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Citation for published version:

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GIANT Consortium, MAGIC Investigators, Voight, BF, Scott, LJ, Steinthorsdottir, V, Morris, AP, Dina, C, Welch, RP, Zeggini, E, Huth, C, Aulchenko, YS, Thorleifsson, G, McCulloch, LJ, Ferreira, T, Grallert, H, Amin, N, Wu, G, Willer, CJ, Raychaudhuri, S, McCarroll, SA, Langenberg, C, Hofmann, OM, Dupuis, J, Qi, L, Segre, AV, van Hoek, M, Navarro, P, Ardlie, K, Balkau, B, Benediktsson, R, Bennett, AJ, Blagieva, R, Boerwinkle, E, Bonnycastle, LL, Bostroem, KB, Bravenboer, B, Bumpstead, S, Burtt, NP, Charpentier, G, Chines, PS, Cornelis, M, Couper, DJ, Crawford, G, Doney, ASF, Elliott, KS, Elliott, AL, Erdos, MR, Fox, CS, Morris, A, Rudan, I, Campbell, H & Wilson, JF 2011, 'Twelve type 2 diabetes susceptibility loci identified through large-scale association analysis (vol 42, pg 579, 2010)', *Nature Genetics*, vol. 43, no. 4, pp. 388-388, https://doi.org/10.1038/ng0411-388b 388. https://doi.org/10.1038/ng0411-388b

#### Digital Object Identifier (DOI):

10.1038/ng0411-388b

#### Link:

Link to publication record in Edinburgh Research Explorer

#### **Document Version:**

Peer reviewed version

#### Published In:

**Nature Genetics** 

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## **Europe PMC Funders Group Author Manuscript**

Nat Genet. Author manuscript; available in PMC 2011 April 21.

Published in final edited form as: *Nat Genet.* 2010 July ; 42(7): 579–589. doi:10.1038/ng.609.

# Twelve type 2 diabetes susceptibility loci identified through large-scale association analysis

Benjamin F Voight<sup>1,3,100</sup>, Laura J Scott<sup>4,100</sup>, Valgerdur Steinthorsdottir<sup>5,100</sup>, Andrew P Morris<sup>6,100</sup>, Christian Dina<sup>7,8,100</sup>, Ryan P Welch<sup>9</sup>, Eleftheria Zeggini<sup>6,10</sup>, Cornelia Huth<sup>11,12</sup>, Yurii S Aulchenko<sup>13</sup>, Gudmar Thorleifsson<sup>5</sup>, Laura J McCulloch<sup>14</sup>, Teresa Ferreira<sup>6</sup>, Harald Grallert<sup>11,12</sup>, Najaf Amin<sup>13</sup>, Guanming Wu<sup>15</sup>, Cristen J Willer<sup>4</sup>, Soumya Raychaudhuri<sup>1,2,16</sup>, Steve A McCarroll<sup>1,17</sup>, Claudia Langenberg<sup>18</sup>, Oliver M Hofmann<sup>19</sup>, Josée Dupuis<sup>20,21</sup>, Lu Qi<sup>22,4</sup>, Ayellet V Segrè<sup>1,2,17</sup>, Mandy van Hoek<sup>25</sup>, Pau Navarro<sup>26</sup>, Kristin Ardlie<sup>1</sup>, Beverley Balkau<sup>27,28</sup>, Rafn Benediktsson<sup>29,30</sup>, Amanda J Bennett<sup>14</sup>, Roza Blagieva<sup>31</sup>, Eric Boerwinkle<sup>32</sup>, Lori L Bonnycastle<sup>33</sup>, Kristina Bengtsson Boström<sup>34</sup>, Bert Bravenboer<sup>35</sup>, Suzannah Bumpstead<sup>10</sup>, Noisël P Burtt<sup>1</sup>, Guillaume Charpentier<sup>36</sup>, Peter S Chines<sup>33</sup>, Marilyn Cornelis<sup>24</sup>, David J Couper<sup>37</sup>, Gabe Crawford<sup>1</sup>, Alex S F Doney<sup>38,9</sup>, Katherine S Elliott<sup>6</sup>, Amanda L Elliott<sup>1,17,40</sup>, Michael R Erdos<sup>33</sup>, Caroline S Fox<sup>21,41</sup>, Christopher S Franklin<sup>42</sup>, Martha Ganser<sup>4</sup>, Christian Gieger<sup>11</sup>, Niels Grarup<sup>43</sup>, Todd Green<sup>1,2</sup>, Simon Griffin<sup>18</sup>, Christopher J Groves<sup>14</sup>, Candace Guiducci<sup>1</sup>, Samy Hadjadj<sup>44</sup>, Neelam Hassanali<sup>14</sup>, Christian Herder<sup>45</sup>, Bo Isomaa<sup>46,47</sup>, Anne U Jackson<sup>4</sup>, Paul R V Johnson<sup>48</sup>, Torben Jørgensen<sup>49,50</sup>, Wen H L Kao<sup>51,52</sup>, Norman Klopp<sup>11</sup>, Augustine Kong<sup>5</sup>, Peter Kraft<sup>22,23</sup>, Johanna Kuusisto<sup>53</sup>, Torsten Lauritzen<sup>54</sup>, Man Li<sup>51</sup>, Aloysius Lieverse<sup>55</sup>, Cecilia M Lindgren<sup>6</sup>, Valeriya Lyssenko<sup>56</sup>, Michel Marre<sup>57,58</sup>, Thomas Meitinger<sup>59,60</sup>, Kristian Midthjell<sup>61</sup>, Mario A Morken<sup>33</sup>, Narisu Narisu<sup>33</sup>, Peter Nilsson<sup>56</sup>, Katharine R Owen<sup>14</sup>, Felicity Payne<sup>10</sup>, John R B Perry<sup>62,63</sup>, Ann-Kristin Petersen<sup>11</sup>, Carl Platou<sup>61</sup>, Christine Proença<sup>7</sup>, Inga Prokopenko<sup>6,14</sup>, Wolfgang Rathmann<sup>64</sup>, N William Rayner<sup>6,14</sup>, Neil R Robertson<sup>6,14</sup>, Ghislain Rocheleau<sup>65,67</sup>, Michael Roden<sup>45,68</sup>, Michael J Sampson<sup>69</sup>, Richa Saxena<sup>1,2,40</sup>, Beverley M Shields<sup>62,63</sup>, Peter Shrader<sup>3,70</sup>, Gunnar Sigurdsson<sup>29,30</sup>, Thomas Sparsø<sup>43</sup>, Klaus Strassburger<sup>64</sup>, Heather M Stringham<sup>4</sup>, Qi Sun<sup>22,23</sup>, Amy J Swift<sup>33</sup>, Barbara Thorand<sup>11</sup>, Jean Tichet<sup>71</sup>, Tiinamaija Tuomi<sup>46,72</sup>, Rob M van Dam<sup>24</sup>, Timon W van Haeften<sup>73</sup>, Thijs van Herpt<sup>25,55</sup>, Jana V van Vliet-Ostaptchouk<sup>74</sup>, G Bragi Walters<sup>5</sup>, Michael N Weedon<sup>62,63</sup>, Cisca Wijmenga<sup>75</sup>, Jacqueline Witteman<sup>13</sup>, The MAGIC Investigators<sup>99</sup>, The GIANT Consortium<sup>99</sup>, Richard N Bergman<sup>76</sup>, Stephane Cauchi<sup>7</sup>, Francis S Collins<sup>77</sup>, Anna L Gloyn<sup>14</sup>, Ulf Gyllensten<sup>78</sup>, Torben Hansen<sup>43,79</sup>, Winston A Hide<sup>19</sup>, Graham A Hitman<sup>80</sup>, Albert Hofman<sup>13</sup>, David J Hunter<sup>22,23</sup>, Kristian Hveem<sup>61,81</sup>, Markku Laakso<sup>53</sup>, Karen L Mohlke<sup>82</sup>, Andrew D Morris<sup>38,39</sup>, Colin N A Palmer<sup>38,39</sup>, Peter P Pramstaller<sup>83</sup>, Igor Rudan<sup>42,84,85</sup>, Eric Sijbrands<sup>25</sup>, Lincoln D Stein<sup>15</sup>, Jaakko Tuomilehto<sup>86,88</sup>, Andre Uitterlinden<sup>25</sup>, Mark Walker<sup>89</sup>, Nicholas J Wareham<sup>18</sup>, Richard M Watanabe<sup>76,90</sup>, Gonçalo R Abecasis<sup>4</sup>, Bernhard O Boehm<sup>31</sup>, Harry Campbell<sup>42</sup>, Mark J Daly<sup>1,2</sup>, Andrew T Hattersley<sup>62,63</sup>, Frank B Hu<sup>22,24</sup>, James B Meigs<sup>3,70</sup>, James S Pankow<sup>91</sup>, Oluf Pedersen<sup>43,92,93</sup>, H-Erich Wichmann<sup>11,12,94</sup>, Inês Barroso<sup>10</sup>, Jose C Florez<sup>1,3,95</sup>, Timothy M Frayling<sup>62,63</sup>, Leif Groop<sup>56,72</sup>, Rob Sladek<sup>65,67</sup>, Unnur Thorsteinsdottir<sup>5,96</sup>, James F Wilson<sup>42</sup>, Thomas Illig<sup>11</sup>, Philippe Froguel<sup>17,97</sup>, Cornelia M van Duijn<sup>13</sup>, Kari Stefansson<sup>5,96</sup>, David Altshuler<sup>1,3,17,40,95</sup>, Michael Boehnke<sup>4</sup>, and Mark I McCarthy<sup>6,14,98</sup>

<sup>1</sup>Broad Institute of Harvard and Massachusetts Institute of Technology (MIT), Cambridge, Massachusetts, USA. <sup>2</sup>Center for Human Genetic Research, Massachusetts General Hospital, Boston, Massachusetts, USA. <sup>3</sup>Department of Medicine, Harvard Medical School, Boston, Massachusetts, USA. <sup>4</sup>Department of Biostatistics, University of Michigan, Ann Arbor, Michigan, USA. <sup>5</sup>deCODE Genetics, Reykjavik, Iceland. <sup>6</sup>Wellcome Trust Centre for Human Genetics, University of Oxford, Oxford, UK. <sup>7</sup>CNRS-UMR-8090, Institute of Biology and Lille 2 University,

Pasteur Institute, Lille, France. 8INSERM UMR915 CNRS ERL3147, Nantes, France. <sup>9</sup>Bioinformatics Program, University of Michigan, Ann Arbor, Michigan, USA. <sup>10</sup>Wellcome Trust Sanger Institute, Hinxton, UK. <sup>11</sup>Institute of Epidemiology, Helmholtz Zentrum Muenchen, Neuherberg, Germany. <sup>12</sup>Institute of Medical Informatics, Biometry and Epidemiology, Ludwig-Maximilians-Universität, Munich, Germany. <sup>13</sup>Department of Epidemiology, Erasmus University Medical Center, Rotterdam, The Netherlands. 14Oxford Centre for Diabetes, Endocrinology and Metabolism, University of Oxford, Oxford, UK, <sup>15</sup>Ontario Institute for Cancer Research, Toronto, Ontario, Canada. 16 Division of Rheumatology, Immunology and Allergy, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts, USA. 17 Department of Molecular Biology, Harvard Medical School, Boston, Massachusetts, USA. 18 Medical Research Council (MRC) Epidemiology Unit, Institute of Metabolic Science, Addenbrooke's Hospital, Cambridge, UK. <sup>19</sup>Department of Biostatistics, Harvard School of Public Health, Boston, Massachusetts, USA. <sup>20</sup>Department of Biostatistics, Boston University School of Public Health, Boston, Massachusetts, USA. <sup>21</sup>National Heart, Lung, and Blood Institute's Framingham Heart Study, Framingham, Massachusetts, USA. <sup>22</sup>Department of Nutrition, Harvard School of Public Health, Boston, Massachusetts, USA. <sup>23</sup>Department of Epidemiology, Harvard School of Public Health, Boston, Massachusetts, USA. <sup>24</sup>Channing Laboratory, Department of Medicine, Brigham and Women's Hospital and Harvard Medical School, Boston, Massachusetts, USA, <sup>25</sup>Department of Internal Medicine, Erasmus University Medical Centre, Rotterdam, The Netherlands. <sup>26</sup>MRC Human Genetics Unit. Institute of Genetics and Molecular Medicine. Western General Hospital. Edinburgh, UK. <sup>27</sup>INSERM, CESP Centre for Research in Epidemiology and Population Health, U1018, Epidemiology of Diabetes, Obesity and Chronic Kidney Disease over the Lifecourse, Villejuif, France. <sup>28</sup>University Paris-Sud 11, UMRS 1018, Villejuif, France. <sup>29</sup>Landspitali University Hospital, Reykjavik, Iceland. 30 Icelandic Heart Association, Kopavogur, Iceland. 31 Division of Endocrinology, Diabetes and Metabolism, Ulm University, Ulm, Germany. 32The Human Genetics Center and Institute of Molecular Medicine, University of Texas Health Science Center, Houston, Texas, USA. 33National Human Genome Research Institute, National Institute of Health, Bethesda, Maryland, USA. 34Research and Development Centre, Skaraborg Primary Care, Sköyde, Sweden, 35Department of Internal Medicine, Catharina Hospital, Eindhoven, The Netherlands. <sup>36</sup>Endocrinology-Diabetology Unit, Corbeil-Essonnes Hospital, Corbeil-Essonnes, France. <sup>37</sup>Department of Biostatistics and Collaborative Studies Coordinating Center, University of North Carolina at Chapel Hill, Chapel Hill, North Carolina, USA. 38 Diabetes Research Centre, Biomedical Research Institute, University of Dundee, Ninewells Hospital, Dundee. UK. <sup>39</sup>Pharmacogenomics Centre, Biomedical Research Institute, University of Dundee, Ninewells Hospital, Dundee, UK. <sup>40</sup>Department of Genetics, Harvard Medical School, Boston. Massachusetts, USA. 41 Division of Endocrinology, Diabetes, and Hypertension, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts, USA. 42Centre for Population Health Sciences, University of Edinburgh, Edinburgh, UK. 43 Hagedorn Research Institute, Gentofte, Denmark. 44Centre Hospitalier Universitaire de Poitiers, Endocrinologie Diabetologie, CIC INSERM 0801, INSERM U927, Université de Poitiers, UFR, Médecine Pharmacie, Poitiers Cedex, France. <sup>45</sup>Institute for Clinical Diabetology, German Diabetes Center, Leibniz Center for Diabetes Research at Heinrich Heine University Düsseldorf, Düsseldorf, Germany. <sup>46</sup>Folkhälsan Research Center, Helsinki, Finland. <sup>47</sup>Malmska Municipal Health Center and Hospital, Jakobstad, Finland. 48 Diabetes Research and Wellness Foundation Human Islet Isolation Facility and Oxford Islet Transplant Programme, University of Oxford, Oxford, UK. <sup>49</sup>Research Centre for Prevention and Health, Glostrup University Hospital, Glostrup, Denmark. <sup>50</sup>Faculty of Health Science, University of Copenhagen, Copenhagen, Denmark. <sup>51</sup>Department of Epidemiology, Johns Hopkins University, Baltimore, Maryland, USA. 52Department of Medicine and Welch Center for Prevention, Epidemiology and Clinical Research, Johns Hopkins University, Baltimore, Maryland, USA. 53Department of Medicine, University of Kuopio and Kuopio University Hospital, Kuopio, Finland. 54Department of General Medical Practice, University of Aarhus,

Aarhus, Denmark. 55Department of Internal Medicine, Maxima Medical Center, Eindhoven, The Netherlands. <sup>56</sup>Department of Clinical Sciences, Diabetes and Endocrinology Research Unit, University Hospital Malmö, Lund University, Malmö, Sweden. <sup>57</sup>Department of Endocrinology, Diabetology and Nutrition, Bichat-Claude Bernard University Hospital, Assistance Publique des Hôpitaux de Paris, Paris, France. 58 INSERM U695, Université Paris 7, Paris, France. 59 Institute of Human Genetics, Helmholtz Zentrum Muenchen, Neuherberg, Germany. 60 Institute of Human Genetics, Klinikum rechts der Isar, Technische Universität München, München, Germany, 61Nord-Trøndelag Health Study (HUNT) Research Center, Department of Community Medicine and General Practice, Norwegian University of Science and Technology, Trondheim, Norway. <sup>62</sup>Genetics of Complex Traits, Institute of Biomedical and Clinical Science, Peninsula Medical School, University of Exeter, Exeter, UK. 63Diabetes Genetics, Institute of Biomedical and Clinical Science, Peninsula Medical School, University of Exeter, Exeter, UK. 64Institute of Biometrics and Epidemiology, German Diabetes Center, Leibniz Center for Diabetes Research at Heinrich Heine University Düsseldorf, Düsseldorf, Germany. 65 Department of Human Genetics, McGill University, Montreal, Canada. 66Department of Medicine, Faculty of Medicine, McGill University, Montreal, Canada. <sup>67</sup>McGill University and Genome Quebec Innovation Centre, Montreal, Canada. <sup>68</sup>Department of Metabolic Diseases, Heinrich Heine University Düsseldorf, Düsseldorf, Germany. <sup>69</sup>Department of Endocrinology and Diabetes. Norfolk and Norwich University Hospital National Health Service Trust, Norwich, UK. 70General Medicine Division, Massachusetts General Hospital, Boston, Massachusetts, USA. 71 Institut interrégional pour la Santé (IRSA), La Riche, France. 72 Department of Medicine, Helsinki University Hospital, University of Helsinki, Helsinki, Finland. 73Department of Internal Medicine, University Medical Center Utrecht, Utrecht, The Netherlands. 74Molecular Genetics, Medical Biology Section, Department of Pathology and Medical Biology, University Medical Center Groningen and University of Groningen, Groningen, The Netherlands. 75 Department of Genetics, University Medical Center Groningen and University of Groningen, Groningen, The Netherlands. 76 Department of Physiology and Biophysics, University of Southern California School of Medicine, Los Angeles, California, USA. 77 National Institute of Health, Bethesda, Maryland, USA. 78Department of Genetics and Pathology, Rudbeck Laboratory, Uppsala University, Uppsala, Sweden, <sup>79</sup>University of Southern Denmark, Odense, Denmark. 80 Centre for Diabetes, Barts and The London School of Medicine and Dentistry, Queen Mary University of London, London, UK. 81 Department of Medicine, The Hospital of Levanger, Levanger, Norway. 82 Department of Genetics, University of North Carolina, Chapel Hill, North Carolina, USA. 83Institute of Genetic Medicine, European Academy Bozen/Bolzano (EURAC), Bolzano, Italy. 84Croatian Centre for Global Health, Faculty of Medicine, University of Split, Split, Croatia, 85 Institute for Clinical Medical Research, University Hospital 'Sestre Milosrdnice', Zagreb. Croatia. 86Department of Public Health, University of Helsinki, Helsinki, Finland. 87South Ostrobothnia Central Hospital, Seinäjoki, Finland. 88 Red RECAVA Grupo RD06/0014/0015, Hospital Universitario La Paz, Madrid, Spain. 89 Diabetes Research Group, Institute of Cellular Medicine, Newcastle University, Newcastle upon Tyne, UK. 90 Department of Preventative Medicine, Keck Medical School, University of Southern California, Los Angeles, California, USA, <sup>91</sup>Division of Epidemiology and Community Health, University of Minnesota, Minneapolis, Minnesota, USA, 92Department of Biomedical Science, Panum, Faculty of Health Science. University of Copenhagen, Copenhagen, Denmark. 93 Faculty of Health Science, University of Aarhus, Aarhus, Denmark. 94Klinikum Grosshadern, Munich, Germany. 95Diabetes Unit, Massachusetts General Hospital, Boston, Massachusetts, USA. 96Faculty of Medicine, University of Iceland, Reykjavík, Iceland. 97Genomic Medicine, Imperial College London, Hammersmith Hospital, London, UK. 98Oxford National Institute for Health Research Biomedical Research Centre, Churchill Hospital, Oxford, UK.

#### Abstract

By combining genome-wide association data from 8,130 individuals with type 2 diabetes (T2D) and 38,987 controls of European descent and following up previously unidentified meta-analysis signals in a further 34,412 cases and 59,925 controls, we identified 12 new T2D association signals with combined  $P < 5 \times 10^{-8}$ . These include a second independent signal at the *KCNQ1* locus; the first report, to our knowledge, of an X-chromosomal association (near *DUSP9*); and a further instance of overlap between loci implicated in monogenic and multifactorial forms of diabetes (at *HNF1A*). The identified loci affect both beta-cell function and insulin action, and, overall, T2D association signals show evidence of enrichment for genes involved in cell cycle regulation. We also show that a high proportion of T2D susceptibility loci harbor independent association signals influencing apparently unrelated complex traits.

Type 2 diabetes (T2D) is characterized by insulin resistance and deficient beta-cell function<sup>1</sup>. The escalating prevalence of T2D and the limitations of currently available preventative and therapeutic options highlight the need for a more complete understanding of T2D pathogenesis. To date, approximately 25 genome-wide significant common variant associations with T2D have been described, mostly through genome-wide association (GWA) analyses<sup>2</sup>-<sup>13</sup>. The identities of the variants and genes mediating the susceptibility effects at most of these signals have yet to be established, and the known variants account for less than 10% of the overall estimated genetic contribution to T2D predisposition. Although some of the unexplained heritability will reflect variants poorly captured by existing GWA platforms, we reasoned that an expanded meta-analysis of existing GWA data would offer augmented power to detect additional common variant signals of modest effect.

#### **RESULTS**

#### **GWA** meta-analysis and replication

We conducted a meta-analysis of eight T2D GWA studies comprising 8,130 T2D cases and 38,987 controls of European descent. We combined case-control data from the Wellcome Trust Case Control Consortium (WTCCC), Diabetes Genetics Initiative (DGI) and Finland-US Investigation of NIDDM genetics (FUSION) scans (the subjects of a previous joint analysis<sup>7</sup>), with those from scans performed by deCODE genetics<sup>6</sup>, the Diabetes Gene Discovery Group<sup>2</sup>, the Cooperative Health Research in the Region of Augsburg group (KORAgen), the Rotterdam study and the European Special Population Research Network (EUROSPAN). The effective sample size (n = 22,044) of stage 1 of the current (hereafter designated 'DIAGRAM+') meta-analysis was more than twice that of the earlier DIAGRAM (DIAbetes Genetics Replication and Meta-analysis) study<sup>7</sup>. After genomic control correction of each component study, we combined association data for 2,426,886 imputed and genotyped autosomal SNPs into a fixed-effects, additive-model meta-analysis using the inverse-variance method (Online Methods, Fig. 1, Supplementary Tables 1 and 2 and Supplementary Note). We observed only modest genomic control inflation ( $\lambda_{gc} = 1.07$ ), suggesting that the observed results were not due to population stratification. After removing SNPs within established T2D loci (Supplementary Table 3), the resulting quantile-quantile plot was consistent with a modest excess of disease associations of relatively small effect (Supplementary Note). Weak evidence for association at HLA variants strongly associated with autoimmune forms of diabetes (Supplementary Table 3 and Supplementary Note) suggested some case admixture involving subjects with type 1 diabetes or latent autoimmune diabetes of adult-hood; however, failure to detect T2D associations at other non-HLA type 1 diabetes susceptibility loci (for example, INS, PTPN22 and IL2RA) indicated that any such misclassification was too modest to drive stage 1 associations outside the HLA. The stage 1 meta-analysis also provided further confirmation of many previously reported signals and, at some of these, refinement of the peak association signal (Fig. 1, Supplementary Table 3 and Supplementary Note).

We selected for stage 2 follow-up the most strongly associated SNP from each of the 23 new autosomal regions showing the most compelling evidence for association (all  $P < 10^{-5}$  in stage 1; Supplementary Table 3). We combined exclusively *in silico* data from three GWA samples (Atherosclerosis Risk in Communities (ARIC) study, Nurses' Health Study and Framingham Heart Study) not included in the primary meta-analysis (2,832 cases and 15,843 controls) with additional (predominantly *de novo*) genotyping in up to 31,580 cases and 44,082 controls, for a maximum possible stage 2 sample size of 34,412 cases and 59,925 controls (effective sample size of 79,246), all of European descent (**Supplementary Tables** 1 and 2).

Stage 2 analyses indicated that the set of 23 signals was enriched for true association signals. In all, 21 showed directional consistency of effect between stage 1 and 2 (binomial test,  $P \sim 3.3 \times 10^{-5}$ ), and for 15, the stage 2 P value was <0.05 (Supplementary Note). In joint analysis of stage 1 and 2 data (up to 42,542 cases and 98,912 controls), 13 autosomal loci exceeded the threshold for genome-wide significance (P ranging from  $2.8 \times 10^{-8}$  to  $1.4 \times 10^{-22}$ ) with allele-specific odds ratios (ORs) between 1.06 and 1.14 (Table 1 and Fig. 2). All signals remained close to or beyond genome-wide significance thresholds (the least significant P value was  $5.2 \times 10^{-8}$ ) when we repeated analyses after implementing a second (post meta-analysis) round of genomic control adjustment within stage 1 data (Supplementary Note).

We extended our search for susceptibility signals to the X chromosome, identifying one further signal in the stage 1 discovery samples meeting our criteria for follow-up (represented by rs5945326, near *DUSP9*,  $P = 2.3 \times 10^{-6}$ ). This SNP showed strong evidence for replication in 8,535 cases and 12,326 controls (OR (allowing for X-inactivation) 1.32 (95% CI 1.16–1.49),  $P = 2.3 \times 10^{-5}$ ), for a combined association P value of  $3.0 \times 10^{-10}$  (OR 1.27 (95% CI 1.18–1.37)) (Table 1 and Fig. 2).

#### Fourteen signals reaching genome-wide significance

Two of the 14 signals reaching genome-wide significance on joint analysis (those near *MTNR1B* and *IRS1*) represent loci for which T2D associations have been recently reported in samples which partially overlap with those studied here<sup>10</sup>, <sup>14</sup>-<sup>16</sup> (Table 1).

A third signal (rs231362) on 11p15 overlaps both intron 11 of *KCNQ1* and the *KCNQ1OT1* transcript that controls regional imprinting  $^{17}$  and influences expression of nearby genes including *CDKN1C*, a known regulator of beta-cell development  $^{18}$ . This signal maps ~150 kb from T2D-associated SNPs in the 3' end of *KCNQ1* first identified in East Asian GWA scans  $^{8}$ ,9. SNPs within the 3' signal were also detected in the current DIAGRAM+ meta-analysis (for example, rs163184,  $P = 6.8 \times 10^{-5}$ ), but they failed to meet the threshold for initiating replication. A SNP in the 3' region (rs2237895) that was reported to reach genome-wide significance in Danish samples  $^{9}$  was neither typed nor imputed in the DIAGRAM+ studies. In our European-descent samples, rs231362 and SNPs in the 3' signal were not correlated ( $r^2 < 0.05$ ), and conditional analyses (see below) establish these SNPs as independent (Fig. 2 and Supplementary Table 4). Further analysis in Icelandic samples has shown that both associations are restricted to the maternally transmitted allele  $^{11}$ . Both T2D loci are independent of the common variant associations with electrocardiographic QT intervals that map at the 5' end of KCNQ1 ( $r^2 < 0.02$ , D' < 0.35 in HapMap European CEU data)  $^{19}$ ,  $^{20}$  (Supplementary Table 5).

Of the remaining loci, two (near *BCL11A* and *HNF1A*) have been highlighted in previous studies<sup>7</sup>,<sup>21</sup>-<sup>23</sup> but are now shown to reach genome-wide significance. Rare mutations in *HNF1A* account for a substantial proportion of cases of maturity onset diabetes of the young, and a population-specific variant (G319S) influences T2D risk in Oji-Cree Indians<sup>24</sup>.

Confirmation of a common variant association at *HNF1A* brings to five the number of loci known to harbor both rare mutations causal for monogenic forms of diabetes and common variants predisposing to multifactorial diabetes, the others being *PPARG*, *KCNJ11*, *WFS1* and *HNF1B*. A T2D association in the *BCL11A* region was suggested by the earlier DIAGRAM meta-analysis (rs10490072,  $P = 3 \times 10^{-5}$ ), but replication was inconclusive<sup>7</sup>; there is only modest linkage disequilibrium (LD) between rs10490072 and the lead SNP from the present analysis (rs243021,  $r^2 = 0.22$ , D' = 0.73 in HapMap CEU).

The remaining nine signals map near the genes *HMGA2*, *CENTD2*, *KLF14*, *PRC1*, *TP53INP1*, *ZBED3*, *ZFAND6*, *CHCHD9* and *DUSP9* (Table 1 and **Figs.** 1 and 2) and represent new T2D risk loci uncovered by the DIAGRAM+ meta-analysis.

#### Understanding the genetic architecture of type 2 diabetes

Combining newly identified and previously reported loci and assuming a multiplicative model, the sibling relative risk attributable to the 32 T2D susceptibility variants described in this paper is ~1.14. With addition of the five T2D loci recently identified by the Meta-Analysis of Glucose and Insulin-related traits Consortium (MAGIC) investigators <sup>12</sup>, <sup>13</sup> and incorporation of estimates of parent-of-origin–specific effect sizes observed at the *KCNQ1* and *KLF14* signals and at a recently described locus on chromosome 11p15 (which confers substantial risk when paternally inherited but is protective when maternally transmitted <sup>11</sup>), this figure rises to ~1.16. Given estimates of sibling relative risk for T2D in Europeans of ~3 (ref. <sup>25</sup>), variant discovery efforts to date have therefore explained only ~10% of observed familial clustering. We used available data to evaluate several mechanisms that might be contributing to that proportion of familiality which reflects residual, unexplained heritability<sup>26</sup>.

**Copy number variants (CNVs)**—We re-examined stage 1 data looking for associations with SNPs known to provide robust, high-LD tags for common CNVs in European populations. After combining four inventories of CNV-tagging SNPs that survey at least 40% of common CNVs genome-wide >1 kb in size, we found no convincing evidence that this class of variants contributes substantially to T2D risk (Supplementary Note).

**Secondary signals revealed by conditional analysis**—If there are additional independent susceptibility variants at the loci identified, total genetic variance attributable to these regions will be underestimated when based on the lead common variants alone. To explore the potential for independent secondary alleles, we repeated the stage 1 meta-analysis after simultaneously conditioning on 30 known and newly discovered autosomal loci (Supplementary Note). Using a cutoff of  $P < 1 \times 10^{-4}$  (to reflect approximate adjustment for the number of independent SNPs in a ~2 Mb interval), we found preliminary evidence for secondary signals at five loci (TP53INP1, CDKN2A, HHEX-IDE and KCNJ11, in addition to that at KCNQ1; Fig. 1, Supplementary Fig. 1 and Supplementary Table 4). At CDKN2A, the secondary signal is consistent with evidence that haplotype-based analyses generate considerably stronger evidence for association than either signal alone<sup>3</sup>, <sup>27</sup>. Further fine-mapping efforts will be required to confirm the secondary signals at TP53INP1, HHEX-IDE and KCNJ11.

The conditional analysis also provided preliminary ( $P < 10^{-5}$ , our stage 1 threshold) evidence for 19 signals outside known loci (Fig. 1 and Supplementary Table 6). The most notable signal (rs1481279, conditional  $P = 8.4 \times 10^{-9}$ ) maps near *NHEDC1* and corresponds to one of the signals of interest following stage 1 (rs7674212,  $P = 1.7 \times 10^{-7}$  unconditioned). Failure to replicate that signal (P = 0.3 in 21,889 stage 2 cases and 39,568 controls) suggests this was a false positive (Supplementary Note). Several regions showed substantial

incremental evidence for association in the conditional analysis as compared to unadjusted analyses and represent potential targets for large-scale replication and gene-gene interaction analyses. Indeed, one of these regions (at rs11708067 near *ADCY5*, unadjusted  $P=1.7\times10^{-4}$ , conditional  $P=2.2\times10^{-6}$ ) has recently been shown, following initial identification through GWA analysis of continuous glucose measures, to have genome-wide significant associations with T2D in large-scale case-control analyses that involved several DIAGRAM + samples  $^{12}, ^{13}$ .

Etiological heterogeneity—To determine whether etiological heterogeneity might have compromised power to detect genuine T2D susceptibility signals, we performed BMI- and age-of-diagnosis (AOD)-stratified analyses within stage 1 data. We compared effect size estimates for all known T2D risk variants in 2,877 obese (defined as BMI > 30 kgm<sup>-2</sup>) and 4,048 nonobese (BMI 30 kgm<sup>-2</sup>) T2D cases when compared to similarly stratified controls (Supplementary Note). Although risk estimates for 23 of the 30 autosomal loci were numerically greater in the nonobese comparison than in the obese comparison (binomial test, P = 0.0018), only TCF7L2 (P < 0.001) and BCL11A (P = 0.02) showed significant (P < 0.001) 0.05) evidence for effect-size heterogeneity. For AOD, we compared risk-locus genotypes for 1,317 cases with AOD <45 years of age and 4,283 cases with AOD >45 years of age, as well as continuous analyses of AOD within all cases (n = 7,104; Supplementary Note), and found no strong evidence of differential effects. Although recognizing that BMI at examination and AOD are imperfect measures of BMI and age at disease onset, we conclude that a focus on more homogeneous subsamples would not have provided more efficient identification of known T2D susceptibility variants. Furthermore, these data argue against the potential for these common variant signals to afford clinically useful subclassification of individuals with T2D.

Overlap with GWA signals for other diseases—We noted that seven of the newly discovered autosomal loci (near BCL11A, ZBED3, KLF14, CHCHD9, HMGA2, HNF1A and *PRC1*) are characterized by strong ( $P < 10^{-6}$ ) associations with phenotypes other than T2D (Supplementary Table 5). In each case, these appear to be distinct and independent signals. For example, variants at the 3' end of HMGA2 (~180 kb distant from the T2D signal) have widely replicated effects on adult height<sup>28</sup> but are weakly correlated with the T2D-associated SNP rs1531343 ( $r^2 < 0.01$ , D' < 0.15 in HapMap CEU). The KLF14 region harbors distinct signals for both T2D and basal cell carcinoma<sup>29</sup>. At *HNF1A*, previous studies have reported a cluster of associations, with phenotypes including low-density lipoptrotein (LDL) cholesterol<sup>30</sup> and circulating C-reactive protein levels<sup>31</sup>-<sup>33</sup>, mapping ~18–72 kb from the peak T2D signal. Though these two sets of *HNF1A* signals maintain appreciable LD in European samples ( $r^2 \sim 0.1$ ,  $D' \sim 1$ ), they are likely to be independent; the T2D association at the lead SNP for lipids (rs2650000) is far weaker than the association at rs7957197 (P = 0.003 compared to  $P = 4.6 \times 10^{-7}$  in stage 1 samples), whereas LDL cholesterol shows a reciprocal pattern of association ( $P = 7 \times 10^{-9}$  at rs2650000 compared to P = 0.73 at rs7957197 in the same lipid meta-analysis data<sup>30</sup>).

If we include the *KCNQ1* associations described above, previous reports at *JAZF1*, *CDKN2A* and *CDKAL1* (refs. <sup>34</sup>-<sup>40</sup>) and other signals identified by systematic analysis of the National Human Genome Research Institute (NHGRI) GWA catalog<sup>41</sup> (Supplementary Table 5 and Supplementary Note), at least 13 of 30 autosomal T2D loci show this pattern of closely approximated (within 500 kb) but distinct associations with traits other than T2D or related anthropometric and glycemic phenotypes. This is in addition to what appear to be coincident signals involving T2D susceptibility variants at *IRS1* (associated with coronary disease), *JAZF1* (associated with height) and *HNF1B* (associated with prostate cancer) (Supplementary Table 5). Simulations conducted using the NHGRI catalog as a reference set indicate that the number of non-T2D signals observed at T2D loci significantly exceeds

expectation ( $P \sim 1.6 \times 10^{-3}$  for non-T2D signals within 500 kb of T2D loci,  $P \sim 7.0 \times 10^{-5}$  (n = 8) for non-T2D signals within 100 kb of T2D loci). Many of these instances of colocalization may represent variants within different regulatory domains that result in tissue- and disease-specific effects mediated through the same genes and pathways.

#### Understanding the biology of T2D-susceptibility loci

This analysis takes the number of independent loci showing genome-wide significant associations with T2D beyond 35. For some, such as those at *KCNJ11* and *SLC30A8*, the molecular mechanisms responsible for the susceptibility effect can be assigned with some confidence<sup>42</sup>. At others, the identities of the causal variants, the genes through which they act and the pathophysiological processes which they influence remain obscure. We used several approaches designed to link DIAGRAM+ and previously reported T2D association signals to biological insights relevant to T2D pathogenesis.

**Physiological analyses**—Variants at FTO are known to influence T2D predisposition through an effect on BMI. In ~21,000 population sample individuals from the GWA meta-analysis of adult BMI completed by the Genetic Investigation of ANthropmetric Traits (GIANT) consortium<sup>43</sup>, no other autosomal T2D susceptibility locus had the property that the T2D risk allele was significantly associated with higher BMI (Supplementary Note). FTO is therefore the only one of the known T2D signals driven by a strong primary causal association with obesity.

We also examined the effect of T2D susceptibility alleles on continuous glycemic measures in up to 46,186 nondiabetic subjects from the MAGIC meta-analysis <sup>12</sup>, <sup>13</sup>. Coefficients for association between the T2D risk allele and higher fasting glucose were positive for 28 of the 31 loci, and 17 of these T2D loci showed significant (P < 0.05) directionally consistent associations with fasting glucose (Fig. 3 and Supplementary Note). However, the magnitudes of effect sizes for fasting glucose and T2D were only weakly correlated (Supplementary Fig. 2 and Supplementary Note), indicating that the mechanisms influencing normal glucose homeostasis and those responsible for the development of T2D are not entirely congruent. T2D risk alleles at four loci (at PPARG, FTO, IRS1 and KLF14) were associated (P < 0.05) with higher fasting insulin, consistent with a primary effect on insulin action, whereas at three other loci (at TCF7L2, CENTD2 and CDKAL1), the association with reduced fasting insulin indicates beta-cell dysfunction (Fig. 3). Indices of beta-cell function (HOMA-B) and insulin sensitivity (HOMA-IR) derived from paired fasting glucose and insulin measures from ~37,000 individuals supported these mechanistic inferences (Fig. 3). In all, risk alleles at ten loci (the previously reported loci at MTNR1B, SLC30A8, THADA, TCF7L2, KCNQ1, CAMK1D, CDKAL1, IGF2BP2 and HNF1B and the newly discovered locus at CENTD2) were associated (P < 0.05) with reduced beta-cell function, and three loci (previously reported loci at PPARG and FTO and the newly discovered locus at KLF14) were associated with reduced insulin sensitivity. The associations with improved insulin sensitivity evident for risk alleles at TCF7L2, IGF2BP2 and CDKAL1 probably reflect truncated ascertainment, as the MAGIC analyses were restricted to nondiabetic individuals. For the previously reported loci, these findings are broadly consistent with those from more detailed physiological studies<sup>6</sup>, <sup>8</sup>, <sup>44</sup> and suggest that, of the newly discovered loci, the risk alleles at CENTD2 modify T2D susceptibility through a detrimental effect on beta-cell function. In contrast, the risk alleles at KLF14 and possibly HMGA2 (ref. 45), along with those at PPARG, IRS1 (ref. 10) and ADAMTS9 (ref. <sup>46</sup>), appear to have a primary effect on insulin action which is not driven by obesity, unlike the alleles at FTO. The MAGIC meta-analysis did not extend to the X chromosome, but analysis of rs5945326, near *DUSP9*, in a sub-set of MAGIC samples (n = 14,644-21,118), revealed no significant (P < 0.05) associations with any fasting glycemic trait. For this

signal, as with the other newly identified loci, more detailed phenotypic analyses will be required to determine how these impact T2D risk. Overall, these data are consistent with the impression that common T2D risk alleles more often act through beta-cell dysfunction <sup>12</sup>, <sup>44</sup>, but they provide further examples of T2D risk variants that exert their primary effects on insulin action.

**Expression analyses—**We used expression data to seek clues to the genes mediating the T2D susceptibility effects we had detected. First, we examined expression-QTL (eQTL) data (in 23,720 transcripts) for subcutaneous adipose tissue (n = 603 with GWA data) and blood (n = 745) samples typed with the Illumina 300K chip<sup>47</sup> (Table 2 and Supplementary Note). Among the newly identified loci, the most compelling signal was at rs972283, strongly associated with expression of KLF14 in adipose tissue and correlated ( $r^2 = 0.3$  in HapMap CEU) with the SNP (rs738134) with the strongest KLF14 cis expression signal. Both the T2D and cis eQTL associations at this locus showed strong parent-of-origin effects<sup>11</sup>. At the TP53INP1 locus, the cis-eQTL data suggest the T2D susceptibility effect is mediated via altered CCNE2 expression. In contrast, the significant cis-eQTL associations at the ZBED3, CENTD2, HNF1A and PRC1 T2D susceptibility signals are likely to be misleading, as the patterns of conditional association indicate that the T2D association and cis eQTL signals are not coincident. At previously reported T2D association signals, we found strong overlap with cis eQTL effects for IRS1 (consistent with data on IRS1 protein expression and function in skeletal muscle<sup>10</sup>), JAZF1 and  $CAMK1D^7$ .

We also explored the tissue expression profiles of 27 autosomal genes mapping to the newly discovered regions of association and performed quantitative RT-PCR analyses across a panel of human tissues relevant to T2D pathogenesis (Supplementary Note). The broad expression of many of the transcripts, including 24 transcripts with evidence of beta-cell transcription (Supplementary Note), limited our ability to prioritize among candidate transcripts on the basis of static patterns of transcript expression.

Pathway and protein-protein interaction analyses—Reasoning that the additional T2D susceptibility loci would amplify our ability to identify over-represented molecular processes<sup>48</sup>, we deployed several complementary approaches to detect evidence of pathway or network enrichment (Supplementary Note). Using GRAIL<sup>49</sup>, we found that genes within T2D-associated regions showed evidence of increased connectivity within PubMed abstracts, though this largely reflects shared roles in monogenic or syndromic diabetes (involving *HNF1A*, *HNF1B* and *WFS1*). We also showed that the extent of protein-protein interaction between the products of genes mapping to the association signals substantially exceeded expectation (Supplementary Note). Pathway enrichment analyses using the PANTHER database<sup>50</sup> uncovered some evidence of over-representation of signal transduction and protein metabolism and modification, and Reactome<sup>51</sup> highlighted a separate set of pathways including metabolism of lipids and lipoproteins, endothelins and beta-arrestins (for details, see Supplementary Note).

The only consistent signal to emerge across multiple analyses involved cell-cycle regulation. Network analyses based on protein-protein interaction data detected (unadjusted  $P \sim 0.004$ ) an 18-member subnetwork characterized by enhanced protein-protein interaction connectivity and highly enriched for genes implicated in cell cycle regulation ( $P = 2.8 \times 10^{-7}$ ). A smaller (five, only partly overlapping genes) cell-cycle network independently emerged from the Reactome analyses, and gene-set enrichment analysis of selected candidate pathways<sup>52</sup> also detected over-representation of association signals ( $P \sim 0.006$ ) among cell-cycle genes (Supplementary Note). Because many genes within these networks are expressed in pancreatic islets and T2D-association effects at several of these loci are

mediated primarily through beta-cell dysfunction<sup>44</sup>, these findings highlight the contribution of regulation of beta-cell mass to the long-term maintenance of normal glucose homeostasis.

In addition, these analyses highlighted notable biological connections between sets of genes within confirmed T2D-association regions. For example, *HMGA2* emerges as a key transcriptional regulator of *IGF2BP2* (refs. <sup>53</sup>,<sup>54</sup>). However, because *Hmga/Hmg1c* knockout mice are deficient in adipocyte differentiation <sup>45</sup>, and the *IGF2BP2* risk allele is associated with reduced beta-cell function <sup>55</sup>, further work is required to establish the relevance of this regulatory interaction to T2D pathogenesis. Our analyses also revealed that *TLE4* (at the *CHCHD9* locus) encodes a homolog of Groucho that forms complexes with TCF proteins, including TCF7L2, to modulate transcription at target sites <sup>56</sup>. Finally, *FURIN*, one of the genes mapping to the newly identified *PRC1* locus, encodes a paired basic amino acid cleaving enzyme; both NOTCH2 and ADAMTS9 (ref. <sup>7</sup>) are known targets of FURIN cleavage <sup>57</sup>-<sup>59</sup>.

Notably, these global approaches failed to provide any consistent support for many other mechanisms previously promoted on the basis of biochemical or physiological evidence as likely contributors to T2D pathogenesis (Supplementary Note). Overall, the relative paucity of signals from these analyses—particularly when contrasted with the compelling patterns of enrichment seen for other complex traits —indicates, either that T2D pathogenesis is characterized by substantial etiological heterogeneity or that the processes critical to T2D development are poorly represented in existing pathway and interaction databases.

#### DISCUSSION

By increasing the discovery sample size, our study has substantially expanded the number of loci for which there is strong statistical evidence indicating a role in T2D predisposition. When combined with recent reports of additional T2D susceptibility loci arising from studies of continuous glycemic traits<sup>12</sup>, <sup>13</sup> and parent-of-origin effects<sup>11</sup>, the number of confirmed loci for T2D currently stands at <sup>38</sup>.

Although these discoveries represent new opportunities to explore the biology of T2D predisposition, the challenges inherent in translating these common variant association signals into biological mechanisms of disease causation are clear. Nevertheless, the analyses we report have generated several mechanistic hypotheses that can direct future efforts at functional evaluation and genetic refinement. At some loci, particularly those near HNF1A, HMGA2 and KLF14, existing biology, coupled with phenotypic and expression data presented here, highlight the named genes as prime candidates for mediating the susceptibility effect. For example, the T2D susceptibility effect near KLF14, which maps within an imprinted region on chromosome 7q32 and which, on the basis of the MAGIC meta-analysis data, appears to be driven by reduced insulin action, is restricted to the maternally transmitted allele<sup>11</sup>. As KLF14 is maternally expressed, and the eQTL association between rs972283 and KLF14 expression (see above) is similarly restricted to the maternal allele, KLF14 (a widely expressed, intronless member of the Krüppel-like family of transcription factors<sup>60</sup>) emerges as the main regional candidate. At the Xchromosome signal, evidence implicating DUSP9 (mitogen-activated protein kinase phosphatase-4) in the regulation of insulin action in mice gives DUSP9 particular salience as an association candidate<sup>61</sup>,<sup>62</sup>. However, as described above, failure to detect associations with continuous glycemic phenotypes (including fasting insulin and HOMA-IR) means that the functional connection with *DUSP9* remains speculative.

In other regions, such as those near *PRC1*, *TP53INP1* and *CHCHD9*, the functional connections and/or eQTL associations of particular genes mapping within or close to the

respective association intervals show *FURIN*, *CCNE2* and *TLE4*, respectively, to be promising biological candidates. At yet other loci, such as those centered around *ZBED3*, *CENTD2* and *ZFAND6*, existing data provide little, if any, basis for strong inferences concerning the genes likely to mediate the T2D susceptibility effect. Accumulation of new data—through deep resequencing of the regions, fine-mapping and functional studies in humans and in animal models—will be required to characterize the specific variants responsible and the genes and pathways through which they execute their effect on T2D risk.

One theme emerging from this work is the high frequency with which loci implicated in T2D susceptibility harbor variants that influence other common traits. This colocalization of common risk variants exceeds chance expectation, often connects diseases with little obvious mechanistic overlap and typically involves statistically independent susceptibility signals. Recent evidence that tissue-specific eQTL signals are preferentially located in regulatory sequences some distance from transcriptional start sites <sup>63</sup>—in common with many complex trait association signals—suggests that further dissection of these regions should improve understanding of the genomic organization of tissue- and/or developmental-stage-specific regulation.

A further conclusion is that common SNP signals are likely to fall short in explaining the observed familial aggregation of T2D, at least in European descent populations. The limited power of our study (Table 1) to detect several of the genome-wide significant variants we report here (based on the stage 1 sample size and stage 2 odds ratios that minimize 'winner's curse' effects) indicates that there are likely to be many additional common variant signals of similar effect that could be detected by further expansion of the GWA meta-analysis approach. However, it seems unlikely that these will explain a substantial proportion of unex-plained heritability. Based on the data presented, the same is likely to be true for common CNVs and for variation on the sex chromosomes. As a result, the attention of researchers in the field is increasingly directed toward evaluation of the contribution of low frequency and rare variants to complex trait susceptibility. Several lines of evidence—the overlap in loci implicated in monogenic and multifactorial diabetes, the congregation of multiple disease signals at a limited number of loci the conditional analyses—point toward the importance of obtaining complete descriptions of causal genetic variation (of all types and frequencies) at the loci uncovered by this and other GWA studies. Such loci are likely to represent hotspots at which the overall contribution to T2D predisposition and biology may be considerably greater than that estimated using the discovered common variants alone.

#### **METHODS**

Methods and any associated references are available in the online version of the paper at http://www.nature.com/naturegenetics/.

#### **Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

#### **Footnotes**

Correspondence should be addressed to D.A. (altshuler@molbio.mgh.harvard.edu), M.B. (boehnke@umich.edu), P.F. (froguel@good.ibl.fr), T.I. (illig@helmholtz-muenchen.de), M.I.M. (mark.mccarthy@drl.ox.ac.uk), K. Stefansson (kstefans@decode.is), C.M.v.D. (c.vanduijn@erasmusmc.nl) or J.F.W. (jim.wilson@hgu.mrc.ac.uk).

**AUTHOR CONTRIBUTIONS Manuscript writing:** B.F.V., L.J.S., V.S., A.P.M., C.D., E.Z., T.F., T.M.F., R. Sladek, U.T., D.A., M.B., M.I.M.

Clinical samples: A.P.M., H.G., C.L., L.Q., M.v.H., P. Navarro, K.A., B. Balkau, R. Benediktsson, R. Blagieva, L.L.B., K.B.B., B. Bravenboer, N.P.B., G. Charpentier, M.C., A.S.F.D., C.S. Fox, C. Gieger, N.G., S.G., S.H., C. Herder, B.I., T.J., P.K., J.K., T.L., A.L., V.L., M.M., T.M., K.M., P. Nilsson, K.R.O., C. Platou, W.R., M.R., M.J.S., B.M.S., G.S., T.S., K. Strassburger, Q.S., B.T., J. Tichet, T.T., R.M.V.D., T.W.V.H., T.V.H., J.V.V.V.-O., C.W., R.N.B., F.S.C., U.G., T.H., G.A.H., D.J.H., K.H., M.Laakso, K.L.M., A.D.M., C.N.A.P., P.P.P., I.R., E.S., J. Tuomilehto, M.W., N.J.W., B.O.B., H.C., A.T.H., F.B.H., J.B.M., J.S.P., O.P., T.M.F., L.G., R. Sladek, U.T., H.E.W., J.F.W., T.I., P.F., C.M.V.D., K. Stefansson, D.A., M.B., M.I.M.

Stage 1 genotyping and analysis: B.F.V., L.J.S., V.S., A.P.M., C.D., E.Z., C. Huth, Y.S.A., G.T., T.F., H.G., N.A., C.J.W., C.L., A.V.S., M.v.H., P. Navarro, K.A., R. Benediktsson, A.J.B., L.L.B., K.B.B., S.B., N.P.B., G. Charpentier, P.S.C., M.C., G. Crawford, M.R.E., M.G., N.G., C.J.G., C. Guiducci, C. Herder, B.I., A.U.J., N.K., T.L., C.M.L., V.L., M.M., T.M., M.A.M., N.N., P. Nilsson, F.P., G.R., R. Saxena, T.S., K. Strassburger, H.M.S., A.J.S., T.T., R.M.v.D., G.B.W., J.W., R.N.B., S.C., F.S.C., U.G., K.L.M., I.R., E.S., J. Tuomilehto, A.U., N.J.W., H.C., F.B.H., T.M.F., L.G., R. Sladek, U.T., H.-E.W., J.F.W., T.I., P.F., C.M.v.D., K. Stefansson, D.A., M.B., M.I.M.

Stage 2 genotyping and analysis: B.F.V., L.J.S., V.S., A.P.M., C.D., C. Huth, Y.S.A., G.T., H.G., N.A., C.J.W., C.L., J.D., L.Q., M.v.H., P. Navarro, K.A., A.J.B., E.B., L.L.B., K.B.B., S.B., N.P.B., P.S.C., M.C., D.J.C., G. Crawford, A.S.F.D., M.R.E., C.S. Franklin, M.G., C. Gieger, N.G., S.G., C.J.G., C. Guiducci, N.H., C. Herder, B.I., A.U.J., T.J., W.H.L.K., N.K., A.K., P.K., J.K., T.L., M. Li, C.M.L., V.L., T.M., K.M., M.A.M., N.N., P. Nilsson, F.P., A.-K.P., C. Proença, I.P., W.R., N.W.R., N.R.R., G.R., M.R., M.J.S., P.S., T.S., K. Strassburger, H.M.S., Q.S., A.J.S., T.T., R.M.v.D., T.W.v.H., J.V.v.V.-O., G.B.W., M.N.W., C.W., J.W., R.N.B., S.C., F.S.C., U.G., T.H., D.J.H., K.H., M. Laakso, K.L.M., A.D.M., C.N.A.P., P.P.P., I.R., E.S., J. Tuomilehto, A.U., N.J.W., H.C., M.J.D., F.B.H., J.S.P., O.P., I.B., J.C.F., T.M.F., L.G., R. Sladek, H.-E.W., U.T., J.F.W., T.I., P.F., C.M.v.D., D.A., M.B., M.I.M.

Analysis group: B.F.V., L.J.S., V.S., A.P.M., R.P.W., C.D., E.Z., C. Huth, Y.S.A., G.T., T.F., H.G., N.A., C.J.W., J.D., M.v.H., M.G., C. Gieger, A.U.J., N.K., A.K., J.R.B.P., A.-K.P., N.W.R., N.R.R., R. Saxena, M.J.D., P.F., M.B., M.I.M.

**Biological analyses:** V.S., G.T., L.J.M., S.A.M., J.D., K.S.E., A.L.E., P.R.V.J., V.L., I.P., A.L.G., J.B.M., U.T., K. Stefansson, M.I.M.

Informatics analyses: B.F.V., V.S., G.W., S.R., O.M.H., A.V.S., T.G., W.A.H., L.D.S. **DIAGRAM consortium management:** B.F.V., L.J.S., V.S., A.P.M., C.D., E.Z., R.N.B., S.C., F.S.C., A.H., K.L.M., E.S., J. Tuomilehto, R.M.W., G.R.A., H.C., M.J.D., A.T.H., T.M.F., L.G., R. Sladek, U.T., H.-E.W., J.F.W., T.I., P.F., C.M.v.D., K. Stefansson, D.A., M.B., M.I.M.

Note: Supplementary information is available on the Nature Genetics website.

**COMPETING FINANCIAL INTERESTS** The authors declare competing financial interests: details accompany the full-text HTML version of the paper at http://www.nature.com/naturegenetics/.

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#### **Acknowledgments**

We acknowledge funding from: the Academy of Finland (no. 124243); Agence Nationale de la Recherche (France); American Diabetes Association (1-05-RA-140, 7-08-MN-OK; 7-06-MN-05); Ardix Medical; Association Diabète Risque Vasculaire; Association de Langue Française pour l'Etude du Diabète et des Maladies Métaboliques; Association Française des Diabétiques; Bayer Diagnostics; British Diabetic Association Research; Becton Dickinson; Broad Institute of Harvard and Massachusetts Institute of Technology; The Burroughs Wellcome Fund; Cardionics; Center for Inherited Disease Research (USA); Centre for Medical Systems Biology (The Netherlands); Centre of Excellence Metabolic Disorders Baden-Wuerttemberg (Germany); Caisse Nationale Assurance Maladie des Travailleurs Salariés (France); Clinical Research Institute HUCH Ltd; Deutsche Forschungsgemeinschaft (DFG GrK 1041, DFG RA459, SFB 518); the Danish Diabetes Association; the Danish Health Research Council; Diabetes UK; Doris Duke Charitable Foundation; Erasmus Medical Center (The Netherlands); the Dutch Diabetes Foundation; European Community (HEALTH-F4-2007-201413, HEALTH-2007-B-223211, LSHG-CT-2006-01947, LSHM-CT-2004-512013, LSHM-CT-2004-005272, LSHM-CT-2006-518153); the European Foundation for the Study of Diabetes; the Federal Ministry of Health (Germany); the Federal Ministry of Education and Research (Germany) (FKZ01GS0823 and DZD e.V.); Fédération Française de Cardiologie; The Finnish Diabetes Research Foundation; The Folkhalsan Research Foundation; The Foundation for Strategic Research (Sweden); The Foundation of Bristol-Myers Squibb; the German National Genome Research Network; Helmholtz Zentrum München-Research Center for Environment and Health; INSERM (France); La Fondation de France; Lilly; The Linnaeus Centre for Bioinformatics (Sweden); the Lundbeck Foundation Centre of Applied Medical Genomics for Personalized Disease Prediction, Prevention and Care; the Medical Research Council UK (G0601261, G0000649; 081696); Munich Center of Health Sciences-LMU Innovativ (Germany); Merck Santé; the Ministry of Health and Department of Educational Assistance, University and Research of the Autonomous

<sup>&</sup>lt;sup>99</sup>A full list of members is provided in the Supplementary Note.

<sup>100</sup> These authors contributed equally.

Province of Bolzano (Italy); the Ministry of Innovation, Science, Research and Technology of the State of North Rhine-Westphalia (Germany); the Ministry of Science, Education and Sport (Croatia); the National Heart, Lung, and Blood Institute (N01-HC-55015, N01-HC-55016, N01-HC-55018, N01-HC-55019, N01-HC-55020, N01-HC-55 HC-55021, N01-HC-55022, N01-HC-25195, R01HL087641, R01HL59367, R01HL086694, N02-HL-6-4278); National Human Genome Research Institute (U01HG004402, U01HG004399, U01HG004171, 1 Z01 HG000024); the National Institute of Diabetes, Digestive and Kidney Diseases (DK078616, K24-DK080140, U54 DA021519, DK58845, DK069922, DK062370, DK073490, K23-DK65978 and DK072193); the US National Institutes of Health (HHSN268200625226C, HHSN268200625226C, 1K08AR055688, UL1RR025005, 1K99HL094535-01A1); the Netherlands Foundation for Scientific Research (175.010.2005.011, 047.017.043); Nord-Pas-de-Calais region (France); Novartis Pharma; Novo Nordisk; the Oxford National Institute for Health Research (NIHR) Biomedical Research Centre (UK); Office National Inter-professionnel des Vins; Peninsula Medical School, Exeter UK; Pfizer, Inc; Pierre Fabre laboratory (France); Programme National de Recherche sur le Diabète (France); Richard and Susan Family Foundation/American Diabetes Association Pinnacle Program Project; Roche; the Royal Society (UK); Russian Foundation for Basic Research (047.017.043); Sanofi-Aventis; Sarnoff Cardiovascular Research; Scottish Government Chief Scientist Office; SenterNovem (IOP Genomics grant IGE05012); Sigrid Juselius Foundation; the Skaraborg Institute, Skövde, Sweden; South Tyrolean Sparkasse Foundation; the Swedish Natural Sciences Research Council; The Swedish Research Council (349 2006-237P); the Association Diabète Risque Vasculaire (France); Topcon; the Wallenberg Foundation; and the Wellcome Trust (072960; 076113; 083270; 088885; 079557; 081682; 086596; 077016; 075491). A more complete list of acknowledgments is provided in the Supplementary Note.

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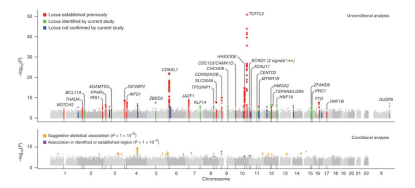


Figure 1. Genome-wide Manhattan plots for the DIAGRAM+ stage 1 meta-analysis. Top panel summarizes the results of the unconditional meta-analysis. Previously established loci are denoted in red and loci identified by the current study are denoted in green. The ten signals in blue are those taken forward but not confirmed in stage 2 analyses. The genes used to name signals have been chosen on the basis of proximity to the index SNP and should not be presumed to indicate causality. The lower panel summarizes the results of equivalent meta-analysis after conditioning on 30 previously established and newly identified autosomal T2D-associated SNPs (denoted by the dotted lines below these loci in the upper panel). Newly discovered conditional signals (outside established loci) are denoted with an orange dot if they show suggestive levels of significance ( $P < 10^{-5}$ ), whereas secondary signals close to already confirmed T2D loci are shown in purple ( $P < 10^{-4}$ ).

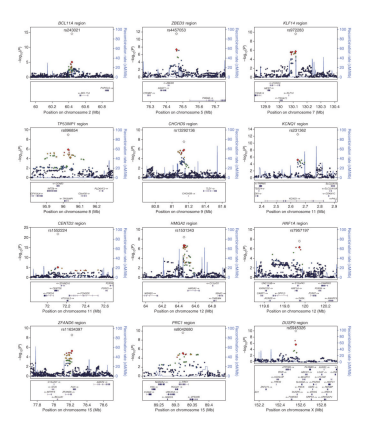


Figure 2. Regional plots of the 12 newly discovered T2D loci. Genotyped and imputed SNPs passing quality control measures across all stage 1 studies are plotted with their meta-analysis P values (as  $-\log_{10}$  values) as a function of genomic position (NCBI Build 36). In each panel, the index association SNP is represented by a diamond, with stage 1 meta-analysis results denoted by a red diamond and the combined stage 1 and stage 2 meta-analysis results denoted with a clear symbol. Estimated recombination rates (taken from HapMap CEU) are plotted to reflect the local LD structure. Color of remaining SNPs (circles) indicates LD with the index SNP according to a scale from  $r^2 = 0$  to  $r^2 = 1$  based on pairwise  $r^2$  values from HapMap CEU (red,  $r^2 = 0.8-1.0$ ; orange,  $r^2 = 0.6-0.8$ ; green,  $r^2 = 0.4-0.6$ ; blue,  $r^2 = 0.2-0.4$ ; black,  $r^2 < 0.2$ ; gray, no  $r^2$  value available). Gene annotations were taken from the University of California Santa Cruz genome browser.

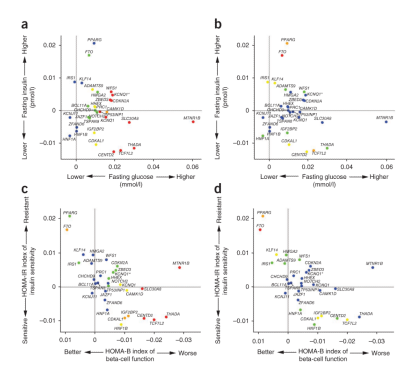


Figure 3. Plots of fasting blood glucose, insulin and derived indices for the established and new T2D loci. (**a,b**) Plots of fasting glucose (x axis) and fasting insulin (y axis). (**c,d**) Plots of HOMA-B (an index of beta cell function; x axis) and HOMA-IR (an index of insulin sensitivity; y axis). Each point refers to a single T2D association signal, with colors denoting the strength of the association to either the x-axis variable (left-hand of each pair of plots) or y-axis variable (right-hand of each pair) (red,  $P < 10^{-3}$ ; orange,  $10^{-3} < P < 10^{-2}$ ; yellow, 0.01 < P < 0.05; green, 0.05 < P < 0.20; blue, P > 0.20). The two KCNQ1 associations are distinguished by the notation KCNQ1 for rs163184 and KCNQ1\* for rs231362. The gene names associated with each signal have been chosen on the basis of proximity to the index SNP and should not be presumed to indicate causality.

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Table 1

Association results for stage 1 + 2 which exceed a genome-wide threshold (overall P value  $< 5 \times 10^{-8}$ )

								Stage 1d		Stage 2d		Stage $1 + 2^d$	
SNP C	Chr.	Position B36 (base pair)	Risk allele $^b$	Nonrisk allele $^b$	Frequency risk allele (Hapmap CEU)	Nearby gene <sup>c</sup>	OR (95%CI)	P value	OR (95% CI)	P value	OR (95% CI)	P value	Power
							up to and 38,9	up to 8,130 cases and 38,987 controls	up to and 59,	up to 34,412 cases and 59,925 controls $^f$		up to 42, and 98,91	up to 42,542 cases and 98,912 controls
New T2D susceptibility loci	ptibili	ty loci											
rs243021	2	60,438,323	Ą	Ö	0.46	BCL11A	1.09 (1.05–1.13)	$8.1\times10^{-6}$	1.08 (1.06–1.10)	$6.2\times10^{-11}$	1.08 (1.06–1.10)	$2.9\times10^{-15}$	09.0
rs4457053	5	76,460,705	Ŋ	A	0.26	ZBED3	1.16 (1.10–1.23)	$4.2\times10^{-8}$	1.07 (1.04–1.10)	$2.7\times10^{-7}$	1.08 (1.06–1.11)	$2.8\times10^{-12}$	0.25
rs972283	7	130,117,394	Ŋ	A	0.55	KLF14	$1.10 \\ (1.06-1.15)$	$1.8\times10^{-6}$	1.06 (1.03–1.09)	$6.4\times10^{-6}$	1.07 (1.05–1.10)	$2.2\times10^{-10}$	0.19
rs896854	∞	96,029,687	Г	C	0.48	TP53INP1	$1.10 \\ (1.06-1.15)$	$1.2\times10^{-6}$	1.05 (1.03–1.08)	$2.2\times10^{-5}$	1.06 (1.04–1.09)	$9.9\times10^{-10}$	0.08
rs13292136	6	81,141,948	C	L	0.93	СНСНD9	1.20 (1.11–1.29)	$1.5\times10^{-6}$	1.08 (1.04–1.13)	$2.4\times10^{-4}$	$1.11 \\ (1.07 - 1.15)$	$2.8\times10^{-8}$	0.02
rs231362	=	2,648,047	G	∢	0.52	KCNQ1	1.11 (1.06–1.16)	$6.4\times10^{-6}$	1.07 (1.05–1.09)	$3.2\times10^{-9}$	1.08 (1.06–1.10)	$2.8\times10^{-13}$	0.38
rs1552224	11	72,110,746	Ą	C	0.88	CENTD2	$1.13 \\ (1.07 - 1.19)$	$7.0\times10^{-6}$	1.14 (1.11–1.18)	$3.2\times10^{-18}$	1.14 (1.11–1.17)	$1.4\times10^{-22}$	0.58
rs1531343 <i>a</i>	12	64,461,161	C	Ö	0.10	HMGA2	1.20 (1.12–1.29)	$1.7\times10^{-7}$	1.08 (1.04–1.12)	$1.1\times10^{-4}$	$1.10 \\ (1.07 - 1.14)$	$3.6\times10^{-9}$	0.07
rs7957197	12	119,945,069	H	A	0.85	HNF1A	1.14 (1.08–1.19)	$4.6\times10^{-7}$	1.05 (1.02–1.09)	$4.6\times10^{-4}$	1.07 (1.05–1.10)	$2.4\times10^{-8}$	0.01
rs11634397	15	78,219,277	Ŋ	A	09.0	ZFAND6	1.11 (1.06–1.16)	$5.1\times10^{-6}$	1.05 (1.03–1.08)	$1.2\times10^{-5}$	1.06 (1.04–1.08)	$2.4\times10^{-9}$	0.07
rs8042680	15	89,322,341	A	C	0.22	PRCI	$1.10 \\ (1.06-1.15)$	$8.2\times10^{-6}$	1.06 (1.03–1.08)	$1.6\times10^{-6}$	1.07 (1.05–1.09)	$2.4\times10^{-10}$	60.0
rs5945326	×	152,553,116	∢	Ü	0.79	DUSP9	1.25 (1.14–1.37)	$2.3\times10^{-6}$	$1.32$ (1.16–1.49) $^f$	$2.3 \times 10^{-5}$	$1.27 \\ (1.18-1.37)$	$3.0\times10^{-10}$	$^{f}66.0$
Previously known	wn												
rs7578326	2	226,728,897	4	Ö	0.64	IRSI	$1.12 \\ (1.07 - 1.17)$	$8.7\times10^{-7}$	1.10 (1.08–1.13)	$2.2\times10^{-15}$	1.11 (1.08–1.13)	$5.4\times10^{-20}$	0.83
rs1387153	11	92,313,476	П	C	0.28	MTNRIB	1.12 (1.07–1.17)	$1.0\times10^{-6}$	1.08 (1.05–1.10)	$4.4\times10^{-10}$	1.09 (1.06–1.11)	$7.8\times10^{-15}$	0.46

OR, odds ratio; CI, confidence interval; Chr., chromosome.

 $^{a}$ Proxy SNP rs2612067 was genotyped in METSIM, FUSION2 and HUNT stage 2 samples.

 $^{\it b}$  Alleles are indexed to the forward strand of NCBI Build 36.

 $^{\mathcal{C}}$  Gene region named for nearest gene (except where there is a very strong positional candidate).

 $d_{\rm All}$  Pvalues reported are two-sided and based on an inverse-variance weighted meta-analysis model (fixed effects).

f Maximum stage 2 sample size for the X chromosome SNP was 8,535 cases and 12,326 controls. Allelic odds ratios calculated assuming X-chromosome inactivation in females. Power to detect SNP given stage 1 sex distribution and X-chromosome inactivation in females, Power to detect autosomal SNPs given the stage 1 sample sizes and stage 2 ORs, the risk AF in Hapmap (CEU), for  $\alpha = 1 \times 10^{-5}$ . Additive model assumed, 8% population prevalence. otherwise as for autosomes. Europe PMC Funders Author Manuscripts

Expression QTL results for T2D-associated variants in blood and adipose tissue

Name   Circ   Rick of page   Circ   Rick of Circ										SNP with str	SNP with strongest correlation with trait	on with trait
65         ZBED3 (N. LOSTI9)         Adipose         0.302 (0.07)         2.8 × 10~5         0.89         nes64520           944         KLFJ4         G         KLFJ4 (N. L. O32367)         Adipose         0.429 (0.068)         1.0 × 10~9         0.011         nes4704389           944         KLFJ4         G         KLFJ4 (N. L. J. 38093)         Adipose         0.225 (0.053)         3.8 × 10~7         0.038         ne373334           46         CFNTD2         A         STARDIQ (N. L. J. 38093)         Blood         0.225 (0.053)         3.8 × 10~7         0.036         ne313434           96         HNFJA         T         CCNZR (N. L. J. 1000017)         Adipose         0.248 (0.067)         3.7 × 10~4         0.29         ne313433           96         HNFJA         T         ACALOS (N. L. J. 1000017)         Adipose         0.248 (0.068)         3.7 × 10~4         0.29         ne313433           96         HNFJA         T         ACALOS (N. L. J. 1002313)         Blood         0.248 (0.068)         3.7 × 10~4         0.29         ne314593           96         HNFJA         T         ACALOS (N. L. J. 1023214)         Blood         0.248 (0.068)         3.7 × 10~4         0.29         ne314593           9         CCOQ5 (N. L. J.			Nearby gene <sup>d</sup>	Risk allele $b$	Gene (transcript)	Tissue	Effect (s.e.m.) $^{\mathcal{C}}$	P value	$^{P}_{ m adj}d$	$SNP(r^2)f$	P value	$P_{ m adj} S$
76,46,715         28ED3         G         PDESB (NM_0032367)         Adipose         0.387 (0.058)         2.8.10-5         0.80         cosession           36,117,394         KLP14         G         KLFF4(NM_138037)         Adipose         0.237 (0.058)         8.1 × 10-1         0.038         0.011         0.010           36,117,394         KLF14         G         KLFF4(NM_138037)         Adipose         0.237 (0.058)         8.1 × 10-7         0.038         0.731           36,029,687         TYSZINPI         T         CCVEZ (NM_1087139)         Blood         0.237 (0.056)         3.3 × 10-7         0.026         0.031           119,945,069         HNPIA         T         CCVEZ (NM_1087133)         Blood         0.234 (0.066)         3.7 × 10-4         0.026         0.031           119,945,069         HNPIA         T         ACALS (NM_1087133)         Blood         0.234 (0.066)         3.7 × 10-4         0.026         0.031           119,945,069         HNPIA         T         ACALS (NM_108733)         Blood         0.234 (0.066)         3.7 × 10-4         0.026         0.031           119,41,104         HNPIA         T         ACALS (NM_108733)         Blood         0.234 (0.066)         3.2 × 10-4         0.026         0.031		ı this study										
380,117.394         KLF/4         G         KLF/4         G         KLF/4         G         KLF/4 (NM_193267)         Adipose         0.429 (0.068)         1.0 × 10^-9         G         G         C		76,460,705	ZBED3	G	PDE8B (NM_003719)	Adipose	0.302 (0.070)	$2.8\times10^{-5}$	0.80	rs6864250 (0.18)	$3.1\times10^{-17}$	$5.8\times10^{-13}$
19.011.394         KLF44         G         KLF4(NML)38693)         Adipose         -0.387 (0.058)         8.1×10-11         0.058         6x33134           96.029.687         7752/KV         1         CCCVZC (NML) 0.006453         Blood         -0.225 (0.053)         3.8×10-5         0.78         6x3733           72.110,746         CEXTDZ         A         STAKDIO (NML) 0.006453         Blood         -0.248 (0.067)         3.7×10-4         0.026         0.5319           119,945,069         HNFIA         T         ACADE (NML) 0.002813         Blood         0.248 (0.067)         3.7×10-4         0.026         0.850-0           119,945,069         HNFIA         T         ACADE (NML) 0.028133         Blood         0.249 (0.066)         3.7×10-4         0.008         0.021           119,945,069         HNFIA         A         ACADE (NML) 0.023133         Blood         0.249 (0.066)         1.3×10-6         0.3         0.03         0.025           110,945,069         HNFIA         A         ACADE (NML) 0.032313         Blood         0.248 (0.066)         1.4×10-6         0.3         0.010         0.03         0.03         0.03         0.03         0.03         0.03         0.03         0.03         0.03         0.03         0.03					ZBED3 (NM_032367)	Adipose	0.429 (0.068)	$1.0\times10^{-9}$	0.011	rs4704389 (0.20)	$3.9\times10^{-16}$	$6.0\times10^{-9}$
96.029.687   TP53INP1   T		130,117,394	KLF14	Ö	KLF14 (NM_138693)	Adipose	-0.387 (0.058)	$8.1\times10^{-11}$	0.058	rs738134 (0.30)	$2.2\times10^{-12}$	0.0014
72.110.746         CEWTD2         A         STARDIO (NML_000645)         Blood         0.337 (0.066)         8.6 x 10 <sup>-7</sup> 0.026         0.031           119.945.069         IMPTA         T         ACADS (NML_000213)         Adipose         0.248 (0.067)         3.7 x 10 <sup>-4</sup> 0.098         0.020           119.945.069         IMPTA         T         ACADS (NML_0002133)         Adipose         0.248 (0.066)         3.7 x 10 <sup>-4</sup> 0.0088         0.020           11.0         ACASL (NML_003733)         Adipose         0.248 (0.066)         1.3 x 10 <sup>-6</sup> 0.139         0.020           11.0         ACASL (NML_003733)         Blood         0.248 (0.066)         1.3 x 10 <sup>-6</sup> 0.31         0.020           11.0         ACASL (NML_032314)         Blood         0.248 (0.066)         1.4 x 10 <sup>-4</sup> 0.048         0.01           11.0         ACAMKKZ (NML_172215)         Adipose         0.0497 (0.066)         1.4 x 10 <sup>-4</sup> 0.048         0.01           22.2.2.341         RRCI         AURKZ (NML_172215)         Blood         0.024 (0.067)         3.4 x 10 <sup>-8</sup> 0.089           22.2.2.8.887         ARGANAL (NML_172215)         Blood         0.024 (0.067)         3.4 x 10 <sup>-8</sup> 0.089         0.039 <tr< td=""><td></td><td>96,029,687</td><td>TP53INP1</td><td>Т</td><td>CCNE2 (NM_057749)</td><td>Blood</td><td>-0.225 (0.053)</td><td><math display="block">3.8\times10^{-5}</math></td><td>0.78</td><td>rs4735339 (0.61)</td><td><math display="block">5.8\times10^{-7}</math></td><td>0.0051</td></tr<>		96,029,687	TP53INP1	Т	CCNE2 (NM_057749)	Blood	-0.225 (0.053)	$3.8\times10^{-5}$	0.78	rs4735339 (0.61)	$5.8\times10^{-7}$	0.0051
19,945,069   HNFA   T		72,110,746	CENTD2	4	STARD10 (NM_006645)	Blood	0.337 (0.066)	$8.6\times10^{-7}$	0.026	rs519790 (0.04)	$2.7\times10^{-24}$	$1.6\times10^{-19}$
R5ADD (NM_002813)         Blood         0.240 (0.065)         3.9 × 10 <sup>-4</sup> 0.0088         n.3741593           A6T (NM_002813)         Adipose         0.318 (0.064)         1.3 × 10 <sup>-4</sup> 0.13         0.1259883           A6T (NM_002733)         Blood         0.218 (0.064)         1.3 × 10 <sup>-4</sup> 0.37         0.2556828           A6T (NM_0027314)         Blood         0.248 (0.064)         1.3 × 10 <sup>-4</sup> 0.92         0.021           A7 (NM_0027314)         Blood         -0.254 (0.064)         1.4 × 10 <sup>-4</sup> 0.048         0.105           A7 (NM_0027314)         Blood         -0.254 (0.064)         1.2 × 10 <sup>-4</sup> 0.048         0.105           A7 (NM_0027315)         Adipose         -0.497 (0.068)         1.4 × 10 <sup>-4</sup> 0.048         0.105           A8,22,23,41         PRCI         A         PRKKA (NM_175215)         Blood         -0.254 (0.064)         1.2 × 10 <sup>-4</sup> 0.048         0.108           A8,22,23,41         PRCI         A         PRKKA (NM_175215)         Blood         -0.254 (0.064)         3.4 × 10 <sup>-6</sup> 0.08         0.108           A8,22,23,41         PRCI         A         PRKA (NM_17568)         Blood         -0.251 (0.057)         3.7 × 10 <sup>-6</sup> 0.55         0.55		119,945,069	HNFIA	Н	ACADS (NM_000017)	Adipose	-0.248 (0.067)	$3.7\times10^{-4}$	0.29	rs9204 (0.02)	$1.3\times10^{-53}$	$5.9\times10^{-50}$
26.728.897         Adipose         0.318 (0.068)         6.4 × 10 <sup>-6</sup> 0.13         18.2259883           1. A. L.					PSMD9 (NM_002813)	Blood	0.240 (0.065)	$3.9\times10^{-4}$	0.0088	rs3741593 (0.00)	$8.3 \times 10^{-8}$	$1.7\times10^{-6}$
2.4.5. (NM_003733)         Blood         0.319 (0.064)         1.3 × 10^6         0.37         84556628           1.2. (NG)         COQ2 (NM_032314)         Blood         0.248 (0.065)         2.1 × 10^-4         0.92         0.021)           1.2. (NG)         COQ2 (NM_0323614)         Blood         -0.254 (0.064)         1.4 × 10^-4         0.048         0.003)           1.2. (NG)         CAMKAZ (NM_172215)         Adipose         -0.497 (0.068)         1.2 × 10^-16         0.18         0.1005           1.2. (NG)         AMKAZ (NM_172215)         Blood         -0.360 (0.063)         3.4 × 10^-8         0.048         0.008)           1.2. (NG)         AMKAZ (NM_172215)         Blood         -0.360 (0.063)         3.4 × 10^-8         0.048         0.008)           1.2. (NG)         AMKAZ (NM_172215)         Blood         -0.371 (0.057)         2.9 × 10^-16         0.039         0.039           1.2. (NG)         AMESTAR (NM_1018668)         Blood         -0.251 (0.059)         3.7 × 10^-8         0.50         0.053           1.2. (NG)         AMESTAR (NM_101869)         AMIPOS         -0.251 (0.059)         3.7 × 10^-8         0.58         0.53945635           1.2. (NG)         AMATS9         AMESTAR (NM_1025320)         Adipose         -0.029 (0.059)					OASL (NM_003733)	Adipose	0.318 (0.068)	$6.4\times10^{-6}$	0.13	rs2259883 (0.19)	$1.1\times10^{-7}$	0.0018
26,728,897         ADAMTS9         COQ5(NM_032314)         Blood         0.248 (0.065)         2.1 × 10 <sup>-4</sup> 0.92         ns 11065504           1, 2, 1, 1, 1         1, 1, 1, 1         1, 1, 1, 1         1, 1, 1, 1         1, 1, 1         1, 1, 1         1, 1, 1         1, 1, 1         1, 1 </td <td></td> <td></td> <td></td> <td></td> <td>OASL (NM_003733)</td> <td>Blood</td> <td>0.319 (0.064)</td> <td><math display="block">1.3\times10^{-6}</math></td> <td>0.37</td> <td>rs4556628 (0.21)</td> <td><math>4.4 \times 10^{-22}</math></td> <td><math display="block">1.4\times10^{-16}</math></td>					OASL (NM_003733)	Blood	0.319 (0.064)	$1.3\times10^{-6}$	0.37	rs4556628 (0.21)	$4.4 \times 10^{-22}$	$1.4\times10^{-16}$
26,222,341         RS/I         ADAMTS9         CAMKAZ (NM_172215)         Adipose         -0.254 (0.064)         1.4 × 10 <sup>-4</sup> 0.048         ns 11065202           26,728,897         ABOMATS9         Adipose         -0.497 (0.068)         1.2 × 10 <sup>-12</sup> 0.18         ns 11065504           26,728,897         ABOMATS9         ABOMATS9         ABOMATS9         Adipose         -0.251 (0.055)         3.4 × 10 <sup>-6</sup> 0.50         ns 11065504           26,728,897         ABOMATS9         ABOMATS9         ABOMATS9         ABOMATS9         Adipose         -0.251 (0.059)         3.7 × 10 <sup>-8</sup> 0.58         ns 2176040           26,680,405         ABOMATS9         A					COQ5 (NM_032314)	Blood	0.248 (0.065)	$2.1\times10^{-4}$	0.92	rs10774561 (0.02)	$8.7\times10^{-39}$	$4.9 \times 10^{-35}$
256,728,897         RSI         Adipose         -0.497 (0.068)         1.2 × 10 <sup>-12</sup> 0.18         ns 11065504 (0.08)           89,322,341         RRI         AMKK2 (NM_175568)         Blood         -0.360 (0.063)         3.4 × 10 <sup>-6</sup> 0.08         ns 11065504 (0.08)           89,322,341         PRCI         A         VPS33B (NM_175568)         Blood         -0.371 (0.055)         3.4 × 10 <sup>-6</sup> 2.0 × 10 <sup>-6</sup> ns 11065504 (0.08)           256,728,897         IRSI         A         VPS33B (NM_18668)         Blood         -0.371 (0.057)         2.9 × 10 <sup>-7</sup> 0.50         ns 12595616 (0.33)           256,728,897         IRSI         A         IRSI (NM_005544)         Adipose         -0.251 (0.059)         3.7 × 10 <sup>-8</sup> 0.58         ns 2176040           12,264,800         PPARC         A         IRSI (NM_014869)         Adipose         -0.630 (0.131)         2.9 × 10 <sup>-6</sup> 1.4 × 10 <sup>-4</sup> ns 10001           64,680,405         ADAMTS9         C         BCD40632 (AK022320)         Adipose         -0.229 (0.056)         7.6 × 10 <sup>-5</sup> 0.28         ns 4521216					UNC119B (NM_032661)	Blood	-0.254 (0.064)	$1.4 \times 10^{-4}$	0.048	rs11065202 (0.09)	$7.8\times10^{-12}$	$2.3\times10^{-9}$
256,728,897         RST         ADAMTS9         CAMKK2 (NM_172215)         Blood         -0.360 (0.063)         3.4 × 10^-8         0.68         ns 11065504           89,322,341         PRCI         A         VPS33B (NM_18668)         Blood         -0.371 (0.057)         2.9 × 10^-10         0.50 (0.03)           226,728,897         IRSI         A         VRS33B (NM_018668)         Blood         -0.251 (0.059)         3.7 × 10^-5         0.89 (0.57)           226,728,897         IRSI         Adipose         -0.251 (0.059)         3.7 × 10^-8         0.89 (0.93)           12,264,800         PPARG         A IQSECI (NM_014869)         Adipose         -0.630 (0.131)         2.9 × 10^-6         1.4 × 10^-4         10.74           64,680,405         ADAMTS9         C         BCD40632 (AK022320)         Adipose         -0.029 (0.056)         7.6 × 10^-5         0.28         10.75					CAMKK2 (NM_172215)	Adipose	-0.497 (0.068)	$1.2\times10^{-12}$	0.18	rs11065504 (0.08)	$2.7\times10^{-117}$	$3.8\times10^{-98}$
89,322,341         PRCI         A         VPS33B (NM_1018668)         Blood         -0.371 (0.057)         3.4 × 10^-6         2.0 × 10^-6         10.25644           226,728,897         IRSI         A         VPS33B (NM_1018668)         Blood         -0.371 (0.057)         2.9 × 10^-10         0.50         10.033)           226,728,897         IRSI         Adipose         -0.251 (0.059)         3.7 × 10^-8         0.89         10.351           12,264,800         PPARG         A         IRSI (NM_0014869)         Adipose         -0.630 (0.131)         2.9 × 10^-6         1.4 × 10^-4         10.74           64,680,405         ADAMTS9         C         BCDM0632 (AK022320)         Adipose         -0.229 (0.056)         7.6 × 10^-5         0.28         10.4 × 10^-6         10.01					CAMKK2 (NM_172215)	Blood	-0.360 (0.063)	$3.4\times10^{-8}$	0.68	rs11065504 (0.08)	$7.0\times10^{-105}$	$5.7\times10^{-94}$
89,322,341         PRC1         A         VPS33B (NM_018668)         Blood         -0.371 (0.057)         2.9 × 10 <sup>-10</sup> 0.50         ns12395616           226,728,897         IRSI         A         IRSI (Contig50189_RC)         Adipose         -0.251 (0.059)         3.7 × 10 <sup>-5</sup> 0.89         ns2943653           12,264,800         PPARG         A         IRSI (NM_005544)         Adipose         -0.031 (0.059)         5.7 × 10 <sup>-8</sup> 0.58         ns2176040           64,680,405         ADAMTS9         C         BCD40632 (AK022320)         Adipose         -0.229 (0.056)         7.6 × 10 <sup>-5</sup> 0.28         ns451216           6002)         ADAMTS9         C         BCD40632 (AK022320)         Adipose         -0.229 (0.056)         7.6 × 10 <sup>-5</sup> 0.28         ns451216					<i>P2RX4</i> (NM_175568)	Blood	0.312 (0.065)	$3.4\times10^{-6}$	$2.0\times10^{-6}$	rs25644 (0.03)	$3.4\times10^{-17}$	$1.9\times10^{-17}$
226,728,897 IRSI A IRSI (Contig50189_RC) Adipose -0.251 (0.059) 3.7×10 <sup>-5</sup> 0.89 1x2943653 (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.93) (0.94) (0.94) (0.94) (0.94) (0.94) (0.94) (0.94) (0.94) (0.95) (0.		89,322,341	PRCI	∢	<i>VPS33B</i> (NM_018668)	Blood	-0.371 (0.057)	$2.9 \times 10^{-10}$	0.50	rs12595616 (0.57)		$4.5 \times 10^{-12}$
IRS1         A         IRSI (Contig50189_RC)         Adipose         -0.251 (0.059)         3.7 × 10^-5         0.89         rs.2943653           PPARG         A         IRSI (NM_005544)         Adipose         -0.331 (0.059)         5.7 × 10^-8         0.58         rs.2176040           PPARG         A         IQSECI (NM_014869)         Adipose         -0.630 (0.131)         2.9 × 10^-6         1.4 × 10^-4         rs.9211           ADAMTS9         C         BCD40632 (AK022320)         Adipose         -0.229 (0.056)         7.6 × 10^-5         0.28         rs.4521216	_	oci										
PPARG         A         IRSI (NM_005544)         Adipose $-0.331 (0.059)$ $5.7 \times 10^{-8}$ $0.58$ $rs.2176040$ PPARG         A         IQSECI (NM_014869)         Adipose $-0.630 (0.131)$ $2.9 \times 10^{-6}$ $1.4 \times 10^{-4}$ $rs.9211$ ADAMTS9         C         BC0440632 (AK022320)         Adipose $-0.229 (0.056)$ $7.6 \times 10^{-5}$ $0.28$ $rs.4521216$ (002)		226,728,897	IRSI	∢	IRS1 (Contig50189_RC)	Adipose	-0.251 (0.059)	$3.7\times10^{-5}$	0.89	rs2943653 (0.93)	$3.4\times10^{-5}$	69.0
PPARG         A $IQSECI$ (NM_014869)         Adipose $-0.630$ (0.131) $2.9 \times 10^{-6}$ $1.4 \times 10^{-4}$ $rs9211$ ADAMTS9         C         BC040632 (AK022320)         Adipose $-0.229$ (0.056) $7.6 \times 10^{-5}$ 0.28 $rs4521216$ (0.02)         C <td></td> <td></td> <td></td> <td></td> <td>IRSI (NM_005544)</td> <td>Adipose</td> <td>-0.331 (0.059)</td> <td><math display="block">5.7\times10^{-8}</math></td> <td>0.58</td> <td>rs2176040 (0.74)</td> <td><math display="block">7.8\times10^{-10}</math></td> <td>0.0042</td>					IRSI (NM_005544)	Adipose	-0.331 (0.059)	$5.7\times10^{-8}$	0.58	rs2176040 (0.74)	$7.8\times10^{-10}$	0.0042
$ADAMTS9 \qquad C \qquad BC040632  (AK022320) \qquad \text{Adipose} \qquad -0.229  (0.056)  7.6 \times 10^{-5}  0.28 \qquad \text{rs}4521216 \qquad \qquad (0.02)  (0.02)$		12,264,800	PPARG	4	<i>IQSEC1</i> (NM_014869)	Adipose	-0.630 (0.131)	$2.9\times10^{-6}$	$1.4\times10^{-4}$	rs9211 (0.01)	$1.1\times10^{-96}$	$7.4 \times 10^{-94}$
		64,680,405	ADAMTS9	C	BC040632 (AK022320)	Adipose	-0.229 (0.056)	$7.6 \times 10^{-5}$	0.28	rs4521216 (0.02)	$3.0\times10^{-13}$	$8.7\times10^{-10}$

										SNP with str	SNP with strongest correlation with trait <sup>e</sup>	on with trait <sup>e</sup>
Chr. Position Nearby Risk Gene (transcript) $ B36 \ (bp)  gene a  \text{allele} b $	Nearby Risk gene $^{a}$ allele $^{b}$	Risk allele $b$		Gene (transcrip	t)	Tissue	Effect (s.e.m.) <sup>C</sup> P value	P value	$^{P}_{ m adj}d$	SNP $(r^2)f$ P value	P value	$^{P}_{ m adj} S$
ADAMTS9 (NM_182920) Adipose	ADAMTS9 (NM.	ADAMTS9 (NM <sub>.</sub>	ADAMTS9 (NM	ADAMTS9 (NM	182920)	Adipose	$-0.274 \ (0.055) \ 1.5 \times 10^{-6}$	$1.5 \times 10^{-6}$	0.036	rs7372321 (0.11)	$1.1\times10^{\text{-9}}$	$2.3 \times 10^{-5}$
7 28.162,747 JAZFI A JAZFI (NM_175061)	(62,747 <i>JAZFI</i> A	∢		JAZFI (NM_17:	5061)	Adipose	$-0.346 (0.055)   1.4 \times 10^{-9}$	$1.4\times10^{-9}$	0.70	rs1635852 (1.00)	$5.2\times10^{-10}$	0.17
10 12,368,016 CAMKID G CAMKID(AL137430)	368,016 CAMKID G	Ö		CAMKID (AL	137430)	Blood	$0.668 \ (0.062) \ 1.6 \times 10^{-25}$	$1.6\times10^{-25}$	0.052	rs11257655 (0.83)	$1.2\times10^{-25}$	0.04
CDC123 (NM_006023)	CDC123 (NM	CDC123 (NM	CDC123 (NM	CDC123 (NM	_006023)	Blood	0.498 (0.064)	$3.9\times10^{-14}$	0.012	rs11257600 (0.57)	$1.4\times10^{-16}$	$4.9\times10^{-5}$
CAMKID (NM_020397)	CAMKID (NIV	CAMKID (NN	CAMKID (NN	CAMKID (NIN	1_020397)	Blood	0.384 (0.064)	$7.6 \times 10^{-9}$	0.44	rs11257655 (0.83)	$3.3 \times 10^{-9}$	0.13
CAMKID (NM_153498)	CAMKID (N)	CAMKID (N)	CAMKID (NI	CAMKID (NI	M_153498)	Blood	0.643 (0.062)	$2.6\times10^{-23}$	n/a	rs12779790 (1.00)	$2.6\times10^{-23}$	n/a
10 94,455,539 HHEXIDE C KIFII (NM_004523)	455,539 HHEXIDE C	C		KIF11 (NM_	.004523)	Adipose	0.202 (0.059)	$9.4 \times 10^{-4}$	0.54	rs10882095 (0.53)	$1.6 \times 10^{-4}$	0.059
KIF11 (NM_004523)	KIFII (NM.	KIF11 (NM <sub>.</sub>	KIF11 (NM	<i>KIF11</i> (NM_	_004523)	Blood	0.185 (0.054)	$9.7\times10^{-4}$	0.62	rs6583826 (0.43)	$7.9 \times 10^{-6}$	0.0025
MARCH5()	MARCH5 ()	MARCH5 ()	MARCH5 ()	MARCH5 ()	MARCH5 (NM_017824)	Adipose	-0.262 (0.060)	$2.5\times10^{-5}$	0.31	rs10748579 (0.04)	$7.4 \times 10^{-36}$	$3.2 \times 10^{-31}$

Chr., chromosome.

 $^{a}$  Gene region named for nearest gene (except where there is a very strong positional candidate).

 $^{b}$ Alleles are indexed to the forward strand of NCBI Build 36.

Effect of T2D risk allele measured in units of standard deviation of the standardized distribution of the mean-log expression ratio.

d Pvalue for the expression trait association at the lead T2D SNP after conditioning on the local SNP that shows the most significant correlation with the expression trait.

The SNP in the 2-Mb window that shows the strongest correlation with the expression trait.

f Correlation  $r^2$  between the T2D-associated SNP and the SNP that shows the strongest correlation with the expression trait (in CEU HapMap).

<sup>g</sup> pvalue for the expression trait association after conditioning on the strongest T2D associated SNP in the region. A more detailed explanation of the table contents and analyses is provided in the Supplementary Note.