Cellular characterisation of the $GCKR$ P446L variant associated with type 2 diabetes risk

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Abstract
Aims/hypothesis Translation of genetic association signals into molecular mechanisms for diabetes has been slow. The glucokinase regulatory protein ($GKRP$; gene symbol $GCKR$) P446L variant, associated with inverse modulation of glucose- and lipid-related traits, has been shown to alter the kinetics of glucokinase ($GCK$) inhibition. As GCK inhibition is associated with nuclear sequestration, we aimed to determine whether this variant also alters the direct interaction between GKRP and GCK and their intracellular localisation.

Methods Fluorescently tagged rat and human wild-type (WT)- or P446L-GKRP and GCK were transiently transfected into HeLa cells and mouse primary hepatocytes. Whole-cell and nuclear fluorescence was quantified in individual cells exposed to low- or high-glucose conditions (5.5 or 25 mmol/l glucose, respectively). Interaction between GCK and GKRP was measured by sensitised emission-based fluorescence resonance energy transfer (FRET) efficiency.

Results P446L-GKRP had a decreased degree of nuclear localisation, ability to sequester GCK and direct interaction with GCK as measured by FRET compared with WT-GKRP. Decreased interaction was observed between WT-GKRP and GCK at high compared with low glucose, but not between P446L-GKRP and GCK. Rat WT-GKRP and P446L-GKRP behaved quite differently: both variants responded to high glucose by diminished sequestration of GCK but showed no effect of the P446L variant on nuclear localisation or GCK sequestration.

Conclusions/interpretation Our study suggests the common human P446L-GKRP variant protein results in elevated hepatic glucose uptake and disposal by increasing active cytosolic GCK. This would increase hepatic lipid biosynthesis but decrease fasting plasma glucose concentrations and provides a potential mechanism for the protective effect of this allele on type 2 diabetes risk.

Keywords Association study · Genetics · Glucokinase · Molecular mechanism

Abbreviations
$GKRP$ Glucokinase regulatory protein
$hGKRP$ Human GKRP
$rGKRP$ Rat GKRP
$GCK$ Glucokinase
$ECFP$ Enhanced cyan fluorescent protein
$EGFP$ Enhanced green fluorescent protein
$EYFP$ Enhanced yellow fluorescent protein
$FRET$ Fluorescence resonance energy transfer
$FRETN$ Sensitised emission-based FRET efficiency
$F6P$ Fructose 6-phosphate
$F1P$ Fructose 1-phosphate
$WT$ Wild-type
$SNP$ Single nucleotide polymorphism
**Introduction**

The past 4 years have seen a rapid expansion in the number of reproducibly associated loci for type 2 diabetes and related traits. However, progress in translating these genetic associations into biological mechanisms for disease has been slow [1, 2]. Contributing factors have included small effect sizes and the mapping of association signals to poorly annotated, often intergenic, sequences [3]. Signals that include an obvious (e.g. non-synonymous) variant in a biologically plausible gene provide tractable targets for functional follow-up and a framework for exploring approaches to variant characterisation [2].

Glucokinase (GCK) regulates glucose storage and disposal in the liver. Hepatic GCK activity is modulated by glucokinase regulatory protein (GKRP; gene symbol GCKR), a competitive inhibitor of GCK. Fructose 6-phosphate (F6P) and fructose 1-phosphate (F1P) enhance and reduce GKRP-mediated inhibition, respectively [4, 5]. A further layer of regulation exists in the form of cellular localisation: GKRP sequesters GCK to the nucleus at low glucose concentrations, further inhibiting GCK activity and glycolysis [6]. At higher glucose concentrations, glucose-mediated dissociation of GCK from GKRP activates GCK and exposes its nuclear export signal [7, 8].

A common GCKR single nucleotide polymorphism (SNP) (rs1260326 [c.1337C>T; p.P446L]) has been reproducibly associated with an inverse relationship between glucose and triacylglycerol levels [9]. This seemingly paradoxical relationship also extends to other glucose- and lipid-related traits. The glucose-lowering leucine allele is also associated with decreased fasting insulin, HOMA-estimated insulin resistance and reduction of type 2 diabetes risk [10] but increased triacylglycerol and total cholesterol levels [11]. This variant is one of only a handful of candidate non-synonymous variants identified to date from type 2 diabetes association studies [2, 12]. We have previously shown that variant P446L-GKRP increases GCK activity as a result of diminished regulation by F6P and have proposed that this results in increased hepatic glycolytic flux and de novo lipogenesis, accounting for the observed associated phenotypes [12]. Interestingly, this kinetic effect was not seen with rat P446L-GKRP [13], likely due to species differences in regulation by F6P [14].

However, kinetic mechanisms alone are not sufficient to explain GKRP regulation of GCK in vivo. For example, Gckr-knockout mice display impaired postprandial glucose handling and paradoxical reductions in GCK protein levels and activity in spite of the removal of GKRP inhibition [15, 16]. This suggests that nuclear interaction with GKRP plays an important role in regulating GCK protein concentration and stability and is essential for the maintenance of glucose homeostasis. In addition, the P446L variant is associated with inverse modulation of fasting and 2 h glucose levels, an observation not readily explained by kinetic data [10]. As the contribution of GKRP and GCK cellular localisation to phenotypes associated with the P446L-GKRP variant protein is currently unexplored, the aims of the current study were to assess the impact of this variant on the ability of GKRP to sequester GCK in the nucleus and to further explore potential differences between rat and human regulatory proteins.

**Methods**

**Cloning of fluorescent fusion plasmids** Human GCK (Mammalian Gene Collection; ATCC, Manassas, VA, USA) was amplified using primers containing BamHI and HindIII restriction sites and subcloned into the pEGFP-c3 vector encoding enhanced green fluorescent protein (EGFP). Generation of enhanced cyan fluorescent protein (ECFP)-tagged GCK (pECFP-GCK) has been described by Baltrus et al. [17]. PCR primers containing Eco47III and EcoRI sites were used to amplify human GCKR (Mammalian Gene Collection) and replace rat Gckr in pEYFP-N1-GCKR [17]. The P446L mutation was introduced into rat and human pEYFP-N1-GCKR plasmids by PCR-based site-directed mutagenesis (Agilent Biotechnologies, Santa Clara, CA, USA). The enhanced yellow fluorescent protein (EYFP) in both rat and human pEYFP-N1-GCKR [17] was replaced by mCherry [18] using AgeI and BsrGI restriction sites. Two independent preparations of all plasmids were generated and verified by sequencing. Primer sequences are available upon request.

**Cell culture and transient transfection of HeLa cells** HeLa cells were grown in DMEM with 5.5 or 25 mmol/l glucose supplemented with 10% (vol./vol.) dialysed fetal bovine serum and penicillin/streptomycin (all Invitrogen, Carlsbad, CA, USA). For quantitative fluorescence microscopy, 4×10⁴ cells were seeded into four-chamber slides (BD Biosciences, San Jose, CA, USA) and cultured for 24 h. Co-transfection of mCherry-GCKR and EGFP-GCK was initially carried out with a range of plasmid DNA concentrations at ten separate GCK/GCKR molar ratios between 40:1 and 1:40. Nuclear localisation of GCK was observed for GCK/GCKR ratios between 1:1 and 1:40. A ratio of 1:4 was used for all future experiments as this is in agreement with the published GCK/GCKR mRNA ratio in liver [12]. Optimal transfection efficiency and fluorescence intensity was observed with transfection of 100 ng total DNA and 0.75 μl of Lipofectamine 2000 (Invitrogen). Transfection was carried out according to the manufacturer’s instructions at 5.5 mmol/l glucose. The medium was replaced 5 h after transfection.
Slides were incubated at 5.5 or 25 mmol/l glucose at 22 h post-transfection and fixed with formaldehyde 24 h post-transfection for mounting with VectaShield Mounting Medium with DAPI (Vector Laboratories, Burlingame, CA, USA). Each preparation condition was transfected two times. For fluorescence resonance energy transfer (FRET) studies, cells were seeded at a density of 2×10^5 cells on 35 mm glass-bottom dishes (MatTek, Ashland, MA, USA) and grown for 24 h. Thereafter, cells were transfected with 1 μg DNA (1:1 ratio of the EYFP and ECFP plasmid) and 2 μl jetPEI (Qbiogene, Montreal, QC, Canada) according to the manufacturer’s instructions. The culture medium was replaced 1 day after transfection, and the cells were incubated for another 24 h with medium containing 5.5 or 25 mmol/l glucose.

Hepatocyte isolation, culture, and transient transfection
Primary hepatocytes were isolated from C57/BL6 mice using a modification of the collagenase perifusion methods of Seglen [19] and Bader et al. [20]. Isolated cells were suspended in Williams’ medium E supplemented with 10 mmol/l glucose, 5% (vol./vol.) fetal calf serum, 1×10^-4 mmol/l dexamethasone, and 1×10^4 pmol/l insulin. Hepatocytes were seeded at a density of 4×10^5 cells on 35 mm collagen-coated glass-bottom dishes (MatTek, Ashland, MA, USA) or on 12-well plates and incubated for 24 h in a humidified atmosphere at 37°C and 5% CO₂. Cells were transfected with 1 μg DNA (either EYFP plasmid alone or 1:1 ratio of the EYFP and ECFP plasmid) and 3.2 μl PromoFectine Hepatocyte (PromoCell, Heidelberg, Germany) according to the manufacturer’s instructions. Thereafter, hepatocytes were cultured for 24 h and finally incubated in medium with either 5.5 or 25 mmol/l glucose for 2 h.

Fluorescence microscopy and image analysis
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×/1.4 DIC objective lens. Excitation wavelengths of 488 nm (2%), 561 nm (5%) and 770 nm (3%) were used for detection of EGFP-GCK, mCherry-GKRP and DAPI, respectively. Confocal images were post-processed using the MediaCybernetics Image-Pro Plus v7.0 software package (Bethesda, MD, USA) with a custom macro designed to calibrate, segment (nuclei touching the image border were not included) and outline the nuclei, outline the cell border, count and record measurements. Measurements included nucleus and cell areas (square micrometres) and fluorescent intensity (minimum, maximum, mean, standard deviation and sum). The nuclei outlines were obtained by using DAPI staining as the template and propagating that outline automatically determined with a gradient method from transmitted light images. Finally, the mean fluorescence was calculated as average grey value in selected regions in the nucleus and cytoplasm in individual cells after background correction. The sensitised emission-based FRET efficiency (FRETN) was calculated from the ECFP nuclear:cytoplasmic ratio, glass-bottom dishes were fixed on the microscope stage and images were obtained with an UPLSAPO 60×1.35 numerical aperture oil-immersion objective (Olympus). For calculation of the EYFP nuclear to cytoplasmic ratio on 12-well plates, an UPLSAPO 20×0.75 numerical aperture objective (Olympus) was used and all motorised devices of the microscope system were synchronised by the scanR acquisition software (Olympus). The image focus was automatically determined with a gradient method from transmitted light images. Finally, the mean fluorescence was calculated as average grey value in selected regions in the nucleus and cytoplasm in individual cells after background correction. The sensitised emission-based FRET efficiency (FRETN) was calculated from the ECFP emission with excitation at 436 nm, EYFP emission with excitation at 436 nm and EYFP emission with excitation at 500 nm, based on the calculation of Vanderklish et al. [22]. FRETN measurements were validated using an ECFP-EYFP fusion construct as described by Langer et al. [21].

Statistical analysis
Statistical analyses were performed by ANOVA followed by Bonferroni’s test for multiple comparison or t tests for independent samples using the Prism analysis program (Graphpad, San Diego, CA, USA). Data are presented as means±SEM. p values <0.05 (two-tailed) were considered significant.

Results
Localisation of GKRP fluorescent fusion proteins in HeLa cells and mouse primary hepatocytes
To investigate the
subcellular localisation of human GKRP (hGKRP) and the effect of the P446L variant on this localisation, mCherry-tagged GCX fusion plasmids were transiently transfected into HeLa cells. HeLa cells were chosen as an established model known to exhibit efficient glucose uptake and lacking endogenous expression of GCKR and GCK [7, 23]. Both wild-type (WT)-hGKRP and P446L-hGKRP localised strongly to the nucleus, but an appreciable increase in cytoplasmic fluorescence intensity was detectable for P446L-hGKRP (Fig. 1a, b). To quantify this apparent difference in localisation, the ratio of nuclear to whole-cell fluorescence in individual cells was calculated. Cells with very low EGFP or mCherry fluorescence intensities (<50,000 arbitrary fluorescence units) were excluded from statistical analysis. However, exclusion of these data did not change the significance of any result. WT-hGKRP (n=20 cells) was associated with significantly increased nuclear localisation when compared with P446L-hGKRP (n=21) (83.4±1.4% nuclear localisation compared with 65.1±3.1%, mean±SEM; p<0.001; Fig. 1c). Previous studies have revealed substantial kinetic differences between human and rat GKRP (rGKRP) [14]. However, no studies have compared cellular localisation between rat and human regulatory proteins. Accordingly, we also generated and transiently transfected mCherry-tagged rat Gckr. The P446L variant was also introduced into the rat Gckr sequence. There was no significant difference in the nuclear-to-whole-cell-fluorescence ratio of WT-rGKRP (62.8±2.3%; n=20) and P446L-rGKRP (56.3±1.5%; n=26; Fig. 1d). Both WT-rGKRP and P446L-rGKRP showed significantly decreased nuclear localisation compared with WT-hGKRP (p<0.001; ANOVA/Bonferroni correction). Localisation of all GKRPs was independent of glucose concentration.

Co-transfection of human fluorescent GCK and GKR fusion plasmids in HeLa cells Transiently transfected human EGFP-GCK localised entirely to the cytoplasm of HeLa cells (Fig. 2a). In accordance with previous in vivo studies, localisation was not affected by glucose concentration (data not shown) [15]. Co-transfection of GCK and human or rat WT- and P446L-GKRP resulted in nuclear sequestration of GCK (Fig. 2b, c). Incubation for 2–16 h at a series of glucose concentrations (1–25 mmol/l) resulted in a mix of nuclear and cytoplasmic localisation for both GKRP and GCK (data not shown). Complete glucose-dependent export of GCK was not observed even at high glucose concentrations, consistent with previous observations in rat hepatocytes [17, 24]. Accordingly, quantitative image analysis was undertaken at two glucose concentrations (5.5 and 25 mmol/l glucose) which have been previously used to investigate the GCK:GKRP interaction as they are either side of the glucose S0.5 (substrate concentration needed for half maximal rate) for GCK (approximately 7.5 mmol/l) [17].

Quantitative analysis was undertaken on a total of 787 cells from two transfections per glucose concentration and from two independent plasmid preparations per variant. In the presence of GCK at 5.5 mmol/l glucose, P446L-hGKRP was again associated with decreased nuclear localisation compared with WT-hGKRP (Fig. 2d; 67.4±2.5% vs 73.6±1.8%

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**Fig. 1** Localisation of GKRP fluorescent fusion proteins at 5.5 mmol/l glucose. Visualisation of mCherry-WT-hGKRP (a) or P446L-hGKRP (b) and overlap with DAPI on transient transfection into HeLa cells. c Quantification of degree of nuclear fluorescence for human and rat GKRPs in HeLa cells. Data are expressed as means±SEM of two individual experiments with a total of 20–26 cells analysed. d Quantification of nuclear/cytoplasmic ratio of human and rat GKRPs on transient transfection into primary mouse hepatocytes. Data are expressed as means±SEM of two individual experiments with a total of 36–52 cells analysed. ***p<0.001 compared with WT-hGKRP (ANOVA/Bonferroni correction)
nuclear fluorescence; \( p=0.04 \)). Human GCK showed significantly decreased nuclear localisation in the presence of P446L-hGKRP (Fig. 2d; 57.5±2.1% vs 80.4±1.8% nuclear fluorescence; \( p<0.001 \)). The nuclear localisation of GCK remained significantly reduced in the presence of P446L-hGKRP compared with WT-hGKRP when individual cells were matched for mCherry, EGFP or mCherry:EGFP fluorescence intensity (\( p<0.05 \) for all groups).

There was no significant difference in GCK sequestration between high and low glucose for either human variant (Fig. 3). The differences in GCK sequestration between WT-hGKRP and P446L-hGKRP observed at 5.5 mmol/l glucose were also detectable at 25 mmol/l glucose (\( p<0.001 \) for GKRP and GCK localisation). By comparison, transient co-transfection of rGKRPs with human GCK resulted in no difference in GCK sequestration between WT-rGKRP and P446L-rGKRP (\( p>0.1 \) for both glucose concentrations). However, there was a significant glucose dependence in GCK sequestration by both rGKRPs, with increased GCK nuclear localisation observed at low glucose (Fig. 3; \( p<0.05 \) for both WT-rGKRP and P446L-rGKRP). These findings are not likely due to differences in rat and human GCK, as previous kinetic studies have extensively characterised the interaction between rat and human GKRP and either rat or human GCK, showing no difference in interaction of GKRP with either rat or human GCK [14].

**Fig. 2** Localisation of co-transfected human GCK and GKRP in HeLa cells at 5.5 mmol/l glucose. **a** Visualisation of human EGFP-GCK and DAPI. **b,c** Co-transfection of mCherry-WT-hGKRP and EGFP-GCK (b) or mCherry-P446L-hGKRP and EGFP-GCK (c): GKRP signal, upper left; GCK signal, upper right; DAPI, lower left; overlay, lower right. **d** Quantification of degree of nuclear fluorescence of GKRP and GCK. Data are expressed as means±SEM of two individual experiments with a total of 77–98 cells analysed. *\( p<0.05 \); ***\( p<0.001 \) \( (t \) test). Black bars, GKRP; white bars, GCK

**Fig. 3** Glucose dependence of GCK localisation in HeLa cells. Degree of nuclear localisation of human GCK transiently transfected in the presence of WT-hGKRP, P446L-hGKRP, WT-rGKRP or P446L-rGKRP at 5.5 mmol/l and 25 mmol/l glucose. Data are expressed as means±SEM of two individual experiments with a total of 18–98 cells analysed. *\( p<0.05 \); ***\( p<0.001 \) (ANOVA/Bonferroni correction). White bars, 5.5 mmol/l glucose; black bars, 25 mmol/l glucose.
Direct interaction of human GCK and GKRP in HeLa cells

FRETN was used to detect the interaction between ECFP-GCK and EYFP, EYFP-WT-hGKRP, or EYFP-P446L-hGKRP in the nucleus of HeLa cells. FRETN measures the transfer of energy from ECFP to EYFP on excitation of ECFP and occurs when these proteins are within close proximity (<10 nm) because of the overlap of the emission spectrum of ECFP and the excitation spectrum of EYFP [22]. There was a significant reduction in interaction with GCK for the P446L-hGKRP variant protein compared with WT-hGKRP at 5.5 mmol/l glucose (Fig. 4). WT-hGKRP showed significantly reduced interaction with GCK at 25 mmol/l glucose compared with 5.5 mmol/l. There was no significant glucose dependence of interaction for P446L-hGKRP.

Discussion

To capitalise on recent advances in our understanding of the genetic basis of diabetes risk, it is necessary to translate association signals into detailed molecular mechanisms that may underlie disease predisposition. We have previously shown that P446L-hGKRP, inversely associated with triacylglycerol levels and glucose levels, affects GCK activity in liver through diminished regulation by F6P [12]. However, the impact of this variant on the cellular localisation of GCK and GKRP has not previously been explored. Recent studies using rat hepatocytes have suggested reliable estimation of GCK localisation can be accomplished by quantification of localisation in multiple fields of cells [25, 26]. We therefore sought to image a large number of cells using a quantitative method to assess nuclear and cytoplasmic fluorescence, and to combine this with assessment of direct interaction of GKRP and GCK by measuring FRETN. We have demonstrated for the first time differences in the degree of nuclear sequestration of GCK between WT-hGKRP and P446L-hGKRP and in the direct cellular interaction of P446L-hGKRP with GCK compared with WT-hGKRP. In spite of the presence of endogenous mouse GCK and GKRP, we were also able to demonstrate these differences in localisation, sequestration and direct interaction in primary mouse hepatocytes.

In combination with our previously described kinetic studies, the P446L-hGKRP variant protein would be expected to result in an increased pool of active cytoplasmic GCK, particularly in the presence of low (i.e. fasting state) glucose concentrations. Our findings suggest a mechanism of action for P446L-hGKRP consistent with the associated clinical phenotypes [9–11]. The reduced ability of P446L-hGKRP to sequester and interact with ECFP-GCK. This may be due to an abundance of EYFP-P446L-hGKRP protein in the cytoplasm preventing GCK from localising to the nucleus even at low glucose concentrations, as suggested by our assessment of GCK localisation in HeLa cells (Fig. 3). As in HeLa cells, we observed significantly decreased interaction between GCK and P446L-hGKRP compared with WT-hGKRP at 5.5 mmol/l glucose in hepatocytes (Fig. 5b). At 25 mmol/l glucose in hepatocytes, accelerated metabolism results in generation of high levels of NADH and NADPH, resulting in autofluorescence. Additionally, there is a reduction in the absolute value of FRETN compared with HeLa cells due to quenching by endogenous mouse GCK and GKRP. Accordingly, the reduced interaction between GCK and GKRP at 25 mmol/l glucose was not significantly detectable above this increased background (Fig. 5b).
GCK in the nucleus in the fasting state would result in increased hepatic glucose disposal, glycolytic flux and generation of precursors for synthesis of molecules such as triacylglycerols and cholesterol. Recent findings associating GCKR with increased VLDL particle concentrations [27] suggest this enhancement of glycolytic flux leads to increased levels of de novo triacylglycerol and cholesterol synthesis and export.

Increased fasting-state hepatic glucose uptake, use and disposal by GCK in the presence of P446L-hGKRP would be predicted to decrease circulating plasma glucose concentrations. Further support for this molecular mechanism comes from at least one cellular study linking over-expression of GCK in liver to reduced glucose levels and raised triacylglycerol levels [28]. We propose that decreased fasting plasma glucose levels could result in decreased basal insulin output by the pancreas, which could ultimately contribute to higher insulin sensitivity and decreased risk of type 2 diabetes in spite of increased hepatic lipid production.

Our results also suggest that, compared with WT-hGKRP, P446L-hGKRP shows reduced glucose dependence in its cellular interaction with GCK (Fig. 4). Accordingly, in spite of higher fasting-state GCK activity in the presence of P446L-hGKRP, GCK may not be activated to the same extent by increased glucose in the presence of this variant. This provides a mechanistic explanation for the observation that the P446L variant is associated with increased 2 h glucose levels but decreased fasting glucose levels [10, 29].

Interestingly, the degree of nuclear localisation of rGKRP is significantly reduced compared with WT-hGKRP in both HeLa cells and hepatocytes. Our studies assessing the localisation of transfected rGKRP in mouse hepatocytes (nuclear/cytoplasmic ratio of 3.37 at 5.5 mmol/l glucose) are highly consistent with our observations of endogenous rGKRP in rat hepatocytes (nuclear/cytoplasmic ratio of 3.44 at 5.5 mmol/l glucose; S. Baltrusch, unpublished observations). However, the abnormality in nuclear GCK sequestration for P446L-GKRP appears to be specific to the human regulatory protein as we did not observe a similar phenomenon with rGKRP. This is consistent with previous studies that have established important species-specific differences in both the regulation of human and rat GKRP by F6P and the impact of the P446L variant on this regulation [12–14]. Taking into account our previous assessment of the interaction of rGKRP with human GCK by FRETN analysis in COS-1 cells [21], direct interaction determined in our experiments in living cells appeared to be strongest between hGKRP and GCK. This is consistent with in vitro studies highlighting hGKRP as a more potent inhibitor of GCK than rGKRP [14]. These studies serve to emphasise the differences between human and rat regulatory proteins and highlight the need for use of appropriate assays to decipher complex mutational mechanisms driving disease associations.

It has previously been reported that glucose-dependent GCK localisation is highly heterogeneous in rat hepatocytes [25], thus necessitating careful analysis in living and fixed cells. Our results comparing the glucose dependence of the nuclear/cytoplasmic ratio for human GCK co-transfected with rGKRP in mouse hepatocytes resulted in a ratio at 25 mmol/l that was 63% of the ratio at 5.5 mmol/l glucose. This is consistent with the ratio calculated from the results of Watanabe et al. [25] analysing endogenous rat GCK (59%) and our previous results measuring the nuclear/cytoplasmic ratio of transfected human GCK in rat hepatocytes.
hepatocytes (61%) [17]. Translocation of human GCK in the presence of hGKRP has not been previously explored, but our results suggest there are important differences between the rat and human GKR translocation systems, and that regulation of translocation in humans will be strongly dependent on the genotype at P446L. Extensive assessment in human hepatocytes of appropriate genotype may provide useful future insight into these questions.

In conclusion, we have demonstrated that the human GCKR P446L variant alters the ability of GKR to sequester GCK in the nucleus. This observation, coupled with our previous kinetic studies, provides a mechanistic link between a common genetic variant associated with diabetes risk and metabolic traits and glucose and triacyl-glycerol metabolism.

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Contribution statement MGR wrote the manuscript, contributed to experimental design, and researched and analysed data. SW analysed data and contributed to the manuscript. JS researched and analysed data and contributed to the manuscript. RW researched and analysed data and contributed to the manuscript. NLB researched and analysed data and reviewed and edited the manuscript. SB researched and analysed data and contributed to the manuscript. All authors approved the final version to be published.

Duality of interest The authors declare that there is no duality of interest associated with this manuscript.

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