

Association Analysis of Dyslipidemia-Related Genes in Diabetic Nephropathy

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Abstract

Type 1 diabetes (T1D) increases risk of the development of microvascular complications and cardiovascular disease (CVD). Dyslipidemia is a common risk factor in the pathogenesis of both CVD and diabetic nephropathy (DN), with CVD identified as the primary cause of death in patients with DN. In light of this commonality, we assessed single nucleotide polymorphisms (SNPs) in thirty-seven key genetic loci previously associated with dyslipidemia in a T1D cohort using a casecontrol design. SNPs (n = 53) were genotyped using Sequenom in 1467 individuals with T1D (718 cases with proteinuric nephropathy and 749 controls without nephropathy i.e. normal albumin excretion). Cases and controls were white and recruited from the UK and Ireland. Association analyses were performed using PLINK to compare allele frequencies in cases and controls. In a sensitivity analysis, samples from control individuals with reduced renal function (estimated glomerular filtration rate < 60 ml/min/1.73 m²) were excluded. Correction for multiple testing was performed by permutation testing. A total of 1394 samples passed quality control filters. Following regression analysis adjusted by collection center, gender, duration of diabetes, and average HbA1c, two SNPs were significantly associated with DN. rs4420638 in the APOC1 region (odds ratio [OR] = 1.51; confidence intervals [CI]: 1.19-1.91; P=0.001) and rs1532624 in CETP (OR=0.82; CI: 0.69-0.99; P = 0.034); rs4420638 was also significantly associated in a sensitivity analysis (P = 0.016) together with rs7679 (P = 0.027). However, no association was significant following correction for multiple testing. Subgroup analysis of end-stage renal disease status failed to reveal any association. Our results suggest common variants associated with dyslipidemia are not strongly associated with DN in T1D among white individuals. Our findings, cannot entirely exclude these key genes which are central to the process of dyslipidemia, from involvement in DN pathogenesis as our study had limited power to detect variants of small effect size. Analysis in larger independent cohorts is required.

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Introduction

Type 1 diabetes mellitus (T1D) has been previously reported to increase the risk of microvascular complications and cardiovascular disease (CVD) [1–3]. In contrast to the reduction in cardiovascular mortality within the general US population, the declining trend is less evident in individuals with diabetes [4]. Despite improved disease management strategies, CVD remains the primary cause of death in patients with T1D [5] and a ten-fold increase in risk is reported in those with diabetic nephropathy (DN) relative to those without it [6]. DN is a complex, multifactorial disease and identifying robust genetic risk factors has proved challenging. Several risk factors are common to both CVD and DN, including hypertension, male gender, smoking and modifiable dyslipidemia [5–11].

Dyslipidemia results from abnormal lipid metabolism with departure from optimum vascular cholesterol and triglyceride levels leading to atherosclerosis, a process of fatty acid plaque deposition in arterial blood vessels. Previous studies reported normal low-density lipoprotein (LDL) and high-density lipoprotein (HDL) cholesterol levels in individuals with T1D, with elevated triglyceride levels more commonly associated with poor glycemic control [12]. This abnormal lipid profile can result from insulin deficiency with subsequent reduction in lipoprotein lipase activity and diminished ability for chylomicron and very low-density lipoprotein (VLDL) clearance [13]. This contrasts with individuals with type 2 diabetes (T2D) who often exhibit reduced HDL levels with a shift in LDL to the more atherogenic dense VLDL particles as a consequence of increased hepatic production. This process is increased by insulin resistance resulting in reduced clearance of VLDL and chylomicrons [14].

Observational studies have identified multiple lipid abnormalities in both incipient and overt DN [15–17], although this has not been consistently reported [18]. While the exact mechanism of effect is not fully understood, dyslipidemia has been associated with DN progression as well as increasing cardiovascular risk [19–

Table 1. Clinical characteristics of diabetic nephropathy (DN) cases and no nephropathy diabetic controls.

Characteristic	DN cases (n = 718)	Controls (n = 749)	P value
Male; n (%)	415 (57.8%)	320 (42.7%)	< 0.001
Age at diagnosis of T1D (yr)	14.8±7.7	15.5±7.9	0.09
Duration of T1D (yr) ^a	33.3±9.4	28.1±9.0	< 0.001
Age at sampling	48.1 ± 10.4	43.6±11.0	< 0.001
HbA1c (%) ^b	9.0±1.9	8.6±1.5	< 0.001
Systolic blood pressure (mmHg) ^b	144.9±20.9	125.0±14.7	< 0.001
Diastolic blood pressure (mmHg) ^b	81.5±11.4	75.4±7.8	< 0.001
Body mass index (kg/m²)	26.3±4.7	26.1±4.2	0.50
Serum cholesterol (mmol/L)	5.34±1.22	5.09±0.91	< 0.001
HDL cholesterol (mmol/L)	1.59 ± 0.55	1.78±0.47	< 0.001
LDL cholesterol (mmol/L)	2.88±0.95	2.80±0.75	0.17
Serum triglycerides (mmol/L) median (interquartile range)	1.4 (1.0-2.2)	1.0 (0.7–1.4)	< 0.001
Serum creatinine (μmol/L); ^c median (interquartile range)	130 (102–183)	92 (79–105)	< 0.001
Estimated glomerular filtration rate (ml/min/1.73m²); ^c median (interquartile range)	48 (33–66)	70 (59–85)	< 0.001
End-stage renal disease; n (%)	193 (26.9%)	NA	NA
Passed quality control criteria; n (%)	684 (95.3%)	710 (94.8%)	0.68

Unless otherwise stated values are mean \pm standard deviation.

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20]. Supporting evidence implicates insulin resistance as pivotal in the development and/or progression of this condition [21–25]. Potential mechanisms contributing to renal injury in DN have included stimulation of pro-inflammatory and pro-fibrotic cytokine production, cell apoptosis, vasoconstriction and modulation of mesangial cell proliferation [26–28]. As such, parallels between mechanisms that underpin atherosclerosis and glomerulosclerosis provide support for investigation of the parameters that contribute to both conditions [29].

While previous evidence demonstrates modulation of lipid profiles through lifestyle changes such as smoking, diet and physical activity, recent studies have also identified common genetic variation as regulators of lipid levels and subsequent dyslipidemia [30-37]. To date, almost 100 genetic loci have been reported in association with serum cholesterol and triglyceride levels [38]. Aulchenko and colleagues highlighted that many of the loci influencing lipid levels and CVD risk had previously been identified in association studies enriched by participants with diabetes [34]. The management of diabetic dyslipidemia, a wellrecognized and modifiable risk factor, is a key element in the multifactorial approach to prevent CVD in individuals with diabetes [23]. In light of the evidence supporting association of these variants with dyslipidemia in individuals with diabetes, we sought to assess the allelic frequency of 53 common single nucleotide polymorphisms (SNPs) in 37 key loci in individuals with DN using a case-control design involving 1467 individuals with T1D. These loci and SNPs were selected on the basis of their functional significance and previous reported association with dyslipidemia [30–37].

Methods

Participants

Research ethics approval was obtained from the South and West Multicentre Research Ethics Committee (MREC/98/6/71) and Oueens University Belfast Research Ethics Committee. Written informed consent was obtained prior to participation. All recruited individuals were white, had T1D diagnosed before 32 years of age and were born in the UK or Ireland. Patients (n = 718) and controls (n = 749) originated from the Warren 3/UKGenetics of Kidneys in Diabetes (GoKinD) and all-Ireland collections [39]. The definition of DN in cases was based on development of persistent proteinuria (>0.5 g protein/24 h) at least 10 years after diagnosis of T1D, hypertension (blood pressure >135/85 mmHg or treatment with antihypertensive agents) and associated diabetic retinopathy. Controls were individuals with T1D for at least 15 years with normal urinary albumin excretion rates and no evidence of microalbuminuria on repeated testing. In addition, control subjects had not been prescribed antihypertensive drug treatment avoiding possible misclassification of diabetic individuals as 'control phenotypes' when the use of antihypertensive treatment may have reduced urinary albumin excretion into the normal range. Individuals with microalbuminuria were excluded from both case and control groups since it was not possible to be confident in assigning case/control status for such individuals whose urinary albumin excretion might either regress or progress over time [40].

SNP selection and genotyping

SNPs (n = 53) were selected on the basis of previously reported association with dyslipidemia [31–34] and of minor allele frequency (MAF) exceeding 0.1 in populations of European descent. Genotyping was performed by MassARRAY iPLEX (Sequenom, San Diego, CA, USA) assays according to the manufacturer's instructions. Quality filters for exclusion of SNPs included call rates below 95% and deviation from HWE (P<0.001). DNA samples were excluded if missing genotypes exceeded 10%. Other quality control measures included parent/offspring trio samples, duplicates on plates, random sample allocation to plates, independent scoring of

^aCalculated from the dates of diagnosis and recruitment.

^bAverage of the three most recent values prior to recruitment.

^cExcludes subjects receiving renal replacement therapy (dialysis or transplant).

Table 2. Minor allele frequencies (MAF) and genotype counts in 684 diabetic nephropathy cases and 710 no nephropathy diabetic controls.

		Genomic	mic				Case		Controls			Confidence	ence	
SNP ID	Ref C/s	C/some Position	ion Variant		Gene	^a Alleles	Counts	MAF	Counts	MAF	bP val	OR Interva	II ^d P val	eP val
rs10903129 3	34 1	25768937	937 intronic		TMEM57	[A/G]	128/319/236	0.42	124/364/220	0.43	0.548	0.92 0.77-1.10	0 0.368	0.261
rs11206510 3	33 1	55496039		intergenic	PCSK9	[7]	32/172/461	0.18	25/217/464	0.19	0.431	0.90 0.72–1.12	2 0.341	0.377
rs1167998 3	34 1	62931632	632 intronic		DOCK7	[C/A]	84/298/298	0.34	83/308/314	0.34	0.719 0	0.98 0.82–1.18	8 0.867	0.957
rs10889353 3	34 1	63118196	196 intronic		DOCK7	[C/A]	81/297/305	0.34	80/307/320	0.33	0.748	0.98 0.82–1.18	8 0.858	0.957
¹ rs12740374 3	32 1	109817590	3'UTR		CELSR2	[1/6]	33/230/421	0.22	36/219/452	0.21	0.494	1.10 0.89–1.36	988 0.388	0.520
rs646776 3	32 1	109818530		intergenic	CELSR2	[G/A]	33/225/424	0.21	37/219/452	0.21	0.678	1.07 0.86–1.32	0.540	0.707
rs2144300 3	31 1	230294916	4916 intronic		GALNT2	[C/J]	105/337/235	0.40	108/330/269	0.39	0.337 1	1.07 0.89–1.28	0.490	0.573
rs4846914 3	32 1	230295691	5691 intronic		GALN72	[G/A]	109/339/236	0.41	110/332/268	0.39	0.320	1.05 0.88-1.26	96 0.578	0.620
rs6754295 3	34 2	21206183		intergenic ,	APOB	[G/T]	27/238/416	0.21	39/239/432	0.22	0.573 1	1.02 0.83–1.27	27 0.833	0.318
rs7557067 3	33 2	21208211		intergenic	APOB	[G/A]	28/241/415	0.22	40/238/432	0.22	0.663	1.03 0.83-1.27	00800 23	0.286
rs673548 3	34 2	21237544	544 intronic		APOB	[A/G]	20/228/430	0.20	30/215/454	0.20	0.951	1.07 0.85–1.34	14 0.564	0.263
rs1260326 3	33 2	27730940	940 missense		GCKR	[1/]	93/329/258	0.38	114/305/290	0.38	0.879	0.99 0.83–1.19	9 0.943	0.642
rs780094 3	34 2	27741237	237 intronic		GCKR	[A/G]	86/311/273	0.36	104/296/294	0.36	0.885	0.98 0.82–1.18	8 0.842	0.798
rs6756629 3	34 2	44065090	090 missense		ABCG5	[A/G]	3/90/591	0.07	4/92/610	0.07	0.947	0.90 0.64-1.27	0950 2	0.449
rs6544713 3	33 2	44073881	881 intronic		ABCG8	[1/]	68/308/307	0.33	86/305/318	0.34	0.525 1	1.03 0.85–1.24	94 0.759	0.811
rs3846662 3	34 5	74651084	084 intronic		HMGCR	[C/J]	104/324/246	0.39	113/323/254	0.40	0.866	0.98 0.82–1.17	7 0.815	0.258
rs3846663 3	33 5	74655726	726 intronic		HMGCR	[1/C]	83/302/297	0.34	92/308/309	0.35	0.831	0.96 0.80–1.16	6 0.694	0.258
rs1501908 3	33 5	156398169		intergenic	TIMD4/HAVCR1	[C/G]	102/313/265	0.38	99/329/279	0.37	0.686	1.10 0.92–1.32	12 0.288	0.200
rs12670798 3	34 7	21607352	352 intronic		DNAH11	[C/J]	37/255/392	0.24	43/254/413	0.24	0.948 (0.99 0.81–1.22	22 0.945	0.759
rs2240466 3	34 7	72856269	.269 intronic		BAZ1B	[1/]	5/175/503	0.14	14/151/545	0.13	0.463	1.13 0.87–1.47	17 0.374	0.109
² rs714052 3	32 7	72864869	869 intronic		BAZ1B	[C/J]	5/173/503	0.13	14/151/543	0.13	0.534 1	1.12 0.86–1.46	16 0.396	0.115
rs7819412 3	33 8	11045161	161 intronic		XKR6	[G/A]	168/322/179	0.49	152/361/196	0.47	0.231	1.05 0.88-1.26	6 0.583	0.420
rs10096633	34 8	19830921		intergenic	THT	[1/C]	11/124/548	0.11	8/145/557	0.11	0.584 (0.82 0.62–1.08	0.154	0.130
rs12678919 3	33 8	19844222		intergenic	TLF	[G/A]	9/5//26	0.08	5/111/593	60.0	0.923	0.84 0.62–1.14	4 0.265	0.219
rs2083637 3	34 8	19865175		intergenic	TbT	[C/J]	50/245/388	0.25	51/271/388	0.26	0.542 (0.90 0.74–1.10	0 0.321	0.191
rs17321515 3	32 8	126486409		intergenic	TRIB1	[G/A]	153/343/188	0.47	153/365/191	0.47	0.949	1.03 0.86–1.23	3 0.717	0.564
³rs2954029 3	32 8	126490972		intergenic	TRIB1	[T/A]	149/333/202	0.46	147/366/197	0.46	0.852	1.02 0.86–1.22	0.797	0.617
rs471364 3	33 9	15289578	578 intronic		TTC398	[G/A]	6/147/529	0.12	11/144/554	0.12	0.967	1.01 0.77–1.33	13 0.951	0.807
rs3905000 3	34 9	107657070	7070 intronic		ABCA1	[A/G]	11/180/492	0.15	17/179/513	0.15	0.863	1.02 0.80–1.31	11 0.882	0.810
rs1883025 3	33 9	107664301	4301 intronic		ABCA1	[A/G]	51/276/356	0.28	55/316/338	0:30	0.168 (0.94 0.78–1.15	5 0.573	0.620
rs7395662 3	34 11	48518893		intergenic	OR4A47	[A/G]	95/326/263	0.38	92/339/279	0.37	0.628	1.08 0.90–1.30	0.415	0.524
rs174547 3	33 11	61570783	783 intronic		FADS1	[C/J]	78/319/287	0.35	70/327/306	0.33	0.402	1.13 0.93–1.37	37 0.212	0.524
	34 11	61597212	212 intronic		FADS2	[1/0]	8/161/508	0.13	10/168/525	0.13	0.817	1.04 0.79–1.35	662.0 98	0.527
rs964184 3	32 11	116648917		intergenic	ZNF259	[d/C]	10/138/535	0.12	10/162/538	0.13	0.314 (0.82 0.63-1.07	0.142	0.172

Table 2. Cont.

SAP ID Ref C/some Position Variant Gene rs2338104 31 12 109895168 intronic KCTD10 rs2338104 31 12 121388962 intergenic HNF1A rs2650000 33 15 58674695 intergenic LPC rs10468017 33 15 58678512 intergenic LPC rs153624 34 16 67902079 intronic LPC rs2271293 34 18 47167214 intronic LPC rs2967605 33 19 1420306 intronic LPCR rs2228671 34 19 11210912 missense LDLR rs2228671 34 19 19407718 intronic 7LP2PB rs232441 35 19 19789528 intronic 7LP2PB rs2325801 34 19 45395619 intronic 7LMM4 rs439401 34 19 45414451		Case	Con	Controls			Confidence		
31 12 109895168 intronic 33 12 121388962 intergenic 31 15 58674695 intergenic 34 16 57005479 intronic 34 16 67902070 intronic 34 18 47167214 intergenic 33 19 8469738 intergenic 34 19 11203306 intronic 32 19 19662220 intronic 34 19 19662220 intronic 34 19 45395266 intronic 34 19 45395266 intronic 34 19 45414451 intergenic 34 19 454295619 intronic 34 19 454242946 intergenic 34 19 45422946 intergenic	Gene	^a Alleles Counts	MAF Counts	nts MAF	F ^b P val	'OR	Interval	^d P val	eP val
33 12 121388962 intergenic 31 15 58674695 intergenic 33 15 58678512 intergenic 34 16 57005479 intronic 34 16 67902070 intronic 33 19 47167214 intergenic 34 19 11202306 intronic 32 19 19407718 intronic 32 19 19662220 intronic 34 19 45395266 intronic 34 19 45395266 intronic 34 19 45414451 intergenic 34 19 45414451 intergenic 34 19 45429546 intergenic 34 19 45414451 intergenic 33 19 45422946 intergenic	KCTD10	[C/G] 149/340/193	0.47 189/	189/329/189 0.50	0.089	0.91	0.76-1.08	0.268	0.172
31 15 58674695 intergenic 33 15 58678512 intergenic 34 16 67902070 intronic 34 16 67902070 intronic 33 19 8469738 intergenic 34 19 11202306 intronic 32 19 19407718 missense 34 19 19662220 intronic 34 19 45395266 intronic 34 19 45395266 intronic 34 19 45414451 intergenic 34 19 45422946 intergenic 34 19 45414451 intergenic 34 19 45414451 intergenic 33 19 45422946 intergenic	HNF1A	[T/G] 76/292/312	0.33 82/3	82/315/307 0.34	0.444	0.88	0.73-1.06	0.180	0.257
33 15 58678512 intergenic 34 16 57005479 intronic 34 16 67902070 intronic 33 19 8469738 intergenic 34 19 11202306 intronic 32 19 11210912 missense 32 19 19407718 intronic 34 19 19662220 intronic 34 19 45395266 intronic 34 19 45395266 intronic 34 19 45414451 intergenic 34 19 45422946 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	TIPC	[C/G] 65/288/328	0.31 68/2	68/284/356 0.30	0.555	1.05	0.87-1.27	0.596	0.348
34 16 57005479 intronic 34 16 67902070 intronic 34 18 47167214 intergenic 33 19 8469738 intergenic 34 19 11202306 intronic 32 19 19407718 intronic 34 19 19662220 intronic 34 19 45395266 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	TIPC	[T/C] 61/290/330	0.30 65/2	65/279/366 0.29	0.403	1.07	0.88-1.29	0.502	0.275
34 16 67902070 intronic 34 18 47167214 intergenic 33 19 11202306 intronic 34 19 11201012 missense 32 19 19407718 intronic 34 19 19662220 intergenic 34 19 45395266 intronic 34 19 45395619 intergenic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	CETP	[T/G] 109/337/233	0.41 149/	149/341/213 0.45	0.015	0.82	0.69-0.99	0.034	0.514
34 18 47167214 intergenic 33 19 8469738 intergenic 34 19 11202306 intronic 32 19 19407718 intronic 34 19 19789528 intronic 34 19 45395266 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 39228784 intergenic	NUTF2	[A/G] 11/131/521	0.12 8/12	8/129/543 0.11	0.470	0.93	0.70-1.25	0.638	0.272
33 19 8469738 intergenic 33 19 11202306 intronic 34 19 19407718 intronic 32 19 19662220 intronic 34 19 19789528 intronic 34 19 4539566 intronic 34 19 45395619 intergenic 34 19 45414451 intergenic 34 19 45422946 intergenic 33 19 39228784 intergenic	TIPG	[T/C] 28/213/437	0.20 26/2	26/212/467 0.19	0.458	1.06	0.85-1.33	0.612	0.585
33 19 11202306 intronic 34 19 11210912 missense 32 19 19407718 intronic 34 19 19789528 intronic 34 19 45395266 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	RAB11B	[A/G] 25/207/446	0.19 24/2	24/214/468 0.19	0.789	1.08	0.86-1.35	0.526	0.717
34 19 11210912 missense 32 19 19407718 intronic 34 19 19789528 intronic 34 19 4539566 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	LDLR	[T/G] 11/145/526	0.12 6/14	6/144/558 0.11	0.313	1.08	0.82-1.43	0.575	0.083
32 19 19407718 intronic 32 19 19662220 intergenic 34 19 4539526 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	TDLR	[T/C] 12/148/521	0.13 8/16	8/160/542 0.12	0.852	1.00	0.77-1.30	0.989	0.289
32 19 19662220 intergenic 34 19 19789528 intronic 34 19 4539566 intronic 34 19 45395619 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	SUGP1	[C/T] 5/91/587	0.07 4/10	4/101/605 0.08	0.778	0.93	0.67-1.29	0.667	0.415
34 19 19789528 intronic 34 19 4539566 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	CILP2PBX4	[T/C] 5/97/582	0.08 6/10	6/107/597 0.08	0.589	0.94	0.68-1.29	0.685	0.382
34 19 45395266 intronic 34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	ZNF101	[G/A] 7/96/571	0.08 5/10	5/106/594 0.08	0.949	0.98	0.72-1.34	0.913	0.487
34 19 45395619 intronic 34 19 45414451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	TOMM40	[G/A] 122/319/242	0.41 121/	121/345/241 0.42	0.873	1.01	0.84-1.21	0.916	0.626
34 19 4541451 intergenic 33 19 45422946 intergenic 33 20 39228784 intergenic	TOMM40	[G/A] 11/163/508	0.14 10/1	10/184/514 0.14	0.522	1.01	0.78-1.30	0.956	0.877
33 20 39228784 intergenic 33 20 39228784 intergenic	APOE/APOC1	[T/C] 96/313/275	0.37 96/3	96/348/266 0.38	0.544	0.99	0.82-1.18	0.882	0.772
33 20 39228784 intergenic	APOC1	[G/A] 24/224/421	0.20	11/199/475 0.16	0.005	1.51	1.19–1.91	0.001	0.0160
	MAFB	[T/C] 68/312/304	0.33 65/3	65/327/318 0.32	0.750	96.0	0.80-1.17	0.713	0.499
rs7679 33 20 44576502 3'UTR <i>PCIF1</i>	PCIF1	[C/T] 29/178/473	0.17 24/2	24/222/462 0.19	0.242	0.91	0.72-1.14	0.389	0.027

^aMinor alleles are presented first followed by major allele.

^bUnadjusted P values were calculated as tests for trend (1 df) across genotypes.

Odds ratios and 95% confidence intervals are calculated on a per allele basis for the minor allele assuming an additive model.

Adjusted P values were calculated as tests for trend (1 df) across genotypes in a logistic regression which included terms for collection center, gender, duration of T1DM and HbA1c category. Associations were no longer significant

after correction for multiple testing performed by permutation test (n = 100,000).

*In a sensitivity analysis (Control samples only with eGFR>60 ml/min/1.73 m², n = 444) adjusted P values were calculated as tests for trend (1 df) across genotypes in a logistic regression which included terms for collection center, gender, duration of T1DM and HbA1c category. Associations were no longer significant after correction for multiple testing performed by permutation test (n = 100,000).

*Proxy for rs599839 ($r^2 = 0.90$).

Table 3. Assessment of gene-gene pair-wise interactions.

SNP 1	Gene 1	SNP 2	Gene 2	¹P valu	ie ² P value
rs3905000	ABCA1	rs7679	PCIF1	0.002	0.014
rs6756629	ABCG5	rs714052	BAZ1B	0.003	0.009
rs2240466	BAZ1B	rs6756629	ABCG5	0.007	0.01
rs2240466	BAZ1B	rs12678919	LPL	0.007	0.115
rs1167998	DOCK7	rs17216525	CILP2PBX4	0.008	0.002
rs10903129	TMEM57	rs6544713	ABCG8	0.009	0.024
rs12678919	LPL	rs714052	BAZ1B	0.009	0.14

The number of significant interactions observed is less than one might expect by chance.

P values for gene-gene interactions were obtained between SNPs using likelihood ratio χ^2 tests in the logistic regression. Data are presented for those which attained significance at the P<0.01 level in an unadjusted model¹. Significance levels are also presented where terms for potential confounders (collection center, gender, duration of T1D and HbA1c) are included in the adjusted model².

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problematic genotypes by two individuals and re-sequencing of selected DNAs to validate genotypes.

Statistical analysis

Clinical characteristics of cases and controls were compared using the z-test for large independent samples and the χ^2 test. Association analyses were performed using PLINK (version 1.07; http://pngu. mgh.harvard.edu/~purcell/plink/). Initially a χ^2 test for trend (1 df) was used with stratification by collection center. Logistic regression analysis was performed on each SNP with terms for potential confounders (collection center, gender, duration of T1D and HbA1c) included in the model. A sensitivity analysis to minimize potential misclassification of case/control status was performed by excluding samples from those control individuals with an estimated glomerular filtration (eGFR) < 60 ml/min/1.73 m². The level of statistical significance was set at 5% and adjustment for multiple testing performed by permutation test (n = 100,000). Potential genegene interactions between SNPs were assessed using likelihood ratio χ^2 tests in the logistic regression with terms for potential confounders (collection center, gender, duration of T1D and HbA1c) included in the model.

Table 4. Study power to detect various odds ratios for selected minor allele frequencies.

Odds	Minor All	ele Frequency (MAF)	
ratio	0.10	0.20	0.30	0.40
1.2	30%	49%	59%	65%
1.3	56%	81%	89%	92%
1.4	79%	96%	99%	99%
1.5	93%	99%	100%	100%

Power calculations are based on 684 cases and 710 controls with odds ratio ranging from 1.2–1.5 for SNPs with a MAF between 0.10 and 0.40 with no correction for multiple testing.

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Results

The clinical characteristics of the DN cases (n=718) and diabetic controls (n=749) genotyped in this study are listed in Table 1. There were more males, higher mean HbA1c and blood pressure values (despite the use of antihypertensive treatment) in the case group compared with the control group. All comparisons were significant at P < 0.001 with the exception of age at diagnosis, LDL cholesterol and body mass index which did not differ significantly between groups. Approximately one quarter of cases (26.9%) had end-stage renal disease (ESRD).

A total of 53 SNPs were genotyped using MassARRAY iPLEX technology in 718 cases and 749 controls (Table 2). We excluded 73 samples (34 cases and 39 controls) from the analysis with ≥10% missing genotypes. The average call rate for all SNPs analysed was 99.3%. The genotype distribution for each SNP did not deviate significantly from HWE in either cases or controls. No duplicate or Mendelian inconsistencies were observed.

Single marker testing stratified by collection center identified two non-coding SNPs (rs1532624 in Cholesteryl ester transfer protein (*CETP*) and rs4420638 in Apolipoprotein C-I *APOCI*) to be significantly associated with DN (Table 2). In logistic regression analysis with adjustment by collection center, gender, duration of T1D, and average HbA1c as covariates, the significance of both SNPs was maintained (rs1532624: odds ratio [OR] = 0.82; confidence intervals [CI]: 0.69–0.99; P = 0.034; rs4420638: OR = 1.51; CI: 1.19–1.91; P = 0.001). The sensitivity analysis (that includes samples only from those controls with eGFR >60 ml/min/1.73 m²) identified two SNPs significantly associated with DN in the fully adjusted model (rs4420638; P = 0.016 and rs7679; P = 0.027). However, no associations were maintained following correction for multiple testing. Subgroup analyses showed no association of any SNP with ESRD status.

With no prior hypotheses or supporting evidence of potential gene-gene interaction, we assumed a more stringent level of significance (P<0.01). Interactions were assessed using likelihood ratio χ^2 tests in the logistic regression with terms for potential confounders (collection center, gender, duration of T1D and HbA1c) included in the model. Seven interaction terms exceeded the minimum threshold set but following correction for multiple testing and examination of the resultant Q-Q plot, none were identified as being worthy of further investigation (Table 3).

Discussion

Dyslipidemia can result through dietary and lifestyle influences or alternatively as a consequence of variation in genes pivotal to lipoprotein metabolism. In persons with diabetes, prolonged elevation of insulin levels often leads to dyslipidemia, a process central to the pathogenesis of atherosclerosis and increasing CVD risk. As previous studies have reported multiple lipid abnormalities in patients with T1D [15-20], we evaluated common polymorphic variation previously associated with dyslipidemia, in persons with T1D, both with and without DN. Univariate analysis identified two SNPs associated with DN (rs1532624 in CETP and rs4420638 in APOCI) both of which remained significant following adjustment for collection center, gender, duration of T1D, and average HbA1c. Interestingly, rs4420638 was also significantly associated with DN in the sensitivity analysis using only those samples from diabetic controls with eGFR >60 ml/min/1.73 m². However, following correction for multiple testing, these associations were no longer significant. Although, published data were available from the US GoKinD genome-wide association study, limited coverage on the Affymetrix 500 K genotyping platform across the genomic locations of both CETP and APOC1, prevented in silico independent replication or meta-analysis of our top SNPs or any potential proxies ($r^2 > 0.8$) [41].

In previously published studies the definition of the DN phenotype has proved challenging. We do not think it is surprising that cases in our study had persistent proteinuria (macroalbuminuria) despite the use of antihypertensive medication. The use of angiotensin converting enzyme inhibitors (ACEi) or angiotensin receptor blockers (ARBs), typically reduces but does not abolish protein excretion in persons with overt diabetic nephropathy [42–44] suggesting that persistent proteinuria is unlikely to be a consequence of suboptimal blood pressure control. The differences in mean blood pressures observed between case and control groups were consistent with findings in clinical practice.

In addition, it has been suggested that some individuals with a very prolonged duration of type 1 diabetes may develop chronic kidney disease (CKD) without albuminuria. Molitch and colleagues [45] identified 89 of the 1,439 individuals recruited to the DCCT/EDIC study that had developed CKD (defined by estimated GFR <60 ml/min/1.73 m²) after almost 20 years of follow up. Of the 89 individuals with CKD, 21 were classified as normoalbuminuric (albumin excretion rate [AER] <30); 14 as microalbuminuric (AER: 30-300); and 54 as macroalbuminuric (AER >300). Of note 43% of the normoalbuminuric individuals with CKD were taking ACEi during the study and 14% were taking ARBs at year 13/14 of the EDIC study [45]. The antihypertensive drugs, ACEi and ARBs, can both lower AER and reduce eGFR which may partly explain why the authors found a small number of individuals with normoalbuminuria and reduced eGFR. The normoalbuminuric patients with reduced eGFR were also 4 years older at time of recruitment than the macroalbuminuric patients (30+/-7 yr vs. 26+/-7 yr [45]). Nevertheless, we did attempt to address this issue of diabetic patients having CKD without albuminuria. In a sensitivity analysis, we excluded all those diabetic patients we had originally recruited as normalbuminuric controls in whom the eGFR was <60 ml/min/1.73 m². We excluded these controls with reduced renal function from our analysis to limit any risk of misclassification of nephropathy status but found this made little difference to the main analysis (Table 2).

CETP is a protein central to the process of dyslipidemia. It acts as a mediator for the transfer of cholesteryl esters from HDL to VLDL or LDL in exchange for triglycerides, reducing serum HDL concentrations [46]. Variation in CETP levels have been correlated with lipid metabolism and insulin resistance in Type 2 diabetes [47], and also in association with the development of obesity [48] and susceptibility to atherosclerosis and other CVD [49]. Recently, Igl and colleagues demonstrated that the genetic influence mediated by rs1532624 could be attenuated by lifestyle factors such as diet or physical activity, highlighting the potential for interaction at this locus [50]. Our study was unable to examine lifestyle influences, as dietary and physical activity measurements were not collected during recruitment. Nonetheless we sought to evaluate the potential for pair-wise gene-gene interaction between the candidate SNPs examined. Several pair-wise interactions which included the CETP and APOB loci were identified but did not remain significant following correction for multiple testing. As no association survived correction for multiple testing, it is unlikely these gene variants play a specific role in the etiology of DN.

Apo C-I is a protein constituent of chylomicrons, VLDL and HDL and while its precise physiological role is not well established,

References

 Libby P, Nathan DM, Abraham K, Brunzell JD, Fradkin JE, et al. (2005) Report of the National Heart, Lung, and Blood Institute: National Institute of Diabetes evidence has demonstrated support for its involvement in HDL metabolism through activation and inhibition of other proteins central to lipid metabolism, including CETP [51]. Association of rs4420638 with DN in T1D in this cohort has been previously reported [52].

Improved therapeutic regimens to lower LDL levels using statins have proved beneficial for patients both with and without diabetes with respect to CVD risk. In addition, increasing evidence suggests statins provide therapeutic benefit independent of cholesterol modulation, by improving endothelial and vascular function and reducing inflammation [53].

Common genetic loci explain only a proportion of the variation observed in lipid levels within the general population. Evidence in support of rare variants with potentially large individual effect size continues to grow, and is likely to make a significant impact on the genetic heritability of this condition [36]. Since our study focused only on common variants, untyped, highly penetrant rare variants in these genes could also contribute to DN. This study has insufficient power to detect rare variants particularly if the effect sizes are small in magnitude, such as the odds ratios of 1.2/1.3 which are more commonly found in common complex diseases (Table 4). Future amalgamation of independent cohorts with similar DN phenotypes will enable a more robust evaluation of such loci. In addition, other factors such as copy number variation or indeed epigenetic mechanisms (e.g. DNA methylation, histone modification and microRNAs) may also attenuate gene function affecting these pathways which modulate disease risk.

Although the SNPs assessed in this study were chosen on the basis of previous associations with dyslipdemia there are a number of inherent limitations associated with using 53 SNPs across 37 genes [54]: (1) identification of association does not necessarily equate to functional significance given the concept of linkage disequilibrium (LD). (2) assessing one or two SNPs per gene may provide inadequate representation of the genetic architecture at that locus. (3) patterns of LD can vary significantly within and between different populations and therefore a significant association in one population may not necessarily translate across all populations.

In conclusion, we found no strong association between common variants in genes involved in dyslipidemia and DN. Further work to investigate lifestyle factors which influence genes may identify potential risk factors for susceptibility to DN.

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Author Contributions

Conceived and designed the experiments: APM DS AJM. Performed the experiments: GL AJM. Analyzed the data: GM CP. Contributed reagents/materials/analysis tools: GL GM CP AJM. Wrote the paper: GM AJM CP APM.

and Digestive and Kidney Diseases Working Group on Cardiovascular Complications of Type 1 Diabetes Mellitus. Circulation 111: 3489–3493.

- Durrington PN (1999) Diabetic dyslipidaemia. Baillieres Best Pract Res Clin Endocrinol Metab 13: 265–278.
- Marcovecchio ML, Dalton RN, Prevost AT, Acerini CL, Barrett TG, et al. (2009) Prevalence of abnormal lipid profiles and the relationship with the development of microalbuminuria in adolescents with type 1 diabetes. Diabetes Care 32:658–663.
- 4. Gu K, Cowie CC, Harris MI (1999) Diabetes and decline in heart disease mortality in US adults. JAMA 281:1291–1297.
- Laing SP, Swerdlow AJ, Slater SD, Botha JL, Burden AC, et al. (1999) The British Diabetic Association Cohort Study, II: cause-specific mortality in patients with insulin-treated diabetes mellitus. Diabet Med 16:466–471.
- Tuomilehto J, Borch-Johnsen K, Molarius A, Forsén T, Rastenyte D, et al. (1998) Incidence of cardiovascular disease in type 1 (insulin-dependent) diabetic subjects with and without diabetic nephropathy in Finland. Diabetologia 41:784-790.
- The Diabetes Control and Complications (DCCT) Research Group (1995) Effect of intensive therapy on the development and progression of diabetic nephropathy in the Diabetes Control and Complications Trial. Kidney Int 47:1703–1720.
- Parving HH, Hommel E (1989) Prognosis in diabetic nephropathy. BMJ 299:230–233.
- Andersen AR, Christiansen JS, Andersen JK, Kreiner S, Deckert T (1983)
 Diabetic nephropathy in type 1 (insulin-dependent) diabetes: an epidemiological study. Diabetologia 25:496–501.
- Sawicki PT, Didjurgeit U, Muhlhauser I, Bender R, Heinemann L, et al. (1994)
 Smoking is associated with progression of diabetic nephropathy. Diabetes Care 17:126–131.
- Mulec H, Johnsen SA, Wiklund O, Bjorck S (1993) Cholesterol: a renal risk factor in diabetic nephropathy? Am J Kidney Dis 22:196–201.
- Molitch ME (2006) Management of dyslipidemias in patients with diabetes and chronic kidney disease. Clin J Am Soc Nephrol;1:1090–1099.
- O'Brien T, Nguyen TT, Zimmerman BR (1998) Hyperlipidemia and diabetes mellitus. Mayo Clin Proc;73:969–976.
- Ginsberg HN (2006) Efficacy and mechanisms of action of statins in the treatment of diabetic dyslipidemia. J Clin Endocrinol Metab;91:383–392.
- Groop PH, Elliott T, Ekstrand A, Franssila-Kallunki A, Friedman R, et al. (1996) Multiple lipoprotein abnormalities in type I diabetic patients with renal disease. Diabetes 45:974–979.
- Jones SL, Close CF, Mattock MB, Jarrett RJ, Keen H, et al., (1989) Plasma lipid and coagulation factor concentrations in insulin dependent diabetics with microalbuminuria. BMJ 298:487–490.
- Watts GF, Naumova R, Slavin BM, Morris RW, Houlston R, et al (1989) Serum lipids and lipoproteins in insulin-dependent diabetic patients with persistent microalbuminuria. Diabet Med 6:25–30.
- Mattock MB, Cronin N, Cavallo-Perin P, Idzior-Walus B, Penno G, et al., (2001) EURODIAB IDDM Complications Study. Plasma lipids and urinary albumin excretion rate in type 1 diabetes mellitus: The EURODIAB IDDM complications study. Diabet Med 18:59–67.
- Thomas MC, Rosengård-Bärlund M, Mills V, Rönnback M, Thomas S, et al (2006) Serum lipids and the progression of nephropathy in type 1 diabetes. Diabetes Care 29:317–322.
- Tolonen N, Forsblom C, Thorn L, Wadén J, Rosengård-Bärlund M, et al., (2008) Relationship between lipid profiles and kidney function in patients with type 1 diabetes. Diabetologia;51:12–20.
- Taskinen MR (2003) Diabetic dyslipidaemia: from basic research to clinical practice. Diabetologia 46: 733–749.
- Krauss RM, Siri PW (2004) Dyslipidemia in type 2 diabetes. Med Clin North Am 88: 897–909.
- Solano MP, Goldberg RB (2006) Management of dyslipidemia in diabetes. Cardiol Rev;14:125–35.
- Chahil TJ, Ginsberg HN (2006) Diabetic dyslipidemia. Endocrinol Metab Clin North Am 35: 491–510.
- Mooradian AD (2009) Dyslipidemia in type 2 diabetes mellitus. Nat Clin Pract Endocrinol Metab; 5:150–159.
- Heeringa P, Tervaert JW (2002) Role of oxidized low-density lipoprotein in renal disease. Curr Opin Nephrol Hypertens; 11: 287–293.
- Keane WF (2000) The role of lipids in renal disease: future challenges. Kidney Int Suppl; 75: S27–S31.
- Nishida Y, Yorioka N, Oda H, Yamakido M (1997) Effect of lipoproteins on cultured human mesangial cells. Am J Kidney Dis; 29: 919–930.
- Kamanna VS, Roh DD, Kirschenbaum MA (1998) Hyperlipidemia and kidney disease: concepts derived from histopathology and cell biology of the glomerulus. Histol Histopathol; 13: 169–179.
- Kooner JS, Chambers JC, Aguilar-Salinas CA, Hinds DA, Hyde CL, et al. (2008) Genome-wide scan identifies variation in MLXIPL associated with plasma triglycerides. Nat Genet; 40:149–151.

- Willer CJ, Sanna S, Jackson AU, Scuteri A, Bonnycastle LL, et al. (2008) Newly identified loci that influence lipid concentrations and risk of coronary artery disease. Nat Genet; 40:161–169.
- Kathiresan S, Melander O, Guiducci C, Surti A, Burtt NP, et al. (2008) Six new loci associated with blood low-density lipoprotein cholesterol, high-density lipoprotein cholesterol or triglycerides in humans. Nat Genet. 2008;40(2):189– 97. Erratum in: Nat Genet; 40:1384.
- Kathiresan S, Willer CJ, Peloso GM, Demissie S, Musunuru K, et al. (2009) Common variants at 30 loci contribute to polygenic dyslipidemia. Nat Genet; 41:56-65.
- Aulchenko YS, Ripatti S, Lindqvist I, Boomsma D, Heid IM, et al. (2009) Loci influencing lipid levels and coronary heart disease risk in 16 European population cohorts. Nat Genet;41:47–55.
- Teslovich TM, Musunuru K, Smith AV, Edmondson AC, Stylianou IM, et al. (2010) Biological, clinical and population relevance of 95 loci for blood lipids. Nature:466:707–713.
- Johansen CT, Wang J, Lanktree MB, Cao H, McIntyre AD, et al. (2010) Excess
 of rare variants in genes identified by genome-wide association study of
 hypertriglyceridemia. Nat Genet;42:684

 –687.
- Waterworth DM, Ricketts SL, Song K, Chen L, Zhao JH, et al. (2010) Genetic variants influencing circulating lipid levels and risk of coronary artery disease. Arterioscler Thromb Vasc Biol; 30:2264–2276.
- Tukiainen T, Kettunen J, Kangas AJ, Lyytikainen LP, Soininen P, et al. (2012)
 Detailed metabolic and genetic characterization reveals new associations for 30 known lipid loci. Hum Mol Genet;21:1444–1455.
- Kavanagh D, McKay GJ, Patterson CC, McKnight AJ, Maxwell AP et al. (2011)
 Association analysis of Notch pathway signalling genes in diabetic nephropathy.
 Diabetologia 54: 334–338.
- Perkins BA, Ficociello LH, Silva KH, Finkelstein DM, Warram JH et al. (2003). Regression of microalbuminuria in type 1 diabetes. N Engl J Med. 348: 2285–2293.
- Pezzolesi MG, Poznik GD, Mychaleckyj JC, Paterson AD, Barati MT et al. (2009) Genome-wide association scan for diabetic nephropathy susceptibility genes in type 1 diabetes. Diabetes 58: 1403–1410.
- Lewis EJ, Hunsicker LG, Bain RP, Rohde RD (1993). The effect of angiotensinconverting-enzyme inhibition on diabetic nephropathy. The Collaborative Study Group. N Engl J Med. 329: 1456–1462. Erratum in: N Engl J Med 1993 330: 152.
- Lewis EJ, Hunsicker LG, Clarke WR, Berl T, Pohl MA et al. (2001).
 Renoprotective effect of the angiotensin-receptor antagonist irbesartan in patients with nephropathy due to type 2 diabetes. N Engl J Med. 345: 851–860.
- 44. Brenner BM, Cooper ME, de Zeeuw D, Keane WF, Mitch WE et al (2001). Effects of losartan on renal and cardiovascular outcomes in patients with type 2 diabetes and nephropathy. N Engl J Med. 345: 861–869.
- 45. Molitch ME, Steffes M, Sun W, Rutledge B, Cleary P et al. (2010). Development and progression of renal insufficiency with and without albuminuria in adults with type 1 diabetes in the diabetes control and complications trial and the epidemiology of diabetes interventions and complications study. Diabetes Care 33: 1536–1543.
- Lagrost L, Gandjini H, Athias A, Guyard-Dangremont V, Lallemant C, et al. (1993) Influence of plasma cholesteryl ester transfer activity on the LDL and HDL distribution profiles in normolipidemic subjects. Arterioscler Thromb;13(6):815–825. 43.
- Weitgasser R, Galvan G, Malaimare L, Derflinger I, Hedegger M, et al. (2004) Cholesteryl ester transfer protein TaqIB polymorphism and its relation to parameters of the insulin resistance syndrome in an Austrian cohort. Biomed Pharmacother; 58: 619–627. 44.
- Terán-García M, Després JP, Tremblay A, Bouchard C (2008) Effects of cholesterol ester transfer protein (CETP) gene on adiposity in response to longterm overfeeding. Atherosclerosis; 196:455

 –460. 45.
- Dullaart RPF, Sluiter WJ (2008) Common variation in the CETP gene and the implications for cardiovascular disease and its treatment: an updated analysis. Pharmacogenomics 9: 747–763. 46.
- Igl W, Johansson A, Wilson JF, Wild SH, Polasek O, et al. (2010) Modeling of environmental effects in genome-wide association studies identifies SLC2A2 and HP as novel loci influencing serum cholesterol levels. PLoS Genet;6(1): e1000798. 49.
- Lahiry P, Cao H, Ban MR, Pollex RL, Mamakeesick M, et al. (2010) APOC1 T45S polymorphism is associated with reduced obesity indices and lower plasma concentrations of leptin and apolipoprotein C-I in aboriginal Canadians. J Lipid Res;51:843–848. 48.
- McKnight AJ, Maxwell AP, Fogarty DG, Sadlier D, Savage DA, et al. (2009) Genetic analysis of coronary artery disease single-nucleotide polymorphisms in diabetic nephropathy. Nephrol Dial Transplant; 24:2473–2476. 47.
- Danesh FR, Kanwar YS (2004) Modulatory effects of HMG-CoA reductase inhibitors in diabetic microangiopathy. FASEB J;18:805–815. 42.
- Cardon LR, Bell JI (2001) Association study designs for complex diseases. Nat Rev Genet. 2(2):91–9.