Candidate Gene Association Study for Diabetic Retinopathy in Persons with Type 2 Diabetes: The Candidate Gene Association Resource (CARe)

Lucia Sobrin,^{1,2} Todd Green,^{2,3,4} Xueling Sim,⁵ Richard A. Jensen,⁶ E. Shyong Tai,^{7,8,9} Wan Ting Tay,¹⁰ Jie Jin Wang,^{11,12} Paul Mitchell,¹¹ Niina Sandholm,^{13,14} Yiyuan Liu,¹⁵ Kustaa Hietala,^{14,16} Sudha K. Iyengar,¹⁷ for the Family Investigation of Nepbropathy and Diabetes-Eye Research Group,¹⁸ Matthew Brooks,¹⁹ Monika Buraczynska,²⁰ Natalie Van Zuydam,¹⁵ Albert V. Smith,^{21,22} Vilmundur Gudnason,^{21,22} Alex S. F. Doney,²³ Andrew D. Morris,²³ Graham P. Leese,²³ Colin N. A. Palmer,¹⁵ for the Wellcome Trust Case Control Consortium 2,²⁴ Anand Swaroop,¹⁹ Herman A. Taylor, Jr,²⁵ James G. Wilson,²⁶ Alan Penman,^{25,27} Ching J. Chen,²⁸ Per-Henrik Groop,^{13,14} Seang-Mei Saw,⁸ Tin Aung,^{10,29} Barbara E. Klein,³⁰ Jerome I. Rotter,³¹ David S. Siscovick,⁶ Mary Frances Cotch,³² Ronald Klein,³⁰ Mark J. Daly,^{2,3,4} and Tien Y. Wong^{10,12,29}

From the ¹Department of Ophthalmology, Harvard Medical School, Massachusetts Eye and Ear Infirmary, Boston, Massachusetts; the ³Program in Medical and Population Genetics, Broad Institute, Cambridge, Massachusetts; the ⁴Center for Human Genetic Research, Massachusetts General Hospital, Boston, Massachusetts; the ⁵Centre for Molecular Epidemiology and the Departments of ⁷Medicine, ⁸Epidemiology and Public Health, and ²⁹Ophthalmology, National University of Singapore, Singapore; the ⁶Cardiovascular Health Research Unit, Department of Medicine, University of Washington, Seattle, Washington; the ⁹Duke-National University of Singapore Graduate Medical School, Singapore; the ¹⁰Singapore Eye Research Institute, Singapore National Eye Centre, Singapore; the ¹¹Centre for Vision Research, University of Sydney, Sydney, New South Wales, Australia; the ¹²Centre for Eye Research Australia, University of Melbourne, Melbourne, Victoria, Australia; the ¹³Division of Nephrology, Department of Medicine, Helsinki University, Helsinki, Finland; the ¹⁴Folkhälsan Institute of Genetics, Folkhälsan Research Center, Biomedicum Helsinki, Helsinki, Finland; the ¹⁵Biomedical Research Institute, University of Dundee, Dundee, Scotland, United Kingdom; the ¹⁶Department of Ophthalmology, Helsinki University Central Hospital, Helsinki, Finland; the Department of Epidemiology and Biostatistics, Case Western Reserve University, Cleveland, Ohio; the ¹⁸Family Investigation of Nephropathy and Diabetes-Eye Research Group (members listed in the Supplementary Material, http://www.iovs.org/lookup/suppl/ doi:10.1167/iovs.11-7510/-/DCSupplemental); the ¹⁹Neurobiology Neurodegeneration and Repair Laboratory and the ³²Division of Epidemiology and Clinical Applications, Intramural Research Program, National Eye Institute, National Institutes of Health, Bethesda, Maryland; the ²⁰Laboratory for DNA Analysis and Molecular Diagnostics, Department of Nephrology, Medical University of Lublin, Lublin, Poland; the ²¹Icelandic Heart Association, Kopavogur, Iceland; the ²²University of Iceland, Reykjavik, Iceland; the ²³Department of Medicine and Therapeutics, Population Pharmacogenetics Group, Ninewells Hospital and Medical School, Dundee, Scotland, United Kingdom; the ²⁴Wellcome Trust Case Control Consortium (members listed in the Supplementary Material, http://www.iovs. org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental); the Departments of ²⁵Medicine, ²⁶Physiology and Biophysics, ²⁷Biostatistics, and ²⁸Ophthalmology, University of Mississippi Medical Center, Jackson, Mississippi; the ³⁰Department of Ophthalmology and Visual Sciences, University of Wisconsin School of Medicine and

Investigative Ophthalmology & Visual Science, September 2011, Vol. 52, No. 10 Copyright 2011 The Association for Research in Vision and Ophthalmology, Inc. Public Health, Madison, Wisconsin; and the ³¹Medical Genetics Institute, Cedar-Sinai Medical Center, Los Angeles, California. ²These authors contributed equally to the work presented here and should therefore be regarded as equivalent authors.

Supported by National Eye Institute Grant K12-EY16335; a Research to Prevent Blindness Career Development Award; the Massachusetts Lions Eve Research Fund; the Sara Elizabeth O'Brien Trust; and Harvard Catalyst/ The Harvard Clinical and Translational Science Center, National Institutes of Health Award Grant UL1 RR 025758 and financial contributions from Harvard University and its affiliated academic health care centers. The National Heart, Lung Blood Institute (NHLBI) Grant N01-HC-65226 has supported genotyping and has created a genotype-phenotype database with data and samples from nine cohorts as part of Candidate Gene Association Resource (CARe). Please see the Supplementary Material (http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DC Supplemental), as well as http://public.nhlbi.nih.gov/Genetics Genomics/home/care.aspx for detailed information about the grants supporting the four cohorts in this study: Atherosclerosis Risk in Communities (ARIC) Study, Cardiovascular Health Study (CHS), Jackson Heart Study (JHS), and Multi-ethnic Study of Atherosclerosis (MESA). Please see the Supplementary Material (http://www.iovs.org/lookup/suppl/doi: 10.1167/iovs.11-7510/-/DCSupplemental) for detailed information about the grants supporting the replication cohorts.

Submitted for publication March 5, 2011; revised June 30 and August 3, 2011; accepted August 5, 2011.

Disclosure: L. Sobrin, None; T. Green, None; X. Sim, None; R.A. Jensen, None; E.S. Tai, None; W.T. Tay, None; J.J. Wang, None; P. Mitchell, None; N. Sandholm, None; Y. Liu, None; K. Hietala, None; S.K. Iyengar, None; M. Brooks, None; M. Buraczynska, None; N. Van Zuydam, None; A.V. Smith, None; V. Gudnason, None; A.S.F. Doney, None; A.D. Morris, None; G.P. Leese, None; C.N.A. Palmer, None; A. Swaroop, None; H.A. Taylor, Jr, None; J.G. Wilson, None; A. Penman, None; C.J. Chen, None; P.-H. Groop, None; S.-M. Saw, None; T. Aung, None; B.E. Klein, None; J.I. Rotter, None; D.S. Siscovick, None; M.F. Cotch, None; R. Klein, None; M.J. Daly, None; T.Y. Wong, None

Corresponding author: Lucia Sobrin, Massachusetts Eye and Ear Infirmary, 243 Charles Street, Boston, Massachusetts 02114; lucia_sobrin@meei.harvard.edu. **PURPOSE.** To investigate whether variants in cardiovascular candidate genes, some of which have been previously associated with type 2 diabetes (T2D), diabetic retinopathy (DR), and diabetic nephropathy (DN), are associated with DR in the Candidate gene Association Resource (CARe).

METHODS. Persons with T2D who were enrolled in the study (n = 2691) had fundus photography and genotyping of single nucleotide polymorphisms (SNPs) in 2000 candidate genes. Two case definitions were investigated: Early Treatment Diabetic Retinopathy Study (ETDRS) grades ≥ 14 and ≥ 30 . The χ^2 analyses for each CARe cohort were combined by Cochran-Mantel-Haenszel (CMH) pooling of odds ratios (ORs) and corrected for multiple hypothesis testing. Logistic regression was performed with adjustment for other DR risk factors. Results from replication in independent cohorts were analyzed with CMH meta-analysis methods.

RESULTS. Among 39 genes previously associated with DR, DN, or T2D, three SNPs in P-selectin (*SELP*) were associated with DR. The strongest association was to rs6128 (OR = 0.43, P = 0.0001, after Bonferroni correction). These associations remained significant after adjustment for DR risk factors. Among other genes examined, several variants were associated with DR with significant P values, including rs6856425 tagging α -Liduronidase (*IDUA*) ($P = 2.1 \times 10^{-5}$, after Bonferroni correction). However, replication in independent cohorts did not reveal study-wide significant effects. The P values after replication were 0.55 and 0.10 for rs6128 and rs6856425, respectively.

CONCLUSIONS. Genes associated with DN, T2D, and vascular diseases do not appear to be consistently associated with DR. A few genetic variants associated with DR, particularly those in *SELP* and near *IDUA*, should be investigated in additional DR cohorts. (*Invest Ophthalmol Vis Sci.* 2011;52:7593-7602) DOI: 10.1167/iovs.11-7510

Diabetic retinopathy (DR) is the leading cause of blindness in working-age Americans^{1,2} and is increasing in prevalence as rates of type 2 diabetes (T2D) soar worldwide.^{3,4} The frequency and severity of DR are heterogeneous within and across ethnic groups,⁵ even with adjustment for risk factors such as duration of diabetes and glycemic control.^{2,6,7} There are people who have a long duration of diabetes without DR and those who have severe DR despite relatively good glycemic control. For these reasons, genetic risk factors are thought to play a role in DR. Heritability has been estimated to be as high as 27% for DR and 52% for proliferative diabetic retinopathy (PDR).⁸⁻¹⁰ However, genetic association studies for DR have been thus far limited mostly to studies of one or a modest number of candidate genes.^{11,12} Most reported associations have not been consistently reproduced.^{11,13,14}

In contrast to DR, genetic association studies for T2D have revealed many consistently associated genes. Genes that increase T2D risk may also predispose to development of retinopathy. In the case of diabetic nephropathy (DN), a *TCF7L2* variant increases the risk of developing DN beyond the risk of diabetes.¹⁵ Because there is evidence that DR shares risk factors and pathophysiological mechanisms with DN and macrovascular diabetic complications,^{6,16-21} genes associated with DN and atherosclerotic vascular disease may also be associated with DR.

The Candidate gene Association Resource (CARe) is a collaboration for association analyses of genotypes and cardiovascular disease phenotypes.²² It comprises >40,000 participants from nine cohorts who have been genotyped for 49,320 single nucleotide polymorphisms (SNPs) from approximately 2,000 candidate genes postulated or known to increase risk of cardiovascular, metabolic, and inflammatory diseases.²³ It includes 2691 T2D participants with fundus photographs of multiple ethnicities. Thus, the CARe framework provides an opportunity to investigate genetic associations for DR with a candidate gene approach. CARe genotyped many genes previously associated with DR,^{24,25} DN,^{25–27} and T2D.^{28–34} The first purpose of this study was to investigate whether these genes are also associated with the presence of DR in CARe. The second purpose was to determine whether the remaining genes included in the CARe genotyping platform, which were also chosen as potential cardiovascular disease genes, are associated with DR.

METHODS

Study Population and Fundus Photography Procedures

Four CARe cohorts have fundus photographs of T2D participants: Atherosclerosis Risk in Communities (ARIC) Study, Cardiovascular Health Study (CHS), Jackson Heart Study (JHS), and Multiethnic Study of Atherosclerosis (MESA).³⁵⁻³⁸ T2D was defined according to the American Diabetes Association 2003 Criteria.³⁹ The fundus photography protocol for each cohort is described in Table 1.⁴⁰⁻⁴² In all studies, except for the JHS, fundus photographs were graded by masked readers at the University of Wisconsin Ocular Epidemiology Reading Center according to the modified Airlie House Classification system.⁴³ Fundus photographs for the JHS were graded by masked JHS ophthalmologist investigators according to the same criteria.

Definition of Diabetic Retinopathy

We examined two DR phenotypes. First we defined cases as participants with an Early Treatment Diabetic Retinopathy Study (ETDRS) grade ≥ 14 in the eye with the higher ETDRS grade or in the only eye photographed, depending on the study's protocol. These analyses were designed to detect associations with the presence of any DR. Our second phenotype defined cases as participants with ETDRS grade ≥ 30 . The latter was intended to reduce misclassification of patients with minimal signs of DR, which may be seen even in persons without diabetes.⁴⁴⁻⁴⁶ For all analyses, controls were defined as T2D participants with an ETDRS grade < 14 (no DR).

Measurement of Other Variables

Data on DR risk factors were obtained from the study examination at which fundus photography was performed. These included duration of diabetes, fasting blood glucose, systolic and diastolic blood pressures, and fasting total cholesterol.⁴⁷⁻⁵⁰ The procedures for measuring these variables are described in online documentation (www.cscc.unc.edu/ aric/, www.chs-nhlbi.org, and www.mesa-nhlbi.org, provided by the National Heart, Lung, and Blood Institute, National Institutes of Health, Bethesda, MD; and www.jsums.edu/jhs, JHS, Jackson State University, Jackson, MS). Some participants were unaware of a diabetes diagnosis and received the diagnosis based on their laboratory values at the study visit at which they also had fundus photography. For these patients, the duration of diabetes was calculated by halving the number of years between their prior study visit (when they did not meet criteria for T2D) and the visit at which they met criteria. If data on a risk factor were not measured at the fundus photography visit, the information was obtained from the visit closest in time to the fundus photography visit.

Genotyping

CARe participant DNA samples were interrogated on a custom genotyping array (iSelect ITMAT-Broad-CARe [IBC] Chip; Illumina, San Diego, CA). Its design is described elsewhere.²³ SNP selection criteria and genotyping quality control (QC) procedures are explained in the Supplementary Methods (http://www.iovs.org/lookup/suppl/doi:10.1167/ iovs.11-7510/-/DCSupplemental).²³

TABLE 1. CAR	te Participants with T2D	TABLE 1. CARE Participants with T2D and DR Grading by Cohort and Ethnicity	t and Ethnicity				
Cohort/ Population	Eyes Photographed per Participant (n)	Fields Photographed per Eye (n)	Size of Each Field Photographed (deg)	Participants with T2D, DR Grading and IBC Chip Genotyping (n)	Participants with ETDRS Grade <14 (n)	Participants with ETDRS Grade ≥14(n)	Participants with ETDRS Grade ≥30 (<i>n</i>)
ARIC EA Afra	One	One	45	885 439	732 315	153 124	91
CHS EA AfrA tHS	One	One	45	193 54	160 35	33 19	20 15
AfrA	Two	Seven	30	55	26	29	22
EA EA Afra Asa Ha	Two	Two	6 5	176 275 79	140 176 54 151	80 5 80 5 7 5 7 80 5 7 80 5 80 5 80 5 80	11 57 46
All EA Afra				-7 1254 872	1032	222	122
ASA HA				231 231	54 151	25 80	107 14 46
EA, Euro	pean American; AfrA, Af	EA, European American; AfrA, African American; AsA, Asian American; HA, Hispanic American	American; HA, Hispanic	American.			

Statistical Analysis

We first investigated genes that have been previously associated with DR, DN, and T2D. For DR, we chose the genes that had the most robust evidence of association from a comprehensive review of the literature²⁴ and a subsequent strong association with the erythropoietin (*EPO*) gene promoter.²⁵ For DN, we chose genes that have shown nominal associations (P < 0.05) with DN or a related quantitative trait.²⁵⁻²⁷ For T2D, we chose genes with SNPs that had met genomewide significance.^{28-31,33} Supplementary Table S1 (http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental) lists the 39 genes included on the IBC chip that met these criteria. In the second phase, we investigated the remaining genes on the IBC chip which were primarily cardiovascular candidate genes.²³

For genetic association testing, we used χ^2 analysis comparing European-American cases (participants with T2D and DR) to controls (participants with T2D and no DR) in each CARe cohort. Results were combined by Cochran-Mantel-Haenszel (CMH) pooling of the odds ratios (ORs).^{51,52} This CMH method is a robust way of maintaining consistency with individual study ORs while estimating a single fixedeffects OR across all cohorts. We report correction for the multiple association tests performed with per gene Bonferroni correction and permutation testing. Per gene Bonferroni correction was performed because it is an intuitive, easily understood correction. This Bonferroni correction is not as conservative as the one that corrects for the total number of unique genetic loci tested, as would be represented by correcting for the total number of tag SNPs that are not in strong LD. For this reason, we also present the empiric permutation testing correction, which, although less intuitive, does account for this LD between SNPs. We defined statistical significance as P < 0.05 after per gene Bonferroni correction. We chose this threshold for the discovery phase of the experiment, although it is a less stringent threshold than one based on a correction that could completely account for LD between SNPs, to minimize type II error (false negatives) in this initial phase. We apply a more stringent threshold for the replication phase (see below), and the final decision of whether an SNP is truly associated is based on its final P value after replication. For SNPs that were statistically significant in the discovery phase, we performed haplotype analyses using the omnibus test and further examined the associations with logistic regression models that included other DR risk factors. Age and duration of diabetes were defined as continuous variables in years. Fasting glucose and total cholesterol were incorporated as continuous variables in milligrams per deciliter. Systolic and diastolic blood pressures were evaluated as continuous variables (in mm Hg). If a participant was taking antihypertension medication, 15 and 10 mm Hg were added to the systolic and diastolic blood pressure values, respectively.53 Sex and study site were also incorporated. All statistical analyses were performed in PLINK.54

Replication

Top significant findings were pursued in the non-European American populations in CARe and in independent Caucasian cohorts with genome-wide genotyping results: Age, Gene/Environment Susceptibility (AGES) study⁵⁵; Blue Mountains Eye Study (BMES)⁵⁶; Genetics of Diabetes Audit and Research Tayside Study (Go-DARTS)57; Finnish Diabetic Nephropathy (FinnDiane) Study9; Family Investigation of Nephropathy and Diabetes-Eye (FIND-Eye) Study¹⁰; Singapore Malay Eye Study (SiMES)⁵⁸; and the Singapore Prospective Study Program (SP2).⁵⁹ The Medical University of Lublin T2D cohort⁶⁰ performed de novo genotyping for replication. The phenotyping protocols for these studies are described in the Supplementary Methods and Supplementary Table S2 (http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental). Because ETDRS grading was not used by all cohorts, phenotype data were harmonized into two categories that were analogous to ETDRS grade ≥ 14 and ETDRS grade ≥ 30 . Metaanalysis of the discovery cohort and replication cohorts results was performed by CMH pooling of ORs. Statistical heterogeneity was assessed with Cochran's Q statistic.^{61,62} The Q statistic calculates a

weighted sum of the square distances of the observed effects from the null hypothesis of equality of the effects. The weight for each study is the inverse of the variance of the effect estimator so that larger and more accurate studies are weighted more heavily. Statistical significance after replication was defined as a meta-analysis $P < 1 \times 10^{-6}$. This threshold was determined empirically on the basis of the number of genes tested, is analogous to the threshold for genome-wide significance of 5.0×10^{-8} for GWAS,⁶³ and corresponds approximately to a P < 0.05, after Bonferroni correction.

This research adhered to the tenets of the Declaration of Helsinki and was approved by the institutional review boards of the Massachusetts Eye and Ear Infirmary and the primary cohorts. Informed consent was obtained from all participants.

RESULTS

Table 1 shows the number of T2D CARe participants by cohort and ethnicity. The prevalence of DR is similar among the cohorts, although it is higher in the JHS African-American population when compared to the other African-American populations. This discrepancy is probably secondary to the more precise phenotyping performed in the JHS with sevenfield photography. Duration of diabetes and fasting glucose levels were not significantly different between the JHS cohort's and the other cohorts' African-American populations.

Table 2 shows the most significant associations for the analysis of the DR, DN, and T2D genes with any DR in European Americans. Only the associations to three SNPs in the P-selectin gene (*SELP*) were significant (P < 0.05, after Bonferroni correction). The three associated SNPs tagged the only associated haplotype (Supplementary Table S3, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental). In logistic regression including other DR risk factors, the associations to rs6128, rs6133, and rs3917779 remained significant (P = 0.026, 0.022, and 0.026, respectively). The mean values for covariates are presented in Supplementary Table S4 (http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental).

Table 3 shows the most significant associations from the analysis of the DR, DN, and T2D candidate genes with DR defined as an ETDRS grade \geq 30 in European Americans. The three SELP SNPs, along with five SNPs in the fat mass and obesity-associated (FTO) gene, were significantly associated (P < 0.05, after Bonferroni correction). Variants rs12708942, rs9806929, and rs4783824 tagged the only associated haplotype (Supplementary Table S3, http://www.iovs.org/lookup/ suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental). Because the effect of FTO on T2D risk is mediated by its effect on obesity, we executed a logistic regression model including age, sex, and body mass index (BMI). BMI was incorporated as a continuous variable (weight in kilograms divided by height in meters squared). Three FTO SNPs continued to be associated with DR (P = 0.002, 0.002, and 0.001 for rs9926180, rs7500562, and rs12149433, respectively). Of note, an EPO variant, rs551238, was associated with this more stringent definition of DR and had a P < 0.05 after permutation correction but not after Bonferroni correction.

We then examined the strength of these associations in the CARe non-European populations. For any DR, the associations with the *SELP* SNPs were not replicated in the African-, Hispanic-, or Asian-American populations (Table 4). When cases were defined as ETDRS grade \geq 30, the SNPs in *SELP* and *FTO* were not associated, and in logistic regression models including age, sex, and BMI, no association with the *FTO* SNPs was detected (data not shown).

In the second phase of the analysis, we examined the remaining genes on the IBC chip in European Americans. The

					ARIC	U U			CHS	-			MESA				CMH Combined Analysis	ombin	ed An	alysis		P Cori	P Correction
					MAF			W	MAF			W	MAF			W	MAF						
SNP C	Chr Gene	Position	Minor Allele		Cases Controls	OR	Ρ	Cases	Cases Controls	OR	ď	Cases (Controls	OR	Ь	Cases (Controls	OR	195	195	ď	Bonferroni	Permutation
rs6128	1 SELP	167829528	ь	0.08	0.16	0.48	0.0007	0.06	0.18	0.30	0.02	0.08	0.20	0.36	0.02	0.08	0.17	0.43	0.31	0.62 3	3.1×10^{-6}	0.0001	0.0001
rs6133	1 SELP	167831970	V	0.05	0.12	0.39	0.0003	0.03	0.13	0.22	0.02	0.07	0.13	0.50	0.16	0.05	0.12	0.38	0.25	0.59 1	$1.1 imes 10^{-5}$	0.0004	0.0001
rs3917779	1 SELP	167837472	V	0.05	0.12	0.40	0.0004	0.03	0.12	0.23	0.03	0.07	0.13	0.50	0.16	0.05	0.12	0.39	0.26		1.6×10^{-5}	0.0006	0.0001
rs12262390 1	0 HHEX	94436103	C	0.12	0.09	1.48	0.05	0.15	0.10	1.61	0.22	0.18	0.09	2.27	0.03	0.14	0.09	1.61	1.18	2.20	0.002	0.10	0.004
rs9356754	6 CDKALI	I 20916721	G	0.51	0.43	1.37	0.01	0.53	0.44	1.43	0.18	0.40	0.38	1.09	0.74	0.49	0.42	1.32	1.08	1.63	0.007	0.29	0.008
rs9465904	6 CDKALI	1 20929641	C	0.37	0.30	1.38	0.01	0.42	0.33	1.49	0.15	0.28	0.26	1.08	0.79	0.36	0.30	1.34	1.08	1.65	0.008	0.33	0.01
rs11466493	3 TGFBR2	2 30661786	G	0.01	0.04	0.14	0.002	0.06	0.05	1.23	0.72	0.04	0.07	0.59	0.41	0.02	0.05	0.41	0.21	0.80	0.009	0.35	0.01
rs17025862	3 TGFBR2	2 30660132	G	0.01	0.04	0.14	0.002	0.06	0.05	1.23	0.72	0.04	0.07	0.59	0.41	0.02	0.05	0.41	0.21	0.80	0.009	0.35	0.01
rs7747989	6 CDKALI	I 20921354	C	0.37	0.30	1.35	0.02	0.44	0.33	1.60	0.09	0.28	0.26	1.08	0.79	0.36	0.30	1.33	1.07	1.65	0.009	0.36	0.01
rs2328572	6 CDKALI	I 21323815	V	0.003	0.02	0.15	0.03	0	0.03	NA	NA	0.01	0.03	0.48	0.48	0.005	0.02	0.19	0.05	0.69	0.01	0.44	0.01
rs10214694	6 CDKALI	I 21321533	Ţ	0.003	0.02	0.15	0.03	0	0.03	NA	NA	0.01	0.03	0.48	0.48	0.005	0.02	0.19	0.05	0.69	0.01	0.45	0.01
rs12190631	6 CDKAL1	I 21020651	G	0.05	0.03	1.38	0.30	0.12	0.02	6.17	0.0001	0.01	0.01	1.29	0.83	0.05	0.03	1.85	1.14	3.01	0.01	0.51	0.02
rs2275729 1	0 HHEX	94442410	G	0.15	0.12	1.32	0.13	0.15	0.12	1.29	0.51	0.21	0.10	2.26	0.02	0.16	0.12	1.44	1.08	1.92	0.01	0.52	0.02
rs1511024	4 FABP2	120459629	Т	0.05	0.03	1.74	0.07	0.05	0.02	3.00	0.12	0.06	0.03	1.76	0.35	0.05	0.03	1.86	1.13	3.06	0.01	0.55	0.01
rs9350294	6 CDKALI	I 20978072	Τ	0.38	0.32	1.26	0.07	0.44	0.32	1.65	0.07	0.33	0.29	1.24	0.45	0.38	0.32	1.31	1.05	1.61	0.01	0.55	0.01

				W	MAF			M	MAF			M	MAF			R.	MAF						
SNP Chr	Gene	BP	Minor Allele	Cases	Cases Controls	OR	Ь	Cases (Controls	OR	Ь	Cases	Controls	OR	Ь	Cases	Controls	OR	£61	195	Ρ	Bonferroni	Bonferroni Permutation
rs6133 1	SELP	167831970	V	0.04	0.12	0.34	0.002	0	0.12	NA	NA	0.09	0.12	0.74	0.69	0.05	0.12	0.32	0.17	0.58	$2.3 imes 10^{-4}$	0.00	0.0002
rs3917779 1	SELP	167837472	Y	0.04	0.12	0.35	0.003	0	0.12	ΝA	NA	0.09	0.12	0.74	0.69	0.05	0.12	0.32	0.17	0.59	$2.9 imes10^{-4}$	0.01	0.0001
rs9926180 16	FTO	52486108	Т	0.33	0.25	1.51	0.01	0.38	0.25	1.81	0.08	0.32	0.17	2.32	0.07	0.34	0.24	1.65	1.24	2.18	$5.0 imes10^{-4}$	0.02	0.001
rs7500562 16	FTO	52488391	C	0.33	0.25	1.51	0.01	0.38	0.25	1.81	0.08	0.32	0.17	2.27	0.08	0.34	0.24	1.64	1.24	2.17	$5.3 imes10^{-4}$	0.02	0.001
rs12149433 16	FTO	52485580	G	0.15	0.09	1.75	0.01	0.20	0.08	2.84	0.01	0.05	0.04	1.07	0.95	0.15	0.0	1.93	1.33	2.81	$5.5 imes10^{-4}$	0.02	0.001
rs6128 1	SELP	167829528	Т	0.08	0.16	0.49	0.009	0.05	0.17	0.26	0.05	0.09	0.18	0.45	0.27	0.08	0.16	0.44	0.27	0.70	$5.6 imes10^{-4}$	0.02	0.001
rs13335343 16	FTO	52488941	Υ	0.15	0.09	1.74	0.01	0.20	0.08	2.84	0.01	0.05	0.04	1.07	0.95	0.15	0.09	1.92	1.32	2.80	$6.1 imes10^{-4}$	0.02	0.001
rs12935710 16	FTO	52500306	T	0.32	0.23	1.55	0.01	0.33	0.25	1.48	0.28	0.27	0.15	2.09	0.14	0.32	0.22	1.61	1.21	2.14	0.001	0.04	0.002
rs10852525 16	FTO	52496582	Y	0.17	0.12	1.52	0.05	0.20	0.10	2.29	0.05	0.09	0.05	2.09	0.34	0.17	0.11	1.71	1.20	2.45	0.003	0.12	0.005
rs9929152 16	FTO	52496904	G	0.33	0.25	1.45	0.03	0.35	0.26	1.53	0.23	0.32	0.20	1.89	0.18	0.33	0.25	1.52	1.15	2.02	0.004	0.14	0.006
rs4783824 16	FTO	52509162	T	0.13	0.09	1.50	0.09	0.20	0.08	2.95	0.01	0.05	0.03	1.69	0.62	0.14	0.08	1.78	1.21	2.64	0.004	0.15	0.005
rs1362570 16	FTO	52491048	C	0.14	0.10	1.52	0.07	0.20	0.09	2.45	0.03	0.05	0.04	1.07	0.95	0.14	0.09	1.70	1.16	2.48	0.007	0.26	0.009
rs4611524 17	FTO	58945384	Т	0.31	0.41	0.67	0.02	0.33	0.42	0.68	0.27	0.36	0.40	0.86	0.74	0.32	0.41	0.69	0.52	0.91	0.009	0.34	0.009
rs12708942 16	FTO	52503705	Υ	0.14	0.10	1.46	0.1	0.20	0.09	2.63	0.02	0.05	0.04	1.15	0.89	0.14	0.09	1.68	1.14	2.47	0.009	0.35	0.01
rs9806929 16	FTO	52507417	Y	0.14	0.10	1.46	0.1	0.20	0.09	2.63	0.02	0.05	0.04	1.15	0.89	0.14	0.0	1.68	1.14	2.47	0.009	0.35	0.01
rs551238 7	EPO	100159464	G	0.34	0.40	0.75	0.08	0.20	0.40	0.37	0.01	0.32	0.36	0.82	0.67	0.31	0.40	0.69	0.52	0.91	0.009	0.37	0.01

TABLE 4. CHM Association Results for *SELP* SNPs in non-European-American CARe Populations, with Cases Defined as ETDRS Grade ≥ 14

		1	MAF				
SNP	Minor Allele	Cases	Controls	OR	L95	U95	P
African American							
rs6128	Т	0.5	0.46	1.17	0.95	1.44	0.14
rs6133	С	0.42	0.45	0.89	0.72	1.09	0.26
rs3917779	G	0.48	0.50	0.94	0.76	1.15	0.55
Hispanic American							
rs6128	Т	0.29	0.26	1.14	0.74	1.75	0.55
rs6133	Α	0.16	0.18	0.91	0.55	1.52	0.72
rs3917779	Α	0.14	0.16	0.89	0.52	1.52	0.67
Asian American							
rs6128	С	0.24	0.27	0.86	0.40	1.87	0.70
rs6133	Α	0	0	NA	NA	NA	NA
rs3917779	Α	0	0	NA	NA	NA	NA

L95, lower 95% CI boundary; U95, upper 95% CI boundary; NA, not available.

top association results for DR defined as ETDRS grade ≥ 14 and ETDRS grade ≥ 30 are shown in Tables 5 and 6, respectively. The lambdas for the quantile-quantile (Q-Q) plots were 1.01 and 1.00, respectively; Fig. 1). Supplementary Table S5 (http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DC Supplemental) shows the results from the principal components analysis. There was no evidence of population stratification. Several variants in the two analyses had associations that were significant after Bonferroni correction. One variant, rs6856425, was significantly associated with DR in all three CARe cohorts for both definitions of DR with $P = 2.1 \times 10^{-5}$ after Bonferroni could not be replicated in the CARe African American cohorts (MAF 18%, OR = 0.94, P = 0.69).

We pursued replication of top findings from Tables 5 and 6 in independent cohorts of European ancestry with a fixedeffects meta-analysis model adjusted for age and sex (Table 7, Supplementary Fig. S1, http://www.iovs.org/lookup/suppl/ doi:10.1167/iovs.11-7510/-/DCSupplemental). None of the variants achieved significance in replication ($P < 1 \times 10^{-6}$). The smallest *P* value was for rs35260 (P = 0.03). For all SNPs examined in replication, there was a significant amount of heterogeneity (P < 0.05 for Q test). We performed a sensitivity analysis by removing the FinnDiane and Go-DARTS cohortsthe former because it had type 1 diabetes participants exclusively and both because they did not use ETDRS grading consistently for phenotyping (Supplementary Table S6, http:// www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/ DCSupplemental). However, none of the associations were statistically significant in this analysis either, and significant heterogeneity remained for all SNPs with the exceptions of rs3917779 (Q test P = 0.06) and rs6856425 for the ETDRS grade ≥ 14 analysis (Q test P = 0.26). Given the significant residual heterogeneity, we then used a random effects model but found no significant difference in the results. We also performed meta-analyses without the CARe cohorts; there was no statistically significant result or any significant heterogeneity in these analyses. Of note, meta-analyses of the CARe cohorts alone also showed no significant heterogeneity.

In addition to the above replication efforts in European cohorts, we pursued replication of the same findings in two Asian cohorts, SiMES and SP2. None of the SNPs was statistically significant in these populations. We also investigated the *FTO* association in Go-DARTS; neither rs9926180 nor rs12935710 was significantly associated (P = 0.84 and 0.23, respectively).

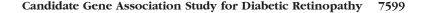
TABLE 3. Top Association Results in European-American Samples with Cases Defined as ETDRS Grade ≥ 30 for Variants in Genes Previously Associated with DR, DN, or T2D

P correction		Bonferroni Permutation	-5 0.0001	-5 0.0001	-	Ŭ	-	-	0.0001	0.0001	0.0001	0.0001	0.0001	0.0002	0.0002	0.0001	0.0001	0.0003	0.0001	0.0002
P 0		Bonferro		4.6×10^{-5}				7.6×10^{-5}	0.0001	0.0003	0.0003	0.0004	0.0005	0.0006	0.001	0.001	0.002	0.002	0.004	0.004
		Ь	1.1×10^{-6}	$1.2 imes10^{-6}$	1.3×10^{-6}	1.3×10^{-6}	$1.8 imes 10^{-6}$	$2.0 imes 10^{-6}$	$3.1 imes 10^{-6}$	7.6×10^{-6}	$8.6 imes 10^{-6}$	1.1×10^{-5}	$1.3 imes 10^{-5}$	1.6×10^{-5}	2.9×10^{-5}	$3.5 imes 10^{-5}$	$5.6 imes 10^{-5}$	6.4×10^{-5}	$9.1 imes 10^{-5}$	9.4×10^{-5}
nalysis		195	0.61	0.61	0.61	0.61	0.62	0.74	0.62	1.98	0.77	0.59	0.61	0.60	4.73	2.53	1.88	8.21	2.38	0.77
oined A		1 T62	3 0.31	4 0.31	4 0.31	4 0.31	4 0.32	0 0.48	3 0.31	1 1.31	2 0.50	8 0.25	1 0.28	9 0.26	8 1.75	8 1.39		1 2.05	8 1.34	9 0.46
CMH Combined Analysis			0.4	0.44	0.44	0.44	0.4	0.60	Ŭ		0.62	0.38	0.41	0.39	2.88	1.88	1.53	4.11	1.78	0.59
CMF	MAF	Cases Controls	0.19	0.19	0.19	0.19	0.18	0.48	0.17	0.33	0.48	0.12	0.14	0.12	0.02	0.09	0.43	0.03	0.10	0.26
		Cases	0.09	0.09	0.09	0.09	0.09	0.35	0.08	0.44	0.36	0.05	0.06	0.05	0.05	0.15	0.53	0.10	0.16	0.17
		Ρ	0.05	0.05	0.05	0.05	0.05	7.4×10^{-5}	0.02	0.13	0.0004	0.16	0.34	0.16	0.19	0.004	0.14	0.001	0.04	0.65
SA		OR	0.46	0.46	0.46	0.46	0.46	0.31	0.36	1.50	0.36	0.50	0.66	0.50	2.28	2.81	1.48	5.88	2.14	0.87
MESA	MAF	Cases Controls	0.21	0.21	0.21	0.21	0.21	0.50	0.20	0.33	0.50	0.13	0.14	0.13	0.03	0.08	0.42	0.02	0.09	0.28
	W	ases (0.11	0.11	0.11	0.11	0.11	0.24	0.08	0.43	0.26	0.07	0.10	0.07	0.06	0.19	0.51	0.10	0.18	0.25
		Р О	0.02	02	0.02	03	0.03	0.07	0.02	0.04	07	02	0.05	0.03	0.0004	0.76	0.02	0.01	0.55	0.94
		OR	0.35 0.	-	-	-	-	0.61 0.	0.30 0.	1.76 0.		0.22 0.	0.32 0.		6.17 0.		1.93 0.	3.33 0.	1.29 0.	
CHS	ιF	ontrols	0.19		0.19			0.48	0.18	0.33		0.13			0.02	0.09	0.43	0.03	0.10	0.25
	MAF	Cases Controls	0.08	0.08	0.08	0.08	0.08	0.36	0.06	0.47	0.36	0.03	0.05	0.03	0.11	0.11	0.59	0.11	0.12	0.24
		Ь	$8.8 imes 10^{-5}$	9.4×10^{-5}	0.0001	$3.8 imes 10^{-5}$	5 0.0001 0	0.0003	0.0007	0.0002	0.005	0.0003	8.9×10^{-5}	0.0004	0.01	0.0006	0.002	NA	0.0007	6.6×10^{-6}
C		OR	0.44 8	0.44 9	0.45	0.44 8	0.45	0.68	0.48	1.60	0.70	0.39	0.37 8	0.40	2.36	1.87	1.47	NA	1.82	0.46 6
ARIC	MAF	Cases Controls	0.18	0.18	0.18	0.18	0.18	0.47	0.16	0.32	0.47	0.12	0.14	0.12	0.02	0.09	0.43	NA	0.10	0.26
	W	Cases	0.09	0.09	0.09	0.09	0.09	0.38	0.08	0.43	0.39	0.05	0.06	0.05	0.04	0.15	0.52	NA	0.16	0.14
		Minor Allele	0	C	Г	c	C	V	Г	V	Г	V	Г	V	c	Г	V	G	C	C
		Position	167794699	167796694	167805168	167797210	167795655	58668304	167829528	99782444	58669348	167831970	167791936	167837472	966918	58654401	99782862	7944506	58588070	114717935
		Gene	5	5	F5	F5	F5	PDE4D	SELP	PCSK6	PDE4D	SELP	F5	SELP	Near IDUA	PDE4D	PCSK6	MTRR	PDE4D	CADMI
		Chr	1	1 F	1 F	1 F	1 F	5 F	1 5	15 F	5 F	1 5	1 F	1 5	4 Λ	5 F	15 F	5 A	5 F	11 0
		SNP	rs9332570	rs7545236	rs2298908	rs1894701	rs6427203	rs35260	rs6128	rs7168655	rs35259	rs6133	rs9332579	rs3917779	rs6856425	rs1824159	rs1006273	rs16879334	rs10036406	rs7105871

The minor allele is the effect allele for the ORs. Chr, chromosome; L95, lower 95% CI boundary; U95, upper 95% CI boundary; NA, not available.

TABLE 6. Top Association Results in European-American Samples with Cases Defined as ETDRS Grade \geq 30 for Variants within All Genes on the IBC Chip

				W	MAF			M	MAF			N	MAF			MAF	F						
SNP Chr	ur Gene	Position	Minor Allele	Cases (Cases Controls OR	OR	d	Cases	Cases Controls	OR	Ь	Cases	Cases Controls	OR	P O	Cases Controls	ontrols	OR	195	U95	Ь	Bonferroni	Bonferroni Permutation
rs6856425 4	4 Near IDUA	966918	0	0.05	0.02	3.01	0.002	0.15	0.02	8.50	$1.8 imes 10^{-5}$	0.09	0.03	3.54 (0.10	0.07	0.02	3.83	2.26	6.46 5.	$5.3 imes 10^{-7}$	$2.1 imes 10^{-5}$	0.0001
rs1555179 6	6 MTHFDIL	151313049	V	0.19	0.35	0.45 3.	$0.45 \ 3.0 \times 10^{-5}$	NA	NA	ΝA	NA	0.18	0.32	0.47 (0.17	0.19	0.34	0.45	0.32	0.65 1	1.3×10^{-5}	0.0005	0.0002
rs7078169 10	10 CUBN	17202320	H	0.09	0.03	2.86	0.0002	0.10	0.03	3.71	0.02	0.05	0.05	0.93 (0.94	0.09	0.04	2.66	1.65	4.27 5.	$5.3 imes 10^{-5}$	0.002	0.0001
rs3901540 5	5 PDE4D	58465490	F	0.15	0.28	0.44	0.0001	0.18	0.29	0.52	0.13	0.32	0.32	0.98 (0.96	0.17	0.29	0.50	0.35	0.70 5	$5.6 imes 10^{-5}$	0.002	0.0002
rs6663966 1	1 GLISI	53842601	C	0.04	0.01	5.16 5.	$5.2 imes10^{-5}$	0.00	0.01	ΝA	NA	0.09	0.02	6.46 (0.01	0.04	0.01	4.15	2.07	8.34 6	6.2×10^{-5}	0.002	0.0009
rs4697843 4	4 HS3ST1	10704856	V	0.21	0.33	0.52	0.0005	0.20	0.33	0.51	0.10	0.18	0.31	0.50	0.21	0.20	0.33	0.52	0.38	0.72 6	6.3×10^{-5}	0.002	0.0001
rs4521758 8	3 PRKDC	48904123	F	0.17	0.11	1.59	0.03	0.30	0.11	3.47	0.0007	0.23	0.09	2.92	0.04	0.20	0.11	1.98	1.41	2.77 7.	$7.3 imes 10^{-5}$	0.003	0.0006
rs10752067 10	D CUBN	17188898	U	0.09	0.04	2.76	0.0003	0.10	0.03	3.73	0.02	0.05	0.05	0.93 (0.94	0.09	0.04	2.59	1.62	4.16 7.	$7.9 imes 10^{-5}$	0.003	0.0001
rs3917462 1	1 Near SELE	167958291	C	0.03	0.01	4.03	0.006	0.05	0.01	9.05	0.009	0.05	0.01	5.16 (0.12	0.03	0.01	4.75	2.17	10.36 9.	$9.3 imes 10^{-5}$	0.004	0.001
rs2508450 11	11 IL10RA	117369039	Н	0.34	0.44	0.65	0.008	0.28	0.49	0.40	0.01	0.23	0.44	0.38 (0.06	0.32	0.45	0.58	0.44	0.76	0.0001	0.004	0.0001
rs2195611 2	2 PDEIA	182964812	C	0.59	0.44	1.83	0.0001	0.58	0.45	1.69	0.12	0.45	0.48	0.91	0.83	0.58	0.45	1.68	1.29	2.19	0.0001	0.005	0.0001
rs11163323 1	1 Near PRKACB	81759046	Г	0.33	0.19	2.05 1.	$1.7 imes10^{-5}$	0.23	0.23	0.98	0.96	0.23	0.19	1.24 (69.0	0.30	0.20	1.76	1.32	2.35	0.0001	0.005	0.0002
rs11155762 6	6 MTHFDIL	151314489	Τ	0.19	0.35	0.45 3.	$3.0 imes10^{-5}$	0.38	0.34	1.17	0.64	0.18	0.32	0.47	0.17	0.22	0.34	0.55	0.40	0.75	0.0001	0.006	0.0005
rs17186372 1	1 NOTCH2	120302474	G	NA	NA	NA	NA	0.25	0.07	4.28	0.0002	0.14	0.07	2.08	0.26	0.21	0.07	3.45	1.81	6.56	0.0002	0.006	0.001
rs6583305 3	3 UBXN7	197603018	Г	0.47	0.36	1.63	0.002	0.55	0.34	2.39	0.008	0.36	0.36	1.00	0.99	0.48	0.35	1.66	1.27	2.16	0.0002	0.007	0.0003
rs6764533 3	3 UBXN7	197572861	V	0.47	0.36	1.63	0.002	0.55	0.34	2.36	0.009	0.36	0.36	1.00	0.99	0.48	0.35	1.65	1.27	2.15	0.0002	0.007	0.0004
rs11545078 8	8 <i>GGH</i>	64101318	V	0.02	0.10	0.15	0.0003	0.08	0.11	0.64	0.47	0.05	0.11	0.37 (0.33	0.03	0.10	0.26	0.13	0.53	0.0002	0.008	0.0002



(Fixed Effects)

Meta-analysis

Lublin

RTS

7. Replication Results in European Samples

TABLE '

620 576 138 0.55

1.03 1.07

> 1.55 0.42 1.39

0.03

0.91

2.24

¥ Z

NA NA 0.16

8

10. NA

2

OR

Score

2

OR

MAF

٩

0.17

0.98 0.88

NA 1.84 NA NA 70.0

> L.03 NA NA

NA 0.08 0.05 0.05 NA 0.05

0.21 0.6 0.45 NA 0.27 NA 0.27 0.27

0.27

1.13 0.9 1.28

NA 00.

0.1

0.68

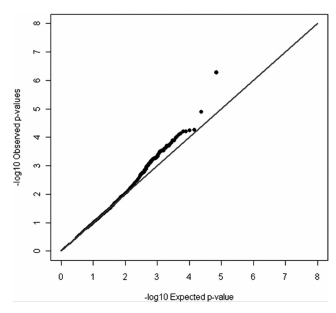


FIGURE 1. Quantile-quantile plot of all single SNPs examined on the IBC chip and their CMH association analysis to diabetic retinopathy, defined as an ETDRS score \geq 30.

DISCUSSION

In this large international collaborative study, genes previously linked with T2D, DR, and DN and vascular diseases were not generally associated with DR. In the CARe European American population, among genes that have been previously associated with DR, DN, and T2D, three SNPs in SELP were associated with DR, even after adjustment for DR risk factors. However, we were unable to replicate this finding in other ethnic groups in CARe or in independent Caucasian cohorts. The SELP SNPs associated with DR in the present study were not in LD with rs6131, the SNP initially associated with diabetic albuminuria (Supplementary Fig. S2, http://www.iovs.org/lookup/suppl/ doi:10.1167/iovs.11-7510/-/DCSupplemental).⁶⁴ P-selectin plays a role in leukocyte adhesion to endothelium during inflammation, and thus there is a biological rationale for its role in both diabetic microalbuminuria and retinopathy.⁶⁵ With regards to FTO, the SNPs associated with DR in CARe were not in significant LD with rs9939609, the SNP associated with T2D (Supplementary Fig. S2, http://www.iovs.org/lookup/suppl/doi:10.1167/ iovs.11-7510/-/DCSupplemental).66

Importantly, we were unable to confirm an association to most genes that have previously been associated with DR and were included on the IBC chip. We note that for several of these genes, the IBC chip did not include SNPs in LD to the previously associated variants because the selection of tag SNPs may not have densely covered those genes (Supplementary Table S1, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs. 11-7510/-/DCSupplemental). For some genes, most notably EPO, we did have excellent proxies to the initially reported variants. The chip included two perfect proxies (rs551238 and rs1734907) for the EPO SNP originally associated to PDR, rs1617640.25 When we defined cases as having an ETDRS grade \geq 30, rs551238 had a significant effect consistent with that found in the previous study, where the minor allele is protective (OR = 0.69, P = 0.009); the P value remains significant after correction by permutation (P = 0.01) but not by the Bonferroni method (P = 0.37). It is possible that with a larger sample size, the association would withstand the Bonferroni correction. We were also unable to detect an association to DR in other genes previously associated with DN and T2D. For the

				ARIC	Ū	CHS	W	MESA		AGES		-	BMES		H	FIND-Eye		Fin	FinnDiane	e)	ġ	Go-DAR
Controls, n				732	1	160	Т	140		249			175			627			570			774
Cases ETDRS ≥14, <i>n</i>	S ≥14, n	-		153	6	33		36		92			67			105			2009			923
Cases ETDRS $\geq 30, n$	S ≥30, n	_		91	7	20		11		37			26			NA			1399			923
SNP	Minor Defin Allele of C	Definition of Cases	OR	Ь	OR	d	OR	Ρ	MAF	OR	Δ	MAF	OR	Ρ	MAF	OR	ď	MAF	OR	d,	MAF	OR
rs9332570	6	ETDRS ≥14	0.43	0.0001	0.33	0.02	0.49	0.06	NA	0.75	0.26	0.21	1.24	0.42	0.19	0.79	0.26	0.27	1.16	0.05	NA	NA
rs35260	V	$ETDRS \ge 14$	0.68	0.002	0.59	0.08	0.29	0.0001	Ν	1.01	0.98	0.49	1.23	0.32	0.48	0.9	0.45	0.5	0.96	0.59	0.44	0.98
rs6128	Т	$ETDRS \ge 14$	0.48	0.0007	0.27	0.02	0.39	0.03	0.09	1.27	0.32	0.21	1.27	0.37	0.17	0.73	0.14	0.24	1.16	0.07	NA	1.13
rs7168655	Υ	$ETDRS \ge \! 14$	1.59	0.0004	1.78	0.04	1.48	0.15	NA	0.91	0.57	0.34	0.96	0.83	0.36	1.08	0.63	0.34	1.02	0.82	0.34	0.96
rs6133	V	ETDRS ≥ 14	0.39	0.0005	0.19	0.03	0.51	0.16	0.15	0.73	0.32	0.14	1.37	0.28	0.13	0.72	0.17	0.09	1.13	0.31	NA	1.08
rs3917779	Υ	ETDRS ≥ 14	0.40	0.0007	0.20	0.03	0.51	0.16	0.09	0.73	0.32	0.14	1.25	0.45	0.12	0.72	0.19	0.09	1.14	0.29	NA	Ν
rs6856425	С	ETDRS ≥ 14	2.48	0.01	5.21	0.004	2.63	0.16	0.04	0.89	0.86	0.02	1.8	0.45	0.02	1.18	0.75	0.03	0.97	0.87	0.02	1.34
rs7105871	С	ETDRS ≥ 14	0.47	$1.7 imes10^{-5}$	0.97	0.91	0.86	0.62	NA	0.81	0.2	0.25	1.16	0.54	0.22	0.88	0.47	0.31	1	0.96	NA	ΥN
rs6856425	C	ETDRS ≥30	3.19	0.003	6.8	0.002	3.25	0.19	NA	ΝA	NA	NA	NA	NA	ΝA	ΝA	NA	0.03	0.79	0.39	0.02	1.34

DN genes, again the IBC chip may not have included SNPs in LD with the previously reported variants. For T2D, however, the IBC chip variants were specifically those previously associated at genome-wide significant levels.

Another explanation for the inability to replicate previous DR associations lies in the heterogeneity among studies regarding DR definitions and participants' mean duration of diabetes. We attempted to mitigate the heterogeneity of DR definitions by examining two different definitions. However, this may not be sufficient to account for all the possible phenotype heterogeneity. The studies from which we selected genes deemed to be previously associated with DR all used controls that were diabetic patients without DR, as we did; however, some of them were performed in type 1 diabetic patients, which is another potential source of heterogeneity. There was also great variability in the duration of T2D among cases and controls in CARe cohorts. In particular, there were participants with short durations of diabetes who were included. There is the potential for misclassification of controls if these participants did not have DR at the time of study inclusion but are at risk for significant DR with longer duration of diabetes. We attempted to correct for this by including duration of disease as a covariate in logistic regression, but these issues could still bias the results toward the null. However, our ability to detect an association with EPO indicates that the amount of control misclassification in CARe is not significant enough to prevent detection of associations of this effect size.^{25,67} Finally, while the current investigation is the largest candidate gene study for DR to date, it still has limited power to detect genetic associations of modest or small effects (Supplementary Table S7, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/ DCSupplemental). In particular, CARe includes a modest number of European-American cases (122) defined as ETDRS grade \geq 30. Examining milder degrees of DR as outcomes may have further decreased our ability to detect associations as the development of early DR has a lower heritability. Of note, the ARIC cohort is larger than the other cohorts, and the top findings in the analyses were often driven by the results in ARIC.

In the second phase of the analysis, we took an unbiased approach at the remaining genes available on the IBC chip. Although we found several strong associations in our discovery cohort, replication in independent samples did not yield variants with consistent effects nor any variants that met the replication significance threshold ($P = 1 \times 10^{-6}$). The rs6856425 association was initially compelling because it was consistent within each CARe cohort. Furthermore, the strength of the association was greater when DR was defined as ETDRS grade ≥ 30 vs. ETDRS grade ≥ 14 , which is in line with the expected greater heritability of more advanced DR phenotypes. The rs35260 variant had the lowest *P* value in the replication meta-analyses (P = 0.03), but this was still far below the replication threshold for significance ($P = 1 \times 10^{-6}$).

Failure to replicate a genetic association can be explained broadly, either as a false positive in the discovery cohort or a false negative in the replication cohort. For rs6856425, a rare variant, the initial estimate was based on limited instances of the minor allele: 9 in cases and 20 in controls. Small numbers of observations can lead to unstable effect estimates that represent chance statistical fluctuations rather than true associations. This underscores the importance of large sample sizes in both discovery and replication cohorts, particularly for rare variants. Another possible reason for false positives is population stratification, but there was no significant population stratification in this study.

False negatives in the replication cohort can be due to a lack of power, genotyping/imputation imprecision or heterogeneity between cohorts. Our aggregate replication cohort sample was well powered (Supplementary Table S7, http://www.iovs.org/lookup/suppl/doi:10.1167/iovs.11-7510/-/DCSupplemental). Of note, imputation was used by most replication cohorts for at least one of the SNPs. Imputation quality scores were greater than 0.90 for all SNPs with the exception of rs6856425 in FIND-Eye where the imputation quality score was 0.67. Errors in imputed genotype calls could lead to false-negative results, particularly for rare SNPs, which are more susceptible to genotyping artifacts. In addition, there was significant heterogeneity among samples that could not be explained by excluding type 1 diabetes participants and cohorts that did not use ETDRS grading consistently. Because the heterogeneity was not present when meta-analysis was restricted to the replication cohorts alone or the CARe cohorts alone, the "winner's curse" effect of large effect sizes in the discovery cohorts likely explains most of this heterogeneity.⁶⁸ Some heterogeneity might also derive from the different DR ascertainment methods and case- control definitions. Cohorts differed in their photography protocols, with some cohorts having one field of only one eye for phenotype determination. This introduces misclassification bias for participants for whom the DR grade in the one or two fields photographed may not accurately represent the DR grade in other fields or the contralateral eye. It is therefore possible that some of the variants associated with DR in CARe may eventually be replicated in larger studies with direct genotyping and better phenotype harmonization.

In summary, in this candidate gene analysis of DR with data from the CARe consortium, with replications in a several large, well-powered samples, we found little evidence of a major DR gene. This is the largest number of candidate genes studied for DR to date. Although no association could be confirmed with a high threshold for significance, the results are hypothesis generating and the genes associated with DR in CARe could be prioritized in studies. The importance of well-powered replication and phenotype harmonization are highlighted by our study. These issues will continue to be important as results from genome-wide association studies for DR become increasingly available.

Acknowledgments

The authors thank Joan W. Miller and David M. Altshuler for their support of the study.

References

- National Institute of Diabetes and Digestive and Kidney Diseases. National diabetes statistics fact sheet: general information and national estimates on diabetes in the United States. 2000. Bethesda, MD: U.S. Department of Health and Human Services, National Institute of Health, Publication No. 02-3892.
- Klein R, Klein BE, Moss SE, Cruickshanks KJ. The Wisconsin Epidemiologic Study of Diabetic Retinopathy, XVII: The 14-year incidence and progression of diabetic retinopathy and associated risk factors in type 1 diabetes. *Ophthalmology*. 1998;105:1801– 1815.
- Klein BE. Overview of epidemiologic studies of diabetic retinopathy. Ophthalmic Epidemiol. 2007;14:179-183.
- Cheung N, Mitchell P, Wong TY. Diabetic retinopathy. *Lancet.* 2010;376:124-136.
- Zhang X, Saaddine JB, Chou CF, et al. Prevalence of diabetic retinopathy in the United States, 2005–2008. JAMA. 2010;304: 649–656.
- Nathan DM. Long-term complications of diabetes mellitus. N Engl J Med. 1993;328:1676–1685.
- Klein R, Klein BE, Moss SE, Cruickshanks KJ. Relationship of hyperglycemia to the long-term incidence and progression of diabetic retinopathy. *Arch Intern Med.* 1994;154:2169–2178.

- Looker HC, Nelson RG, Chew E, et al. Genome-wide linkage analyses to identify loci for diabetic retinopathy. *Diabetes*. 2007; 56:1160-1166.
- 9. Hietala K, Forsblom C, Summanen P, Groop PH. Heritability of proliferative diabetic retinopathy. *Diabetes.* 2008;57:2176-2180.
- Arar NH, Freedman BI, Adler SG, et al. Heritability of the severity of diabetic retinopathy: the FIND-Eye study. *Invest Ophthalmol Vis Sci.* 2008;49:3839-3845.
- Abhary S, Hewitt AW, Burdon KP, Craig JE. A systematic metaanalysis of genetic association studies for diabetic retinopathy. *Diabetes.* 2009;58:2137-2147.
- Liew G, Klein R, Wong TY. The role of genetics in susceptibility to diabetic retinopathy. *Int Ophtbalmol Clin.* 2009;49:35–52.
- Lohmueller KE, Pearce CL, Pike M, Lander ES, Hirschhorn JN. Meta-analysis of genetic association studies supports a contribution of common variants to susceptibility to common disease. *Nat Genet.* 2003;33:177-182.
- Hirschhorn JN, Lohmueller K, Byrne E, Hirschhorn K. A comprehensive review of genetic association studies. *Genet Med.* 2002;4: 45-61.
- Kottgen A, Hwang SJ, Rampersaud E, et al. TCF7L2 variants associate with CKD progression and renal function in populationbased cohorts. J Am Soc Nepbrol. 2008;19:1989–1999.
- Krentz AJ, Clough G, Byrne CD. Interactions between microvascular and macrovascular disease in diabetes: pathophysiology and therapeutic implications. *Diabetes Obes Metab.* 2007;9:781–791.
- 17. Ohno T, Kinoshita O, Fujita H, et al. Detecting occult coronary artery disease followed by early coronary artery bypass surgery in patients with diabetic retinopathy: report from a diabetic retinocoronary clinic. *J Thorac Cardiovasc Surg.* 2010;139:92–97.
- Kofler B, Mueller EE, Eder W, et al. Mitochondrial DNA haplogroup T is associated with coronary artery disease and diabetic retinopathy: a case control study. *BMC Med Genet.* 2009;10:35.
- Holman RR, Paul SK, Bethel MA, Matthews DR, Neil HA. 10-year follow-up of intensive glucose control in type 2 diabetes. *N Engl J Med.* 2008;359:1577–1589.
- 20. Cheung N, Wong TY. Diabetic retinopathy and systemic vascular complications. *Prog Retin Eye Res.* 2008;27:161–176.
- Kawasaki R, Cheung N, Islam FM, et al. Is diabetic retinopathy related to subclinical cardiovascular disease? *Ophthalmology*. 2011;118:860-865.
- 22. Musunuru K, Lettre G, Young T, et al. Candidate gene association resource (CARe): design, methods, and proof of concept. *Circ Cardiovasc Genet.* 2010;3:267–275.
- Keating BJ, Tischfield S, Murray SS, et al. Concept, design and implementation of a cardiovascular gene-centric 50 k SNP array for large-scale genomic association studies. *PLoS One.* 2008;3:e3583.
- Uhlmann K, Kovacs P, Boettcher Y, Hammes HP, Paschke R. Genetics of diabetic retinopathy. *Exp Clin Endocrinol Diabetes*. 2006;114:275-294.
- 25. Tong Z, Yang Z, Patel S, et al. Promoter polymorphism of the erythropoietin gene in severe diabetic eye and kidney complications. *Proc Natl Acad Sci U S A.* 2008;105:6998-7003.
- 26. Ng DP, Tai BC, Lim XL. Is the presence of retinopathy of practical value in defining cases of diabetic nephropathy in genetic association studies?—the experience with the ACE insertion/deletion polymorphism in 53 studies comprising 17,791 subjects. *Diabetes*. 2008;57:2541-2546.
- Freedman BI, Bostrom M, Daeihagh P, Bowden DW. Genetic factors in diabetic nephropathy. *Clin J Am Soc Nepbrol.* 2007;2: 1306-1316.
- Frayling TM. Genome-wide association studies provide new insights into type 2 diabetes aetiology. *Nat Rev Genet.* 2007;8:657– 662.
- Sladek R, Rocheleau G, Rung J, et al. A genome-wide association study identifies novel risk loci for type 2 diabetes. *Nature*. 2007; 445:881–885.
- Saxena R, Voight BF, Lyssenko V, et al. Genome-wide association analysis identifies loci for type 2 diabetes and triglyceride levels. *Science*. 2007;316:1331–1336.
- Frayling TM, Timpson NJ, Weedon MN, et al. A common variant in the FTO gene is associated with body mass index and predisposes to childhood and adult obesity. *Science*. 2007;316:889-894.

- Zeggini E, Scott LJ, Saxena R, et al. Meta-analysis of genome-wide association data and large-scale replication identifies additional susceptibility loci for type 2 diabetes. *Nat Genet.* 2008;40:638– 645.
- Sandhu MS, Weedon MN, Fawcett KA, et al. Common variants in WFS1 confer risk of type 2 diabetes. *Nat Genet*. 2007;39:951–953.
- 34. Winckler W, Weedon MN, Graham RR, et al. Evaluation of common variants in the six known maturity-onset diabetes of the young (MODY) genes for association with type 2 diabetes. *Diabetes.* 2007;56:685–693.
- Bild DE, Bluemke DA, Burke GL, et al. Multi-ethnic study of atherosclerosis: objectives and design. *Am J Epidemiol.* 2002;156: 871-881.
- The Atherosclerosis Risk in Communities (ARIC) Study: design and objectives. The ARIC investigators. *Am J Epidemiol.* 1989;129: 687–702.
- 37. Fried LP, Borhani NO, Enright P, et al. The Cardiovascular Health Study: design and rationale. *Ann Epidemiol.* 1991;1:263–276.
- Wilson JG, Rotimi CN, Ekunwe L, et al. Study design for genetic analysis in the Jackson Heart Study. *Ethn Dis.* 2005;15:86-30-37.
- 39. Report of the expert committee on the diagnosis and classification of diabetes mellitus. *Diabetes Care* 2003;26(suppl 1):S5-S20.
- Wong TY, Klein R, Islam FM, et al. Diabetic retinopathy in a multi-ethnic cohort in the United States. *Am J Ophthalmol.* 2006; 141:446-455.
- 41. Cheung N, Wang JJ, Rogers SL, et al. Diabetic retinopathy and risk of heart failure. *J Am Coll Cardiol.* 2008;51:1573-1578.
- 42. Klein R, Marino EK, Kuller LH, et al. The relation of atherosclerotic cardiovascular disease to retinopathy in people with diabetes in the Cardiovascular Health Study. *Br J Ophthalmol.* 2002;86:84–90.
- 43. Diabetic retinopathy study. Report Number 6. Design, methods, and baseline results. Report Number 7. A modification of the Airlie House classification of diabetic retinopathy. Prepared by the Diabetic Retinopathy. *Invest Ophthalmol Vis Sci* 1981;21:1–226.
- 44. Ojaimi E, Nguyen TT, Klein R, et al. Retinopathy signs in people without diabetes the multi-ethnic study of atherosclerosis. *Oph-thalmology.* 2011;118:656-662.
- Colagiuri S, Lee CM, Wong TY, Balkau B, Shaw JE, Borch-Johnsen K. Glycemic thresholds for diabetes-specific retinopathy: implications for diagnostic criteria for diabetes. *Diabetes Care.* 2011;34: 145–150.
- 46. Wong TY, Liew G, Tapp RJ, et al. Relation between fasting glucose and retinopathy for diagnosis of diabetes: three population-based cross-sectional studies. *Lancet.* 2008;371:736–743.
- Guillausseau PJ, Massin P, Charles MA, et al. Glycaemic control and development of retinopathy in type 2 diabetes mellitus: a longitudinal study. *Diabet Med.* 1998;15:151-155.
- Stratton IM, Kohner EM, Aldington SJ, et al. UKPDS 50: risk factors for incidence and progression of retinopathy in type II diabetes over 6 years from diagnosis. *Diabetologia*. 2001;44:156–163.
- Tight blood pressure control and risk of macrovascular and microvascular complications in type 2 diabetes: UKPDS 38. UK Prospective Diabetes Study Group. *BMJ*. 1998;317:703–713.
- Ferris FL 3rd, Chew EY, Hoogwerf BJ. Serum lipids and diabetic retinopathy. Early Treatment Diabetic Retinopathy Study Research Group. *Diabetes Care*. 1996;19:1291–1293.
- 51. Cochran WG. Some methods for strengthening the common X2 tests. *Biometrics*. 1954;10:417-451.
- Mantel N, Haenszel W. Statistical aspects of the analysis of data from retrospective studies of disease. *J Natl Cancer Inst.* 1959;22: 719–748.
- Tobin MD, Sheehan NA, Scurrah KJ, Burton PR. Adjusting for treatment effects in studies of quantitative traits: antihypertensive therapy and systolic blood pressure. *Stat Med.* 2005;24:2911– 2935.
- Purcell S, Neale B, Todd-Brown K, et al. PLINK: a tool set for whole-genome association and population-based linkage analyses. *Am J Hum Genet.* 2007;81:559–575.
- Qiu C, Cotch MF, Sigurdsson S, et al. Retinal and cerebral microvascular signs and diabetes: the age, gene/environment susceptibility-Reykjavik study. *Diabetes*. 2008;57:1645–1650.

- Mitchell P, Smith W, Wang JJ, Attebo K. Prevalence of diabetic retinopathy in an older community. The Blue Mountains Eye Study. *Ophthalmology.* 1998;105:406-411.
- 57. Doney AS, Leese GP, Olson J, Morris AD, Palmer CN. The Y402H variant of complement factor H is associated with age-related macular degeneration but not with diabetic retinal disease in the Go-DARTS study. *Diabet Med.* 2009;26:460-465.
- Wong TY, Cheung N, Tay WT, et al. Prevalence and risk factors for diabetic retinopathy: the Singapore Malay Eye Study. *Ophthalmol*ogy. 2008;115:1869-1875.
- 59. Yim-Lui Cheung C, Wong TY, Lamoureux EL, et al. C-reactive protein and retinal microvascular caliber in a multiethnic Asian population. *Am J Epidemiol.* 2010;171:206–213.
- Buraczynska M, Baranowicz-Gaszczyk I, Tarach J, Ksiazek A. Tolllike receptor 4 gene polymorphism and early onset of diabetic retinopathy in patients with type 2 diabetes. *Hum Immunol.* 2009;70:121-124.
- 61. Cochran WG. Problems arising in the analysis of a series of similar experiments. *J R Stat Soc.* 1937;4:102–118.

- Cochran WG. The combination of estimates from different experiments. *Biometrics*. 1954;10:101–129.
- 63. Pe'er I, Yelensky R, Altshuler D, Daly MJ. Estimation of the multiple testing burden for genomewide association studies of nearly all common variants. *Genet Epidemiol.* 2008;32:381–385.
- 64. Liu Y, Burdon KP, Langefeld CD, et al. P-selectin gene haplotype associations with albuminuria in the Diabetes Heart Study. *Kidney Int.* 2005;68:741-746.
- Tedder TF, Steeber DA, Chen A, Engel P. The selectins: vascular adhesion molecules. *FASEB J.* 1995;9:866-873.
- 66. Freathy RM, Timpson NJ, Lawlor DA, et al. Common variation in the FTO gene alters diabetes-related metabolic traits to the extent expected given its effect on BMI. *Diabetes.* 2008;57:1419-1426.
- Abhary S, Burdon KP, Casson RJ, Goggin M, Petrovsky NP, Craig JE. Association between erythropoietin gene polymorphisms and diabetic retinopathy. *Arch Ophthalmol.* 2010;128:102–106.
- Kraft P. Curses—winner's and otherwise—in genetic epidemiology. *Epidemiology*. 19:649–651, 2008; discussion 57–58.