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Large-scale study reveals genetic risk of diabetes in the Japanese population



Manhattan plot of genome-wide association results of the meta-analysis in 36,614 cases and 155,150 controls. Association signals that reached genome-wide significance (P^{-8}) are shown in green if novel and blue if previously reported. Credit: Suzuki et al. (2019) *Nature Genetics*

The genetic and genomic revolutions have led to an abundance of data about the genetic factors that confer a predisposition to type 2 diabetes (T2D), alongside environmental and lifestyle-related causes. However, most of the studies were based on individuals of European descent, meaning that the findings, and any treatments based on them, may not be optimal for other ethnic groups.

A new study performed by researchers from Osaka University, The



University of Tokyo, RIKEN, and others, and published in the journal *Nature Genetics* has shed more light on the genetics of <u>diabetes</u> in the Japanese population by analyzing data on over 36,000 sufferers of T2D and over 150,000 controls of Japanese ancestry. Their work has revealed 28 novel <u>genomic regions</u> associated with T2D, as well as related molecular pathways and cells, with some of these associations being specific to individuals of Japanese descent.

The team conducted a type of study called a <u>meta-analysis</u>, which involves combining the data from a number of independent studies in order to increase the amount of available data and thus the statistical power. As shown by this new study, this approach can potentially provide novel findings that aren't unearthed by constitutive studies, each with a smaller sample size.

"We incorporated the data on links between type 2 diabetes and over 12 million variants across the whole genome from four different genomewide association studies in the Japanese population," Yukinori Okada says. "We found 88 genomic regions significantly associated with this disease, including 28 new ones, some of which are not found in European populations in previous reports."





Regional association plots of GLP1R region in (a) the Japanese (n = 191,764) and (b) the European (n = 159,208) T2D GWAS. $-\log_{10}(P \text{ value})$ is shown on the y axis, and variant position appears on the x axis. Each variant is colored according to the strength of LD (r^2) with the reference variant (colored in purple). (c) A three-dimensional ribbon model of GLP1R, as viewed parallel to the cell membrane. (d) A snake plot of GLP1R. Credit: Suzuki et al. (2019) *Nature Genetics*



The group then looked in more detail at the identified genes, the effects of mutations on them and the proteins they encode, along with the associated pathways and cells. Examples of <u>genetic factors</u> linked to T2D include mutations in the GLP1R, involved in glucose-dependent insulin secretion, and in the genes CPA1 and GP2, known to help certain pancreatic cells transform into insulin-producing beta cells. Although the mutant forms of these genes were linked to T2D in this study, these mutations actually do not exist or are extremely rare in previous reports on European populations.

"Our findings indicate that some of the genetic underpinnings and molecular pathways of type 2 diabetes in the Japanese <u>population</u> may differ from those in European populations, which is unsurprising considering that, when comparing individuals of the same body mass index, Japanese are more prone to this disease," lead author Ken Suzuki says. "Our work could lead to Japanese- or Asian-specific therapeutic measures being developed to more effectively prevent or treat diabetes in this ethnic group."



Database	Pathway name	FDR-q of pathways	
		Japanese	Europeans
Pathways with FDR- <i>q</i> < 0.05 in both Japanese and Europeans			
KEGG	Maturity onset diabetes of the young	6.3 × 10 ⁻¹⁰	1.2 × 10 ⁻³
REACTOME	Regulation of beta cell development	6.1 × 10 ⁻⁶	0.028
REACTOME	Developmental biology	8.6 × 10 ⁻³	0.016
KEGG	Prostate cancer	0.016	0.031
REACTOME	G1 phase	0.046	0.043
Pathways with FDR- <i>q</i> < 0.05 only in Japanese			
REACTOME	Regulation of gene expression in beta cells	4.0 × 10 ⁻⁴	0.084
REACTOME	Signaling by NOTCH	7.6 × 10 ⁻³	0.17
REACTOME	Integration of energy metabolism	8.6 × 10 ⁻³	0.41
REACTOME	NOTCH1 intracellular domain regulates transcription	8.6 × 10 ⁻³	0.41
REACTOME	Regulation of insulin secretion	8.6 × 10 ⁻³	0.29
REACTOME	Negative regulation of FGFR signaling	0.010	0.48
KEGG	Phosphatidylinositol signaling system	0.011	0.54
KEGG	Chronic myeloid leukemia	0.022	0.18
REACTOME	Signaling by NOTCH1	0.022	0.61
KEGG	Pathways in cancer	0.046	0.16
REACTOME	Cell cycle mitotic	0.046	0.10
KEGG	NOTCH signaling pathway	0.047	0.27
Pathways with	h FDR- <i>q</i> < 0.05 only in Europeans		
KEGG	Type II diabetes mellitus	0.11	0.011
KEGG	ABC transporters	0.70	0.028
REACTOME	Pre NOTCH transcription and translation	0.20	0.028
REACTOME	Fatty acid triacylglycerol and ketone body metabolism	0.26	0.031
REACTOME	Pre NOTCH expression and processing	0.17	0.031
REACTOME	Diabetes pathways	0.24	0.037
REACTOME	PPARA activates gene expression	0.30	0.043
KEGG	Tight junction	0.72	0.050

Pathway analysis of T2D GWAS results for Japanese and Europeans. Credit: The University of Tokyo



More information: Identification of 28 new susceptibility loci for type 2 diabetes in the Japanese population, *Nature Genetics* (2019). <u>DOI:</u> 10.1038/s41588-018-0332-4, www.nature.com/articles/s41588-018-0332-4

Provided by Osaka University

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