

A Common Haplotype of the Glucokinase Gene Alters Fasting Glucose and Birth Weight: Association in Six Studies and Population-Genetics Analyses

Michael N. Weedon, Vanessa J. Clark, Yudong Qian, Yoav Ben-Shlomo, Nicholas Timpson, Shah Ebrahim, Debbie A. Lawlor, Marcus E. Pembrey, Susan Ring, Terry J. Wilkin, Linda D. Voss, Alison N. Jeffery, Brad Metcalf, Luigi Ferrucci, Anna Maria Corsi, Anna Murray, David Melzer, Bridget Knight, Bev Shields, George Davey Smith, Andrew T. Hattersley, Anna Di Rienzo, and Tim M. Frayling

Fasting glucose is associated with future risk of type 2 diabetes and ischemic heart disease and is tightly regulated despite considerable variation in quantity, type, and timing of food intake. In pregnancy, maternal fasting glucose concentration is an important determinant of offspring birth weight. The key determinant of fasting glucose is the enzyme glucokinase (GCK). Rare mutations of *GCK* cause fasting hyperglycemia and alter birth weight. The extent to which common variation of *GCK* explains normal variation of fasting glucose and birth weight is not known. We aimed to comprehensively define the role of variation of *GCK* in determination of fasting glucose and birth weight, using a tagging SNP (tSNP) approach and studying 19,806 subjects from six population-based studies. Using 22 tSNPs, we showed that the variant *rs1799884* is associated with fasting glucose at all ages in the normal population and exceeded genomewide levels of significance ($P = 10^{-9}$). *rs3757840* was also highly significantly associated with fasting glucose ($P = 8 \times 10^{-7}$), but haplotype analysis revealed that this is explained by linkage disequilibrium ($r^2 = 0.2$) with *rs1799884*. A maternal A allele at *rs1799884* was associated with a 32-g (95% confidence interval 11–53 g) increase in offspring birth weight ($P = .002$). Genetic variation influencing birth weight may have conferred a selective advantage in human populations. We performed extensive population-genetics analyses to look for evidence of recent positive natural selection on patterns of *GCK* variation. However, we found no strong signature of positive selection. In conclusion, a comprehensive analysis of common variation of the glucokinase gene shows that this is the first gene to be reproducibly associated with fasting glucose and fetal growth.

Blood glucose concentration is tightly regulated in humans, despite considerable variation in quantity, type, and timing of food intake. The appropriate release of insulin and other hormones prevents the damaging effects caused by hyper- or hypoglycemia. In the absence of food, blood glucose levels return to a tightly defined level of 3.5–5.5 mmol/liter in young adults.

Variation of fasting glucose levels is clinically important. Even in the normal range, the fasting glucose level is associated with future risk of type 2 diabetes and ischemic heart disease.^{1–3} During pregnancy, maternal fasting glucose concentration is an important determinant of offspring birth weight.^{4,5} Maternal hyperglycemia stimulates the release of fetal insulin, an important fetal growth factor. Factors such as age and obesity explain some of the variation in fasting glucose levels within a population, but, when these are accounted for, considerable variation between individuals still remains. Fasting glucose level is strongly heritable, and much of the interindividual differ-

ences in fasting glucose concentration can be explained by genetic variation.^{6–8}

One of the principal regulators of fasting glucose concentrations is the enzyme glucokinase (GCK [MIM 138079]).⁹ GCK is predominantly expressed in the pancreas and liver and catalyzes the first rate-limiting reaction in glycolysis.⁹ Its lack of inhibition by its product and its sensitivity to changes in glucose concentrations within the physiological range make it ideally suited for its role as the pancreatic β -cell glucose sensor.⁹

The key role of *GCK* in glucose homeostasis is demonstrated by the impact of rare gene mutations that cause a subtype of diabetes known as “maturity-onset diabetes of the young” (MODY [MIM 606391]).^{10–12} *GCK* MODY is characterized by mild, stable hyperglycemia during fasting.¹³ *GCK* MODY mutations also affect birth weight. Children born to mothers with a *GCK* mutation are ~600 g heavier at birth¹⁴; conversely, children who have a *GCK* MODY mutation but whose mothers do not are ~500 g

From the Institute of Biomedical and Clinical Science, Peninsula Medical School, Exeter, United Kingdom (M.N.W.; A.M.; D.M.; B.K.; B.S.; A.T.H.; T.M.F.); Department of Human Genetics, University of Chicago, Chicago (V.J.C.; Y.Q.; A.D.R.); Department of Social Medicine (Y.B.-S.; N.T.; D.A.L.; G.D.S.) and Avon Longitudinal Study of Parents and Children (M.E.P.; S.R.), University of Bristol, Bristol, United Kingdom; Department of Epidemiology and Population Health, London School of Hygiene & Tropical Medicine, London (S.E.); Department of Endocrinology and Metabolism, Peninsula Medical School, Plymouth, United Kingdom (T.J.W.; L.D.V.; A.N.J.; B.M.); Clinical Research Branch, National Institute on Aging, National Institutes of Health, Baltimore (L.F.); Laboratory of Clinical Epidemiology, Geriatric Department, National Institute of Research and Care on Aging, Florence (A.M.C.)

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Address for correspondence and reprints: Dr. Tim M. Frayling, Peninsula Medical School, Magdalen Road, Exeter EX1 2LU, United Kingdom. E-mail: Tim.Frayling@pms.ac.uk

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Table 1. Initial tSNP Association Results for Fasting Glucose in the EFS

SNP	Alleles (1/2)	Mean (SD) Glucose Level [No. of Subjects] for Genotype			ANOVA <i>P</i>
		11	12	22	
<i>rs3217943</i>	C/T	4.48 (.47) [84]	4.53 (.44) [602]	4.53 (.43) [783]	.594
<i>rs2537184</i>	G/A	4.53 (.44) [372]	4.53 (.44) [744]	4.51 (.44) [377]	.985
<i>rs882020</i>	T/C	4.68 (.36) [16]	4.53 (.46) [324]	4.53 (.43) [1,117]	.371
<i>rs2268575</i>	C/T	4.67 (.44) [37]	4.52 (.45) [438]	4.52 (.43) [976]	.120
<i>rs2971679</i>	T/C	4.53 (.47) [42]	4.53 (.45) [403]	4.53 (.44) [1,039]	.995
<i>rs2971677</i>	A/C	4.51 (.45) [39]	4.52 (.45) [398]	4.53 (.43) [1,020]	.870
<i>rs2971676</i>	A/G	4.46 (.36) [17]	4.52 (.45) [232]	4.53 (.44) [1,240]	.765
<i>rs4724290</i>	T/C	4.49 (.43) [108]	4.53 (.44) [590]	4.53 (.44) [735]	.714
<i>rs2041547</i>	C/T	4.55 (.43) [390]	4.53 (.45) [744]	4.49 (.41) [355]	.212
<i>rs2971671</i>	C/T	4.60 (.46) [68]	4.56 (.44) [477]	4.51 (.43) [887]	.054
<i>rs2284777</i>	C/T	4.52 (.34) [19]	4.52 (.43) [336]	4.53 (.43) [1,144]	.890
<i>rs758988</i>	C/A	4.52 (.44) [1,028]	4.54 (.45) [421]	4.51 (.37) [44]	.727
<i>rs2284770</i>	A/T	4.50 (.45) [460]	4.54 (.44) [673]	4.52 (.41) [320]	.345
<i>rs2284769</i>	G/C	4.67 (.29) [19]	4.51 (.45) [291]	4.53 (.44) [1,190]	.234
<i>rs1799884</i>	T/C	4.53 (.51) [55]	4.58 (.44) [449]	4.50 (.43) [988]	.010
<i>rs1476891</i>	A/G	4.53 (.44) [683]	4.52 (.45) [614]	4.55 (.45) [158]	.775
<i>rs3757840</i>	G/T	4.48 (.42) [350]	4.53 (.43) [714]	4.57 (.48) [407]	.016
<i>rs34976798^a</i>	G/T	4.68 (.28) [6]	4.49 (.44) [173]	4.53 (.44) [1,270]	.347
<i>rs2908287</i>	G/A	4.52 (.43) [842]	4.52 (.44) [546]	4.58 (.46) [100]	.408
<i>rs2331003</i>	A/G	4.53 (.42) [940]	4.53 (.44) [490]	4.56 (.42) [67]	.847
<i>rs1003573</i>	C/T	4.53 (.43) [549]	4.52 (.43) [656]	4.54 (.46) [233]	.919
<i>rs917791</i>	G/A	4.55 (.46) [193]	4.51 (.43) [698]	4.54 (.40) [551]	.311

NOTE.—Glucose is measured in mmol/liter.

^a *r*² = 1 with *rs13239289*.

lighter at birth.¹⁴ If both the mother and the child have a *GCK* mutation, the effects cancel each other out and the baby is of normal birth weight.¹⁴

In the past, low-birth-weight babies had a high level of neonatal mortality, which was more marked before modern medical support was available. In contrast, high birth weight can pose a danger to the mother and to the child during birth. Although it is likely that natural selection has had a strong influence on birth-weight variation, several different models of selection are plausible. For example, it may be that variation influencing birth weight evolves under purifying selection, which eliminates alleles that increase or decrease birth weight beyond the optimal value for a given environment. Alternatively, if the population-average birth weight is not close to the optimal value, new variants that improve birth weight will be advantageous. Hence, variants that affect birth weight are interesting candidates for population-genetics studies aimed at understanding the evolution of birth weight.

The extent to which common variation of *GCK* explains

normal variation of fasting glucose and birth weight is not known. We recently described an association between a variant of the *GCK* promoter, *GCK-30* (*rs1799884* [dbSNP]), and fasting glucose in young adults and pregnant women.¹⁵ We also demonstrated an association of the minor allele with increased offspring birth weight.¹⁵ It is not known whether further variation in the *GCK* gene alters fasting glucose or birth weight or the extent to which natural selection influenced variation of this gene. In the present study, we aimed to comprehensively define the role of variation of *GCK* on fasting glucose and birth weight, using a tagging SNP (tSNP) approach and studying 19,806 subjects from six different studies.

Material and Methods

Association Study

Determining the Linkage-Disequilibrium (LD) Structure of the GCK Gene Region.—To determine the LD structure of *GCK* (45 kb) and the surrounding region (5 kb upstream and 19 kb downstream of *GCK* isoform 1), we first sequenced the exons of 43 unrelated

Table 2. Characteristics of Subjects Examined in the Fasting Glucose Study

Characteristic ^a	Finding for					
	EFS	PEB Adults	BCG	InChianti	BWHHS	ALSPAC Children
No. (% male)	1,552 (50)	356 (52)	626 (53)	1,022 (43)	2,652 (0)	839 (54)
Median age, in years (IQR)	32.0 (7.0)	34.1 (7.0)	25.0 (1.0)	71.0 (12.0)	68.0 (9.0)	8.0 (.0)
Median BMI, in kg/m ² (IQR)	24.8 (5.6)	25.9 (5.4)	24.3 (7.3)	27.0 (5.5)	27.6 (5.0)	16.5 (4.0)
Median fasting glucose, in mmol/liter (IQR)	4.5 (.6)	4.7 (.6)	4.6 (.6)	4.8 (.7)	5.6 (.8)	4.9 (.7)
Median fasting blood insulin, in pmol/liter (IQR)	55.3 (38.5)	56.7 (36.8)	39.6 (25.8)	66.4 (48.3)	Not applicable	37.7 (44.6)

NOTE.—Only results of successfully genotyped subjects are included.

^a IQR = interquartile range.

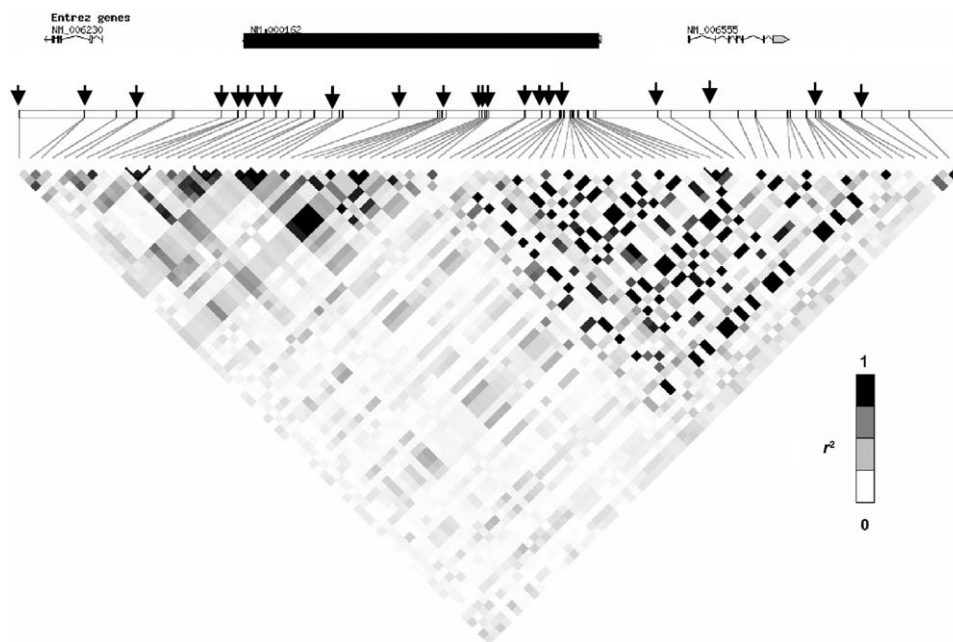


Figure 1. LD structure of glucokinase. r^2 values between the 84 SNPs across a 116-kb region are presented. Arrows indicate tSNPs used in association study.

U.K. white subjects. No common (>5% minor-allele frequency) coding variants were identified. We then used a directed resequencing approach, centering our PCR amplicons on SNPs reported in the November 2002 version of the Santa Cruz genome Web site. We initially focused on common SNPs in regions of conservation among humans, mice, and rats (using mVista), and we defined a cut-off of >100 bp that showed >75% matches. Next, to increase the average SNP density to 1 SNP per 2 kb, we chose additional amplicons randomly. This study was initiated before the availability of HapMap data.

Choice of tSNPs.—We used the Tagger program to choose tSNPs.¹⁶ We forced in *rs1799884*, because of the previous association evidence, and *rs34976798*, which was discovered in our population-genetics survey (see table 1). The latter was chosen because it tags a high-frequency–derived haplotype in the non-African samples. A pairwise tSNP approach was equally efficient as a multimarker approach, so only single-marker tSNPs were used.

Fasting glucose initial study: 1,552 subjects from the Exeter Family Study (EFS).—We genotyped 1,552 subjects (median age 33 years) from the population-based EFS, to test for nominal associations ($P < .05$) of tSNPs. The EFS has been described in detail elsewhere.^{17,18} The clinical characteristics of these and the replication studies' subjects are presented in table 2. Subjects from all studies with fasting glucose over the normal range (>6 mmol/liter) were excluded. Briefly, the EFS is a study of newborn babies and their parents from a geographically defined region of Exeter, United Kingdom.^{17,18} Fasting glucose was measured using dry-slide technology on Vitros 950 analyzers (Ortho Clinical) before December 2001 and with manufacturer's standard reagents on Modular analyzers (Roche Diagnostics) thereafter.

Fasting glucose replication studies.—We used five population-based studies for replication of nominally significant tSNPs. We studied 356 parents (median age 35 years) from the Plymouth EarlyBird study (PEB).¹⁹ The PEB is a nonintervention prospective study of

school-aged children and their parents. Fasting glucose was measured on a Cobas Integra 700 analyzer (Roche Diagnostics).

We studied 626 subjects (median age 25 years) from the Barry Caerphilly Growth study (BCG).²⁰ These subjects are from the follow-up of consecutively born infants, from the towns of Barry and Caerphilly in South Wales, born between 1972 and 1974. Between 1997 and 1999, all the original children who had completed the 5-year follow-up were traced, and they completed a questionnaire and attended a screening clinic, where growth and oral glucose tolerance test measurements were taken.

We studied 1,022 subjects (median age 71 years) from the Invecchiare in Chianti study (InChianti), a prospective population-based cohort of elderly Italians from two towns (Greve in Chianti and Bagno a Ripoli) from the Chianti region of Tuscany. The study includes randomly selected participants aged 65–102 years and 30 men and women from each age decade between 20 and 70 years. The data collection started in September 1998 and was completed in March 2000. Fasting glucose concentration was determined by an enzymatic colorimetric assay, with use of a modified glucose-oxidase-peroxidase method and a Roche-Hitachi 917 analyzer (Roche Diagnostics).

We studied 2,652 subjects (median age 68 years) from the British Women's Health and Heart Study (BWHHS). The BWHHS is a cohort of women aged 60–79 years, randomly selected from 23 British towns.²¹ Baseline data were collected between April 1999 and March 2001.

We studied 839 children (median age 8 years) from the Avon Longitudinal Study of Parents and Children (ALSPAC) (the only subjects from the ALSPAC study for whom fasting glucose measurements were available). Glucose was measured by the glucose-oxidase method on a YSI 2300 stat plus analyzer. Further details of the study are described in the "Birth-weight study" section. Blood samples were taken after a minimum 6-h fast. Plasma glu-

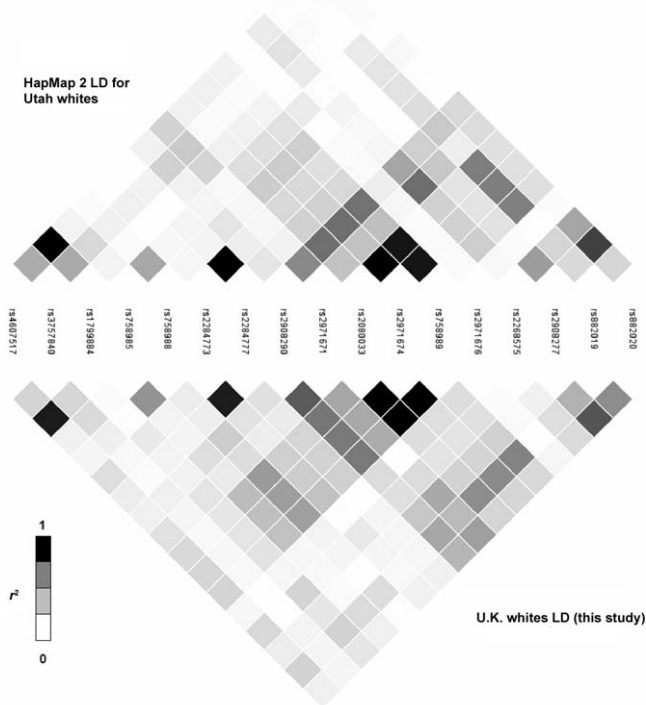


Figure 2. Comparison of LD structure determined in this study with that determined from HapMap 2 data. The LD structures are based on the r^2 values between the 17 SNPs that are present in both data sets.

cose was measured by a glucose oxidase Trinder method,²² with use of a Hitachi Modular analyzer.

Birth-weight study.—We examined ALSPAC subjects for the association of maternal and offspring *GCK* polymorphisms with offspring birth weight. ALSPAC is a prospective study of pregnancies recruited from all pregnant women in three Bristol-based District Health Authorities with expected delivery dates between April 1991 and December 1992. We have reported elsewhere the association of one *GCK* polymorphism with offspring birth weight in 1,159 of the ALSPAC mother-child pairs.¹⁵ In the present study, we extended this earlier analysis by including >6,500 mother-child pairs. Birth weights were taken from hospital records, and gestational age was estimated from the last menstrual period and ultrasound data. For the present study, subjects born after gestation of <36 wk, nonwhites, and twins were excluded from the analysis.

Association-study statistics.—All standard statistical analyses were performed using Stata SE version 9 (StataCorp). We used analysis of variance (ANOVA) and linear regression to test for association of fasting glucose and birth weight with genotype. For LD statistics and tSNP identification, we used Tagger.¹⁶ For the haplotype analyses, we used qtpphase from the unphased suite of programs.²³ Fixed-effects meta-analysis statistics and plots were generated using Stats Direct version 2.4.2 (Sale), with a weighted-mean-difference approach. For the association of genotype with fasting glucose, we included only normal, glucose-tolerant subjects in our analyses and therefore excluded anyone with a fasting glucose >6 mmol/liter. For the studies of elderly subjects, this meant exclusion of a high proportion of subjects (13.9% from InChianti

and 36.5% from the BWHHS). We therefore also performed the analyses excluding only World Health Organization–defined diabetic subjects (fasting glucose >7 mmol/liter). For the birth-weight studies, we analyzed (1) the effect of maternal genotype on birth weight both with and without correction for fetal genotype and (2) the effect of fetal genotype on birth weight both with and without correction for maternal genotype.

Power.—We used Quanto for power calculations.²⁴ For the initial study of all tSNPs, with the assumption of an additive model and an allele frequency of 20%, we had ~80% power to detect a per-allele effect on fasting glucose of >0.055 mmol/liter, at $P < .05$. We did not correct for the testing of multiple SNPs; instead, we included all nominally significant SNPs ($P < .05$) in the replication studies. For birth weight, with the assumption of an additive model and an allele frequency of 20%, the ALSPAC cohort provided ~80% power to detect per-allele differences of >26 g for both maternal and fetal genotype analyses at the $P < .05$ level of significance.

Genotyping method.—All genotyping was performed by Kbioscience. Details of the company and the KASPar genotyping method can be found at the Kbioscience Web site.

Genotyping quality control.—In the initial study of the EFS subjects, we analyzed 23 tSNPs, and all were in Hardy-Weinberg equilibrium (HWE) ($P > .01$) except *rs2537185* ($P = 4 \times 10^{-9}$). The overall success rate for the 23 tSNPs in the initial study was 94.6%;

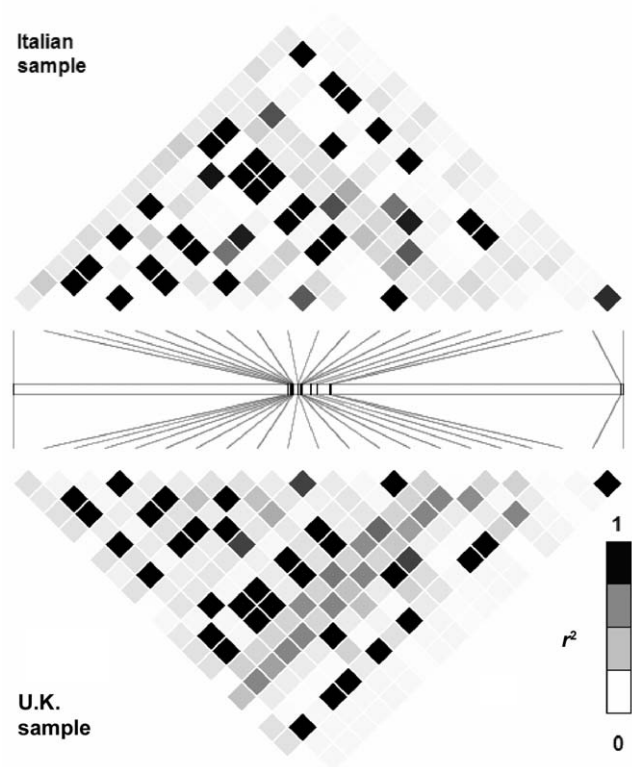


Figure 3. Comparison of LD structure determined for 16 Italians from the Natural Selection study with that determined from U.K. samples used for the tSNP-association study. The LD structure is based on the r^2 and values between 20 SNPs that are present in both data sets.

Table 3. Fasting Glucose Results for *rs1799884* and *rs3757840* for All Cohorts in the Present Study

Study	Mean (SD) Glucose Level [No. of Subjects] for								Linear Trend <i>P</i>
	<i>rs1799884</i>				<i>rs3757840</i>				
	GG	GA	AA	Dominant <i>P</i>	CC	CA	AA		
ALSPAC Children	4.90 (.32) [506]	4.98 (.32) [207]	4.95 (.35) [21]	.008	4.87 (.32) [174]	4.94 (.32) [368]	4.95 (.32) [180]		.02
BCG	4.57 (.38) [426]	4.64 (.37) [180]	4.48 (.45) [18]	.13	4.57 (.41) [127]	4.59 (.36) [324]	4.64 (.40) [166]		.10
EFS	4.50 (.43) [988]	4.58 (.44) [449]	4.53 (.51) [55]	.01	4.48 (.42) [350]	4.53 (.43) [714]	4.57 (.48) [407]		.02
PEB	4.65 (.69) [224]	4.75 (.41) [94]	4.66 (.48) [14]	.08	4.61 (.44) [81]	4.69 (.45) [172]	4.79 (.48) [76]		.01
BWHHS	5.49 (.39) [1,671]	5.54 (.37) [678]	5.60 (.28) [69]	.001	5.48 (.41) [594]	5.51 (.38) [1,238]	5.53 (.35) [578]		.03
InChianti	4.77 (.49) [657]	4.82 (.51) [321]	4.95 (.47) [44]	.03	4.76 (.51) [178]	4.79 (.49) [456]	4.80 (.50) [293]		.40

NOTE.—Glucose is measured in mmol/liter. The data are presented in ascending order of the median age of the cohort. We chose to present the data this way because of the relatively few AA genotypes for *rs1799884* and because our previous data¹⁵ and the initial cohort suggested a dominant model. However, given the high frequency of the *rs3757840* minor allele and the data from the EFS, an additive model appeared to fit best for this SNP.

the minimum success rate was 92.1% for *rs2971671*. The overall genotype error rate in the initial part of the study was 0.6% (18 errors from 3,243 duplicate genotypes). Six of these duplicate errors were for SNP *rs2537185* (6/141 = 4.3% error rate). Given this error rate and the departure from HWE, *rs2537185* was excluded from further analysis. For the genotyping of *rs1799884* and *rs3757840*, all cohorts were in HWE (all *P* > .10), and the LD statistics between the two SNPs were similar across all studies (*r*² range 0.18–0.22; *D'* range 0.98–1.00).

Population-Genetics Study

DNA samples.—We used a separate set of samples to test for evolutionary neutrality in the *GCK* region. We surveyed sequence variation in samples from three major ethnic groups: 16 Hausa from Yaounde (Cameroon), 16 Han Chinese from Taiwan, and 16 individuals from central Italy; we used a European sample separate from that used in the study SNP survey, because these same population samples, together with the Hausa and Han Chinese samples, were surveyed elsewhere at 50 unlinked noncoding (therefore, putatively neutrally evolving) regions^{25,26} and at other candidate diabetes genes.²⁷ This is important because it helps distinguish between differences in patterns of sequence variation due to demography alone and those due to selection and demography. SNPs associated with fasting glucose or with possible evidence of selection were genotyped in the Human Genome Diversity Project (HGDP) panel of 1,052 individuals (of whom 927 are unrelated) from 52 populations. Sequence data were also collected from one common chimpanzee, one gorilla, and one orangutan, to determine the putative ancestral allele at each polymorphic site.

PCR amplification and DNA sequencing.—Primers for PCR amplification and sequencing were designed using PRIMER.²⁸ The survey included a segment of 17,355 bp centered on *rs1799884* in

the *GCK* promoter and spanning the entire gene. PCR primers were designed to amplify a 2,000–2,400-bp fragment, with at least 200 bp overlap between amplicons. Sequencing was performed using internal sequencing primers with the ABI Big Dye Terminator v. 3.1 Cycle Sequencing kit and with ABI 3700 or 3730 automated sequencers. The Phred-Phrap-Consed package was used to assemble and analyze all sequences and score genotypes.²⁹

Genotyping of HGDP.—On the basis of the fasting glucose association results and the survey of population-genetics variation, three SNPs were genotyped in the HGDP panel: *rs3757840*, *rs1799884*, and *rs13239289*. SNPs *rs1799884* and *rs13239289* were genotyped using the 5' nuclease assay (TaqMan [Applied Biosystems]). SNP *rs3757840* was genotyped by RFLP analysis with *Bgl*III. Primer sequences and PCR conditions for both the 5' nuclease assays and the RFLP assay are available from the authors.

Data analysis.—Summary statistics of population variation were calculated using the online applications SLIDER and MAXDIP.^{25,30} The HKA test was performed using an online application. Haplotypes were inferred, using PHASE2,³¹ in each population sample separately. Coalescent simulations for the haplotype test were performed with the assumption of values of 4*Nr* estimated from the data for each population sample.³² For all tests, gene conversion was assumed to occur twice as often as crossover, and the mean conversion tract length was 500 bp.

Results

LD Structure of the Glucokinase Region

The LD structure of the 116-kb region of *GCK* that is based on 84 SNPs genotyped for 43 unrelated U.K. white subjects is shown in figure 1. We compared the LD structure determined by our resequencing of the *GCK* region with data from HapMap 2 and from the Italian sample of the pop-

Table 4. Haplotype Results for the Association of Fasting Glucose against *rs1799884* and *rs3757840* for All Cohorts

Study	Mean (SD) Glucose Level [No. of Subjects] for Haplotype				Global <i>P</i>	Difference in Glucose (95% CI) [<i>P</i>] for Pairwise Haplotype Comparison ^a		
	GC	GA	AA			GA and GC	AA and GA	AA and GC
ALSPAC Children	4.91 (.01) [707]	4.92 (.01) [455]	4.99 (.02) [241]	.009	.014 (−.02, .05) [.46]	.063 (.01, .11) [.013]	.077 (.03, .12) [.001]	
BCG	4.58 (.02) [571]	4.62 (.02) [439]	4.61 (.03) [208]	.21	.041 (−.01, .09) [.08]	−.01 (−.07, .05) [.70]	.03 (−.03, .09) [.31]	
EFS	4.50 (.01) [1,293]	4.55 (.01) [886]	4.58 (.02) [511]	.002	.04 (.00, .08) [.03]	.08 (.03, .12) [.0008]	.04 (−.01, .08) [.15]	
PEB	4.67 (.03) [277]	4.72 (.04) [178]	4.78 (.05) [103]	.09	.055 (−.03, .14) [.21]	.06 (−.05, .02) [.31]	.11 (.06, .16) [.03]	
BWHHS	5.50 (.01) [2,301]	5.51 (.01) [1,487]	5.55 (.01) [786]	.008	.01 (−.02, .04) [.64]	.05 (.01, .08) [.005]	.05 (.02, .08) [.0007]	
InChianti	4.79 (.02) [817]	4.78 (.02) [666]	4.86 (.03) [384]	.03	−.01 (−.07, .04) [.66]	.08 (.02, .15) [.0109]	.07 (.01, .13) [.03]	

NOTE.—Glucose is measured in mmol/liter.

^a The first allele is *rs1799884*, and the second allele is *rs3757840*. The A allele of both SNPs is the relative glucose-raising allele.

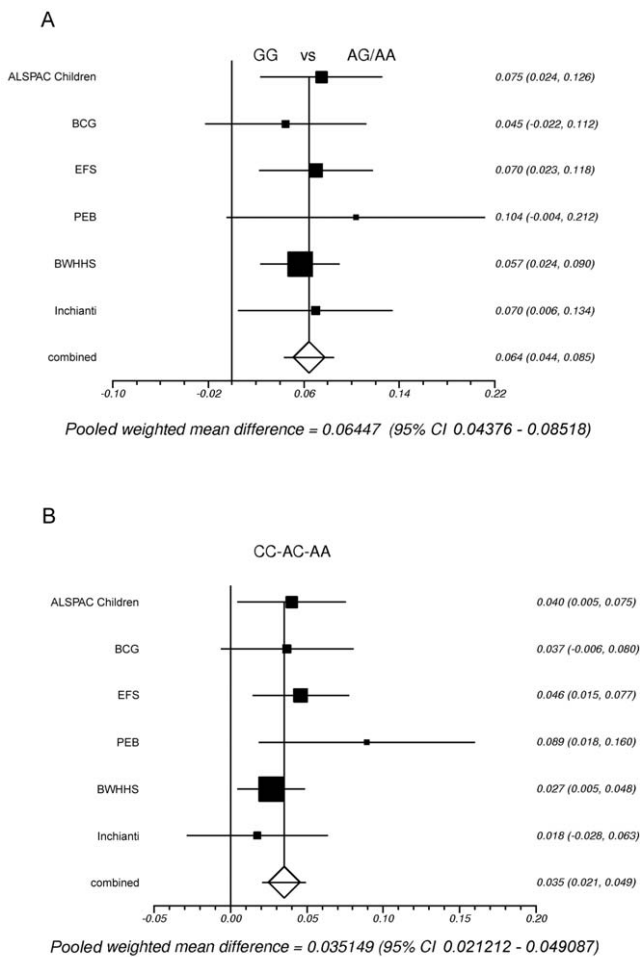


Figure 4. A, Meta-analysis of the fasting glucose association results of *rs1799884* AA and AG versus GG for all studies. Studies are ordered by ascending age of the cohort. Heterogeneity $P = .94$; combined $P = 1 \times 10^{-9}$. B, Meta-analysis of the fasting glucose association results of *rs3757840*. The effect size per allele is presented. Heterogeneity $P = .56$; combined $P = 8 \times 10^{-7}$.

ulation-genetics survey. Seventeen SNPs were in both the HapMap 2 and our data set; figure 2 compares the data sets. Twenty SNPs were identified by both the tSNP association study and the population-genetics survey of the GCK region; figure 3 compares the data sets. Patterns of LD were very similar across all three sets of samples.

tSNP Performance

Of the 23 genotyped tSNPs, 22 passed our stringent quality-control criteria. These 22 SNPs captured all 84 SNPs, with a mean r^2 of 0.837. Of the captured SNPs, 73% SNPs were captured with an $r^2 > 0.8$, 89% were captured with an $r^2 > 0.5$, and 98% were captured with an $r^2 > 0.2$.

Initial tSNP Fasting Glucose Association Analysis

The results for the initial tSNP association analysis are shown in table 1. Two SNPs, *rs3757840* and *rs1799884*, were nominally significant.

Fasting Glucose Replication Studies

We next attempted to replicate the associations in an additional 5,495 subjects from five further studies. The results for both SNPs in the individual cohorts are shown in table 3 and figure 4. Overall, there was strong replication of the SNPs with fasting glucose in these cohorts (overall combined P values: for *rs1799884*, $P = 1 \times 10^{-9}$; for *rs3757840*, $P = 8 \times 10^{-7}$). The result was very similar when samples with impaired fasting glucose (i.e., a cut-off of 7 mmol/liter rather than 6 mmol/liter) were included.

Are There Two Independent Effects?

The SNPs *rs1799884* and *rs3757840* are 2,148 bp apart and are in LD ($D' = 1$; $r^2 = 0.2$). To determine whether these SNPs have independent effects on fasting glucose, we performed haplotype analyses using the unphased program.²³ The haplotype results are presented in table 4 and figure 5. Alleles from haplotypes formed by these two SNPs are given as *rs1799884*, then *rs3757840*. The pairwise comparison of the GA and the GC haplotypes (a test of *rs3757840*) was only marginally significant (fig. 5A) (combined effect size = 0.02 mmol/liter; combined $P = .03$). In contrast, the AA versus GA comparison (a test of *rs1799884*) was highly significant (fig. 5B) (combined effect size = 0.05 mmol/liter; $P = 1 \times 10^{-5}$), and the GC versus AA comparison (a test of both SNPs) was also significant (fig. 5C) (combined effect size = 0.06 mmol/liter; $P = 1 \times 10^{-11}$). This strongly suggests that the *rs1799884* SNP statistically explains the association of the *rs3757840* variant with fasting glucose. Regression analyses, which analyzed the effect of one SNP on fasting glucose while adjusting for the other, produced similar results (data not shown).

Birth-Weight Analysis for *rs1799884*

We next examined the effect of *rs1799884* on birth weight, using 6,366 mothers and 7,232 offspring from the ALSPAC study. Results are shown in table 5. We have replicated our previous results, which shows that the presence of a maternal A allele at *rs1799884* is associated with an increase of 27 g in offspring birth weight ($P = .02$). Correction for fetal genotype did not affect the magnitude of the association (26 g); we still found no evidence of a fetal genotype effect, either before ($P = .72$) or after ($P = .56$) correction for maternal genotype. When we combined this result with our data published elsewhere,¹⁵ the maternal genotype effect size was 32 g (95% CI 11–53 g) greater birth weight for A allele carriers at *rs1799884* ($P = .002$).

Population-Genetics Analyses

The resequencing survey of 48 unrelated individuals from three major ethnic groups, centered on the fasting glucose-associated SNP *rs1799884*, yielded 112 polymorphisms: 105 biallelic SNPs and 7 insertion/deletions.

Summary statistics of sequence variation are presented in table 6. Polymorphism levels are summarized in terms

of the number of polymorphic sites (S) and nucleotide diversity per bp (π). Information about LD levels is summarized by ρ_{H01} , which is a composite likelihood estimator of the population crossover-rate parameter ($4Nr$, where N is the effective population size and r is the crossover rate between adjacent nucleotides).^{25,30} Tajima's D summarizes information about the spectrum of allele frequencies in each sample³³; a value near zero is expected under an equilibrium-neutral model, whereas negative and positive values indicate, respectively, an excess of rare and intermediate frequency variants.

Contrary to the pattern observed in most studies of human variation,^{26,34–39} GCK polymorphism levels in the Hausa sample from sub-Saharan Africa are lower than those observed in both non-African samples. We compared these results with those of the 50 noncoding regions surveyed in the same samples.²⁶ Five of these regions were more variable in the Italian than in the Hausa sample, but none of them showed a difference as large as that observed at GCK . When the Hausa and the Chinese samples were compared, only one noncoding region showed a difference as large as that observed at GCK for the same samples. Hence, polymorphism levels at GCK in the Hausa sample appear to be unusually low relative to those observed in the non-African samples, as might be expected if positive natural selection drove a variant to fixation in the Hausa

Table 5. ALSPAC Study Birth-Weight Association Results for the $rs1799884$ Genotype

Genotype	Mean Birth Weight (SE) [No. of Subjects] for $rs1799884$			P^a
	GG	GA	AA	
Maternal	3,476 (6.7) [4,269]	3,501 (10.1) [1,905]	3,522 (31.8) [192]	.02
Offspring	3,500 (6.3) [4,896]	3,507 (9.6) [2,090]	3,518 (30.9) [246]	.72

NOTE.—Birth weight is measured in grams and is adjusted for gestational and sex.
^a The P values are for a dominant model for the $rs1799884$ A allele.

but not in the other samples. To test whether polymorphism levels in the Hausa are low relative to the expectations of a standard neutral model, we used the HKA test⁴⁰ to compare levels of polymorphism and sequence divergence between human and chimpanzee at GCK with those observed at 50 noncoding regions surveyed in the same samples. This test detected a nonsignificant deficit of polymorphism levels in the Hausa ($P = .148$) and a nonsignificant excess of polymorphism levels in the Italians and Chinese ($P = .068$ and $P = .105$, respectively). Likewise, the allele-frequency spectrum is not unusual, compared with the 50 noncoding regions in the same samples and compared with the expectations of the standard neutral model.

Visual inspection of the inferred haplotypes for each population sample revealed a haplotype class defined by at least seven SNPs ($rs13306391$, $rs34976798$, 65255 [nucleo-

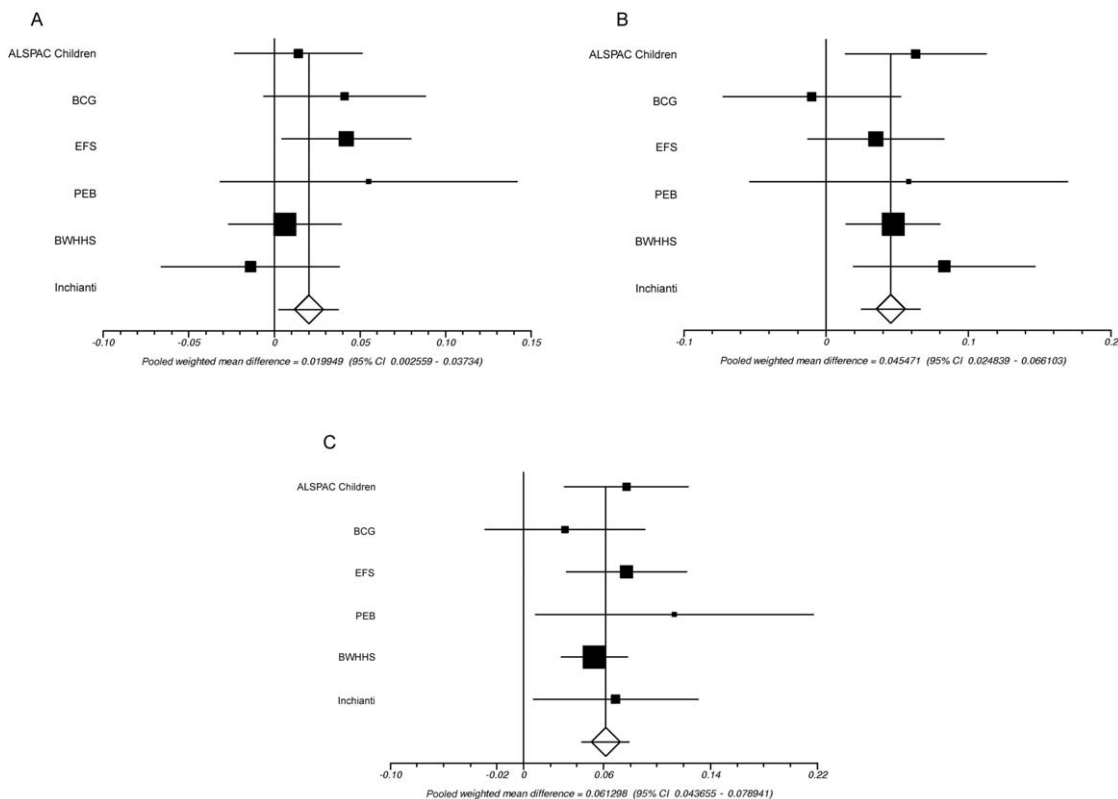


Figure 5. Meta-analysis of the pairwise differences between haplotypes of $rs1799884$ and $rs3757840$ across fasting glucose. *A*, GA versus GC. *B*, AA versus GA. *C*, AA versus GC. Heterogeneity $P = .40$, $.41$, and $.65$, respectively.

Table 6. Summary Statistics of Observed Variation in the Resequencing Survey

Population	S ^a	π^b (%)	Tajima's <i>D</i> ^c	ρ_{H01}^d (%)
Hausa	47	.060	-.491	.0397
Italians	81	.122	.159	.0212
Chinese	75	.116	.268	.0270

^a Number of variable sites.

^b Nucleotide diversity per bp.

^c Summary statistic of the allele frequency spectrum.^{25,30}

^d Composite likelihood estimator of the population recombination rate parameter (*4Nr*) based on a model of gene conversion in which conversion events are twice as common as crossover events and the conversion tract length is geometrically distributed with mean 500 bp.

tion position from start of AC006454, submitted to dbSNP], *rs13239289*, *rs34807880*, *rs34380942*, and *rs34780996*) at high frequency in the non-African samples (27 of 32 inferred haplotypes) and fixed in the Hausa sample. To determine whether positive natural selection acted on this haplotype class, we used the haplotype test,³² which determines whether there is a deficit of variation within a haplotype class, given its frequency and relative to neutrality expectations. Briefly, given the number of variable sites in a population sample, coalescent simulations under neutrality are performed to estimate the probability that a haplotype subset of a given size has as few or fewer polymorphic sites than in the observed haplotype distribution. This test did not detect a significant departure in either the Italian or the Chinese data (*P* > .2).

We also used the results of a recent analysis of the phase I data of the International HapMap Project to assess the evidence of positive natural selection (Haplotter).^{41,42} The analysis is based on a test statistic, called the “integrated haplotype score” (IHS), which measures how unusual the haplotypes around a given SNP are, relative to the genome-wide HapMap data; the larger the absolute value of IHS, the more unusual the haplotypes. The *GCK* gene does not show an unusual clustering of SNPs with large IHS values (*P* = .99 in all HapMap population samples). We also searched for three SNPs of interest: *rs1799884* and *rs3757840*, which are associated with variation in birth weight and fasting glucose, and *rs13239289*, which defines the high-frequency haplotype in the Italians and Chinese. Because *rs3757840* and *rs13239289* are not in the HapMap phase I data, we used surrogate SNPs *rs7810158* and *rs3757838* (*r*² = 0.74 and *r*² = 1, respectively). None of the SNPs is associated with an unusual haplotype structure on a genome-wide scale.

It was shown elsewhere that birth weight is inversely correlated with heat stress, which raises the possibility that genetic variation in birth weight may be shaped by selective pressures related to climate variables.⁴³ To assess whether such spatially varying selective pressures acted on variation in the *GCK* promoter, we genotyped three SNPs in the HGDP panel. Two of the SNPs (*rs1799884* and *rs3757840*) were chosen because they show a signif-

icant association with variation in birth weight and fasting glucose. The third SNP (*rs13239289*) was chosen among those that define the haplotype fixed in the Hausa sample and at near-fixation frequency in the non-African samples. We tested the null hypothesis of no correlation between SNP allele frequency in each HGDP population and a set of climate-related variables for the same populations by using the Spearman rank correlation test. However, a correlation with some of these variables might be expected simply as a result of the history of human migrations and may not necessarily reflect the action of spatially varying selection. To determine whether the observed correlations are unusual compared with empirical genomewide patterns, we compared the rank correlation score for each SNP with the distribution of rank correlation scores for a set of microsatellite loci genotyped in the HGDP panel for each climate variable, as described in the work of Thompson et al.⁴⁴; the proportion of microsatellites with more-extreme rank correlation scores can be considered an empirical *P* value for the correlation observed at the SNPs. As shown in table 7, *rs13239289* has a strong correlation with temperature variables, which is unusual in comparison with the microsatellite distribution. However, this SNP—as determined by genotyping *rs34976798*, which defines the same haplotype and has an *r*² of 1.0 with *rs13239289* in the 48 samples—is not associated with fasting glucose (*P* = .35) in the tSNP analysis (table 1.)

Discussion

We performed a tSNP association analysis of the glucokinase gene to identify common genetic variants associated with fasting glucose and birth weight. Using six studies, we consistently replicated the association of a haplotype, defined by *rs1799884*—previously called “*GCK(-30)*”—with fasting glucose and birth weight. Previous smaller studies of *GCK* by ourselves and others^{15,45,46} that analyzed only *rs1799884* have found a similar increase in fasting glucose associated with the A allele. Our results provide evidence that this variant alters fasting glucose at genomewide levels of significance. Not all individual studies provided sig-

Table 7. Correlation between Allele Frequency and Climate Variables in the HGDP Panel

Climate Variable ^a and Season	Findings for SNP					
	<i>rs13239289</i>		<i>rs3757840</i>		<i>rs1799884</i>	
	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>	<i>r</i>	<i>P</i>
Minimum:						
Summer	-.277	.2176	-.076	.6648	-.081	.7916
Winter	-.542	.0209	-.161	.4034	-.103	.7580
Maximum:						
Summer	-.223	.4184	-.239	.1761	-.146	.5966
Winter	-.549	.0126	-.223	.3068	-.029	.9445
Surface:						
Summer	-.234	.2915	-.200	.2159	-.140	.5664
Winter	-.564	.0126	-.211	.3182	-.096	.7748

^a Mean temperature.

nificant evidence of association, and some provided stronger evidence for *rs3757840*, a SNP partially correlated with *rs1799884*, with an r^2 of 0.22. Our results highlight the importance of using multiple replication studies to establish the true effect of common variants on common phenotypes. Our results also indicate that association signals may be detected with SNPs with r^2 values as low as 0.22.

Our tSNP analysis found no evidence that *GCK* variation, other than the variation in the haplotype that includes *rs1799884*, affects *GCK* activity. The tSNP with the highest r^2 with *rs1799884*—*rs3757840*—was associated with fasting glucose, but the association is explained statistically by LD with *rs1799884* ($r^2 = 0.22$). We also note that the tSNP *rs2971671*, with the second highest r^2 with *rs1799884* ($r^2 = 0.19$), was close to reaching nominal significance in our initial study ($P = .054$). The identification of the signal of *rs1799884* with use of a SNP with a low r^2 highlights the potential of using a tSNP approach for association studies in identification of common variants for complex traits. This is important, since the utility in identification of common disease variants of the HapMap project and whole-genome association studies has been questioned.⁴⁷

We have replicated our previous finding that the maternal *rs1799884* genotype influences offspring birth weight. The association of maternal *rs1799884* with offspring birth weight (27 g) is, as predicted, from the correlation between maternal glucose concentration and offspring birth weight—on the basis of regression of maternal pregnant glucose and birth weight in the EFS mothers, a 0.08 mmol/liter increase in fasting glucose would be expected to increase birth weight by 21 g. We found no evidence that fetal genotype is associated with offspring birth weight (A allele effect size = 4 g; $P = .72$).

The population-genetics analyses of *GCK* was centered on the *rs1799884* SNP associated with fasting glucose and revealed a few interesting patterns. An extended haplotype defined by the derived allele at 17 SNPs occurs at high frequency (84%) in the two non-African samples and is fixed in the Hausa sample. Accordingly, the Hausa sample exhibits lower polymorphism levels compared with the non-African samples. This could be consistent with a complete selective sweep in the Hausa sample and a nearly complete sweep in the non-African samples. However, a statistical test aimed at determining whether the deficit of polymorphism in the Hausa is inconsistent with evolutionary neutrality failed to reject the null hypothesis ($P = .148$). Likewise, the haplotype test did not yield significant results in the non-African samples. Although these results may be due to limited power of the tests, we conclude that there is no evidence in the resequencing data to indicate recent positive natural selection. Interestingly, however, one of the SNPs defining the haplotype fixed in the Hausa and at high frequency in the non-African samples shows a strong correlation with climate variables. Given the reported correlation between birth weight and indices of heat stress,⁴⁵ it is tantalizing to spec-

ulate that *GCK* variation linked to this haplotype may have been shaped by spatially varying selective pressures related to climate.

We also asked whether the variants contributing to fasting glucose and birth-weight variation in our association studies were associated with a selective advantage. However, neither our resequencing data nor an analysis of the HapMap phase I data suggests that this is the case. Given the small phenotypic effect of *rs1799884* on birth weight (i.e., ~32 g), it is perhaps not surprising that no fitness effect appears to be associated with this variant.

Other important conclusions can be drawn from our study. The effect of *rs1799884* on fasting glucose is constant in groups of subjects whose median age varied from 8 years to 71.5 years (heterogeneity $P = .94$) when a cutoff of 6 mmol/liter is taken (i.e., normal glucose-tolerant subjects only).

In conclusion, we have comprehensively assessed the role of the glucokinase gene in determining fasting glucose level and birth weight. Birth weight is a trait strongly influenced by evolutionary pressure, but a comprehensive analysis of *GCK* did not reveal any consistent signals of recent positive selection. We have reproducibly, and at genomewide levels of significance, identified a common haplotype of the glucokinase gene that affects fasting glucose and birth weight.

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Web Resources

The URLs for data presented herein are as follows:

dbSNP, <http://www.ncbi.nlm.nih.gov/SNP/>
Haplotter, <http://hg-wen.uchicago.edu/selection/haplotter.htm>
HKA, <http://genapps.uchicago.edu/hka/index.html>
Kbioscience, <http://www.kbioscience.co.uk/>
MAXDIP, <http://genapps.uchicago.edu/maxdip/index.html>
mVista, <http://genome.lbl.gov/vista/>
Online Mendelian Inheritance in Man (OMIM), <http://www.ncbi.nlm.nih.gov/Omim/> (for *GCK* and *MODY*)

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