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Type 2 Diabetes Genetics: Beyond GWAS

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Abstract

The global epidemic of type 2 diabetes mellitus (T2D) is one of the most challenging problems of the 21st century and the fifth leading cause of death worldwide. Substantial evidence suggests that T2D is a multifactorial disease with a strong genetic component. Recent genome-wide association studies (GWAS) have successfully identified and replicated nearly 75 susceptibility loci associated with T2D and related metabolic traits, mostly in Europeans, and some in African, and South Asian populations. The GWAS serve as a starting point for future genetic and functional studies since the mechanisms of action by which these associated loci influence disease is still unclear and it is difficult to predict potential implication of these findings in clinical settings. Despite extensive replication, no study has unequivocally demonstrated their clinical role in the disease management beyond progression to T2D from impaired glucose tolerance. However, these studies are revealing new molecular pathways underlying diabetes etiology, gene-environment interactions, epigenetic modifications, and gene function. This review highlights evolving progress made in the rapidly moving field of T2D genetics that is starting to unravel the pathophysiology of a complex phenotype and has potential to show clinical relevance in the near future.

The global epidemic of type 2 diabetes (T2D) is a major public health problem of 21st century and the fifth leading cause of death worldwide [1]. The disease is also a leading cause of morbidity and contributes to development of premature coronary heart disease (CHD), stroke, peripheral vascular disease, renal failure, and amputation. According to latest statistics released by the International Diabetes Federation, the number of people living with diabetes is expected to rise from 366 million in 2011 to 552 million by 2030; 80% of these people with diabetes will live in developing countries (http://www.idf.org/diabetesatlas/5e/ the-global-burden). According to these predictions, in three leading countries with diabetes populations USA, India, and China the approximate estimate of 23.7, 61.3 and 90 million people with diabetes in US, India, and China in 2011 will increase to 29.6, 101.2, 129.7 million by 2030 (Figure 1). The global health expenditure on diabetes is expected to increase from 376 billion dollars in 2010 to 490 billion in 2030 [2].

Substantial evidence suggests that T2D is a multifactorial disease with a strong genetic component. High concordance rate obtained in monozygotic twins (96%) supports a substantial contribution of genetic factors to T2D [3-7]. Furthermore, 40% of first-degree relatives of T2D patients develop diabetes as compared to 6% in the general population [8]. Segregation analysis also points to the polygenic nature of T2D, in addition to the existence of a major genetic component [9,10]. The general estimates of heritability (h²) of T2D is 0.49 and the relative recurrence risk for a sib of an affected person (λ_s) to develop T2D is 3.5 [11,12]. However, the findings and the results of recent genome-wide association studies (GWAS) have significantly underestimated these heritability estimates and presents a challenge for ongoing and future investigation.

Clinical Phenotype of T2D

Unlike simple characterization of type 1 diabetes (T1D) which is primarily due to autoimmune mediated destruction of pancreatic beta cells resulting in insulin deficiency, the pathogenesis of T2D is more complex and remains a matter of debate. Hyperglycemia in T2D is a consequence of complex interplay between insulin resistance (sensitivity) and abnormal insulin secretion [13]. It is initially characterized by compensatory insulin secretion associated with insulin resistance. However, the β -cell's response is inadequate for the increased demand during progressive resistance to insulin-mediated glucose disposal,



jections for 2030 in USA, China, and India according to the latest projections by International Diabetes Federation (http://www.idf.org/diabetesatlas/5e/the-global-burden).

eventually resulting in β -cell failure and overt diabetes. There is also accumulating evidence that the β -cell is adversely influenced by influx of fatty acids [14] and cholesterol, which accumulates and exerts toxic effects when efflux by HDL is limited [15].

The progression of insulin resistance often associated with obesity and leading to mild glucose intolerance preceding an increase in glu-

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cose levels above defined thresholds for diabetes, has made it difficult to define the true phenotype, possibly because of early involvement of multiple tissues such as muscle, fat and pancreatic β -cells. Furthermore, the insulin resistant state, usually preceding diabetes by several years, is associated with a cluster of co-morbidities including obesity, dyslipidemias, and elevated blood pressure [16,17], and the presence of three or more traits is currently recognized as the metabolic syndrome which is also predictive of diabetes [18].

More than 80% of diabetic subjects are obese and these individuals typically have an android body type (upper body obesity) manifesting as an increased waist circumference [19]. Obesity, the metabolic syndrome, and T2D are becoming increasingly prevalent even in children and adolescents living in rapidly developing countries in different parts of world [20-24]. The prevalence of T2D in the US is higher among minorities and ethnic populations like African Americans, Native Americans, Hispanic Americans, Asian Americans, and Pacific Islanders than in the general population [25-28]. High prevalence is even seen in children belonging to ethnic minorities [29-32]. Although environmental factors play an important role in determining the risk of disease, overwhelming data support that genetic factors influence the disease susceptibility [33]. Lifestyle modifications (weight reduction and physical activity) delay or even prevent the development of T2D [34,35], such changes are exceedingly difficult to sustain outside the research setting and seem to have contributed little to control the diabetes epidemic [36].

T2D remains undiagnosed for many years because hyperglycemia is usually not severe enough to provoke noticeable symptoms, unlike T1D which often presents with keto-acidosis requiring admission to hospital for correction and initiation of treatment. But hyperglycemia can cause significant pathological and functional changes, which can cause organ damage before the diagnosis of T2D is made. The long term effects of diabetes include micro-vascular and macro-vascular complications and those who have genetic susceptibility are at greater risk [37]. Micro-vascular complications are progressive development of disease in fine capillaries supplying blood to the kidneys and retina of the eye that results in blindness. Macro-vascular complications include hypertension, coronary artery disease, peripheral vascular disease, cerebral vascular disease, and hyperlipidemia. Diabetic patients also have neuropathy, which may lead to foot ulcers, amputations, sexual dysfunction, and non-healing skin wounds. Certain infections such as staphylococcal sepsis is more common in diabetics and infections of the ear, nose, throat as well as reactivation of tuberculosis associated with high rate of mortality and morbidity are more likely with poor blood glucose control. Consequently, diabetes is a disproportionably expensive disease. The economic impact of diabetes in the US is enormous, and is expected to increase from \$113 billion to \$336 billion [38].

Linkage and Candidate Gene Association Studies

Progress in identifying the genetic basis of simple Mendelian (monogenic) diseases during the past decade has been substantial. More than 3000 monogenic disorders have been successfully mapped by linkage and family based studies (http://www.ncbi.nlm.nih.gov/ omim). On the other hand, complex diseases like T2D do not segregate in simple Mendelian fashion but rather are affected by multiple genetic and environmental factors. Linkage and candidate-gene focused studies were successful in identifying some rare familial forms of T2D presenting at young ages called maturity onset diabetes of young (MODY), mitochondrial diabetes and neonatal diabetes. However, linkage and association studies on the common form of T2D provided inconsistent results and failed replication in multiple populations [39]. Only *PPARG, KCNJ11* and *TCF7L2* were identified as established genes associated with common forms of T2D [40]. Apparently, inconsistencies across populations were due to the heterogeneity of the disease itself or its pathogenesis, incorrect candidate selection because of incomplete knowledge of molecular mechanisms, variation in study design, sample size, population-specific linkage disequilibrium, choice, analytical methods, or over-interpretation of results [40-43].

Genome-Wide Association Studies (GWAS)

Completion of the Human Genome Project in 2003 [44] led to subsequent advances in biomedical research. Since 2007, a new technology in the form of 'genome-wide chips' has facilitated remarkable progress in T2D genetic research with the first publication of five large GWA scans within the span of four months, showing that more than 500,000 SNP markers distributed across the genome [45-49]. This approach has been successful in locating genes for other diseases besides T2D and obesity [40] namely, type 1 diabetes [50], prostate cancer [51], rheumatoid arthritis [52], Crohns disease [53,54], and cardiovascular disease [55] and is being applied to other complex disorders. Use of this 'hypothesis-free' approach involved in GWAS has opened new areas of biology to explore as discoveries of more than seventy entirely new T2D loci clearly suggest that associations are not limited to candidate genes and by applying GWAS and re-sequencing approaches, new genes involved in disease pathogenesis can be identified [56] (Table 1).

The number of risk alleles for complex diseases identified by GWAS since 2007 exceeded those identified in the entire preceding decade, and these studies can effectively detect multiple common variants with small effects with odds ratios (ORs) <1.2 [36,39,57] but offers limited power to detect rare variants with stronger effects [58]. However, GWAS have the best chance of detecting and identifying additional genes for T2D and are being pursued in different population groups, to provide a more realistic 'genetic risk landscape' of the disease [59], and account for variation in population-specific environmental interactions. Notably, most immediate advances from GWA studies are not disease prediction or prevention, rather these studies are providing a better understanding of the disease pathophysiology since further studies can lead to defining the role of the newly identified genes in key metabolic pathways involved in pathogenesis. Also GWAS, together with targeted resequencing of the identified loci allows recovery of a more complete inventory of the sequence variation leading to identification of novel metabolic pathways and proteins as potential targets for interventions [57]. Since substantial numbers, maybe thousands, of genes with small effects are operative in human disease and can be identified by this approach, their combined effect may be substantial [56]. Furthermore, gene-gene interaction may not only be additive but effects of a mutation can be more than doubled when compounding mutations are present. The phenomenon known as epistasis was first shown to occur in a rodent knockout model in which the insulin receptor when combined with IRS-1 knockout resulted in 40% increase in diabetes onset compared to isolated knock-outs [60]. The phenomenon could contribute to effect size and in part account for the low heritability accounted for by classic GWAS. It follows that multi locus association testing could be effective in the presence of epistasis. With this in mind the Welcome Trust Case Control Consortium used twolocus tests of association and detected epistatic signaling of association at 79 SNP pairs [61]. However the large number of possible associations in GWAS requires filtration strategies based on knowledge of the biochemistry and known interacting pathways [62]. However, several models for analysis have been proposed and consensus on the optimal methodology has not been reached [63].

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		Gene			Risk Allele Freg in			First Author Journal: Published
SNP	Gene	Region	Population	Trait	Controls	OR/Beta [95% CI]	P-Value	Year
rs10923931	NOTCH2, ADAM30	1p12	European	T2D	T[0.11]	1.13[1.08-1.17]	4x10-8	Zeggini et al. Nat Genet; 2008
rs243021	BCL11A	2p16.1	European	T2D	A[NR]	1.08[1.06-1.10]	3x10-15	Voight et al. Nat Genet; 2010
rs7578326	IRS1	2q36.3	European	T2D	A[NR]	1.11[1.08-1.13]	5x10-20	Voight et al. Nat Genet; 2010
rs2943641	LOC64673, IRS1	2q36.3	European	T2D/Other traits	C[0.63]	1.19[1.13-1.25]	9x10-12	Rung et al. Nat Genet; 2009
rs7593730	RBMS1, ITGB6	2q24.2	European	T2D	?[0.78]	1.11[1.08-1.16]	4x10-8	Qi et al. Hum Mol Genet; 2010
rs7578597	THADA	2p21	European	T2D	T[0.9]	1.15[1.10-1.20]	1x10-9	Zeggini et al. Nat Genet; 2008
rs4607103	ADAMTS9	3p14.1	European	T2D	C[0.76]	1.09[1.06-1.12]	1x10-8	Zeggini et al. Nat Genet; 2008
rs4402960	IGF2BP2	3q27.2	European	T2D	T[0.3]	1.14[1.11-1.18]	9x10-16	Scott et al. Science; 2007
rs358806	NR	3p14	European	T2D	?[0.8]	1.16[1.03-1.33]	3x10-6	WTCC Consortium Nature; 2007
rs1801282	PPARG	3p25.2	European	T2D	C[0.82]	1.14[1.08-1.20]	2x10-6	Scott et al. Science; 2007
rs17036101	SYN2, PPARG	3p25.2	European	T2D	G[0.93]	1.15[1.10-1.21]	2x10-7	Zeggini et al. Nat Genet; 2008
rs1801214	WFS1	4p16.1	European	T2D	T[NR]	1.13[1.08-1.18]	3x10-8	Voight et al. Nat Genet; 2010
rs4689388	WFS1, PPP2R2C	4p16.1	European	T2D/Other traits	T[0.57]	1.16[1.10-1.21]	1x10-8	Rung et al. Nat Genet; 2009
rs12518099	LOC72901, CETN3	5q14.3	European	T2D/Other traits	C[0.23]	1.16[1.10-1.22]	7x10-7	Rung et al. Nat Genet; 2009
rs4457053	ZBED3	5q13.3	European	T2D	G[NR]	1.08[1.06-1.11]	3x10-12	Voight et al. Nat Genet; 2010
rs7754840	CDKAL1	6p22.3	European	T2D	C[0.36]	1.12[1.08-1.16]	4x10-11	Scott et al. Science; 2007
rs896854	TP53INP1	6q22.1	European	T2D	T[NR]	1.06[1.04-1.09]	1x10-9	Voight et al. Nat Genet; 2010
rs9472138	VEGFA	6p21.1	European	T2D	T[0.28]	1.06[1.04-1.09]	4x10-6	Zeggini et al. Nat Genet; 2008
rs864745	JAZF1	7p15.1	European	T2D	T[0.5]	1.1[1.07-1.13]	5x10-14	Zeggini et al. Nat Genet; 2008
rs972283	KLF14	7q32.3	European	T2D	G[NR]	1.07[1.05-1.10]	2x10-10	Voight et al. Nat Genet; 2010
rs13266634	SLC30A8	8q24.11	European	T2D	C[0.61]	1.12[1.07-1.16]	5x10-8	Scott et al. Science; 2007
rs564398	CDKN2A, CD- KN2B	9p21.3	European	T2D	T[0.56]	1.13[1.08-1.19]	1x10-6	Zeggini et al. Science; 2007
rs7018475	CDKN2B	9p21.3	European	T2D	?[NR]	1.35[1.18-1.56]	3x10-8	Huang J et al. Eur J Hum Genet; 2012
rs13292136	CHCHD9	9q21.31	European	T2D	C[NR]	1.11[1.07-1.15]	3x10-8	Voight et al. Nat Genet; 2010
rs12779790	CDC123,CAMK1D	10p13	European	T2D	G[0.18]	1.11[1.07-1.14]	1x10-10	Zeggini et al. Nat Genet; 2008
rs1111875	HHEX	10q23.33	European	T2D	C[0.52]	1.13[1.09-1.17]	6x10-10	Scott et al. Science; 2007
rs7903146	TCF7L2	10q25.2	European	T2D	T[0.26]	1.38[1.31-1.46]	2x10-31	Saxena et al. Science; 2007
rs1552224	ARAP1	11q13.4	European	T2D	A[NR]	1.14[1.11-1.17]	1x10-22	Voight et al. Nat Genet; 2010
rs5219	KCNJ11	11p15.1	European	T2D	T[0.46]	1.14[1.10-1.19]	7x10-11	Scott et al. Science; 2007
rs231362	KCNQ1	11p15.5	European	T2D	G[NR]	1.08[1.06-1.10]	3x10-13	Voight et al. Nat Genet; 2010
rs1387153	MTNR1B	11q14.3	European	T2D	T[NR]	1.09[1.06-1.11]	8x10-15	Voight et al. Nat Genet; 2010
rs1153188	DCD	12q13.2	European	T2D	A[0.73]	1.08[1.05-1.11]	2x10-7	Zeggini et al. Nat Genet; 2008
rs1531343	HMGA2	12q14.3	European	T2D	C[NR]	1.1[1.07-1.14]	4x10-9	Voight et al. Nat Genet; 2010
rs7957197	HNF1A	12q24.31	European	T2D	T[NR]	1.07[1.05-1.10]	2x10-8	Voight et al. Nat Genet; 2010
rs7961581	TSPAN8, LGR5	12q21.1	European	T2D	C[0.27]	1.09[1.06-1.12]	1x10-9	Zeggini et al. Nat Genet; 2008
rs8042680	PRC1	15q26.1	European	T2D	A[NR]	1.07[1.05-1.09]	2x10-10	Voight et al. Nat Genet; 2010
rs11634397	ZFAND6	15q25.1	European	T2D	G[NR]	1.06[1.04-1.08]	2x10-9	Voight et al. Nat Genet; 2010
rs8050136	FTO	16q12.2	European	T2D	A[0.4]	1.23[1.18-1.32]	7x10-14	Zeggini et al. Science; 2007
rs4430796	HNF1B	17q12	European	T2D	G[NR]	1.14[1.08-1.20]	2x10-6	Voight et al. Nat Genet; 2010
rs5945326	DUSP9	Xq26	European	T2D	G[NR]	1.27[1.18-1.37]	3x10-10	Voight et al. Nat Genet; 2010
rs7542900	SLC44A3, F3	1p21.3	African Amer.	T2D	C[0.56]	1.16[1.09-1.25]	6x10-6	Palmer ND et al. PLoS One; 2012
rs7560163	RBM43, RND3	2q23.3	African Amer.	T2D	C[0.86]	1.33[1.19-1.49]	7x10-9	Palmer ND et al. PLoS One; 2012
rs2722769	GALNTL4, LOC729013	11p15.3	African Amer.	T2D	C[0.53]	1.35[1.19-1.54]	2x10-6	Palmer ND et al. PLoS One; 2012
rs7107217	TMEM45B, BARX2	11q24.3	African Amer.	T2D	C[0.91]	1.18[1.10-1.27]	3x10-7	Palmer ND et al. PLoS One; 2012
rs17045328	CR2	1q32.2	Asian	T2D	G[0.3]	1.38[1.20-1.59]	7x10-6	Sim X et al. PLoS Genet; 2011
rs12027542	PCNXL2	1q42.2	Asian	T2D	A[0.61]	1.41[1.23-1.61]	4x10-7	Sim X et al. PLoS Genet; 2011
rs7630877	PEX5L	3q26.33	Asian	T2D	A[0.17]	1.32[1.17-1.49]	7x10-6	Sim X et al. PLoS Genet; 2011
rs3773506	PLS1	3q23	Asian	T2D	C[0.06]	1.81[1.39-2.35]	9x10-6	Sim X et al. PLoS Genet; 2011
rs831571	PSMD6	3p14.1	Asian	T2D	c[0.61]	1.09[1.06-1.12]	8x10-11	Cho YS et al. Nat Genet; 2011
rs2063640	ZPLD1	3q12.3	Asian	T2D	A[0.17]	1.23[1.13-1.34]	3x10-6	Sim X et al. PLoS Genet; 2011
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rs6815464	MAEA	4p16.3	Asian	T2D	C[0.58]	1.13[1.10-1.16]	2x10-20	Cho YS et al. Nat Genet; 2011
rs1048886	C6orf57	6q13	Asian	T2D	G[0.18]	1.54[1.32-1.80]	3x10-8	Sim X et al. PLoS Genet; 2011
rs1535500	KCNK16	6p21.2	Asian	T2D	T[0.42]	1.08[1.05-1.11]	2x10-8	Cho YS et al. Nat Genet; 2011
rs9470794	ZFAND3	6p21.2	Asian	T2D	C[0.27]	1.12[1.08-1.16]	2x10-10	Cho YS et al. Nat Genet; 2011
rs7636	ACHE	7q22.1	Asian	T2D	A[0.06]	1.85[1.42-2.41]	5x10-6	Sim X et al. PLoS Genet; 2011
rs6467136	GCC1,PAX4	7q32.1	Asian	T2D	G[0.79]	1.11[1.07-1.14]	5x10-11	Cho YS et al. Nat Genet; 2011
rs7041847	GLIS3	9p24.2	Asian	T2D	A[0.41]	1.1[1.07-1.13]	2x10-14	Cho YS et al. Nat Genet; 2011
rs17584499	PTPRD	9p24.1	Asian	T2D	T[0.06]	1.57[1.36-1.82]	9x10-10	Tsai et al. PLoS Genet; 2010
rs6583826	KIF11	10q23.33	Asian	T2D	G[0.26]	1.18[1.10-1.27]	7x10-6	Sim X et al. PLoS Genet; 2011
rs10741243	TCERG1L	10q26.3	Asian	T2D	G[0.93]	1.75[1.38-2.23]	5x10-6	Sim X et al. PLoS Genet; 2011
rs1359790	SPRY2	13q31.1	Asian	T2D	G[0.71]	1.15[1.10-1.20]	6x10-9	Shu et al. PLoS Genet; 2010
rs1436955	C2CD4B	15q22.2	Asian	T2D	C[0.73]	1.13[1.08-1.19]	7x10-7	Shu et al. PLoS Genet; 2010
rs16955379	CMIP	16q23.2	Asian	T2D	C[0.8]	1.08[1.05-1.12]	3x10-7	Cho YS et al. Nat Genet; 2011
rs17797882	WWOX	16q23.2	Asian	T2D	T[0.32]	1.08[1.05-1.12]	9x10-7	Cho YS et al. Nat Genet; 2011
rs391300	SRR	17p13.3	Asian	T2D	G[0.62]	1.28[1.18-1.39]	3x10-9	Tsai et al. PLoS Genet; 2010
rs10460009	LPIN2	18p11.31	Asian	T2D	C[0.6]	1.35[1.18-1.54]	9x10-6	Sim X et al. PLoS Genet; 2011
rs472265	PAPL	19q13.2	Asian	T2D	G[0.22]	1.39[1.20-1.61]	9x10-6	Sim X et al. PLoS Genet; 2011
rs3786897	PEPD	19q13.11	Asian	T2D	A[0.56]	1.1[1.07-1.14]	1x10-8	Cho YS et al. Nat Genet; 2011
rs2833610	HUNK	21q22.11	Asian	T2D	A[0.57]	1.17[1.09-1.24]	4x10-6	Sim X et al. PLoS Genet; 2011
rs3923113	GRB14	2q24.3	South Asian	T2D	A[0.74]	1.09[1.06-1.13]	1x10-8	Kooner JS et al. Nat Genet; 2011
rs16861329	ST6GAL1	3q27.3	South Asian	T2D	G[0.75]	1.09[1.06-1.12]	3x10-8	Kooner JS et al. Nat Genet; 2011
rs1802295	VPS26A	10q22.1	South Asian	T2D	A[0.26]	1.08[1.05-1.12]	4x10-8	Kooner JS et al. Nat Genet; 2011
rs2028299	AP3S2	15q26.1	South Asian	T2D	C[0.31]	1.1[1.07-1.13]	2x10-11	Kooner JS et al. Nat Genet; 2011
rs7178572	HMG20A	15q24.3	South Asian	T2D	G[0.52]	1.09[1.06-1.12]	7x10-11	Kooner JS et al. Nat Genet; 2011
rs4812829	HNF4A	20q13.12	South Asian	T2D	A[0.29]	1.09[1.06-1.12]	3x10-10	Kooner JS et al. Nat Genet; 2011
rs2237897	KCNQ1	11p15.4	Japanese	T2D	C[0.61]	1.33[1.24-1.41]	1x10-16	Unoki et al. Nat Genet; 2008
rs2237892	KCNQ1	11p15.5	Japanese	T2D	C[0.61]	1.4[1.34-1.47]	2x10-42	Yasuda et al. Nat Genet; 2008
rs7172432	C2CD4A,C2CD4B	15q22.2	Japanese	T2D	?[0.58]	1.11[1.08-1.14]	9x10-14	Yamauchi et al. Nat Genet; 2010
rs730570	C14orf70	14q32.2	Mexican	T2D	G[NR]	1.14[1.08-1.21]	8x10-6	Parra EJ et al. Diabetologia; 2011

Text in bold indicates highly replicated loci in multiple ethnic populations

Table 1: Summary of type 2 diabetes genes discovered in GWA studies (in order of gene region).

The common strategy used for elucidating the inherited components of complex disease is based on common disease-common variant hypothesis [64,65]. The hypothesis has held true in the case of putative causal variants in APOE, including APOE £4, association with Alzheimer's disease, interleukin 23 receptor association with Crohns disease (the at-risk allele has a frequency 93% in the general population), and PPARG2 association with T2D (at-risk (pro) allele frequency ~85% in general population). The assumption with the use of common variants is that either they will identify the association with the common disease, or a quantitative trait (QT) directly, or indirectly being in linkage disequilibrium (LD) with a functional variant. Most of the T2D loci identified using GWAS approach are common variants with small effects. The first GWAS in T2D was published by a French group of investigators who identified a zinc transporter and member of solute carrier family SLC30A8 and HHEX along with confirming the association of TCF7L2 and KCNJ11 with T2D [49]. Three additional GWAS in their jointly published findings confirmed TCF7L2, KCNJ11, PPARG, SLC30A8, and HHEX. They also discovered novel T2D loci called CD-KAL1 and a variant near CDKN2A-B [47,48,66]. These findings were also simultaneously confirmed by GWAS performed on an Icelandic sample by the deCODE researchers [67]. Association of FTO with T2D was also first discovered by GWAS for obesity [OR for T2D being 1.27, $p < 10^{-8}$ [68] and was later confirmed in replication studies [46].

Additionally, a joint meta-analysis of three previously conducted T2D GWA scans by the Diabetes Genetics Replication And Meta-analysis (DIAGRAM) consortium [69] has detected an additional six novel loci with strong evidence of association; JAZF1, CDC123, TSPAN8, THADA, ADAMTS9, and NOTCH2 [69] in 10,128 individuals using 2.2 million SNPs. Also, another European study reported a strong association of an intronic SNP (rs560887) in G6PC2 with fasting plasma glucose levels by screening 392,935 SNPs in 654 non-T2D participants using GWAS [70]. Association of G6PC2 with fasting plasma glucose levels was replicated in the FUSION study [71]. Further, a variant near MTNR1B has been identified to increase FBG levels in two separate GWA studies [72,73]. Association of MTNR1B variants (rs10830963 and rs1387153) with FBG or T2D in our Punjabi cohort from India has not been confirmed. Instead a less common variant on 5' UTR (rs1374645) of MTNR1B revealed a strong association with low FBG levels in normoglycemic individuals with low BMI (<25 kg/m²). Furthermore, our data showed strong interaction of this variant with BMI with respect to FBG levels [74]. Most recently, a meta-analysis was performed on 21 GWAS comprising 46,186 participants to identify loci influencing glycemic traits. They performed follow-up studies on 25 loci in up to 76,558 additional subjects and identified nine loci associated with fasting glucose and HOMA-B (ADCY5, MADD, ADRA2A, CRY2, FADS1, GLIS3, SLC2A2, PROX1 and C2CD4B) and one influencing fasting insulin and HOMA-IR (IGF1) in European populations [75].

Another study analyzed combined GWAS data from 8,130 individuals with T2D and 38,987 controls of European descent, and in follow-up meta-analysis found signals in a further 34,412 cases and 59,925 controls and identified 12 new T2D associations including new signals at *KCNQ1*, *KLF14*, *DUSP9*, *CENTD2*, *HMGA2*, *HNF1A* and *PRC1* [76].

Genetic Diversity among Populations

The majority of GWAS so far (96%) have been predominantly performed in Caucasians [77,78]. Given the existence of a strong genetic diversity among the world communities due to biological traits, cultural histories, languages, caste system, physical appearance, food habits etc., the information from GWA findings on Caucasians should not be transferred and used to predict risk in other populations [79,80]. Even the largest and best executed GWAS is unlikely to provide a complete assessment of "the genetic risk landscape" of the disease [59]. These loci neither account for all cases of T2D, nor do they sufficiently account for the variation in quantitative sub-phenotypes of T2D. Extending the use of new methodologies such as GWA to study non-European ancestries with different mutational spectra and demographic and cultural histories is important for identifying the population-specific patterns for allele frequency and LD relationship of the susceptibility loci, and population-specific environmental factors for disease risk or protection [57]. Several GWAS in other ethnic groups are currently ongoing and will be capable of revealing novel susceptibility loci. GWAS have proven more successful in small homogenous populations like Finland, Iceland and Costa Rica, and have potential to detect regionally common variants which may be missed in outbred populations [78]. It is of interest to note that many of the common loci originally associated with diabetes in European populations have not been replicated in other non-European populations [49,67,77,81]. For example, loci associated with Crohns disease in Caucasian GWAS were not associated in Japanese [81], also the association of CDKAL1 locus with T2D was not seen in African Americans [67]. As discussed earlier, the common intron 1 FTO variant (rs9939609) was initially associated with diabetes in Europeans, and phenotypic interactions appear to be diabetogenic, but the associations were due to obesity [68,69]. However, the question has been explored in independent association and meta-analyses of South Asian populations in whom BMI and waist association with FTO is similar to that seen in Europeans, but a strong association with diabetes is only partly accounted for by BMI [82,83]. A recent large scale meta-analysis conducted on 96,551 individuals from East and South Asia confirmed that the association of rs9939609 with T2D was independent of obesity [84].

There is now a strong consensus that a single population is not sufficient to uncover all the variants underlying complex diseases [77]. Recent GWAS studies in non-European populations have yielded intriguing new variants, for instance, the KCNQ1 signal (first noticed in Japanese) provided insight into novel etiological pathways leading to insulin resistance and T2D. Joint meta-analysis of Caucasian GWA studies later discovered a second independent signal at the KCNQ1 locus (rs231362) and an overlap between loci implicated in monogenic and multifactorial forms of diabetes at HNF1A [76]. The new Caucasian signal at KCNQ1 was later confirmed in our study on Asian Indians [85]. These findings provide new insights into the pathophysiology of this extremely complex disease by discovering previously unknown variants and often replicating the findings but not always. Association of UBE2E2 with T2D was first described in East Asians [86]. Recent larger meta-analysis using East Asian GWAS cohorts yielded eight new loci for T2D in (GLIS3, PEPD, FITM2-R3HDML-HNF4A, KCNK16, MAEA, GCC1-PAX4, PSMD6 and ZFAND3) [87]. Six novel loci for T2D (*GRB14*, *ST6GAL1*, *VPS26A*, *HMG20A*, *AP3S2* and *HNF4A*) were first discovered in our South Asian population GWAS by Kooner et al. [88]. To date, GWAS and meta-analyses in T2D and related quatitative traits have together identified approximately 75 susceptibility variants which together explain about 10% of the observed familial aggregation (Table 1). Although these effect sizes detected so far do not reflect biological or clinical significance of these variants, they highlight a particular genomic region likely to be associated with the trait. Of all the complex traits studied by GWAS , Crohns disease, has so far had the strongest association with genetic components of risk; up to 20% being explained by variants detected by GWAS [89]. Furthermore, GWAS provided important biological insight with direct clinical relevance adding to significant understanding of the disease compared to what was known about Crohns disease five years back.

Hidden Heritability and Rare Variant Hypothesis

Despite these successes, results of multiple GWAS and meta-analysis studies (based on the common disease-common variant hypothesis [64,65]) show that common variants discovered in these studies explain only a small fraction with odds ratios (ORs) ranging from 1.1-1.2 (p values <10⁻⁸) of the heritable component of T2D risk, and these studies have not pinpointed with certainty the causal variants at the associated loci [90]. Importantly, commercial arrays used in GWAS have captured much less than 60% of common SNP information in the majority of the GWAS conducted to date [91]. One approach could be to expand the sample sizes to detect more variants of moderate effects, however, the GWAS approach is usually underpowered to locate those variants with a frequency of <5% [57], and is considered unlikely to yield more information on the heritable control of disease than what has already been revealed by GWAS [92]. Moreover, association studies, based on the common disease-common variant hypothesis would have less relevance if the disease is caused by less common mutations at multiple sites and each arising on a different ancestral haplotype [93].

Many studies have suggested the large role of rare variants in complex diseases [93,94], and that the genes carrying common variants with small effects also are likely to carry rare variants with large effects [69,90] which can only be detected by targeted resequencing [57,95]. Also, several independently and dominantly acting rare variants from different genes, together can confer a detectable relative risk for a given "common disease-rare variant" hypothesis [96]. These low frequency variants with stronger penetrance would be expected if trait alleles have undergone purifying selection [94]. Resequencing of certain candidate genes have shown that certain rare variants can be identified in individuals with extreme phenotypes [97]. These rare variants when clustered together can produce a significant genetic effect in a complex disease [56]. For instance, rare functional recessive mutations contribute to HDL and triglyceride levels [98,99], suggesting the possibility of discovering such variants in a common disease such as T2D [100]. Fearhead and colleagues [101] detected several rare functional variants in five genes among 124 subjects with multiple adenomatous polyps showing combined ORs of 2.2; p = 0.0001. Furthermore, 25% of these individuals had a single rare variant and none of them had two or more. Their findings suggested that mainly missense and possibly also nonsense, promoter, and splicing variants can collectively explain a significant portion of inherited susceptibility to colorectal adenomas. It is realistic to expect similar findings in other chronic diseases [94,96] including diabetes. It is also important to note that rare variants are often founder variants caused by genetic drift and can be population-specific and less cosmopolitan. For instance, the rare variants discovered in the APC gene for colorectal adenomas in Ashkenazi Jews were not present

in Koreans [102]. These findings therefore suggest that the search for genetic variation in diabetes-susceptible populations should be carried out in multiple ethnicities in carefully chosen candidate genes, and in clearly phenotyped disease groups. Further use of whole genome rare variants from the 1000 Genomes Project (www.1000genomes.org) and T2D-GENES Consortium with data from multiple ethnic groups will substantially improve power of these studies and capture heritability associated with rare variants from specific regions or even from the other regions of the genome, especially those (deletions/insertions, duplications, inversions) that have escaped detection in GWAS [103]. Additionally, imputation from 1000 Genomes Project will allow discovery of new association signals from the missing "dark matter". It will also assist in post-GWAS fine mapping and functional characterization of known associations and enhance our ability to fine map signals of local adaptation as has been successfully achieved for refining the locus for smoking quantity at 15q25 (P = 9.4×10^{-19}) using imputed data from 1000 Genomes which increased marker density to five-fold compared to HapMap2 [104].

In addition to major gene mutations and SNPs, changes in the normal genome sequence can result in significant miscoding with disease association [105]. Submicroscopic deletions, duplications, insertions, inversions and translocations can occur in a small DNA segment and are known as copy number variations (CNV) [106]. Since methodology for CNV detection is relatively new [107], few studies have looked for CNV associations with T2D. Since copy number variation in the leptin receptor gene (LEPR) is associated with T2D [108], it is possible that genome-wide detection of CNV could improve association with T2D as shown for insulin resistance in the HyperGEN study, in which joint tests of association for SNPs and CNVs were performed resulting in detection of association in the T-cell receptor gene (TCRVB) in African Americans [109]. Furthermore, generation of CNV maps [110] supports potential for studying CNV for missing variation in T2D and possibly accounting for the inter-ethnic variation found among diabetes-prone populations [111]; however, common CNVs may be tagged by SNPs and have been indirectly explored in previous SNP-based studies [112]. Further, the event resulting in a CNV may also disrupt regional genes and their regulatory elements accounting for disease association [106].

Role of Epigenetics in Missing Heritability

The assumption of SNP-based GWAS has been that genomic sequence variations account for disease phenotypes, however recent developments in the field of epigenetics has shown altered expression of genes that could lead to enhanced prediction of obesity and diabetes [113]. The modifications include DNA methylation and histone modifications such as ubiquitination, adenosine diphosphate ribosylation, phosphorylation, acetylation and methylation. Since the modifications are often observed during cellular growth and differentiation, it follows that the changes may influence embryological development of cells involved in carbohydrate homeostasis such as the β -cells, adipocytes and myocytes. For example, DNA methylation occurs in the leptin gene promotor and in GLUT4 in 3T3-L1 cells [114]. The epigenetic mechanism mediate the influence of the environment on gene expression [115]. For instance, leptin plays an important role in programming of obesity-related traits [116,117], change in expression could be involved in the pathogenesis of T2D, in part supporting epidemiological findings that fetal growth and early nutrition in childhood influence adult disease incl, HNF1B (rs 4430796 uding T2D [118]. For example, increased methylation of PDX-1 results in decreased expression in pancreatic islets from patients with T2D suggesting that modification of *PDX*-1 plays a role in T2D pathogenesis [119]. Besides, epigenetic modifications are sensitive to change by nutrients, hormones and toxins that can operate during developmental windows according to tissue-specific and sex-specific mechanisms [120], which potentially can be reversed with treatment and prevention [121,122].

Gene-Environment Interaction

Evidence from the epidemiology of T2D overwhelmingly supports a strong environmental influence interacting with genetic predisposition in a synergistic fashion as has been recently reviewed [123], however current state-of-the-art methods for measuring environmental effects lack precision and can result in changes in statistical power to detect interaction [123,124]. Since lifestyle factors are important in preventing diabetes [125,126], interaction of gene variants with measures of dietary intake and exercise have been selected for studies on gene-environment interaction. For example, HNF1B (rs 4430796) was shown to interact with exercise; low levels of activity enhanced the risk of T2D in association with absence of the risk allele, but there was no protective effect of exercise when the allele was present. It follows that subgrouping by genotype may serve to enhance risk prediction while considering gene-environment interaction as has been done for exercise [127]. Also lifestyle including exercise modified the effect of a CDKN2A/B variant on 2-hour glucose levels in the Diabetes Prevention Program [128] but was not confirmed in the HERITAGE study using different measurements and phenotypes involving insulin sensitivity and β -cell function [129]. The pro12ala *PPARG* variant also interacts with physical activity for effect on 2-hour glucose levels [130], which was confirmed in the smaller HERITAGE study [129]. In addition, a relationship of dietary fat intake with plasma insulin and BMI differs by the pro12ala PPARG genotype [131].

It is possible that there are genes that because of their known metabolic involvement are likely to interact with specific nutrients. For example, SLC30A8 which encodes a zinc transporter localized in secretory granules, interacted with dietary zinc to effect fasting insulin levels [132]. However, the majority of GWAS variants have not shown interaction with environmental factors for effect on diabetes or related traits. Therefore, it is likely that prospective future studies will utilize improved assessment methods to increase power and avoid false interpretation [133,134]. This could be enhanced by prioritizing variants that are most likely to have effects [135] or selective sampling according to extremes of the environmental factor could reduce the requirement for sample size [136]. These and other strategies such as meta-analysis, nested case control and genotype-based studies have been recently reviewed [123,133] and the difficulties in measuring environmental exposures have been emphasized, including the application of analyses based on logistic regression [124] and problems with instruments such as physical activity questionnaires [137]. Validated food frequency questionnaires are popular instruments for evaluation diabetes risk and are often used in conjunction with food analysis software [138,139]. Similar methodology has been adapted to assess two predominant food consumption patterns by Prudent and Western [140], and demonstrated synergistic interaction with genotype and a less healthy Western dietary pattern in determining male risk for T2D by showing that the gene-diet interaction was higher in men with a high genetic risk score determined by a gene counting method [141]. Also the effects of diet may predominate at specific developmental periods [142] suggesting that age and associated physiological changes are important as well as differences between genders. It has also been observed that homogeneity of an environmental factor such as physical activity in an Asian Indian study, may reduce ability to detect interaction, but could be solved by subgrouping by the level of activity [143], but increased recruitment would be needed to maintain power.

It is also speculated that several rapidly developing communities lived through centuries of famine and starvation and over the years have adapted to survival under conditions of unreliable food supply because they have "thrifty genes" which enable efficient storage of energy during periods of food deprivation [144]. The genes that once represented a survival advantage, now, in the modern environment with continuous low physical activity and dietary excess, result in obesity and T2D [145,146]. In addition, early protein deprivation, as indicated by low birth weight is considered as an important contributor to the high prevalence of T2D especially in people living in the under-developed world [147-150] possibly when followed by rapid catch-up growth in infancy and early childhood [151]. The "thrifty phenotype" hypothesis was based on data showing a positive association of low birth weight with cardiovascular disease and diabetes in adulthood [152], but the metabolic basis of gestational programming of the fetus is poorly understood. In Europeans, the risk of T2D increases with increase in birth weight whereas in Pima Indians and in South Asians T2D rates increase with both low and high birth weight [153] suggesting that the effective birth weight distribution is bimodal. Poor intra-uterine nutrition also may be associated with abnormal β-cell development and differentiation [154-156]. Therefore, it is imperative to develop a deeper understanding of genetic, epigenetic, environmental, cultural, and social factors that may interact to cause progression to T2D, and are responsible for the declining health of the fastest developing nations of the third world.

Prospects of Translation of Genetic Findings into Clinical Practice

As discussed in more detail in previous sections, investigations on genetic determinants of T2D have progressed substantially following increased characterization of the genome and use of GWAS, however a large portion of the heritability is unaccounted for, and many of the genes commonly found in GWAS have small effects [157-160]. Thus, state-of-the-art clinical use of genotyping is currently restricted to investigation. Also the use SNPs to predict disease onset or response to treatment is limited but can be done with some monogenic forms of diabetes [160]. Furthermore, metabolic pathways that have been responsive to treatment with pharmaceutical agents or exercise, or have been associated with disease have rarely contained genes that have been identified by large scale studies such as GWAS.

However, pharmacogenetic studies on association with the effects of anti-diabetes drugs have provided a significant clinical link with epidemiological findings [161], and frequently used agents in the treatment of T2D such as metformin, sulfonylureas, and thiozolidinediones have been shown to have gene associations. The principal metforminassociated pathway involves adenosine-monophosphate-activated kinase (AMPK), a regulator of glucose levels [162], and therefore has potential as a drug target [163] for the control of T2D by metformin and possibly other agents. It serves as a sensor of cellular energy status via its phosphorylation and activation. Liver AMPK controls hepatic glucose production by inhibiting expression of phosphoenolpyruvate carboxykinase (PEPCK) and glucose-6-phosphatase (G6Pase) and this action is down-regulated by the drug, metformin. Candidate genes encoding enzymes, cofactors and transcription factors involved in AMPK-associated pathways have potential for studies on therapeutic responses. Since gene variants coding for steps in metabolic pathways involving metformin's action may lead to defining treatment responses, investigators in the United Kingdom conducted a meta-analysis using the glycemic response to metformin as the phenotype. A variant at a locus containing ATM, the ataxia telangiectasia mutated gene (rs11212617) was associated with the response [164]. The importance of the variant was emphasized by showing that specific inhibition of ATM attenuated the phosphorylation and activation of AMPK in response to metformin in a rat hepatoma cell line [164]. Although the effect attributed to the variant was small, the study identified an important upstream AMPK regulator adding information on how metformin works, and in addition identified a possible mechanistic link with DNA repair and cancer prevention. Shu et al. have investigated polymorphisms in SLC22A1 which encodes organic cation transporter 1(OCT-1) which functions to facilitate absorption of metformin into hepatocytes and identified association with reduced responsiveness to metformin's effect on glucose levels during an oral glucose tolerance test [165]. Also association with HbA1c was shown with a variant in SL-C47A1 encoding the multidrug and toxin extrusion protein 1 involved in the excretion of metformin into bile and urine [166]. Furthermore, the two genes both acting to increase intracellular metformin, have an interactive effect [167]. The roles of both genes were supported by the Diabetes Prevention Program investigators [168] who confirmed association of the glucose-lowering response to metformin and variants in SLC47A1, and interactions with SLC22A1 and the AMPK genes STK11, PRKAA1, and PRKAA2.

Genes encoding the receptor and pathways for PPAR γ agonists constituting the thiozolidinediones, are often required for glucose control because of their role in promoting insulin sensitivity. Of 133 PPARG variants tested in the TRIPOD study, eight showed evidence for association with response to troglitazone therapy defined as change in tolerance to intravenous glucose (IVGTT) [169] but no association with fasting glucose suggesting that the response shown in the IVGTT represents insulin resistance whereas fasting glucose is more likely to be a measure of β -cell failure.

In studies where overt T2D has been the phenotype the majority of associated polymorphisms have encoded proteins known to be involved in β-cell metabolism; for example TCF7L2, KCNJ11 and HHEX have shown robust association [170,171]. This suggests that these genes could prove useful in predicting $\beta\mbox{-cell}$ preservation during the course of T2D. The glucokinase gene (GCK) coding for the initial glucosesensing step in the β -cell can have activating mutations causing hypoglycemia that might provide structural and functional models leading to drug targets for treating T2D [172]. In the GoDARTs study, investigators examined the medication response of metformin and sulphonylurea based on the TCF7L2 variants mainly affecting the β -cell. The carriers of the at risk 'T' allele responded less well to sulphonylurea therapy than metformin [173]. Also it is of significant public health interest that in the Diabetes Prevention Program, lifestyle modifications were shown to reduce the risk of diabetes conferred by risk variants of TCF7L2 at rs7093146, and in placebo participants who carried the homozygous risk genotype (TT), there was 80% higher risk for developing diabetes compared to the lifestyle intervention group carrying the same risk genotypes [35]. These findings could herald significant future progress in the field of T2D pharmacogenomics, possibly leading to the development and use of agents tailored on the basis of genotype.

Predictive Power of Genetics beyond Traditional T2D Risk Factors

Traditional clinical assessment includes a complete history of the presenting traits including family history of the traits, and followed

by laboratory studies that present either as continuous variables or dichotomous values according to a cut-point. From this basic clinical information it has been possible to derive risk scores that predict T2D such as the Finnish Diabetes Risk Score and the Diabetes Risk Calculator [174,175], which have been used to assess predictive ability using receiver operating characteristic curves or the area under the curve (AUC) as an index of discriminative accuracy. Using these scores prediction has only been considered to be modest [176], leading to the question whether genetic risk prediction might be superior. Studies have been based on genetic variants alone and in combination with environmental factors, but studies have differed in the number and choice of genes and environmental factors. However, disappointingly the genetic risk models had lower prediction capacity than the clinical environment-based models [176]. Differences in populations, BMI, age and duration of follow-up may offer some explanation [176,177], but improving genetic methodologies may be the main issue needing to be addressed.

Conclusions

The escalating current and predicted world-wide prevalence of T2D underscores the need to develop effective strategy for early detection, prevention and targeted treatments. In this regard the field of genetic epidemiology has potential to progress to effective translation to patient care possibly within the next decade [178]. Current stateof-the-art approaches that may lead to this goal include defining the pathogenesis of T2D and the phasic nature of the pre-diabetes phenotype beginning in utero and progressing to overt T2D associated with β-cell failure. Thus far, genes identified by GWAS using diagnosed T2D as the phenotype have mainly been associated with β -cell failure but have only accounted for less than 5% of the heritability. Although the greater than 60% heritability appears clinically obvious, environmental interactions, biochemical DNA modifications (epigenetic), and the combined effect of rare variants and other genes present potential challenges in defining risk association. However, the contribution of GWAS over the past decade should not be minimized because of low contribution to heritability, since the genes showing association have been pointers to key metabolic pathways involved in pathogenesis and in the action of drugs used for T2D treatment. Future functional studies on model organisms or cell lines will enable functional characterization of identified risk loci and will provide better understanding of the biological mechanisms of disease development. Improved SNP- and expression-array technology, sequencing, and targeted re-sequencing are likely to enhance current understanding and progress. In addition, inclusion of "at risk" populations in the developing world is likely to improve understanding of gene-environment interaction and is more likely to lead to detection of rare variants and novel variants that associate with unique phenotypes and disease associations.

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