variants.

ClinSeq sequencing, combined with previous studies and emerging 1000 Genomes data, suggests it is unlikely there are additional common nonsynonymous *GCKR* variants in the general western European population (11, 47, 48). Follow-up genotyping confirmed the rarity of individual variants, but highlighted the importance of considering ethnicity for replication of rare variants. Accordingly, our findings supported a collective analysis of rare variants to explore relationships with phenotypes. However, functional characterization revealed that not every variant is likely to have the same biochemical characteristics and therefore the same phenotypic consequences (Table 2). This is an important limitation of *in silico* predictions, as



bioinformatic methods often cannot distinguish between different types of functional variants. The separation of phenotypes based on functional classification in the ClinSeq cohort suggests the mutational load of low-frequency *GCKR* variants may still play a significant role in heritability of human glucose and lipid traits.

Some of the challenges highlighted by this study are likely to be amplified as more sequencing data becomes available. Resequencing studies will identify a large number of rare variants per individual, many of which may be novel. Phenotyping a large number of individuals for each rare variant will often not be practical. Unless there are reliable computational, cell biological, or biochemical methods for determining the functional consequences of a variant (as we have been able to do here with *GCKR*), it will generally be difficult to interpret the significance of rare sequence changes. While the era of complete genome sequencing holds much promise for identifying heritable risk factors and ushering in the era of personalized medicine, the leap from sequence discovery to functional inference and medical consequence will often not be trivial.

Methods

Subject recruitment. ClinSeq participants were evaluated at the NIH Clinical Center or Suburban Hospital (Bethesda, Maryland, USA). The target enrollment was 1,000 nonsmokers between the ages of 45 and 65. Participants were selected to represent the full spectrum of risk for developing coronary artery disease (CAD) using the Framingham score (61). Participants were divided into groups of 250 participants with a Framingham 10-year CAD risk of less than 5%, 5%–10%, or more than 10% as well as an additional group with known CAD.

Clinical and laboratory analysis. Participants were evaluated by an abbreviated medical history focused on cardiovascular disease. Clinical assessment included measurement of height, weight, blood pressure, head and abdominal circumferences, and electrocardiogram, echocardiogram, and measurement of coronary artery calcification (by multidetector computed tomography). Blood and urine were collected to assess baseline lipids, glucose metabolism, renal function, and specific markers for CAD risk. Genomic DNA was isolated by standard techniques for PCR-based Sanger sequencing of *GCKR*. Bidirectional sequence generation and analysis using PolyPhred followed by manual review were performed as previously described (46). Primer sequences are presented in Supplemental Methods.

The cohort consisted of 800 participants enrolled in the ClinSeq study between January 5, 2007, and April 21, 2010. Fifty-seven individuals were excluded from clinical analysis on the basis of T2D or extremely high triglycerides, leaving 743 individuals for analysis. ClinSeq participants were subdivided into genotypic subgroups for comparison of baseline clinical parameters. The mutational status of rs1260326 (c.1337C>T, p.Pro446Leu) was determined by Sanger sequencing or TaqMan assay (for participants with low-quality Sanger sequencing at this nucleotide) for all subjects.

Cloning of fluorescent plasmids. The pcDNA-DEST53 vector (Invitrogen) was modified to express the YFP variant ZsYellow1 or CFP variant AmCyan1 (Clontech) (Supplemental Methods). Plasmids were verified by sequencing. The Mammalian Gene Collection full-length cDNA of human *GCKR* and Xenopus Gene Collection full-length cDNA of *X. laevis* *gckr* (ATCC) were amplified with Phusion DNA polymerase (New England Biolabs) according to the manufacturer's instructions using primers containing attB1 and attB2 recombination sequences (Supplemental Methods). PCR products were recombined in frame for N-terminal fusions into the Gateway entry vector pDONR-201 using Gateway cloning technology (Invitrogen). The Mammalian Gene Collection full-length cDNA of

human β cell *GCK* (ATCC) was also PCR amplified and cloned in frame for N-terminal fusions into pDONR-201. The liver-specific exon 1 and shared exon 2 of *GCK* were synthesized (Integrated DNA Technologies) and cloned into this N-terminal vector, replacing the β cell-specific exon 1 using a SacII restriction site. Mutations in *GCKR* were introduced into the Gateway entry vector using PCR-based site-directed mutagenesis according to the manufacturer's instructions (Agilent Biotechnologies; Supplemental Methods). WT or mutant *GCKR* were recombined into the YFP-expressing Gateway vector. Two independent preparations were generated for each variant; all mutations were verified by sequencing. *GCK* was recombined into the CFP gateway expression vector.

The CFP-*GCK* promoter, ORF, and poly(A) signal were cloned into the YFP Gateway destination vector so that the CMV promoters upstream of CFP and YFP were in opposite orientations (Supplemental Methods). The *GCKR* coding sequence (either WT or mutant) was recombined into this vector to generate a plasmid expressing both YFP-GKRP and CFP-*GCK*. Two independent preparations were generated for each variant and verified by sequencing.

Cell culture and transfection. HeLa cells were cultured in DMEM with 25 mM glucose (Invitrogen) supplemented with 10% FBS and penicillin/streptomycin. 4×10^4 cells were seeded into 4-chamber slides (BD Biosciences) and cultured for 24 hours. Transfection was carried out with a set amount of plasmid DNA (200 ng for YFP-GKRP plasmids; 1 μ g for dual-expression plasmids) and 3 μ l of lipofectamine 2000 (Invitrogen) according to the manufacturer's instructions. Medium was replaced 5 hours after transfection. Slides were fixed with formaldehyde 24 hours after transfection and mounted with VectaShield Mounting Medium (Vector Laboratories). Confocal images were acquired using a Zeiss LSM 510 NLO Meta System mounted on a Zeiss Axiovert 200M microscope with an oil immersion Plan-Apochromat 63 \times /1.4 DIC objective lens. Each preparation was transfected at least 2 times.

Kinetic characterization. Recombinant glutathione S-transferase-tagged (GST-tagged) human pancreatic *GCK* and WT and variant FLAG-tagged GKRP were prepared as described previously (13, 62). Purity and concentration were measured by the Agilent 230 Protein Kit (Agilent Technologies) and Bio-Rad Bradford reagent assay (Bio-Rad Laboratories), respectively. GKRP preparations ranged in yield between 54 and 100 μ g/l broth, and *GCK* yields averaged 1.5 mg/l broth.

GKRP inhibition of *GCK* activity was determined spectrophotometrically using glucose 6-phosphate dehydrogenase-linked (G6PDH-linked) assays (Sigma-Aldrich) as described previously (13). Independent WT GKRP preparations were generated for each variant preparation. Three preparations were generated for p.Gln234Pro, 2 for p.[Pro383Thr;Pro446Leu], and 1 each for p.Arg612Cys, p.Val103Met, p.Ile219Val, p.Ile500Ser, p.Pro446Leu, and p.Pro383Thr. Except for p.Val103Met, assays were standardized by matching WT and variant GKRP activity (1 unit; corresponding to the amount required for half-inhibition of 10 mU/ml *GCK* at 5 mM glucose). On average, one GKRP unit resulted from 3 μ g/ml of WT GKRP. Relative activities are listed in Supplemental Table 6.

Glucose-dependence assays were carried out over a glucose concentration range of 0 to 100 mM. F1P and F6P assays (0–500 μ M) were performed as described previously (13). Both phosphate esters were purchased from Sigma-Aldrich.

Genotyping. *GCKR* variants were genotyped using the iPLEX Sequenom MassARRAY platform on samples of German and/or Finnish origin. The study design and phenotypic characteristics of these samples have been previously described (31, 55–57, 63). All genotyped SNPs had a genotyping call rate of more than 95%, and polymorphic variants had an HWE *P* value of more than 0.001. p.[Ser183CysfsX34;Ala519Thr] and p.Val103Met follow-up genotyping were performed using TaqMan tech-



nology (Applied Biosystems) in samples of Ashkenazi and Mexican-American origin, respectively (Supplemental Methods and ref. 58). c.1555G>A (p.Ala519Thr) was queried to determine p.[Ser183CysfsX34;Ala519Thr] genotype. Two-step PCR was performed on an ABI7900HT machine, before post-read allelic discrimination using SDS2.3 software.

Statistics. HWE was confirmed using a χ^2 test with 1 degree of freedom. HWE calculation was made with the Online Encyclopedia for Genetic Epidemiology Studies HWE calculator (<http://www.oege.org/software/hwe-mr-calc.shtml>). Nonparametric, unpaired (2-tailed) tests were performed to compare baseline clinical measurements in GCKR subgroups. Mann-Whitney U test was used for continuous variables and Fisher's exact test for categorical variables using GraphPad InStat v3.1a. Results are presented as mean \pm SD. Effect estimates of the GCKR Leu446 allele and rare variant subgroups on baseline laboratory measurements were derived from P trend linear regression. Results are shown as estimate \pm SEM and P value for trend. Stepwise multivariate (race, sex, age, BMI, genotype status) logistic regression was used to identify covariates that have a significant effect on baseline clinical laboratory values. Least-square means were derived to adjust for the effects of confounding covariates (Supplemental Methods). Results are shown as mean \pm SEM. P trend tests, stepwise logistic regressions, and calculations of least-square means were performed with SAS v.9.2. For all tests, 2-tailed P values of less than 0.05 were considered significant. Statistical analysis for kinetic experiments utilized paired 2-tailed t tests, with a cutoff for significance of $P < 0.05$.

Study approval. The ClinSeq protocol was approved by the Institutional Review Boards of the National Human Genome Research Institute, NIH, and Suburban Hospital. All study participants gave informed consent upon enrollment.

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