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# A Low-Frequency Inactivating *AKT2* Variant Enriched in the Finnish Population Is Associated With Fasting Insulin Levels and Type 2 Diabetes Risk

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To identify novel coding association signals and facilitate characterization of mechanisms influencing glycemic traits and type 2 diabetes risk, we analyzed 109,215 variants derived from exome array genotyping together with an additional 390,225 variants from exome sequence in up to 39,339 normoglycemic individuals from five ancestry groups. We identified a novel association between the coding variant (p.Pro50Thr) in AKT2 and fasting plasma insulin (FI), a gene in which rare fully penetrant mutations are causal for monogenic glycemic disorders. The low-frequency allele is associated with a 12% increase in FI levels. This variant is present at 1.1% frequency in Finns but virtually absent in individuals from other ancestries. Carriers of the FI-increasing allele had increased 2-h insulin values, decreased insulin sensitivity, and increased risk of type 2 diabetes (odds ratio 1.05). In cellular studies, the AKT2-Thr50 protein exhibited a partial loss of function. We extend the allelic spectrum for coding variants in AKT2 associated with disorders of glucose homeostasis and demonstrate bidirectional effects of variants within the pleckstrin homology domain of AKT2.

The increasing prevalence of type 2 diabetes is a global health crisis, making it critical to promote the development of more efficient strategies for prevention and treatment (1). Individuals with type 2 diabetes display both pancreatic  $\beta$ -cell dysfunction and insulin resistance. Genetic studies of surrogate measures of these glycemic traits can identify variants that influence these central features of type 2 diabetes (2), highlighting potential pathways for therapeutic manipulation. Comprehensive surveys of the influence of

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This article contains Supplementary Data online at http://diabetes. diabetesjournals.org/lookup/suppl/doi:10.2337/db16-1329/-/DC1. common genetic variants on fasting plasma glucose (FG) and fasting plasma insulin (FI) have highlighted defects in pathways involved in glucose metabolism and insulin processing, secretion, and action (3). Recent studies have identified type 2 diabetes–associated alleles that are common in one population but rare or absent in others (4–6). These associations were observed either due to an increase in frequency of older alleles based on population dynamics and demography (5) or the emergence of population-specific alleles (4,6).

We set out to identify and characterize low-frequency allele (minor allele frequency [MAF] <5%) glycemic trait associations by meta-analysis of exome sequence and exome array genotype data in a multiancestry sample. We also performed in vitro functional studies of protein expression, localization, and activity to understand the consequences of our novel findings.

## **RESEARCH DESIGN AND METHODS**

#### **Genetic Association Studies**

#### Study Samples

The Genetics of Type 2 Diabetes (GoT2D) and Type 2 Diabetes Genetic Exploration by Next-generation sequencing in multi-Ethnic Samples (T2D-GENES) Consortia were initially designed to evaluate the contribution of coding variants to type 2 diabetes risk (7). We performed a discovery association analysis to find novel coding variants associated with fasting glycemic traits in 14 studies from GoT2D that contributed exome array information on 33,231 individuals without diabetes of European ancestry. Further discovery analysis was performed with GoT2D and

A.L.G. and C.M.L. jointly directed this research. A.Man., H.M.H., J.G., X.S., T.Tuk., and P.Fo. contributed equally to this work.

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T2D-GENES with exome sequence data (average  $80 \times$  coverage) in five ancestral groups comprised of 12,940 individuals (6,504 with type 2 diabetes, 6,436 without) with measured FG or FI levels available in 2,144 European, 508 South Asian, 1,104 East Asian, 844 Hispanic, and 508 African American individuals without diabetes. We performed a replication analysis and an assessment of allele frequency distributions in 5,747 individuals from four Finnish cohorts: The Cardiovascular Risk in Young Finns Study (YFS) (8), Helsinki Birth Cohort Study (HBCS) (9), Health 2000 GenMets Study (GenMets) (10), and National FINRISK Study 1997 and 2002 (11). We also assessed the allele frequencies of novel findings in 46,658 individuals from the Cohorts for Heart and Aging Research in Genomic Epidemiology (CHARGE) Consortium with available exome array data (12), although none of the studies passed our quality control filter of a minor allele count (MAC) greater than 5 for inclusion in our replication analysis. See Supplementary Table 1 for study details, sample characteristics, ascertainment criteria, detailed genotype calling, and quality control procedures for each cohort. The relevant institutional review boards, conducted according to the Declaration of Helsinki, approved all human research, and all participants provided written informed consent. A detailed description of ethics permissions is provided in the Supplementary Data.

#### Phenotypes

For the discovery and replication analysis, we excluded individuals from the analysis if they had a diagnosis of type 2 diabetes, were currently receiving oral or injected diabetes treatment, had FG measures  $\geq$ 7 mmol/L, had 2-h postload glucose measures  $\geq$ 11.1mmol/L, or had HbA<sub>1c</sub> measures  $\geq$ 6.5% (48 mmol/mol). Additional exclusions occurring at the study level included pregnancy, nonfasting at time of exam, type 1 diabetes, or impaired glucose tolerance. See Supplementary Table 1A for details. Within each study, we adjusted FG and log-transformed FI levels for age, sex, BMI, and additional study-specific covariates. We applied rank-based inverse-normal transformations to study- or ancestry-specific residuals to obtain satisfactory asymptotic properties of the exome-wide association tests.

We tested for genetic associations with type 2 diabetes, hypertension (HTN), and other related quantitative traits in the Finnish discovery and replication cohorts. We analyzed lipid levels (total cholesterol, HDL cholesterol, LDL cholesterol, and triglycerides), blood pressure (systolic [SBP] and diastolic [DBP] blood pressure and HTN), height, BMI, central adiposity measures (waist-to-hip ratio, waist circumference, hip circumference), adiponectin level, 2-h insulin level, and Matsuda index, which is known to correlate with whole-body insulin sensitivity as measured by the hyperinsulinemic-euglycemic clamp (r = 0.7,  $P < 1.0 \times$  $10^{-4}$ ) (13). For quantitative traits and HTN, we adjusted for age, sex, BMI (for glycemic, blood pressure and central adiposity measures), stratified by type 2 diabetes status and sex (for central adiposity measures) within study. We adjusted LDL and total cholesterol for use of lipid-lowering medication, by dividing total cholesterol by 0.8 if on lipidlowering medication, prior to calculating LDL cholesterol using the Friedewald equation (14). SBP and DBP were adjusted for use of blood pressure-lowering medication by adding 15 mmHg to SBP and 10 mmHg to DBP measurements if an individual reported taking blood pressurelowering medication (15). The Matsuda index was log transformed and analyzed in individuals without diabetes only. After adjusting for covariates, traits were inversenormalized within strata. In addition to studying these metabolic outcomes, we used ICD codes to query electronic medical records in the METSIM (METabolic Syndrome In Men) study and FINRISK 1997 and 2002 cohorts (in all individuals regardless of type 2 diabetes status) and categorized affection status for lipodystrophy, polycystic ovary disease, and ovarian or breast cancer.

## Statistical Analysis

Discovery Analysis. We performed association analyses within each study for the exome array data sets and within ancestry for the exome sequence data sets. We used linear mixed models implemented in EMMAX (16) to account for relatedness. Within each study/ancestry, we required variants to have a MAC greater than or equal to five alleles for single variant association tests. We meta-analyzed the single variant results from the (European ancestry) exome array studies using the inverse-variance meta-analysis approach implemented in METAL (17) and combined these with the European ancestry exome sequence results. Then, we metaanalyzed summary statistics across ancestries. We used P < $5 \times 10^{-7}$  as exome-wide statistical significance thresholds for the single variant tests (18). We used the binomial distribution to assess enrichment of previously reported associations with FG or FI by calculating a P value for the number of nonsignificant variants with consistent direction of effects.

Gene-Based Association Analysis. We performed genebased association tests using variants with MAF <1%(including rare variants with MAC  $\leq$ 5), annotating and aggregating variants based on predicted deleteriousness using previously described methods (7). Briefly, we defined four different variant groupings: "PTV-only," containing only variants predicted to severely impair protein function; "PTV+missense," containing protein-truncating variants (PTV) and nonsynonymous (NS) variants with MAF <1%; "PTV+NS<sub>strict</sub>," composed of PTV and NS variants predicted damaging by five algorithms (SIFT, LRT, MutationTaster, PolyPhen-2 HDIV, and PolyPhen-2 HVAR); and "PTV+NSbroad," composed of PTV and NS variants with MAF <1% and predicted damaging by at least one prediction algorithm above. We used the sequence kernel association test (SKAT) (19) and a frequency-weighted burden test to conduct exome array meta-analyses in an unrelated subset of individuals using RAREMETAL (20). We conducted exome sequence gene-based analyses within ancestry using a linear mixed model to account for relatedness and combined results across ancestries with MetaSKAT

(21), which accounts for heterogeneous effects. We further combined gene-based results from exome array and exome sequences using the Stouffer method with equal weights. For gene-based tests, we considered  $P < 2.5 \times 10^{-6}$  as exome-wide significant, corresponding to Bonferroni correction for 20,000 genes in the genome (18).

**Replication Analysis.** The *AKT2* p.Pro50Thr variant was observed at sufficient frequency in the independent Finnish cohorts to perform single variant association test of association with FI. We tested association in SNPTEST (22) (v.2.4.0) in each study with the same additive linear model used in the discovery analysis. Covariate adjustments for FI levels were sex, age, and 10 principal components, and models were run with and without adjustment for BMI.

**Estimate of Effect on Raw FI Level and Variance Explained.** To characterize the association between *AKT2* p.Pro50Thr and FI, we examined full regression models with raw FI in three studies (Finland-United States Investigation of NIDDM Genetics [FUSION], METSIM, and YFS). We estimated the raw effect on log-transformed FI levels with a fixed-effects meta-analysis. The variance in log-transformed FI explained by *AKT2* p.Pro50Thr was estimated by a weighted average of the narrow-sense heritability of *AKT2* p.Pro50Thr seen in these three studies.

**Population Genetics and Constraint.** We used the Exome Aggregation Consortium (ExAC) for constraint metrics and allele frequencies (23). We obtained sequence alignments for AKT proteins and mRNAs in 100 vertebrates from the University of California, Santa Cruz Genome Browser (24), used Shannon entropy (normalized K = 21) as a conservation score (25), and plotted the sequence logos in R using the RWebLogo library (26).

Associations With Other Traits. We conducted association tests for traits other than FI and FG within studies for both discovery studies as well as the independent Finnish studies used for replication. P values for type 2 diabetes and HTN came from EMMAX (16) or the Wald test from logistic regression (Finnish replication data sets) and meta-analyzed using an N-weighted meta-analysis (17). Odds ratios (ORs) were obtained from logistic regression adjusting for age, sex, with and without BMI, and principal components and meta-analyzed using an inverse-variance meta-analysis. Trait Distributions and Phenotype Clustering. We examined distributions of traits among AKT2 missense allele carriers (p.Pro50Thr, p.Arg208Lys, and p.Arg467Trp) in the T2D-GENES exome sequencing data set. We used nonparametric rank-based methods (kruskal.wallis and permKS functions in R) on both the inverse-normalized covariateadjusted traits used in the genetic association studies and normalized raw trait values (scale function in R). We clustered AKT2 missense allele carriers on scaled trait values (pheatmap function in R).

## In Vitro Functional Studies

## Plasmids and Cell Lines

The generation of the *AKT2* allelic series was initiated by the production of pDONR223-AKT2 through PCR of the

human AKT2 open reading frame with the integration of terminal attR sites using primers (see below). HeLa, HuH7, and 293T cells were obtained at The Broad Institute and maintained in 10% FBS DMEM, 100 units/mL penicillin and 100  $\mu$ g/mL streptomycin, and documented mycoplasma-free. HeLa and HuH7 cells were starved for 18 h and stimulated for 15 min with 100 nmol/L insulin for activation analyses.

## Primers for Functional Work

The generation of the AKT2 allelic series was initiated by the production of pDONR223-AKT2 through PCR of the human AKT2 open reading frame with the integration of terminal attR sites using primers FWD: 5'-GGGGACAAGT-TTGTACAAAAAGTTGGCACCATGAATGAGGTGTCTGTCA-TC-3', REV: 5'-GGGGACCACTTTGTACAAGAAAGTTGG-CAACTCGCGGATGCTG-3' and subsequent Gateway BP reaction into pDONR223 obtained from The Broad Institute Genetic Perturbation Platform. Site-directed mutagenesis was then performed to generate AKT2.E17K (AKT2. Lys17), AKT2.P50T (AKT2.Thr50), AKT2.R208K (AKT2. Lys208), AKT2.R274H (AKT2.His274), AKT2.R467W (AKT2.Trp467) with the following primers: AKT2.E17K: FWD: 5'-GGCTCCACAAGCGTGGTAAATACATCAAGACCT-GG-3', REV: 5'-CCAGGTCTTGATGTATTTACCACGCTTG-TGGAGCC-3'; AKT2.P50T: FWD: 5'-AGGCCCCTGATCA-GACTCTAACCCCCTTAAAC-3', REV: 5'-GTTTAAGGGG-GTTAGAGTCTGATCAGGGGCCT-3'; AKT2.R208K: FWD: 5'-GTCCTCCAGAACACCAAGCACCCGTTCC-3', REV: 5'-GGAACGGGTGCTTGGTGTTCTGGAGGAC-3'; AKT2. R274H: FWD: 5'-GGGACGTGGTATACCACGACATCAAGCT-GGA-3', REV3'REV: 5'-TCCAGCTTGATGTCGTGGTATAC-CACGTCCC-3'; and AKT2.R467W: FWD: 5'-GGAGC-TGGACCAGTGGACCCACTTCCC-3', REV: 5'-GGGAAG-TGGGTCCACTGGTCCAGCTCC-3'. COOH-terminal, V5tagged lentiviral pLX304-AKT2.E17K, pLX304-AKT2.P50T, pLX304-AKT2.R208K, pLX304-AKT2.R274H, and pLX304-AKT2.R467W were each generated by subsequent Gateway LR reactions with pDONR223-AKT2.E17K, pDONR223-AKT2.P50T, pDONR223-AKT2.R208K, pDONR223-AKT2. R274H, and pDONR223-AKT2.R467W, respectively, and pLX304 obtained from The Broad Institute Genetic Perturbation Platform. Control plasmid pLX304-empty vector was additionally acquired from The Broad Institute Genetic Perturbation Platform.

## Antibodies

Anti-Akt (#4685), anti–phospho-Akt S473 (#4060), anti– phospho-Akt T308 (#9275), anti– $\beta$ -actin (#4970), anti-GSK3 $\beta$  (#9315), anti–phospho-GSK3 $\beta$  (#9336), anti-GST (#2625), and anti-V5 (#13202) were purchased from Cell Signaling Technologies. Horseradish peroxidase–conjugated anti-rabbit and anti-mouse IgG antibodies were purchased from Millipore.

## 3-D Modeling

The 3-D structure of AKT2 with the full allelic series was predicted using IntFOLD (27) and visualized in PyMOL (28).

#### In Vitro Kinase Assays

Following lentiviral infection and subsequent  $5\mu$ g/mL blasticidin selection, V5-AKT2, V5-AKT2.Lys17, V5-AKT2. Thr50, V5-AKT2.Lys208, V5-AKT2.His274, and V5-AKT2. Trp467 variants were each isolated from HeLa cell lysate with V5 agarose beads (Sigma-Aldrich) and incubated with 150 ng GST-GSK3 $\beta$  substrate peptide (Cell Signaling Technologies) and 250 mmol/L cold ATP in kinase assay buffer (Cell Signaling Technologies) for 35 min at 30°C.

# **Proliferation Assay**

Lentiviral pLX304 control or V5-AKT2 variant infected HuH7 cells were cultured in 24-well plates for 72 h in 10% FBS /phenol red-free DMEM for 72 h. We added WST-1 (Takara Clontech) to each well at the manufacturerrecommended 1:10 ratio and incubated for 4 h at 37°C prior to absorbance measurement at 450 nm with BioTek Synergy H4 plate reader.

#### Immunoblots

We washed cells with PBS and lysed in EBC buffer (120 mmol/L NaCl, 50 mmol/L Tris-HCl [pH 7.4], 50 nmol/L calyculin, cOmplete protease inhibitor cocktail [Roche], 20 mmol/L sodium fluoride, 1 mmol/L sodium pyrophosphate, 2 mmol/L ethylene glycol tetraacetic acid, 2 mmol/L ethylenediaminetetraacetic acid, and 0.5% NP-40) for 20 min on ice. To preclear cell lysates, we centrifuged at 12,700 rpm at 4°C for 15 min. We measured protein concentration with Pierce BCA protein assay kit using a BioTek Synergy H4 plate reader. We resolved lysates on Bio-Rad Any kD Mini-PROTEAN TGX polyacrylamide gels by SDS-PAGE and transferred by electrophoresis to nitrocellulose membrane (Life Technologies) at 100 V for 70 min. We blocked membranes in 5% nonfat dry milk/TBST (10 mmol/L Tris-HCl, 150 mmol/L NaCl, 0.2% Tween 20) buffer pH 7.6 for 30 min. We incubated blots with indicated antibody overnight at 4°C. The membrane was then washed in TBST, three times at 15-min intervals, before a 1-h secondary horseradish peroxidaseconjugated antibody incubation at room temperature. We again washed nitrocellulose membranes in TBST, three times for 15 min, prior to enhanced chemiluminescent substrate detection (Pierce).

## Statistical Analysis

The quantified results of the in vitro kinase and proliferation assays were normalized to internal control values for each replicate. We used generalized linear models of the quantified assay results to assess effects of variants within and across replicate rounds, allowing for interaction by replicate. The graphical representation was produced using functions in the effects (v 3.0-3) package in R.

#### **Gene Expression Studies**

#### Study Samples

We compared the expression pattern of *AKT2* to the two other members of the *AKT* gene family, *AKT1* and *AKT3*, using multitissue RNA sequencing (RNA-seq) data from the pilot phase of the Genotype-Tissue Expression (GTEx) project (dbGaP accession number: phs000424.v3.p1) in 44 tissues with data from more than one individual. Detailed procedures for sample collection, RNA extraction, RNAseq, and gene and transcript quantifications have been previously described (29). Using data from the Identifying Biomarkers of Ageing using whole Transcriptome Sequencing (EuroBATS) project, samples from photo-protected subcutaneous adipose tissue from 766 twins were extracted (130 unrelated individuals, 131 monozygotic and 187 dizygotic twin pairs) and processed as previously described (30,31). Using data from METSIM, subcutaneous fat biopsy samples were obtained from a sample of 770 participants and processed as previously described (32).

#### Phenotypes

We studied the association of age, BMI, and FI levels with gene expression levels and with expression-associated SNPs (expression quantitative trait loci [eQTL]) in the *AKT2* region. Age and sex were available for the GTEx study samples. In additional to age and BMI, FI level was measured at the same time point as the fat biopsies in the EuroBATS sample data, following a previously described protocol (33). Baseline age, BMI, and FI levels were used for the METSIM participants (34).

## Statistical Analysis

The comparison of expression levels of *AKT2* versus *AKT1* and *AKT2* versus *AKT3* was performed using log<sub>2</sub>transformed reads per kilobase per million mapped reads (RPKMs). We studied BMI, age, and FI (not available in GTEx data) associations with *AKT2* expression using linear mixed models as implemented in the lme4 package in R. The gene expression RPKM values were inverse-variance rank normalized for these analyses. Covariates included study-specific fixed- and random-effects (see Supplementary Data for additional details on each cohort), using sex, BMI, and age as additional fixed-effects as appropriate. The eQTL analysis was performed on single nucleotide polymorphisms (SNPs) within a 1 Mb of *AKT2* using linear mixed models to assess the association of the SNPs with the inverse-normalized RPKM expression values.

#### RESULTS

#### **Genetic Association Studies**

We tested the association of FI and FG with 390,225 variants from exome sequence data (GoT2D and T2D-GENES) and 109,215 variants derived from exome array genotyping (GoT2D) (7) (individual study genomic inflation factor [ $\lambda_{GC}$ ] <1.06; Supplementary Fig. 1). We examined variants that had been previously associated with FG and FI (3,18). Of 28 FG and 14 FI loci with the reported SNPs or close proxies in our data set, 13 FG and 4 FI showed directionally consistent significant associations. Among the remaining genome-wide association study loci not significant in our data, we observed directionally consistent associations in 14 of 15 FG and 9 of 10 FI loci ( $P_{\text{enrichment}} = 5 \times 10^{-4}$  for

FG and 0.01 for FI) (Supplementary Data, Supplementary Table 2).

In addition, we identified a novel significant single variant association between rs184042322 and FI (MAF 1.2%,  $P = 1.2 \times 10^{-7}$ ), a coding variant in *AKT2* (*V-AKT Murine Thymoma Viral Oncogene Homolog 2*) where amino acid Pro50 is substituted with a threonine (NP\_001617.1: p.Pro50Thr) (Fig. 1, Supplementary Fig. 1). The same allele drove a significant FI signal for *AKT2* in gene-based analysis ( $P = 6.1 \times 10^{-7}$ ), in which we discovered two additional significant gene-based associations between *GIMAP8* and FG ( $P_{\text{PTV}} = 2.3 \times 10^{-6}$ ) and between *NDUFAF1* and FI ( $P_{\text{PTV+NSBroad}} = 9.2 \times 10^{-7}$ ) (Supplementary Fig. 2, Supplementary Table 2D).

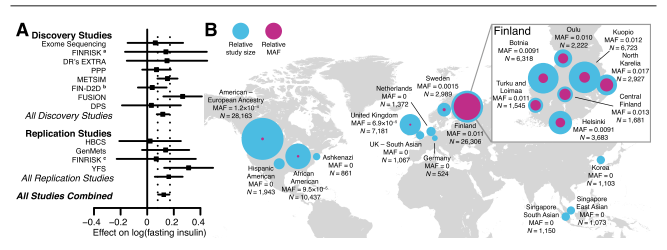
In an effort to replicate the single variant association of AKT2 Pro50Thr with FI, we aggregated the allele frequency estimates of AKT2 Pro50Thr in our data with data from the CHARGE Consortium and the four Finnish studies. In ExAC, rs184042322 is multiallelic (p.Pro50Thr and p.Pro50Ala) but Pro50Ala is observed only twice in the Latino population sample and not seen in our exome sequencing data, which includes 1,021 individuals of Hispanic ancestry. AKT2 Pro50Thr was observed at a much higher frequency in Finnish individuals (MAF 1.1%) than other non-Finnish pooled European (MAF 0.02%), African American (MAF 0.01%), Asian (MAF <0.01%), or Hispanic (MAF <0.01%) individuals (Fig. 1). We replicated the association between FI and AKT2 Pro50Thr by meta-analysis of the association in the four Finnish studies ( $P = 5.4 \times 10^{-4}$ , N = 5,747) with the discovery studies (P\_{\rm combined} = 9.98  $\times$  $10^{-10}$ , N = 25,316). We observed no evidence of effect-size heterogeneity between studies ( $P_{\text{heterogeneity}} = 0.76$ ). The

minor T allele was associated with a 12% (95% CI 7–18) increase in FI levels in the discovery and replication studies, a per allele effect of 10.4 pmol/L (95% CI 6.6–14.3).

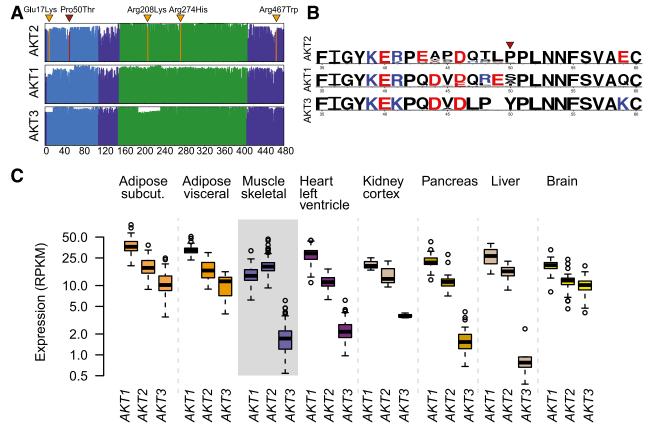
The serine/threonine protein kinases AKT1, AKT2, and AKT3 are conserved across all vertebrates (Fig. 2). Pro50 and the seven preceding residues in the pleckstrin homology (PH) domain appear to be specific for the AKT2 isoform. Population genetic studies show a strong intolerance to missense and loss-of-function (LoF) variation in *AKT2* (Supplementary Data, Supplementary Fig. 3, Supplementary Fig. 4, Supplementary Table 3). Notably, in ExAC data, *AKT2* contains fewer missense variants than expected (the missense constraint metric, Z = 3.5, is in the 94th percentile of all genes) and extreme constraint against LoF variation (estimated probability of being LoF intolerant = 1).

AKT2 is a primary transducer of PI3K signaling downstream of the insulin receptor and is responsible for mediating the physiological effects of insulin in tissues including liver, skeletal muscle, and adipose. *Akt2* null mice are characterized by hyperglycemia and hyperinsulinemia, and some develop diabetes (35,36). In humans, highly penetrant rare alleles in *AKT2* cause familial partial lipodystrophy and hypoinsulinemic hypoglycemia with hemihypertrophy (Glu17Lys) (37,38) and a syndrome featuring severe insulin resistance, hyperinsulinemia, and diabetes (Arg274His) (39). Additional rare alleles have been observed in individuals with severe insulin resistance (Arg208Lys and Arg467Trp), but no variant has been associated with glycemic traits at the population level (40).

Given the spectrum of diseases and traits associated with *AKT2* (41), we hypothesized that *AKT2* Pro50Thr would be



**Figure 1**—*AKT2* Pro50Thr association with FI levels. *A*: For each study, the square represents the estimate of the additive genetic effect for the association of the *AKT2* Pro50Thr allele with log-transformed FI levels and the horizontal line gives the corresponding 95% CI of the estimate. Inverse-variance meta-analyses were performed for all discovery studies, all replication studies, and all studies combined. The vertical dashed lines indicate the 95% CI for the estimate obtained in the meta-analysis of all studies combined. DPS, The Finnish Diabetes Prevention Study; DR's EXTRA, Dose-Responses to Exercise Training study; FIN-D2D, National Diabetes Prevention Programme in Finland; PPP, Prevalence, Prediction and Prevention of Diabetes (PPP)-Botnia study. *B*: MAF for each available region and ancestry. Across countries of the world, the MAF ranges from 0 to 1.1%. The relative sample sizes (*N*) for each region/ancestry are displayed with the blue circles and the rale with the size of the circles showing comparative differences. Within Finland (inset), where the MAF ranges from 0.9 to 1.7%, birthplace and study center data were used to show the allele distribution across the country. <sup>a</sup>FINRISK 2007, <sup>b</sup>FIN-D2D 2007, <sup>c</sup>FINRISK 1997 and 2002.



**Figure 2**—Expression and conservation properties. *A*: Amino acid alignment and conservation of the three AKT proteins in vertebrates. The *x*-axis gives the amino acid position and the height of the lines shows the conservation score across 100 vertebrate genome alignments. The functional domains are the PH domain (blue) and the kinase domain (green). The position of AKT2 Pro50Thr is shown in red and the locations of the other *AKT2* disease-causing mutations (37–40) are shown in orange: Glu17Lys, Arg208Lys, Arg274His, and Arg467Trp. *B*: WebLogo plots of amino acids 35–60 are shown for AKT2, AKT1, and AKT3, contrasting the homology of the three isoforms. The height of letters gives the relative frequency of different amino acids across the 100 vertebrate species, with the colors showing amino acids with similar charge. *C*: Expression of *AKT1*, *AKT2*, and *AKT3* in eight insulin-sensitive tissues using RNA sequencing data from the GTEx Consortium. subcut., subcutaneous.

associated with features of metabolic syndrome or lipodystrophy. In quantitative trait analysis in the initial discovery and replication cohorts, we did observe a constellation of features indicative of a milder lipodystrophy-like phenotype associated with the rare allele: associations with increased 2-h insulin values (effect = 0.2 SD of log-transformed 2-h insulin, 95% CI 0.1–0.4,  $P = 7.9 \times 10^{-8}$ , N = 14,150), lower insulin sensitivity (effect = -0.3 SD of the log-transformed Matsuda index, 95% CI -0.5 to -0.2,  $P = 1.2 \times 10^{-6}$ , N =8,566), and increased risk of type 2 diabetes (OR 1.05, 95% CI 1.0–1.1,  $P = 8.1 \times 10^{-5}$ , 9,783 individuals with type 2 diabetes and 22,662 without diabetes), with no effects on FG, postprandial glucose, or fasting lipid levels ( $P \ge 0.01$ ) (Supplementary Table 4). In the T2D-GENES exome sequencing data where FG and FI levels were available in individuals with diabetes, we observed one individual who was homozygous for the P50T allele with FI and FG levels in the 99.8th and 98.8th percentiles, respectively. There was a significant difference in trait distributions by P50T genotype (FI, P = 0.002; FG, P = 0.02) (Supplementary Fig. 5,

Supplementary Table 4). Next, we used electronic health records available in the Finnish METSIM and FINRISK cohorts to characterize the impact of *AKT2* Pro50Thr on disease risk. We found no evidence for association with any cancer, polycystic ovary disease, or acanthosis nigricans (Supplementary Table 5); however, these tests are underpowered due to the low number of cases and potential for misclassification. Nor did we find evidence for enrichment of low-frequency associations in any *AKT2*-related pathways or genes implicated in monogenic forms of glycemic disease (Supplementary Data, Supplementary Table 6, Supplementary Table 7, Supplementary Fig. 6, Supplementary Fig. 7).

#### In Vitro Functional Studies

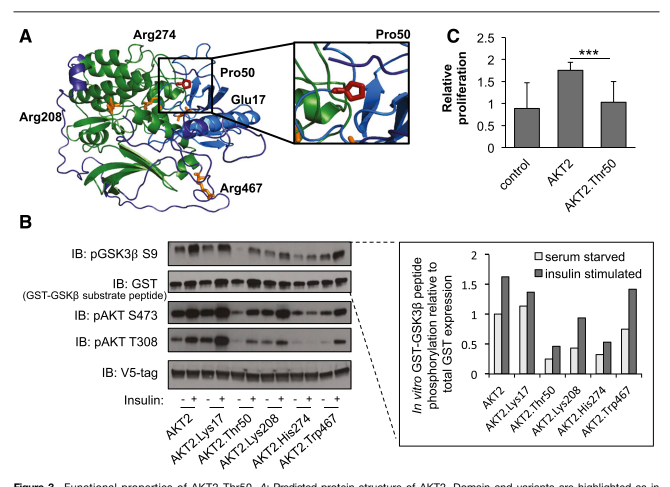
To understand the functional consequences of the *AKT2* Pro50Thr variant on the protein, we investigated protein expression, activation, kinase activity, and downstream effector phosphorylation.

First, we used in silico classifiers that predict potential functional consequences of alleles on protein function. Two

of the five classifiers predicted *AKT2* Pro50Thr to be deleterious (Supplementary Table 3). Second, we used 3-D models of *AKT2* viewed in the PyMol software, which predicted that the Pro50Thr variant causes a change in the conformations of the lipid binding pleckstrin homology (PH) domain (Fig. 3, Supplementary Fig. 8). We hypothesized that the variant protein is inefficiently recruited to the plasma membrane thereby impacting *AKT2* phosphorylation and downstream activity.

To assess the molecular and cellular consequence of the *AKT2* Thr50 variant on protein function, we performed a comparative analysis of AKT2-Thr50 with inactivating and activating alleles implicated in monogenic disorders of insulin signaling. Analysis of AKT2-Thr50 expression showed that while AKT2 protein levels remained unchanged, there was a partial loss of AKT2-Thr50 phosphorylation at its activation sites (Thr308 and Ser473) in HeLa cells,

suggesting impaired AKT2 signaling (Fig. 3, Supplementary Fig. 9). Similar effects were observed in human liverderived HuH7 cells (Supplementary Fig. 10). AKT2-Thr50 also showed a reduced ability to phosphorylate its downstream target GSK3β. These defects in AKT2-Thr50 activity were confirmed through an in vitro kinase assay (P < 0.01) (Fig. 3). AKT2-Thr50 showed a similar decrease in kinase function to the lipodystrophy-causing AKT2-His274 variant. Using a 4-h time course analysis of AKT2 activity, we verified a reduction in both maximally phosphorylated Thr308 and Ser473 in AKT2-Thr50 (Supplementary Fig. 11). To understand how this loss of activity could manifest as a defect in a known cellular function of AKT2 (42), we determined the impact of AKT2-Thr50 on cell proliferation in HuH7 cells. While the addition of AKT2 stimulated hepatocyte proliferation, the response to AKT2-Thr50 was reduced (effect =  $-1.2, P < 1.0 \times 10^{-3}$ ) (Fig. 3*C*, Supplementary Fig. 12).



**Figure 3**—Functional properties of AKT2-Thr50. *A*: Predicted protein structure of AKT2. Domain and variants are highlighted as in Fig. 2*A*. The relative spatial positioning of the AKT2-Pro50 residue is magnified within the inset. *B*: HeLa cells were infected with lentiviral V5-AKT2, V5-AKT2-Lys17, V5-AKT2-Thr50, V5-AKT2-Lys208, V5-AKT2-His274, or V5-AKT2-Trp467; starved for 18 h (white bar); and stimulated for 20 min with 100 nmol/L insulin (gray bar). V5-tagged AKT2 was isolated from cell lysates with anti-V5 agarose beads and incubated with GSK3β-GST peptide in an in vitro kinase assay. Quantification of phosphorylated substrate peptide (pGSK3β) relative to total peptide (GST-GSK3β) is shown at the inset. Immunoblots and quantification shown are representative of three independent replicates. Linear model statistical analyses across all three independent replicates are available in Supplementary Fig. 9. The in vitro kinase was immunoblotted (IB) with the indicated antibodies. *C*: HuH7 cells were infected with lentiviral V5-AKT2, V5-AKT2, V5-AKT2, Thr50, or control pLX304. At 72 h, relative cellular proliferation was determined with WST-1 assay of HuH7 cells. Error bars represent SD. \*\*\**P* = 4.5 × 10<sup>-5</sup>.

## **Gene Expression Studies**

We queried RNA sequencing data from the GTEx Project and found that, in agreement with previous studies (43), *AKT2* is highly and ubiquitously expressed across all tissues (44 tissue types, 3–156 individuals/tissue). Notably the *AKT2* Pro50Thr containing exon is expressed in all tissues and individuals (Supplementary Fig. 13), suggesting that the PH domain is important to AKT2 function (44). Of the three *AKT* homologs, *AKT2* had 1.4-fold higher expression in skeletal muscle than *AKT1* ( $P = 1.5 \times 10^{-19}$ ) and 11-fold higher expression than *AKT3* ( $P = 7.8 \times 10^{-91}$ ). Skeletal muscle was the only tested tissue displaying such pronounced *AKT2* enrichment (Fig. 2, Supplementary Data, Supplementary Fig. 14, Supplementary Table 8).

Motivated by the age-related loss of adipose tissue in Akt2 null mice (35,36) and the growth and lipodystrophy phenotypes in carriers of fully penetrant alleles (37-40), we examined associations of expression levels of AKT2 with BMI, FI, and age in the three adipose tissue data sets (Supplementary Table 9). We found an association between lower BMI levels and higher AKT2 expression in two cohorts (EuroBATS: effect = -0.07 SD,  $P = 6.1 \times 10^{-28}$ ; METSIM: effect = -0.06 SD,  $P = 8.1 \times 10^{-8}$ ) and also observed that higher AKT2 expression was associated with lower log-transformed FI (EuroBATS: effect = -0.04 SD,  $P = 1.1 \times 10^{-3}$ ; METSIM: effect = -0.4 SD,  $P = 3.3 \times$  $10^{-11}$ ). We next tested for gene eQTL and found an eQTL in the 5'UTR of AKT2 (rs11880261, MAF 35%,  $r^2 = 0.002$ , D' = 0.47 in the Finnish 1000 Genomes samples) with the common allele associated with lower AKT2 expression levels (METSIM:  $P = 6.9 \times 10^{-14}$ ; EuroBATS:  $P = 2.3 \times 10^{-8}$ ; GTEx: P = 0.08) (Supplementary Fig. 15). No association was detected between rs11880261 and FI levels, suggesting that the common variant eQTL does not drive the initial FI association (Supplementary Data, Supplementary Table 10).

#### DISCUSSION

Meta-analyses of exome sequence and array genotyping data in up to 38,339 normoglycemic individuals enabled the discovery, characterization, and functional validation of a FI association with a low-frequency *AKT2* coding variant. Rare, penetrant variants in genes encoding components of the insulin-signaling pathway, including *AKT2*, cause monogenic but heterogeneous glycemic disorders (45). In parallel, common alleles in or near many of these genes impact FI levels—the *AKT2* Pro50Thr association shows an effect 5–10 times larger than those of these previous published associations (3). This discovery expands both the known genetic architecture of glucose homeostasis and the allelic spectrum for *AKT2* coding variants associated with glucose homeostasis into the low-frequency range and highlights the effects of both locus and allelic heterogeneity (Fig. 4).

Individuals of Finnish ancestry drove the *AKT2* Pro50Thr association signal. This demonstrates the value of association studies in different ancestries where frequencies of rare alleles may increase due to selective pressure or

stochastic changes from population bottlenecks and genetic drift. The allele associated with increased FI most likely rose to a higher frequency due to genetic drift and exists within the spectrum of rare and low-frequency variation observed in Finland, the excess of which facilitates the study of complex trait associations (46).

Although the *AKT2* Pro50Thr allele shows a strong effect on all of the insulin measures and modest increased type 2 diabetes risk (OR 1.05), we see no effect on any of the glucose measures in individuals without diabetes. Due to the effects of both type 2 diabetes and its treatment on glucose homeostasis, we have not tested genetic associations of FG and FI in individuals with type 2 diabetes, although we observed an individual with diabetes homozygous for P50T with extreme FI and FG levels. The mechanism for such heterogeneous effects is unclear and detailed in vivo physiological studies are needed.

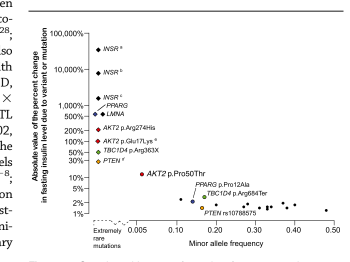


Figure 4-Genetic architecture of rare, low-frequency, and common variants associated with FI levels. In this plot, the absolute values of the percent change in FI level due to rare monogenic mutations (diamonds) and common genetic variants (circles) are plotted against the MAF of the variant. The extremely rare monogenic mutations (above the dashed line to the left of the x-axis) were observed in 2-18 individuals (3,37-40,48,53,54), with the height of the point indicating the percent change in FI levels of mutation carriers from 40 pmol/L, an estimate of population mean FI level. Mutations in INSR and AKT2 p.Arg274His cause compensatory hyperinsulinemia, individuals with TBC1D4 p.Arg363Ter show normal FI levels but postprandial hyperinsulinemia, and mutations in PTEN cause enhanced insulin sensitivity providing protection against type 2 diabetes. For common variants, the percent change in FI levels per insulin-increasing allele is plotted above the solid horizontal axis. These observations are from sequencing (6) and array-based genome-wide association studies (3). For several genes, the effects from rare mutations can be compared with the effects of common variants in or near the gene: PPARG (blue), TBC1D4 (green), PTEN (orange), and AKT2 (red). <sup>a</sup>Donohue syndrome: biallelic LoF mutations in INSR (54). <sup>b</sup>Rabson-Mendenhall syndrome: biallelic LoF mutations in INSR (54). <sup>c</sup>Postpubertal severe insulin resistance: heterozygous or homozygous LoF mutations in INSR (54). <sup>d</sup>LoF PTEN mutations cause Cowden syndrome in which carriers exhibit a lowered FI level (mean 29 pmol/L) compared with matched control subjects (3). Carriers with the AKT2 p.Glu17Lys mutation were described with hypoinsulinemic hypoketotic hypoglycemia and hemihypertrophy with undetectable serum insulin (37,38).

We leveraged similar findings to generate hypotheses for future work on AKT2 and downstream targets to further illuminate tissue-specific mechanisms. All reported carriers of the lipodystrophy-causing AKT2 Arg274His allele are hyperinsulinemic, and three of the four carriers have diabetes (39). These observations are similar to the ones made for TBC1D4 (which encodes a protein that acts as a substrate immediately downstream of AKT2 in the PI3K pathway). In TBC1D4, a population-specific, protein-truncating variant (Arg684Ter) is associated with increased type 2 diabetes risk (OR 10.3), increased postprandial glucose and insulin levels, and a modest decrease in FI and FG levels (6) (Fig. 4). Arg363Ter, another stop codon allele in *TBC1D4*, is rare (not observed in ExAC) and has been reported with a modest elevation in FI levels but extreme postprandial hyperinsulinemia and acanthosis nigricans (47). Small interfering RNA-mediated gene knockdown of AKT2 in human primary myotubes completely abolishes insulin action on glucose uptake and glycogen synthesis (48), which highlights the importance of an intact AKT2-TBC1D4 signaling pathway in the regulation of insulin sensitivity in humans. TBC1D4 is ubiquitously expressed with adipose and skeletal muscle tissue ranking among the tissues with highest expression in GTEx. TBC1D4 Arg363Ter seems to have an effect in adipocytes (47), whereas Arg684Ter falls in an exon that is exclusively expressed in skeletal and heart muscle (6,49). This is a likely cause of the TBC1D4 Arg684Ter tissue specificity, which appears to differ from the other TBC1D4 Arg363Ter variant as well as the AKT2 variants.

The phenotypes exhibited by carriers of rare, penetrant AKT2 alleles reflect differential AKT2 activation with kinetically inactivating variants, resulting in hyperinsulinemia and lipodystrophy, whereas kinetically activating variants lead to hypoglycemia (37-39). The decrease of cellular proliferation we observe demonstrates that the downstream signaling changes caused by AKT2-Thr50 are sufficient in hepatocytes to impair AKT2 function at the cellular level while maintaining varying portions of regulatory capacity. Along with the observed association with increased FI levels in human populations, these results support AKT2 Pro50Thr as a partial LoF variant. The inactivating AKT2 Pro50Thr variant contrasts with the known activating AKT2 Glu17Lys mutation and showcases bidirectional effects within the PH domain of AKT2. Although the Pro50 residue is conserved in AKT2 throughout all vertebrates, the variant lies within the PH domain that is not conserved between AKT isoforms (Fig. 2). These residues, harboring the Pro50 variant, may functionally distinguish AKT2 from AKT1 and AKT3. Although AKT isoforms are activated in the same mechanism within the PI3K pathway downstream of insulin, the  $Akt2^{-/-}$  mouse is the only knockout of the gene family to be characterized by insulin resistance and diabetes (35,50-52). A deeper understanding of what makes the AKT2 isoform distinct could offer potential sites for therapeutic intervention and enable more targeted approaches to disease prevention.

# Appendix

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