Differential gene expression profiling of the sciatic nerve in type 1 and type 2 diabetic mice

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Abstract. Diabetic peripheral neuropathy (DPN) is a common complication of diabetes mellitus (DM). The pathogenic mechanisms of DPN and the therapeutic interventions required may be distinct between type 1 (T1) and type 2 (T2) DM. However, the molecular mechanisms underlying the pathogenesis of DPN in both types of diabetes remain unclear. The aim of the current study was to identify the changes in genes and pathways associated with DPN in sciatic nerves of T1- and T2DM mice using bioinformatics analysis. The microarray profiles of sciatic nerves of T1DM (GSE11343) and T2DM (GSE27382) mouse models were downloaded from the Gene Expression Omnibus database to identify differentially expressed genes (DEGs) in each. DEGs in the two types of DM (with fold change ≥ 2 and P<0.05) were identified with BRB-ArrayTools. Gene Ontology (GO) term and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses were performed using the Database for Annotation, Visualization and Integrated Discovery. A protein-protein interaction (PPI) network was constructed using the Search Tool for the Retrieval of Interacting Genes/Proteins and visualized using Cytoscape. Compared with control samples, 623 and 1,890 DEGs were identified in sciatic nerves of T1- and T2DM mice, respectively. Of these, 75 genes were coordinately dysregulated in the sciatic nerves of both models. Many DEGs unique to T1DM mice were localized to the nucleoplasm and were associated with regulation of transcription processes, while many unique to T2DM mice were localized at cell junctions and were associated with ion transport. In addition, certain DEGs may be associated with the different treatment strategies used for the two types of DM. This analysis provides insight into the functional gene sets and

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pathways operating in sciatic nerves in T1- and T2DM. The results should improve understanding of the molecular mechanisms underlying the pathophysiology of DPN, and provide information for the development of therapeutic strategies for DPN specific to each type of DM.

Introduction

Diabetic peripheral neuropathy (DPN) is a common complication of diabetes mellitus (DM). More than half of patients with DM worldwide suffer from DPN during the course of the disease (1,2). DPN is characterized by distal-to-proximal nerve damage leading to neuropathic pain and loss of sensation (3). It is also associated with notably high morbidity and mortality: Previous studies reported that the mortality rate approximated to 25-50% within 5-10 years following onset of diabetic neuropathy (4,5). Strict maintenance of normal glycemic level is the only effective treatment available for DPN at present (6).

Though the occurrence of DPN is common to both type 1 (T1) and type 2 (T2)DM, a number of studies have suggested that its pathogenic mechanism may differ between the two (7,8). For example, more abnormalities at the molecular, functional and morphometric levels including increased frequencies of denervated Schwann cells and significant fiber loss have been observed in the peripheral nerve of T1DM mice compared with in T2DM mice (8,9). In addition, different structural changes may lead to variation in nerve conduction velocity for DPN in the two diabetic models (10). These fundamentally different mechanisms for DPN may lead to varying effects of the common treatments, such as in control of glucose, which has been suggested to be more beneficial for DPN in T1DM than in T2DM (7). Therefore, determining the distinct molecular mechanisms underlying DPN in T1- and T2DM is of paramount importance for the development of successful therapeutic interventions.

In the last decade, the development of high-throughput platforms including microarray technology has allowed researchers to concurrently determine the expression levels of several thousands of differentially expressed genes (DEGs) in diseases (11,12). In addition, the comparison of gene expression profiles of DEGs through microarray technology using bioinformatics analysis has determined distinct pathophysiological mechanisms in different diseases or syndromes including neuropathic pain and chronic radicular

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pain (13). For instance, Zhang et al (14) screened a number of DEGs in a control (healthy) group of mice and a group with streptozotocin (STZ)-induced diabetes, and identified genes co-regulated by both STZ and rosiglitazone, which may be potential targets in the treatment of DPN. However, there are few studies that have compared the gene expression profiles of DEGs between DPN in T1- and T2DM through microarray profiling. In the present study, the aim was to compare the DEGs between the sciatic nerves of T1- and T2DM mouse models by microarray profiling. Furthermore, the distinct biological processes and pathways associated with DPN in T1- or T2DM were analyzed and compared based on gene ontology (GO) and pathway enrichment analyses. This was hoped to provide novel insights into the distinct pathophysiological mechanisms and implicate drug therapies for DPN specific to the different types of DM.

Materials and methods

Source of microarray data. The gene expression profiles of GSE11343 and GSE27382 were obtained from the Gene Expression Omnibus (GEO) database (http://www.ncbi.nlm. nih.gov/geo/). The annotation platforms for GSE11343 and GSE27382 were GPL1261 and GPL9746, respectively.

For GSE11343 submitted by Wiggin *et al* (15), included datasets were of five sciatic nerve samples from mice with T1DM induced by STZ (GSM286169, GSM286173, GSM286176, GSM286178 and GSM286430) and four sciatic nerve samples from normal mice (GSM286159, GSM286160, GSM286163 and GSM286165).

For GSE27382 submitted by Pande *et al* (16), included datasets were of six sciatic nerve samples from db/db mice with T2DM (GSM677112-GSM677117) and seven sciatic nerve samples from db/+ (normal) mice (GSM677105-GSM677111).

Pre-processing of microarray data and identification of DEGs. Pre-processing for the cell intensity (CEL) files including conversion into expression measures, background correction and quartile data normalization was performed with BRB-ArrayTools (version 4.5.1) (17). The univariate t-test with a fold change ≥ 2 and nominal significance level of 0.05 was applied in BRB-ArrayTools to identify the DEGs between the diabetic and normal groups.

Gene ontology and pathway enrichment analyses. To identify the DEGs determined with BRB-ArrayTools, GO and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathway enrichment analyses were performed via the Database for Annotation, Visualization and Integrated Discovery (DAVID 6.8; http://david.abcc.ncifcrf.gov/) (18,19). GO terms [categorized into molecular function (MF), biological process (BP) and cellular component (CC)] and KEGG pathways with P<0.05 were considered significantly enriched by the DEGs.

Construction of protein-protein interaction (PPI) network and module analysis. With the purpose of evaluating the relationships among DEGs from the perspective of protein interaction, a protein-protein interaction (PPI) network was constructed with the Search Tool for the Retrieval of Interacting Genes/Proteins (STRING 10.5; http://www.string-db.org) and visualized using



Figure 1. Numbers of (A) upregulated and (B) downregulated differentially expressed genes in the sciatic nerve of T1- and T2DM mouse models. T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Cytoscape software (version 3.4.0; http://cytoscape.org/) (18). A combined score >0.4 was selected to determine significant interactions among DEGs. In accordance with a previous analysis (20), the connectivity degree of a protein, namely the number of proteins it connected with, was considered to indicate importance of the protein in the PPI network.

Module analysis was performed in the plugin ClusterONE (version 1.0) in Cytoscape with a threshold of P<0.001. In addition, function and pathway enrichment analyses were performed for DEGs in the modules with a threshold of P<0.05.

Results

Identification of DEGs. Based on the criteria of a nominal significance level of 0.05 and fold change ≥ 2 , 623 and 1,890 DEGs were identified in sciatic nerves of T1DM (GSE11343) and T2DM (GSE27382) mice, respectively, as compared with the control samples. Among them, 75 genes were identified to be coordinately dysregulated in sciatic nerves of both models, with 20 genes upregulated and 55 genes downregulated (Fig. 1 and Table SI). Meanwhile, 160 upregulated and 388 downregulated DEGs were unique to T1DM (Fig. 1 and Table SII), and 721 upregulated and 1,094 downregulated DEGs were unique to T2DM (Fig. 1 and Table SIII).

GO and pathway enrichment analyses. There were 60 coincident enriched GO terms for DEGs in T1- and T2DM, the top 20 of which are listed in Table I. In addition, the top 20 enriched GO terms for DEGs that were unique to sciatic nerves from the T1- or T2DM model are presented in Table II. Besides these,

Table SI. List of genes coordinately regulated in type 1 and type 2 diabetes mellitus.

Fable SII. List	of genes	specific to	o type 1	diabetes	mellitus.
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Gene Gene Ger count symbol cou		Gene count	Gene symbol	Gene count	Gene symbol
Upregi	ulated				
1	C1qtnf9	8	Mvo5b	15	Hipk3
2	Hspb7	9	Tef15	16	Hif3a
3	Fmo2	10	Ltbp2	17	Pde2a
4	Cyp1a1	11	Tlr7	18	Fkbp5
5	Gpihbp1	12	Map3k6	19	8430436 N08Rik
6	Mbd1	13	Angptl4	20	Dnah11
7	Gm4876	14	D8Ertd82e		
Down	egulated				
1	Kcna2	20	Pcyox11	39	Tmem229a
2	Kif5b	21	Rapgef4	40	Rian
3	Cct4	22	Sptssb	41	C1ql3
4	Fchsd2	23	5930427L02Rik	42	Chic1
5	Fgf14	24	Efcab14	43	Cntn3
6	Fndc5	25	Frzb	44	Map1b
7	Kcna1	26	Cemip	45	Slc35f1
8	Slc25a27	27	Zic2	46	Sox9
9	Galnt3	28	Zic1	47	Gpm6b
10	Ncald	29	Slc1a3	48	Slc47a1
11	Senp8	30	Snhg14	49	Elavl2
12	Asph	31	Bmp5	50	Ptn
13	Pdia3	32	Lamc3	51	Dcc
14	Bicd1	33	Nipal1	52	Atp13a5
15	Elovl4	34	C78859	53	Нрса
16	Npas3	35	Glipr2	54	Sox8
17	Opcml	36	Enah	55	Zic5
18	Nr2f2	37	Gm19935		
19	Ntrk3	38	Paqr8		

21 and 27 KEGG pathways were enriched for DEGs in T1- and T2DM, respectively. Among them, 4 pathways were coincident in both models (Table III), while 17 and 23 pathways were unique to T1- and T2DM respectively (Table IV).

PPI network. A total of 419 nodes and 1,343 edges were involved in the PPI for T1DM, whereas 1,416 nodes and 11,077 edges were involved in that for T2DM (data not shown). The top 10 proteins in each of T1- and T2DM with a relatively high degree of connectivity in the PPI network are presented in Table V.

Extent of enriched function and topological structure analysis for the modules of the PPI networks. Based on the module analysis by ClusterONE, a single module was determined in the PPI for T1DM (Fig. 2A) and 7 modules for T2DM (Fig. 2B-H). The information on the modules, including node number, P-value, protein with highest degree of connectivity, and numbers of GO terms and KEGG pathways are shown in Table VI.

Discussion

DPN is among the most frequent complications for T1- and T2DM. It is generally considered that DPN results from damage by hyperglycemia, regardless of the type of DM (3). In

Gene	Gene	Gene	Gene	Gene	Gene
count	symbol	count	symbol	count	symbol
Upreg	ulated				
1	Myh2	55	Wdr63	108	Bbd9
2	Doc2b	56	Gm17455	109	Klk1b24
3	Atxn711	57	Per2	110	Acer2
4	Tbata	58	Gimap8	111	Angpt2
5	Rph3al	59	Cpa3	112	BC003965
6	Apoc3	60	C77370	113	Rtp3
7	Nr5a2	61	Rgs3	114	Iffo1
8	Alpk3	62	Oosp1	115	Lhx6
9	Tall	63	Dbp	116	Tmem88
10	Aldh3b2	64	DII4 Manla12	11/	4930539E08Rik
11	Kasu2 Jign1	66	D7Bwg0826e	110	Aucyap111 Ananc11
13	Etnppl	67	Myct1	120	Clca3a2
14	Colec11	68	Ranbp17	121	Cvvr1
15	Ifi44	69	Adcy4	122	Lrpap1
16	Krt18	70	Shisa6	123	Cdh13
17	Gimap4	71	Asx11	124	Abcg1
18	Adgrf5	72	Ppp1r3d	125	Syt7
19	Spns2	73	Camk2n1	126	Jag2
20	Cabp4	74	Mboat1	127	Odf3b
21	D7Wsu130e	75	Arrdc4	128	Rilp
22	Ms4a2	/6	Ctsg	129	SOX /
23	EXOC312 Ms/182	78	PVr	130	9430030IN1/KIK
24 25	Csf2rb2	70	4833438C02Rik	131	Usp3
26	Ace2	80	Raserf2	133	Vps51
27	Itih4	81	Tnfsf13os	134	Keng1
28	Robo4	82	Gm11772	135	Krt20
29	Gm11716	83	4833411I10Rik	136	Rabgap11
30	4921507P07Rik	84	Mfng	137	Sphk2
31	Gp1bb	85	Sox17	138	1810021B22Rik
32	Vwf	86	D930003E18Rik	139	4930447M23Rik
33	Erg	87	Rtn4rl2	140	Arhgef15
34	Eng	88	Kank3	141	Cpsfl
35	Mettl22	89	Ptprb David2	142	Foxp4
27	Lamas 5720459M16D:1	90	Kpusu2	145	A otr1b
38	4833403116Rik	91	Sigiri Trim47	144	Hed17b11
39	Tceanc2	93	Adamdec1	146	Myrin
40	Fam107a	94	Cgrrf1	147	Gimap5
41	Slc27a2	95	Tigd3	148	Prkcg
42	Zfpm1	96	Spn	149	Abi3
43	Clec14a	97	Car7	150	Bmp4
44	Oasl1	98	Mettl13	151	Mgst2
45	Anxa8	99	Rap1gap	152	Ppp1r131
46	D130037M23Rik	: 100	2210019G11Rik	153	Selo
47	Fam189a1	101	6330403A02Rik	154	Sema3f
48	Clca3a1	102	Tmprss2	155	Mocs1
49 50	Mmrn2 Chfa2t3	103	AK8 Cyp11e1	150	NOS3 Pold1
51	Notab4	104	Ushbn1	157	AU022252
52	She	105	Slc6a7	150	D630045112Rik
53	Soat2	107	6330416G13Rik	160	Ngef
54	Unc45bos				- 8
Down	regulated				
1	Ccnt2	131	9530029O12Rik	260	Dag1
2	Col4a5	132	A330075M08Rik	x 261	Itm2a
3	Crkl	133	Abce1	262	Luc712
4	Dnajc18	134	Anp32a	263	Marcks
5	Ebf2	135	Arhgef6	264	Nktr
6	Lamb1	136	Chl1	265	Saa1
7	Pbrm1	137	Ср	266	Senp6
8	K3hdm4	138	Ctse	267	Serinc I
9	Kadiu	139	DCDId2	208	SITPI

Table SII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
10	Rbfox2	140	Eif4a1	269	Slc25a24	76	Epha7	206	Zfp871	335	Camk1d
11	Rsl24d1	141	Epb4112	270	Snx18	77	Fam49a	207	Araf	336	Gm2a
12	Sat1	142	Far1	271	Snx5	78	Gcc2	208	Brd4	337	Mirg
13	Slc25a3	143	Gatm	272	Tbl1xr1	79	Glg1	209	Cwf1912	338	Mob1a
14 15	Smarce1	144	Gpbp111	273	Z1C4	80	Kdelr2	210	Decr2	339	Peli2 Dimov1
15	Spry2 Suz12	145	HDa-a2 Hmger	274	Akap13	81 82	Lipep Mcf2	211	Hox0308 Hed17b12	340 341	PKIIOX I These 1
17	Timp?	140	Inngel Iof1r	276	Dtx31	82	Pcdhb15	212	Krit1	342	Trove?
18	Zfp626	148	Klhl28	277	Hsph1	84	Pum1	213	Larp4	343	Dyrk1a
19	Actb	149	Mboat7	278	Lin7c	85	Rhou	215	Mapre1	344	Efnb2
20	Amd1	150	Pddc1	279	Peg3	86	Ssbp2	216	Phip	345	Fgfr2
21	Arf1	151	Ppp1r10	280	Ppp4c	87	Stoml2	217	Rbms3	346	Gmfb
22	Arhgap29	152	Prpf39	281	Pum2	88	Syne2	218	Rhobtb3	347	Lcor
23	Atp6v1b2	153	Pura	282	Rad51c	89	Sypl	219	Tspyl1	348	Slc16a6
24	C030046G05	154	Rbm25	283	Scd2	90	Tetl	220	UstI	349	Sox6
25	CDX5	155	Knd3	284	SICI 2a2	91	Tripi Triad7	221	Apool	350 251	Srrm2
20	Cryzl1	150	Trps1	285	Sp7 Stk25	92	Trn53	222	Aseco Cask	352	Ogloa Arhgef11
28	D2Ertd640e	158	Ttc14	287	Txnip	94	Usp4	223	Cd247	353	Braf
29	Elavl1	159	Vma21	288	Usp48	95	Ywhaz	225	Cdh11	354	Dhcr24
30	Hist1h1c	160	Vps13a	289	Zfp825	96	Zbtb20	226	Cdkn2c	355	Golt1b
31	Il6st	161	Yipf4	290	AA409587	97	Zcchc3	227	Cpxm1	356	Otud4
32	Katnbl1	162	Add3	291	B4galt6	98	1810014B01Rik	228	Cyp20a1	357	Pfdn2
33	Lats2	163	Atxn7l3b	292	D9Ertd292e	99	Adamts15	229	Ercc4	358	Saa2
34	P4ha1	164	BC052688	293	Dusp3	100	Aplnr	230	Fbx13	359	Smg1
35	Pi4kb	165	C230076A16Rik	294	Klhl24	101	Arl8a	231	Ggact	360	Igf2
36	Plekhgl	166	Col8a1	295	Msn Daw 2 alt	102	Btbd/	232	Hnrnpl	361	Lsm12
3/	Polb	162	Cpeb2	296	Ppp3cb	103	Crim1 Eubr1	233	HOOK3	362	Malti
30	Rhhp4	160	Dab?	297	Canx	104	Fubpi Kansli	234	Msi2	364	Tial1
40	Rnf13	170	Hngds	299	Ccl2	105	LOC552902	236	Rab14	365	Zfp451
41	Slc25a30	171	Lrrc41	300	Ccl7	107	Lpp	237	Rbm12b1	366	Cxadr
42	Srsf11	172	Lrrk1	301	Dnm11	108	Mau2	238	Smek1	367	9430020K01Rik
43	Stard3nl	173	Nrk	302	Fbn2	109	Mmgt1	239	Vti1a	368	Gpc3
44	Strn3	174	Pbx1	303	Jmjd1c	110	Mtpn	240	1300017J02Rik	369	Serinc3
45	Tmem39a	175	Prkx	304	Nlgn3	111	Ntrk2	241	E030016H06Rik	370	E230029C05Rik
46	Upk1b	176	Rab23	305	Prkacb	112	Nuded1	242	Eif4ebp2	371	Rbm5
47	AI314180	177	Rbms2	306	Selk	113	Peak1	243	Gid4	372	Tgfbrl
48	Ccdcoo Chat12	170	Secola2	202	Smuri2 Smof1	114	PhI14 Dolb	244	HSG1/D/ Kat2h	3/3	Veen
49 50	Crist15	1/9	Srgap2 Steep4	300	SISI I Ser1	115	Polli Pom1k	243	Kif1b	375	v can Irf2bp2
51	Ddit3	181	Taf15	310	Cd209a	117	Psmb2	240	Man3k2	376	Dapl1
52	Gpc6	182	Tmtc4	311	Dhdh	118	Rrm2b	248	Nek4	377	Pdia4
53	Hrsp12	183	Usp9x	312	Fn1	119	Sfpq	249	Ppig	378	Uba6
54	Ikzf2	184	Wac	313	Hacd3	120	Sgcd	250	Rbm41	379	Wif1
55	Mbnl1	185	Actr2	314	Nbeal1	121	Sp3	251	Tmem45a	380	Slc22a8
56	Nit2	186	Cdk6	315	Ncaph2	122	Srpx	252	Vamp3	381	Ubfd1
57	Npr3	187	Cflar	316	Ss18	123	Stxbp4	253	March7	382	Eya1
58	Pja2	188	Ddi2	317	Zfp266	124	Tcf/12	254	9430053009Rik	383	SIC38a5
59 60	STOUPDP	189	Dnajc13	318 210	ZIP3611	125	Vps13b Wat5a	200	Abi2	384	Nnat
61	Siglech Sny27	190	Dyrk2 Etnk1	320	Adam10	120	VVIII.Ja Zeche24	250	Arglu1	386	Asgi i Ranbn3l
62	Stk3	192	Gadd45a	321	Arnc4	127	Zfp207	258	Cspg5	387	Sall1
63	Taok1	193	Khsrp	322	Atp8a1	129	Zkscan8	259	Cybb	388	Fabp7
64	Tgfbr3	194	Mpdz	323	Ddx6	130	5830407P18Rik		5		1
65	Tpbg	195	N4bp212	324	Malat1						
66	Trak2	196	Otub1	325	Nid1						
67	Ttc5	197	Rab31	326	Snhg11						
68	Whsc111	198	Sorbs1	327	Cx3cr1	recen	t decades disti	nct me	lecular function	ns and	1 morphometric
69 70	Zeb2	199	Tacc1	328	DIk1	ahno	rmalities betw	een T)PN in $T1_{-2n}$	d T71	DM have been
70 71	Atrx	200	1 mem80	329 220	EtV3 Mior1	renor	ted (78) In ed	dition	it has been pro	u 121	that the there
/1 72	Cupa Cup1	201	Wnt16	33U 321	Rsrc2	repor	a intorrection	for D	n nas ocen pro	posec	and on the the
73	Dmtf1	202	Zeche7	337	Sdf211	peuti	c interventions	IOL D	EIN SHOUID DITT		sed on the type
74	Dnaib6	204	Zdhhc2	333	Slc9a2	OI d18	abetes (/). For	UPN 1	n 11DNI, it has	been	suggested that
75	E130308A19Rik	205	Zfand5	334	Atic	ımpro	oved glycemic o	contro	may preserve i	nerve	runction and/or

Table SIII. List of genes	specific to type	2 diabetes mellitus.
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Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count
Upreg	ulated					66	Zranb3	307	Pik3r5	548
1	Sept2	242	Alcam	48	Bex1	67	Orm2	308	Abcc3	549
2	Tph2	243	Duoxa1	484	Cables1	68	Krt79	309	Fbp2	550
3	Lipf	244	Lrrc3	485	Ccl4	69	Cxcl2	310	Gm15498	551
4	Ighm	245	Tcerg11	486	Cnr2	70	Cdca5	311	Kiflb	552
5	Irdn Dm/22	246	Irx3	487	Epgn	/1	Adam8	312	Lpxn Dmana 1	553 554
07	PII 52 Ovtr	247	Lep Sla16a10	400	Lpcat5	72	Rub1	313	Slc16a12	555
8	Cna?	240	Clenn	409	Slc24a3	74	Agtr?	315	Stil	556
9	Gpr50	250	Cteflos	491	Tnfrsf23	75	Destamp	316	Nup210	557
10	Mbn13	251	Uhrf1	492	Bcl2l13	76	Myl4	317	Gdf5	558
11	Chil3	252	Ccl2	493	Cd320	77	C430042M11Rik	318	Serpina1b	559
12	Ucp1	253	Stc2	494	Pagr9	78	Stra61	319	Bche	560
13	Ubd	254	Gm6484	495	Pih1d2	79	Tshr	320	Ncaph	561
14	Atp1a4	255	Trav9d-3	496	Tmem252	80	S100a8	321	Stom	562
15	Ighg	256	Gm7694	497	Zwilch	81	Ncan	322	Hist1h3d	563
16	Dusp9	257	Iqgap3	498	1700013H16Rik	82	S100g	323	Tmem178	564
17	Cdkl4	258	Otop1	499	Dsg1b	83	Kenj15	324	2610020C07Rik	565
18	Hmga2	259	Ocstamp	500	Fabp5	84	Pkp2	325	Rps6ka3	566
19	Atp6v0d2	260	Mylk	501	Rhod	85	1600025M17Rik	326	Fam20c	567
20	Tfr2	261	Ube2t	502	Trim25	86	Gas2l2	327	Trabd2b	568
21	Gabrr2	262	C3ar1	503	Wfdc21	87	D830036C21Rik	328	Astlb	569
22	2010309G21Rik	263	Dhx32	504	Histlhle	88	SIc3/a2	329	Edem I	570
23	Mmp12	264	Cd200r1	505	Lmanl	89	Cenb2	330	Cdt1	571
24	111rn 0020610D08D:1	265	Cc122 Chat	507	Rtn2	90	Angpt13	222	Cenpa	572 572
23	9030019P08RIK	200	A 520020C20Dil	502	HOXCIS Ket222	91	Gnumb	332	AF231703	575
20	Alg1 Olr1	268	A530020G20RIK	500	NILZZZ Dfb1	92	Cldn15	334	Hn11	575
28	Slc5a7	260	AJ50055022Kik	510	Pik3an1	94	Cen55	335	Nepn	576
20	Ptchd4	209	4933427G17Rik	511	Sh3tc1	95	Abch4	336	Sel11	577
30	F7	271	Lat2	512	Srbx2	96	Gprin3	337	Cvp4b1	578
31	ll7r	272	Prkag3	513	Tmem120b	97	Car6	338	Muc13	579
32	Apoc4	273	Ddias	514	Tmem144	98	Brca1	339	Rhof	580
33	Spc25	274	Dgat2	515	Card11	99	Igkv15-103	340	Mcm10	581
34	Mogat2	275	Trim67	516	Etf1	100	Tinag	341	Tnfrsf11a	582
35	Lppr4	276	Cenpe	517	Plet1os	101	Trim29	342	Ppih	583
36	Crisp2	277	Cd164l2	518	Cd68	102	Cela1	343	Rasgrp2	584
37	Npr3	278	Mest	519	Cdca2	103	Mki67	344	Srp54b	585
38	Rnase2a	279	Mgll	520	Ccnd2	104	Cdk1	345	Gzme	586
39	Rgs1	280	Slc8al	521	Dep2	105	Lrrc39	346	Fam126b	587
40	4930502E18Rik	281	Gdf3	522	Fgd4	106	Dtl LL t2 2	347	Heph G1200-4	588
41	KII14 Certa a 1	282	V S1g4	525	Foxm1	107	Ugisaz Duen2	240 240	Ca200f4	389 500
42	Sympo2	200	Adoral Illaro2	524 525	GIII05//	100	Tmem182	349	Chinin2 Ghr	501
43	Synpoz Itgay	285	Tmem37	525	Ullars Il?ra	110	Acov1	351	Cennw	592
45	Sez6l	286	Cd1d1	527	Llol2	111	Chrna2	352	Mcm5	593
46	Cdca7l	287	Kif2c	528	Nckap11	112	Apoc2	353	Ska3	594
47	Serpine1	288	Rrm2	529	Serpina3m	113	Fzd4	354	Mrgprg	595
48	Hmmr	289	Cks2	530	Snta1	114	Fads3	355	Pcdh7	596
49	Iglv1	290	Pdcd11g2	531	Hpdl	115	Arhgap25	356	2810408I11Rik	597
50	Fam110c	291	Ffar2	532	Knstrn	116	Dock8	357	E2f8	598
51	Kcnj14	292	Stap1	533	Nedd1	117	Ccnb1	358	5430437J10Rik	599
52	Prr16	293	Pm20d1	534	Nf2	118	Gm6277	359	Kif18b	600
53	Cldn23	294	Clec4a2	535	Pitpnm2	119	4930486L24Rik	360	Akr1c13	601
54	Cyp2r1	295	Fmr1nb	536	Pole	120	Heph11	361	Cd300a	602
55	Mc2r	296	Blnk	537	Thbs2	121	Tpx2	362	Cd84	603
56	Grtp1	297	Tnfrsf12a	538	Cd300lb	122	Slpi	363	Lrrc27	604
57	4930480G23Rik	298	Vegfc	539	Cdc42ep1	123	Sirpbla	364	Mcempl	605
58	Dest1	299	Acaalb	540	Gm30173	124	IIrI3	305	Adam12	606
39 60		300 201	AIS2CT12	541 542	igiais Vot7	125	0120204L05D	200 267	CKap2 Eam101b	00/ 600
00 61	Aauac 23100021.00D 11-	301	Cat	542 542	Kann ^A	120	>130204L03KlK	360	i amitutu Jasef6	600
62	2310002L09Klk	302	Upi Haver?	545 547	Pmch	127	Rhm28	360	Pramef12	610
62 63	Cyp2C70 Nek2	303	Marco	544 545	Fet13	120	Dnmt31	370	Theref??	611
64	Dnaih13	305	4931406C07Rib	546	Car4	130	Sucnr1	371	Ttk	612
65	Gsdma2	306	Dnmt3a	547	Cd180	131	Atp6v0a1	372	Car12	613
					-	-	ĩ	-		-

Gene

symbol

548 Comt 549

Fsd2

Itgb1

Plk4

Spc24

Atp2a3

Myo1f

Pla2g12a

Pou2af1

Gm15972

Rp1391

Usp29 Arl4a

Ercc61

Shmt1

Atp10a Glb112

Slc2a10

Smoc1

Zfp106

Arfgef2

Gent1

Hesx1

Mrc2

Plk1

Neurl2

Zfp426

Clip1

Dyrk2

E2f7

Pex16

Qpctl Slc22a4

Tmem132a

D17H6S56E-5

Adam23

Ece2

Lipa

Plek

Tgfb1

Fgfrl1

Mutyh

Nim1k

5830403F22Rik

Ptprk

Aurka

Fignl1

Isoc1

Tgm2

Hvcn1

Mpeg1

Sphk1

Tpsab1

Rab20

Skil

Trappc13 Wipi1

Copg2os2

Lss

Adamts9

AI844685 Ckap2l

Ccl7

1700017B05Rik

A930018P22Rik

Tns1

Table SIII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
132	Irak2	373	Depdc1a	614	Gatb	198	Apol6	439	2010003K11Rik	680	Nlrc3
133	Prss27	374	Fam64a	615	Gm11827	199	Rrbp1	440	Plekhg6	681	Gm9899
134	Birc5	375	Hpgd	616	Ptpn22	200	Pex51	441	Kif22	682	Phtf2
135	Mis18bp1	376	Dbf4	617	Stx17	201	F10	442	Ndc80	683	Tead4
136	Spaca1	377	Igsf21	618	Trib3	202	Cena2	443	NIrp10	684	Tfcp211
137	Adh6b	378	Kcna5	619	E2f2	203	Tmem135	444	Ptges31	685	Ttpi2 Mon245
138	AIP Th1	3/9	AI481207	620 621	Itsn1 C2ad2	204	Gm8350	445	Abbd15	687	Map2K3 Sh2h2
140	Cxcl3	381	Gla	622	Cd44	205	Nus1	447	Acsl1	688	Spint2
141	Sdsl	382	Hrct1	623	Hfe	200	Tm4sf5	448	Ltc4s	689	Wdr62
142	Tsc22d2	383	Naip1	624	Mmp19	208	Cdr2	449	Crk	690	4930511E03Rik
143	Klk1	384	Slc18a1	625	2500002B13Rik	209	Aurkb	450	Naip2	691	B230307C23Rik
144	Fosl1	385	2210406H18Rik	626	2700099C18Rik	210	Lonrf3	451	Peg3	692	Cib2
145	Rnf128	386	3930402G23Rik	627	Ano9	211	Bcar3	452	4430402I18Rik	693	Olfm2
146	Cdca8	387	Adap2	628	Ccne1	212	Dsg1a	453	Fgf21	694	Zfp655
147	Timp4	388	Palmd	629	Lox	213	Peg10	454	Lrrc28	695	Cenpk
148	Zc3n12d Dlin5	389	Vidir Mfga8	030 631	Dyniro2 Env2	214	Clecke	455	Piekiii2 Rad51ap1	690 607	DgKu Hena12a
149	Ttc39b	390	Mpp7	632	Gm3912	215	Galnt6	450	Sgol2a	698	Hspa12a
151	AA467197	392	Sgol1	633	Lipg	217	Gvs2	458	Ehd2	699	Morn5
152	Tenm4	393	Mmp13	634	Maff	218	Melk	459	St8sia4	700	Rbp7
153	Nuf2	394	Ms4a6d	635	Milr1	219	Spag5	460	Dcbld1	701	Chpt1
154	8430408G22Rik	395	Tatdn2	636	Tmem82	220	Gsdma	461	Mtor	702	Ehd3
155	Cdca3	396	Pigt	637	2600006K01Rik	221	Glipr1	462	Plaur	703	Hk3
156	Hilpda	397	Sbk1	638	Acvr1c	222	Pde7a	463	Akap2	704	LOC545086
157	Esco2	398	Lrg1	639	Bcl2110	223	Top2a	464	Cenf	705	Mta3
158	SIc9a6	399	Oit3	640	Itih5 Distations 1	224	Rab38	465	LTT1 Not1	706	Spry4
159	Irem2 Pkp1	400	Qsox1	641 642	Plekhm I Slo7a8	225	Adn4 Cxcl10	400	Nat1 Gm31718	707	Gpr65
161	Tex33	401	v wao Atp8b4	643	Tes	220	Kif11	468	Kif20a	708	Rel
162	Aap7	403	B430306N03Rik	644	Cd6	228	Hrasls	469	Optn	710	Stk17b
163	Fmr1	404	Msr1	645	Gsg2	229	Kcnk3	470	Plod1	711	Zfp69
164	Fggy	405	Nfe2l2	646	Gm2011	230	2810417H13Rik	471	Rad54b	712	Ccr7
165	BE949265	406	Nkd1	647	Incenp	231	1700029I15Rik	472	AU021933	713	Cenpn
166	Ube2c	407	Ptx3	648	Pold3	232	Serpina3a	473	Ccrl2	714	Fbxo42
167	Pbk	408	Cd22	649	Bst1	233	Lamb3	474	Piezo1	715	Ttpa
168	Wif1	409	Galns	650	Opte	234	PletI	475	Stab2	716	Vac14
109	Inedd9	410 411	Aimia Cd300lg	652	Smado Tmem206	235	RID Btc	470	C5ar1 Hof	718	Nc11 Vbx2
170	Slc7a10	411	Ermp1	653	7fp787	230	Cede125	478	Kbtbd12	719	Cd3001d
172	Spd11	413	Fcgr4	654	Cdc20	238	Rbpms	479	Map7	720	Gk
173	Tmed5	414	Shcbp1	655	Ripk3	239	Mettl24	480	Rad18	721	Xrcc3
174	Slc15a3	415	Knop1	656	Sdcbp2	240	Tnc	481	Sdk2		
175	Clec4d	416	Serpina12	657	Myo19	241	Tpcn2	482	Tlr8		
176	Serpinb9b	417	Larp1b	658	Tinagl1	Down	regulated				
177	Cux2	418	Thbs1	659	Tmem106a	1	March2	366	Tspan2	731	St6gal2
178	Cdk18	419	Seh11	660	1rim16	2	March4	367	Ttyh1	732	Sytl2
1/9	C0 Lhfnl2	420	Lgaiss Nok6	001 662	5055450115K1K	3	March5	368	Unc13a	733	Tnfrsf19
181	Scel	421	Ctse	663	Dolk	4	March6	369	Ztp612	734	Agr2
182	Glyctk	423	Hsd11b2	664	Laptm5	5	2900055J20KIK Roll11b	370	0330184L 24Dik	736	$C_{03003}/E_{1}/P_{1}$
183	Mms221	424	Itgb2	665	Lats2	7	Cdk7	372	Aak1	737	Costo Costa
184	Tcp11x2	425	Slamf7	666	Lbp	8	Esr1	373	Adamts11	738	Dcdc2a
185	Cacnale	426	3110027N22Rik	667	Ormd13	9	Gm13446	374	Aif11	739	E130002L11Rik
186	Hipk2	427	Dmrt2	668	Pde1b	10	Gpr85	375	Ank3	740	Fam131a
187	Ccl3	428	Hspb6	669	Prelp	11	Hoxb9	376	Atl1	741	Fam167a
188	Fabp12	429	Ptbp3	670	Serpina3c	12	Htr7	377	B930095G15Rik	742	Fam169a
189	Ncapg	430	SIC22a12	671	Hr VIL125	13	Kctd14	378	C030039L03Rik	743	Gap43
190 101	rupiaz Dopa3	431 432	Tic/ Cenni	072 673	Shb	14	Mapk4	379	Cacnb3	744	Gng4
192	Elov13	433	Cxcl9	674	Vav1	15 16	Ncdn Ochrol2	380 201	Calb2	745	Kendl Vætden
193	Prss46	434	Itga7	675	6030422H21Rik	10 17	Osupio Pfn2	201 202	Cdb8	740 777	Krituap I rrn1
194	Mmp8	435	Lcn2	676	Ccdc86	18	Pou3f2	383	Clasp?	748	Lysmd4
195	Folh1	436	Pid1	677	Cenpp	19	Slc14a1	384	Dlg4	749	Mmd2
196	Cenpf	437	Prc1	678	Gch1	20	Tardbp	385	Epb4114b	750	Nacad
197	Npy	438	1700112E06Rik	679	Ms4a6b	21	4833423F13Rik	386	Fbxo30	751	Necab1

Table SIII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
22	Angel1	387	Gpr61	752	Nrxn1	88	R3hdm1	453	Chd5	818	Nrsn1
23	BC030500	388	Itgb3bp	753	Nxpe3	89	Runx1t1	454	Dgkg	819	Pcsk1
24	Cacng4	389	Lcat	754	Parm1	90	Sall4	455	E130309D14Rik	820	Pcyt1b
25	Chd9	390	Lect1 Magi1	755	Ptprt	91 02	SIc29a4 S1c35f2	456 457	Hapin2 Myt11	821	Rnf15/ Serpini1
20	Drys13	391	Mag11 Meg3	750	Pvalo Rem2	93	Trank1	458	Ntsr2	822	Slain1
28	Ece1	393	Mmp9	758	Rps6ka6	94	Zdhhc18	459	Oprl1	824	Slc35f4
29	Eif5a2	394	Neto2	759	Sphkap	95	Zfp318	460	Prdm8	825	Slc4a10
30	Endod1	395	Otud7a	760	Syt11	96	Zfp37	461	Prr18	826	Vps13c
31	Epha5	396	Phactr1	761	Th	97	2900009J20Rik	462	Psmf1	827	Wt1
32	Evi5	397	Prkcz	762	Thy1	98	A930041H05Rik	463	Rab3a	828	Adamts16
33	Gjd2	398	Psd2 Dref120	763	Tmem/4	100	Ak5	464	Rasget1a	829	Ano3
34 35	Gpr88	399 400	Kni150 Somo3b	765 765	Trim3 /	100	Arg2 Bank1	465	Rgillo Rgs7bp	831	Arlgels
36	Hsosi2 Htr1d	400	Sema4g	766	Winf3	101	Ccdc28a	467	Robol	832	Cntn1
37	Icall	402	Slc2a3	767	Byes	102	Cdkn1b	468	Rph3a	833	Ddx25
38	Kalrn	403	Smr2	768	Ccdc92	104	Clec2l	469	Slc2a12	834	Dzank1
39	Kenmb2	404	Srr	769	Elfn1	105	Csmd1	470	Slc6a2	835	Fam19a3
40	Kcnv1	405	Trpm3	770	Erich3	106	Dip2a	471	Sstr4	836	Gprin2
41	Mag	406	Zfp385b	771	Galnt5	107	Dlgap1	472	Syt7	837	Hsbp111
42	Nxph1	407	A830010M20Rik	772	Gca	108	F5	473	Tdo2	838	Htr4
43	Osbpl6	408	AI661384	773	Ggt7	109	Fzd3	474	Tmem108	839	Jakmip1
44	Pax3 Dodb0	409	A1854703	//4 775	Grial Hawh5	110	Htria	475	vwaba Zfr2	840 841	Jakmip2
45	Pcdn9 Pcp/11	410	Aw040457 Adam22	776	HOXD3 Kenhl	112	Klhl32	470	9530077C05Rik	842	Lix1
40	Pter	412	R3gat1	777	Kenin1	112	L1cam	478	Acsl6	843	Maneal
48	Rgs7	413	Bajap211	778	Kenmb1	114	Lrrtm2	479	Aqp4	844	Npy2r
49	Scmh1	414	C530044C16Rik	779	Lrfn5	115	Map6d1	480	Atcay	845	Rab3c
50	Slc1a6	415	Chrnb2	780	Mab2112	116	Mfsd6	481	Ccdc68	846	Reep2
51	Smco3	416	Cntnap5a	781	Necab3	117	Mrpl35	482	Cdk5r1	847	Rufy2
52	Sostdc1	417	Cplx2	782	Nyap1	118	Nedd41	483	Cgref1	848	Sh3gl2
53	Stxbp1	418	Crh	783	Sptbn2	119	Pcsk6	484	Cpb1	849	Vstm5
54 55	Stxbp5	419	Csdc2	784	Syn1 Tmom151a	120	PIXIDI Pop1r2	485	Cund2	850 851	1810041L15K1K
55	Then2290	420	D2Frtd282e	786	Tillelli151a	121	Rbm33	487	Dnah7a	852	Aldoc
57	Upp2	422	Dclk1	787	Tro	123	Rfx4	488	Dzip3	853	Asic1
58	Zfp788	423	Eras	788	Trpa1	124	Sacs	489	Fndc4	854	BC059841
59	5330417C22Rik	424	Erich6	789	Agtr1b	125	Serpina11	490	Fxyd7	855	Car10
60	A730089K16Rik	425	Gm36529	790	Calca	126	Shd	491	Gabra5	856	Ccdc172
61	Abhd10	426	Greb11	791	Camta1	127	Tcte2	492	Glrb	857	Cldn10
62	Adgral	427	Grm4	792	Crmpl	128	Tial T2	493	Lrchl	858	D130043K22Rik
64	C/9242	428	Hopx	793	Fam131b	129	Ttbk1	494 705	Mir124a-Ing Neam2	859 860	Gabrg2 Gm16551
65	Cen97	429	Nalen	794 795	Gabra?	130	Ttc3	496	Plk5	861	Kenin2
66	Cpsf2	431	Nebl	796	Kend3	131	Vamp4	497	Pogz	862	Kenip4
67	Cpsf6	432	Nova1	797	Lmo3	133	2900056L01Rik	498	Prox1	863	Kif3c
68	Csmd3	433	Pcp4	798	Map3k9	134	2900064F13Rik	499	Rasl10b	864	Mrgpra3
69	Cyfip2	434	Plekha6	799	Mpped2	135	Arhgap20	500	Rnf208	865	Pappa2
70	Eml2	435	Rab6b	800	Ngfr	136	Cede112	501	Scin	866	Rab39b
71	Fat3	436	Rbfox3	801	Nmnat2	137	Cfap69	502	Scn8a	867	Rassf6
72	Fg118 Calaa7h	437	Rbm11	802	Prkar1b Dime2	138	Cmtm5	503 504	SCITI Slitela	808 860	KDIOX1 Pufy2
75	Golga70 Gpr173	438	Rgso Rtn/	803 804	Rundc3b	139	Dua Dusp15	504	Susd5	870	Sez612
75	Grin1	440	Sarm1	805	Scn4b	140	Ephx4	506	Tmem169	871	Soga3
76	Hs3st1	441	Sema3e	806	Slc5a5	142	Faim2	507	Tmem196	872	Stac
77	Irf4	442	Shisa4	807	Tesc	143	Gabrb3	508	Ttc39c	873	Sv2a
78	Klf12	443	Slc6a17	808	Trpv1	144	Galnt14	509	Tyrp1	874	Tmem179
79	Lancl1	444	Slco1c1	809	Tst	145	Gm12371	510	Unc5c	875	Tspyl4
80	Lrba	445	Synm	810	Ube2ql1	146	Gm13889	511	Vwc2l	876	Aplp1
81	Lrrn4cl	446	Tubg2	811	Zfp365	147	Gprasp1	512	5930412G12Rik	877	Atp6v1g2
82 82	LITUM4	44/ 110	Wari / 02201921 06D:1-	812 812	ZIP804a	148 140	rispa4i Josf11	515 517	Abcg4	0/8 870	rarz Megf11
03 84	Pagr5	448 449	aifm3	013 814	∠700040A07KIK AF529169	149	Itih3	514	Acsm3	880	Nso2
85	Plch1	450	Bhlha9	815	B3galt5	151	Kenip3	516	Arhgap12	881	Pclo
86	Prokr2	451	Cd200r3	816	Bean1	152	Kcnk1	517	Cabp7	882	Pou4f2
87	Prr51	452	Cdh1	817	F2rl2	153	Lingo1	518	Ccdc30	883	Ptprn

Table SIII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
154	Mkx	519	Clip3	884	Rasgrp1	220	Ism1	585	Dok4	950	Pacsin1
155	Nipa1	520	Clip4	885	Rit2	221	Jam3	586	Epb4113	951	Rims3
156	Phex	521	Dgkb	886	Slc1a2	222	Klf7	587	Exph5	952	Rspo2
157	Pnkd	522	Diras1	887	Them5	223	Ltbp1	588	Fez1	953	Rundc3a
158	Ppm11 Spink10	523 524	Elovi2 Gm10700	888	Tmem151b Zwint	224	MapK8ip1	500	Fmn2 Cria2	954	Scg5
160	St3gal6	525	Gpr149	890	6430604M11Rik	225	Nrin3	590	Kif5c	955	Sicoars Srrm3
161	St6galnac3	526	Has2	891	Bsn	227	Phyhip	592	Mast1	957	St8sia3
162	Stk33	527	Hoxd10	892	C530008M17Rik	228	Prepl	593	Ms4a3	958	Tram111
163	Tfap2a	528	Hs3st4	893	Cadps2	229	Rasgrf1	594	Ndrg4	959	Zswim5
164	Wdr31	529	Iqsec3	894	Cbln2	230	Serpind1	595	Pirt	960	2900011008Rik
165	4930519N06Rik	530	Kazn	895	Ctnna2	231	Sox1	596	Prmt8	961	Astn1
167	Acsog1	532	Kcn11 Kif3a	890 807	Cyp4x1 Elavl3	232	Sox2ot Spock2	508	Rab33a Rab37	962 063	Atp202 Bend6
168	Asx13	533	Lepr	898	Elavis Ebxo2	234	Susd2	599	Slc25a12	964	Bex2
169	Atad1	534	Lingo2	899	Gm10419	235	Syndig11	600	Srgap3	965	Cacna2d3
170	B3galt1	535	Mapt	900	Igfbp2	236	Zfhx2	601	Tmem163	966	Cckar
171	B630019K06Rik	536	Mycl	901	Jph3	237	A730054J21Rik	602	Tmem79	967	Cend1
172	Ccdc138	537	Neto1	902	Jph4	238	Adam11	603	Ttc9b	968	D930028M14Rik
173	Cdr2l	538	Nipal3	903	Klhdc8a	239	Adcy1	604	4930524007Rik	969	Dnm3
174	Cur61	539 540	NW02 Ddzd4	904 005	Nrcam Pren2e	240	Adgr13	605 606	A330050F15R1K	970 071	Fam155a Gnal
176	Dhh	541	Plexd3	906	Ppm1i	242	Appa1	607	Arhgan28	972	Inw
177	Edil3	542	Plp1	907	Ptgs2	243	Apcdd1	608	Arhgap44	973	Kcna4
178	Fign	543	Ppfia2	908	Tcte1	244	B3gnt5	609	Celf4	974	Kcnj10
179	Gabbr1	544	Ptbp2	909	Tmem150c	245	Bmp7	610	Cntnap4	975	Lsamp
180	Gabra1	545	Scrn1	910	Trim9	246	Crym	611	Elovl7	976	Mapk8ip2
181	Gm13629	546	Slc25a18	911	Acpp	247	Dbndd2	612	Ferls	977	Mtcl1
182	GIII3089 Gnaz	547 548	Steodal	912 913	Augros Anks1b	248 240	Eerraz Fam81a	614	Gpro7	978	Phil24 Snap25
184	Gpr62	549	Svop Svngr1	914	Ccdc184	250	Fbx116	615	Ibsp	980	Tmem591
185	Hcn1	550	Tmem246	915	Chga	251	Fbxo41	616	Ints6	981	Atp2b3
186	Kenk5	551	Tmem88b	916	Ckmt1	252	Flrt1	617	Lgi3	982	Col25a1
187	Kenq5	552	Arpp21	917	Galnt13	253	Gkn3	618	Lrrtm1	983	Dpysl5
188	Lrrc49	553	Atp1b1	918	Gng3	254	Hey2	619	Magee1	984	Grip1
189	Mbp Dala1	554	Cd209g	919	Kcnabl Konal	255	Kend2	620	Pcdh20	985	Hapini
190	Pax9	555	Cua Chst5	920 921	Kilf5a	250	L in7a	622	Plekhh1	980	I hfpl4
192	Pcdh10	557	Cnnm1	922	Napb	258	Lrrtm3	623	Prkg2	988	Nap113
193	Praf2	558	Dleu2	923	Nrxn3	259	Ncam1	624	Prune2	989	Ppm1e
194	Prss12	559	Dscam	924	Ptgds	260	Nrsn2	625	Rcan2	990	Prph
195	Ptchd2	560	Dusp26	925	Ptprn2	261	Pak7	626	Scn2b	991	Rnf182
196	Rimklb	561	Hebp2	926	Rgs17	262	Pcolce2	627	Slc7a14	992	Rtn1
197	SgtD Sh3d10	563	II31ra L hfp13	927	IOX3	263	Ppp2r2c	628	StX1b Sult4e1	993	SIC39a12
199	Slc16a9	564	Lmpi5 Lm8	929	Zmat4	265	S100b	630	Svt14	995	Zechc18
200	Smpd3	565	Mfsd4	930	Csrnp3	266	Samd14	631	Tmem132e	996	Ano4
201	Sorcs3	566	P2rx3	931	Dpp6	267	Sh3g13	632	Ubash3b	997	Arhgef7
202	Stmn2	567	Pigz	932	Fam184b	268	Slc6a11	633	A830018L16Rik	998	BC048546
203	Usp31	568	Pnma2	933	Gdap1	269	Trio	634	Adcy8	999	Gabrg1
204	Vamp1 Vat11	569 570	Rab39	934	Kctd8	270	2610100L16Rik	635	Cdh10	1000	Hecw1
205	Zdbf2	570	Rlbn1	935	Rnf112	271	Another Apple 1	637	Chrna6	1001	Map7d2
207	Aldh1a2	572	Sema4f	937	Syt2	273	Ccl27a	638	Eno2	1002	Mog
208	Atp2b4	573	Slitrk5	938	Tmem200c	274	Ccp110	639	Fgf4	1004	Pcsk2
209	Cacnb4	574	Smarcal1	939	3632451006Rik	275	Cd24a	640	Frrs11	1005	Pgbd5
210	Camk1g	575	Sntg1	940	9430021M05Rik	276	Clvs2	641	Gm5124	1006	Rprm
211	Chn1 Dio2cc	576	Stmn3	941	AW060742	277	Cntn6	642	Gpr22	1007	Sgpp2
212	D10308 Ekbn1b	570 570	Susd4 Tmom117	942 042	Anki Asio3	278	CxcII3	643 644	IIIr2 Lanal3	1008	SIC1/a/ S122627
215 214	Gib1	570 579	Vwa7	743 944	Brsk2	∠19 280	Dende5	645	Ogfrl1	1010	Spock1
215	Gna14	580	1700019D03Rik	945	Cldn11	281	Dkk2	646	Olfm3	1011	Adrb3
216	Herc3	581	A2m	946	Faxc	282	Gabrb2	647	Phactr3	1012	Akap6
217	Hnrnpr	582	Brsk1	947	Fstl5	283	Gm15663	648	Reep1	1013	C030017B01Rik
218	Igsf1	583	C030011L09Rik	948	Lhfpl5	284	Hs3st2	649	Rgs4	1014	Calb1
219	Irf6	584	Cabp1	949	Nptx2	285	104	650	Kimbp2	1015	Cpne4

Table SIII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
286	I118	651	Skida1	1016	Dync1i1
287	Map1a	652	Slc6a1	1017	Isl1
288	Mllt11	653	Slitrk1	1018	Isl2
289	Nfasc	654	Tmem56	1019	Kndc1
290	Pak3	655	Trnp1	1020	Map4
291	Pgm211	656	Tspy15	1021	Mapk10
292	Picd4	007 658	v stm2a	1022	Pnynipi Pou4f1
293	Rasaiz Rltpr	659	A330068G13Rik	1023	Rims1
295	Sec62	660	Aard	1025	Sv2b
296	Snph	661	Acsl3	1026	Syt1
297	Stk32c	662	Atp8a2	1027	Syt4
298	Syngr3	663	Cacna1b	1028	Tubb4a
299	Tbc1d30	664	Cdh9	1029	Amph
300	Trim2	665	Ceslf	1030	Elavl4
301	1110032F04R1k	666	Col4a6	1031	Ermn
302	Ank2	00/ 668	Dner En 3k	1032	Grap
304	Apc2 Apoh	669	Gm16532	1033	Mlc1
305	Asphd1	670	Itga8	1035	Mup10
306	BB319198	671	Kif1a	1036	Nrg1
307	Cacna2d2	672	L1td1	1037	Scn10a
308	Cadm3	673	Megf10	1038	Tagln3
309	Clgn	674	Mtus2	1039	Vgf
310	Cmah	675	Nipal2	1040	Add2
311	Ddn	676	Nrxn2	1041	Cdhrl
312 212	Dhrs2 Efeeb1	679	NtrK1 Dodb8	1042	Chrna/
313	Elcaul Fam180a1	679	Pla2g3	1043	Dlg2
315	Gm32444	680	Rab9b	1044	Drg2 Dpp10
316	Gpr158	681	Sowahb	1046	Fam19a4
317	Gpr75	682	Trpc3	1047	Grik1
318	Grm8	683	Trpc6	1048	Htr3a
319	Inadl	684	Tvp23a	1049	Plekhd1
320	Kenq2	685	Vwa5b2	1050	Ppp1r1c
321	Ketd4	686	A730017C20Rik	1051	Ppp2r2b
322	Kif2la Kat27	687	Amer3	1052	Kalyl
323 324	Kft2/	680	Arngel4	1055	Stosiai Stmp/
325	Lonrf?	690	Ceser1	1054	Synpr
326	Lztfl1	691	Cntnap2	1055	2900052N01Rik
327	Mfsd2a	692	Cplx1	1057	Chgb
328	Mro	693	Crygs	1058	Fam19a2
329	Pcnxl2	694	Dock3	1059	Resp18
330	Prkg1	695	Dpp4	1060	Slc17a6
331	Srd5a1	696	Fam189b	1061	Spock3
332	Srs112 Tmom231	697 608	Gdap111 Gm2115	1062	V snl l Noff
334	Tnik	699	Gnrasn?	1063	Scn9a
335	Trim36	700	Hoxb6	1065	Nts
336	Ttbk2	701	Lgil	1066	Sult1e1
337	Aox3	702	Mobp	1067	Avil
338	Ccdc47	703	Mogat1	1068	2810037O22Rik
339	Celsr2	704	Mtmr7	1069	Nefh
340	Cldn9	705	Nkx2-2	1070	Cdk5r2
341	Dbndd1	706	Omg	1071	Ttc9
342	Esd Exud2	707	Pdelc Dterral	1072	Adcyap1
345	Greb1	708	Sep1a	1073	Slc24a2
345	Gria4	710	Slc35f3	1074	Nefm
346	Kenn2	711	Spire2	1076	Panx2
347	Kent1	712	Stxbp51	1077	Syt9
348	Lynx1	713	Tenm1	1078	Rimkla
349	Ndp	714	Timd4	1079	Gm2102
350	Nkain2	715	6330563C09Rik	1080	Scg2
351	Pi16	716	Ap3b2	1081	Tac1

Table SIII. Continued.

Gene count	Gene symbol	Gene count	Gene symbol	Gene count	Gene symbol
352	Pip5kl1	717	Arhgap36	1082	Prrx11
353	Pla2g2d	718	Caln1	1083	AI593442
354	Ppl	719	Celf6	1084	Calcb
355	Prokr1	720	Cntn4	1085	Shh
356	Rab27b	721	Corin	1086	Tmem130
357	Rragd	722	Epb41	1087	Cyp2f2
358	Rsrp1	723	Fosb	1088	Snap91
359	Scrt2	724	Kenj3	1089	Nap112
360	Slc35d3	725	Kcnk10	1090	Scn11a
361	Spink2	726	Lrrc75b	1091	Myt1
362	Stk39	727	Morn4	1092	Grm7
363	Synrg	728	Onecut2	1093	Tubb3
364	Tmem255a	729	Pcsk1n	1094	Cadps
365	Tshz2	730	Scg3		

decrease the likelihood of developing DPN (6). By contrast, for DPN in T2DM, not only glycemic control, but treatments for other risk factors including obesity may be equally critical in preventing DPN (6). Therefore, since the molecular mechanisms underlying the pathogenesis of DPN in each type of DM may differ, understanding the mechanisms may lead to novel therapeutic approaches for prevention or treatment.

In the present study, there were marked differences between the gene expression profiles, GO terms and KEGG pathways of DEGs in the sciatic nerves of T1- and T2DM mouse models. This was in accordance with previous studies revealing considerable alterations in global gene expression profiles of several tissues including skeletal muscles and adipose tissue (21), and organs such as the intestine (22), liver (23) and brain (24) in the two types of DM. It is established that T1DM is characterized by destruction of pancreatic islet cells by autoimmunity, with loss of pancreatic insulin production, while T2DM is a metabolic disease with high pancreatic insulin production in the setting of insulin resistance (6). Therefore, the different insulin level and insulin signaling pathways between the two types of DM may converge on and modulate the transcription of genes (25); it thus seemed reasonable to search for altered gene expression patterns in DPN between T1- and T2DM models characterized by absolute or relative insulin deficiency.

A number of DEGs unique to the sciatic nerve of T1DM mice were associated with cell proliferation, as well as the mitogen-activated protein kinase signaling pathway, which is a critical pathway for cell proliferation (26). In neuropathic pain, a correlation has been determined between the proliferation of nerve cells including microglia and astrocyte and the abnormal pain responses (27,28). In addition, previous study has demonstrated that the proliferation of glia cells including astrocyte was associated with structural changes in the nervous system, such as axonal growth (29). Furthermore, with loss of C-fibers, increased frequencies of denervated Schwann cells and regenerating fibers have been identified in T1DM mice but not in T2DM mice (8). Therefore, the DEGs associated with cell proliferation may serve a crucial role during the pathological changes of DPN in T1DM.

Insulin resistance is among the major factors that leads to the development and progression of complications in T2DM (30).

		T1DI	М	T2DM		
GO ID	GO term	Gene count	P-value	Gene count	P-value	
GO:0005515 MF	Protein binding	57	3.1E-6	76	9.2E-20	
GO:0005737 CC	Cytoplasm	21	6.7E-6	123	3.8E-14	
GO:0001525 BP	Angiogenesis	14	9.6E-6	89	7.3E-18	
GO:0000166 MF	Nucleotide binding	45	2.4E-5	46	5.5E-10	
GO:0007399 BP	Nervous system development	21	3.2E-5	86	7.3E-11	
GO:0008201 MF	Heparin binding	13	4.6E-5	57	5.9E-14	
GO:0006468 BP	Protein phosphorylation	11	5.4E-5	61	4.3E-12	
GO:0005578 CC	Proteinaceous extracellular matrix	24	5.5E-5	66	9.1E-20	
GO:0016310 BP	Phosphorylation	13	6.2E-5	78	8.8E-16	
GO:0009986 CC	Cell surface	12	6.5E-5	54	5.6E-17	
GO:0016020 CC	Membrane	17	8.9E-5	23	3.2E-15	
GO:0005794 CC	Golgi apparatus	12	2.4E-4	45	2.5E-17	
GO:0004672 MF	Protein kinase activity	14	3.1E-4	24	4.2E-16	
GO:0030335 BP	Positive regulation of cell migration	15	4.5E-4	19	8.5E-17	
GO:0005783 CC	Endoplasmic reticulum	6	5.3E-4	32	5.2E-16	
GO:0019933 BP	cAMP-mediated signaling	18	5.8E-4	30	7.5E-13	
GO:0048471 CC	Perinuclear region of cytoplasm	16	6.3E-4	21	5.2E-12	
GO:0016301 MF	Kinase activity	19	6.7E-4	26	7.1E-15	
GO:0004674 MF	Protein serine/threonine kinase activity	32	1.3E-3	24	2.1E-18	
GO:0006629 BP	Lipid metabolic process	22	1.6E-3	42	7.6E-22	

Table I. Top 20 coincident enriched GO terms in T1- and T2DM.

GO, Gene ontology; BP, biological process; MF, molecular function; CC, cellular component; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.



Figure 2. Protein modules in the PPI networks for T1- and T2DM. The size of a protein was determined by the degree of its connection to other proteins, and the width of the edge connecting two proteins by the combined score of those proteins. (A) Protein module in the PPI network for T1DM; (B-H) the 7 modules in the PPI network for T2DM. PPI, protein-protein interaction; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

TDM GO:0005654 CC Nucleoplasm 92 5.38-7 GO:0007062 CC Extracellular exosome 116 1.38-6 GO:000506829 CC Cytosol 84 2.8E-6 GO:00050680 BP Negative regulation of prihelial cell proliferation 12 6.0B-6 GO:00050680 BP Negative regulation of canonical Wnt signaling pathway 13 3.7E-5 GO:00050750 CF Focal adhesion 27 4.7B-5 GO:0005156 BP Cell flate commitment 11 3.7E-5 GO:0005205 CF Focal adhesion 27 4.7B-5 GO:0005052 CC Focal adhesion 27 4.7B-5 GO:0005042 CC Nucleus 208 2.4E-4 GO:0005416 BP Cell differentiation 7 8.6B-4 GO:000542 CC Nucleus 208 2.4E-4 GO:000542 CC Nucleus 208 2.4E-4 GO:000542 CN Nucleus 208 2.4E-4 GO:000542 CC Nucleus 208 2.4E-4 GO:0003525 MP Negative regulation of	GO ID	GO term	Gene count	P-value
G0:0005654 CCNucleoplasm925.8E.7GO:0070062 CCExtracellular exosome1161.3E.6G0:0005829 CCCytosol842.8E.6G0:005893 BPPositive regulation of transcription, DNA-templated384.3E.6G0:0050560 BPNegative regulation of epithelial cell proliferation126.0E.6G0:0005050 PNRegative regulation of canonical Wnt signaling pathway133.5E.5G0:0005050 CFBasement membrane113.7E.5G0:0005050 CFFocal adhesion277.47E.5G0:0005052 CFFocal adhesion282.4E.4G0:0005052 CFFocal adhesion282.4E.4G0:0005032 CCNucleus2082.4E.4G0:0005032 CCNucleus2082.4E.4G0:0005032 MFmRNA 3'-UTR AU-rich region binding53.0E.4G0:0005032 MFNegative regulation of ell proliferation253.1E.4G0:0003025 MFNegative regulation of clarascription from RNA polymerase II promoter371.1E.3G0:000312 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3G0:000312 CCExtracellular matrix191.7E.3G0:0003512 CCCell junction468.7E-21G0:0003512 CCCell junction468.7E-21G0:0003512 CCCell junction468.7E-21G0:0003512 CCCell junction312.4E-44G0:0003512 CCCell junction468.7E-21G0:0000	T1DM			
GO:0070062 CC Extracellular exosome 116 1.3E-6 GO:0005829 CC Cytosol 84 2.8E-6 GO:0005893 BP Positive regulation of transcription, DNA-templated 38 4.3E-6 GO:0005808 DP Negative regulation of pithelial cell proliferation 12 6.0E-6 GO:0005109 C Balamer regulation of canonical Wnt signaling pathway 13 3.5E-5 GO:00051519 C GO:00051561 C Fate comminment 11 3.7E-5 GO:0005925 CC Focal adhesion 27 4.7E-5 GO:00050544 CP Positive regulation of transcription from RNA polymerase II promoter 50 60.6000553 GO:00050534 CC Nucleus 288 2.4E-4 GO:000587 BP Organ morphogenesis 12 2.9E-4 GO:000587 BP Organ morphogenesis 3.0E-4 GO:000887 BP 78.8E-4 GO:0003818 BP Negative regulation of transcription from RNA polymerase II promoter 7 7.8E-4 GO:0003818 BP Regulation of alternative mRNA splicing, via spliceosome 7 7.8E-4 GO:000312 CC Extracellular matr	GO:0005654 CC	Nucleoplasm	92	5.3E-7
G0:0005829 CC Cytasol 84 2.8E.6 G0:00050880 BP Positive regulation of transcription, DNA-templated 38 4.3E.6 G0:0005080 BP Negative regulation of epithelial cell proliferation 12 6.0E.6 G0:0005080 BP Cellular response to transforming growth factor β stimulus 11 8.3E.6 G0:0005090 BP Negative regulation of canonical Wrt signaling pathway 13 3.5E.5 G0:0005025 CF Focal adhesion 27 4.7E.5 G0:0005052 CF Focal adhesion 27 4.7E.5 G0:0005054 CN Nucleus 2.9E.4 60:0005634 CN Nucleus 2.9E.4 G0:0005054 CN Nucleus 2.8 2.4E.4 60:0005825 MF mRNA 3'-UTR AU-rich region binding 5 3.0E.4 G0:00005216 PN Organ morphogenesis 12 2.9E.4 60:0000827 MF 3.8 4.8E.4 G0:00005216 PN Negative regulation of call proliferation 7 3.8E.4 60:0000318 BP Regulation of alternative mRNA splicing, via spliceosome 7 7.8E.4 G0:0000316 CC Cell	GO:0070062 CC	Extracellular exosome	116	1.3E-6
G0:0043593 BPPositive regulation of transcription, DNA-templated384.3E-6G0:0050600 BPNegative regulation of epithelial cell proliferation126.0E-6G0:0071560 BPCellular response to transforming growth factor β stimulus118.3E-6G0:0005004 CCBasement membrane131.7E-5G0:000525 CFocal adhesion274.7E-5G0:000525 CCFocal adhesion274.7E-5G0:000525 CCFocal adhesion2082.4E-4G0:000534 CCNucleus2082.4E-4G0:000584 PDOrgan morphogenesis122.9E-4G0:000587 BPOrgan morphogenesis122.9E-4G0:000587 BPNegative regulation of transcription from RNA polymerase II promoter503.0E-4G0:000587 BPNegative regulation of cell proliferation73.6E-4G0:000587 BPNegative regulation of cell proliferation73.6E-4G0:000381 BPRegulation of allernative mRNA splicing, via spliceosome77.8E-4G0:000381 BPRegulation of transcription from RNA polymerase II promoter371.1E-3T2DMT125.1E-433.6E-4G0:0000521 CCCell junction468.7E-213G0:0000521 GMFIon channel activity481.8E-4G0:0005216 MFIon channel activity481.8E-4G0:0005216 MFIon channel activity405.0E-13G0:0005216 MFIon channel activity405.0E-13G0	GO:0005829 CC	Cytosol	84	2.8E-6
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	GO:0045893 BP	Positive regulation of transcription, DNA-templated	38	4.3E-6
GO:0071560 BP Cellular response to transforming growth factor β stimulus 11 8.3E.6 GO:0005064 CC Basement membrane 13 1.7E.5 GO:000525 CC Focal adhesion 27 4.7E.5 GO:000534 PC Cell differentiation 43 6.9E.5 GO:0005054 CC Nucleus 208 2.4E.4 GO:000584 PC Organ morphogenesis 12 2.9E.4 GO:000584 PC Nucleus 208 2.4E.4 GO:000587 PC Organ morphogenesis 12 2.9E.4 GO:000887 BP Organ morphogenesis 12 2.9E.4 GO:000887 BP Organ morphogenesis 12 2.9E.4 GO:0008870 BP Oligodendrocyte differentiation 7 3.6E.4 GO:0008870 BP Negative regulation of transcription from RNA polymerase II promoter 37 1.1E.3 GO:0003818 BP Regulation of transcription from RNA polymerase II promoter 37 1.1E.4 GO:0003054 CC Cell junction 147 1.4E.26 GO:0003054 CC Cell junction 147 1.4E.	GO:0050680 BP	Negative regulation of epithelial cell proliferation	12	6.0E-6
GO:0005604 CCBasement membrane131.7E.5GO:000906 BPNegative regulation of canonical Wnt signaling pathway133.5E.5GO:0005165 BPCell fate commitment113.7E.5GO:0005156 BPCell fate commitment274.7E.5GO:00050154 BPCell differentiation436.9E.5GO:00050154 BPOrgan morphogenesis2082.4E.4GO:0005634 CCNucleus2082.4E.4GO:0005875 BPOrgan morphogenesis122.9E.4GO:0005875 BPNegative regulation of cell proliferation253.1E.4GO:0008285 BPNegative regulation of cell proliferation73.6E.4GO:000381 BPRegulation of atternative mRNA splicing, via spliceosome77.8E.4GO:00031012 CCExtracellular matrix191.7E.3T2DM01.6E.33.1E.24GO:000324 CCCell junction468.7E.21GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFIon channel activity483.8E-14GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFIon channel activity481.8E-14GO:0003210 MFColamber593.5E-12GO:0003210 MFIon channel activity405.0E-13 <td>GO:0071560 BP</td> <td>Cellular response to transforming growth factor β stimulus</td> <td>11</td> <td>8.3E-6</td>	GO:0071560 BP	Cellular response to transforming growth factor β stimulus	11	8.3E-6
GO:00090090 BPNegative regulation of canonical Wnt signaling pathway133.5E.5GO:0045165 BPCell fate commitment113.7E.5GO:000525CCFocal adhesion274.7R.5GO:000525CCFocal adhesion436.9E.5GO:000548 BPCell differentiation436.9E.5GO:000547CCNucleus2082.4E.4GO:00053475 BPOrgan morphogenesis122.9E.4GO:00053525 BFmRNA 3'-UTR AU-rich region binding53.0E.4GO:0005825 BPNegative regulation of ellopoliferation253.1E.4GO:000381 BPRegulation of alternative mRNA splicing, via spliceosome77.8E.4GO:00031012 CCExtracellular matrix191.7E.3T2DMT1.4E.2660:0003142 CCAxon981.9E.26GO:0003195 CCCell junction1471.4E.261.4E.26GO:0003195 CCTerminal bouton468.7E.211.125.1E.14GO:0003195 CCTerminal bouton468.7E.211.125.1E.14GO:0003214 CCAxon981.9E.203.6E.132.4E.14GO:0003214 CCPresynaptic membrane312.4E.143.6E.13GO:0003214 CCPresynaptic membrane312.4E.143.6E.13GO:0005216 MFIon channel activity405.0E.133.6E.13GO:0005214 MFVoltage-gated ion channel activity405.0E.133.6E.13GO:0005509 MFCalcium ion binding111<	GO:0005604 CC	Basement membrane	13	1.7E-5
GO:0045165 BPCell fate commitment113.7E.5GO:005925 CCFocal adhesion274.7E.5GO:0005925 CCFocal adhesion274.7E.5GO:0005925 BPCell differentiation436.9E.5GO:0005634 CCNucleus2082.4E.4GO:0005634 CCNucleus2082.4E.4GO:0005825 BPOrgan morphogenesis122.9E.4GO:0008285 BPNegative regulation of cell proliferation253.0E.4GO:0008285 BPNegative regulation of cell proliferation73.6E.4GO:0008285 BPNegative regulation of atternative mRNA splicing, via spliceosome77.8E.4GO:0000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3GO:001012 CCExtracellular matrix191.7E.3T2DMCCCell junction1471.4E.26GO:0003054 CCCell junction1471.4E.26GO:0003054 CCCell junction1125.1E.18GO:00031012 CCFerminal bouton981.9E.26GO:0003744 CCAxon981.9E.26GO:0003745 BPRegulation of in transmembrane transport423.8E.14GO:0003765 BPRegulation of in transmembrane transport223.8E.44GO:0003765 BPRegulation of in transmembrane transport423.8E.14GO:0005216 MFIon channel activity405.0E.13GO:0005509 MFCalcium ion binding1112.9E.12G	GO:0090090 BP	Negative regulation of canonical Wnt signaling pathway	13	3.5E-5
GO:0005925 CCFocal adhesion274.7E-5GO:0005925 CCFocal adhesion436.9E-5GO:000544 BPPositive regulation of transcription from RNA polymerase II promoter501.6E-4GO:0005634 CCNucleus2082.4E-4GO:0005925 MFmRNA 3':UTR AU-rich region binding53.0E-4GO:000525 MFmRNA 3':UTR AU-rich region binding53.1E-4GO:0008285 BPNegative regulation of cell proliferation253.1E-4GO:000811 BPRegulation of alternative mRNA splicing, via spliceosome77.8E-4GO:00031012 CCExtracellular matrix191.7E-3T2DM1471.4E-26468.7E-21GO:000324 CCCell junction1471.4E-26GO:000315 CCTerminal bouton468.7E-21GO:000312 BPIon channel activity481.8E-14GO:0003216 MFIon channel activity481.8E-14GO:0003216 MFIon channel activity405.0E-13GO:0003510 CCSynaptic membrane312.4E-14GO:0005216 MFIon channel activity405.0E-13GO:0005216 MFVoltage-gated ion channel activity405.0E-13GO:0005216 MFVoltage-gated ion channel activity405.0E-13GO:0005216 MFVoltage-gated jon channel activity405.0E-13GO:0005216 MFVoltage-gated jon channel activity405.0E-13GO:0005216 MFColcinnion binding1112.9E-12<	GO:0045165 BP	Cell fate commitment	11	3.7E-5
GO:0030154 BPCell differentiation436.9E.5GO:0045944 BPPositive regulation of transcription from RNA polymerase II promoter501.6E.4GO:000563 CCNucleus2082.4E.4GO:000587 BPOrgan morphogenesis122.9E.4GO:0005875 MFmRNA 3'-UTR AU-rich region binding53.0E.4GO:0005825 BPNegative regulation of cell proliferation253.1E.4GO:0008285 BPNegative regulation of cell proliferation73.6E.4GO:0000121 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3GO:000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3GO:000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3GO:003012 CCExtracellular matrix191.7E.3T2DMTerminal bouton468.7E.21GO:0030424 CCAxon981.9E.26GO:0043155 CCTerminal bouton468.7E.21GO:005216 MFIon transport412.4E.14GO:0005216 CFPresynaptic membrane312.4E.14GO:0005214 CCSynaptic vesicle393.6E.13GO:0005242 MFVoltage-gated ion channel activity405.0E.13GO:0005244 MFVoltage-gated ion channel activity405.0E.13GO:0005265 PFRegulation of ion transmembrane5093.5E.12GO:0005265 MFClaicum ion binding1112.9E.12<	GO:0005925 CC	Focal adhesion	27	4.7E-5
GO:0045944 BPPositive regulation of transcription from RNA polymerase II promoter501.6E-4GO:0005634 CCNucleus2082.4E-4GO:0005878 POrgan morphogenesis122.9E-4GO:0008285 BPNegative regulation of cell proliferation253.1E-4GO:0008285 BPNegative regulation of cell proliferation73.6E-4GO:0008285 BPNegative regulation of alternative mRNA splicing, via spliceosome77.8E-4GO:000318 PRegulation of alternative mRNA splicing, via spliceosome71.1E-3GO:00031012 CCExtracellular matrix191.7E-3T2DMT468.7E-21GO:0003142 CCAxon981.9E-26GO:000424 CCAxon981.9E-26GO:0004215 CCTerminal bouton468.7E-21GO:0005216 MFIon channel activity481.8E-14GO:0003217 MCPresynaptic membrane312.4E-14GO:0003214 CCSynaptic vesicle393.6E-13GO:0003216 MFIon channel activity405.0E-13GO:0005214 MFVoltage-gated ion channel activity405.0E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005269 MFCalcium ion binding1112.9E-12GO:0005886 CCPlasma membrane593.5E-12GO:0005810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmision441.4E-11GO:0007756 C <t< td=""><td>GO:0030154 BP</td><td>Cell differentiation</td><td>43</td><td>6.9E-5</td></t<>	GO:0030154 BP	Cell differentiation	43	6.9E-5
GO:0005634 CCNucleus2082.4E-4GO:0009887 BPOrgan morphogenesis122.9F-4GO:0005925 MFmRNA 3'-UTR AU-rich region binding53.0E-4GO:0008285 BPNegative regulation of cell proliferation253.1E-4GO:0008285 BPOligodendrocyte differentiation73.6E-4GO:000318 DPRegulation of alternative mRNA splicing, via spliceosome77.8E-4GO:000122 DPNegative regulation of transcription from RNA polymerase II promoter371.1E-3GO:0030054 CCCell junction1471.4E-26GO:0030054 CCCell junction468.7E-21GO:0030054 CCCell junction468.7E-21GO:003195 CCTerminal bouton468.7E-21GO:0005216 MFIon transport1125.1E-18GO:0005216 MFIon transport123.8E-14GO:0005214 CCSynaptic membrane312.4E-14GO:0005214 MFVoltage-gated ion channel activity405.0E-13GO:0005214 MFVoltage-gated ion channel activity405.0E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005886 CCPlasma membrane593.5E-12GO:0007268 BPChemical synaptic transmission441.4E-14GO:0007726 BPChemical synaptic transmission441.4E-14GO:0007726 BPChemical synaptic transmission441.4E-14GO:0007726 BPChemical synaptic transmission441	GO:0045944 BP	Positive regulation of transcription from RNA polymerase II promoter	50	1.6E-4
GO:0009887 BPOrgan morphogenesis122.9E.4GO:0035925 MFmRNA 3'-UTR AU-rich region binding53.0E.4GO:0008285 BPNegative regulation of cell proliferation253.1E.4GO:0048709 BPOligodendrocyte differentiation73.6E.4GO:0000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E.3GO:001012 CCExtracellular matrix191.7E.3T2DMT74.6E.4GO:0003054 CCCell junction1471.4E.26GO:003054 CCCell junction1471.4E.26GO:0030424 CCAxon981.9E.26GO:0043195 CCTerminal bouton468.7E-21GO:0042734 CCPresynaptic membrane312.4E.14GO:0042734 CCPresynaptic membrane312.4E.14GO:00021CSynaptic vesicle393.6E.13GO:0002540 MFVoltage-gated ion channel activity4050.61.3GO:0002544 MFVoltage-gated ion channel activity4050.61.3GO:0005244 MFVoltage-gated ion channel activity4050.61.3GO:0005269 MFCalcium ion binding1112.9E.12GO:0007268 BPChemical synaptic transmission441.4E.11GO:000776 CCVoltage-gated potassium channel complex272.2E.10GO:000776 CCChromosome, centromeric region387.1E.11GO:000776 CCVoltage-gated potassium channel complex272.2E.10GO:0007	GO:0005634 CC	Nucleus	208	2.4E-4
GO:0035925 MFmRNA 3'-UTR AU-rich region binding53.0E-4GO:0008285 BPNegative regulation of cell proliferation253.1E-4GO:0008285 BPOligodendrocyte differentiation73.6E-4GO:0000122 BPRegulation of tanscription from RNA spliceosome77.8E-4GO:0003054 CCExtracellular matrix191.7E-3T2DM981.9E-26GO:0003054 CCCell junction468.7E-21GO:0003054 CCCell junction468.7E-21GO:0003054 CCCell junction468.7E-21GO:0003054 CCCell junction468.7E-21GO:0003195 CCTerminal bouton468.7E-21GO:0005216 MFIon transport1125.1E-18GO:0005216 MFIon channel activity481.8E-14GO:0005205 MFRegulation of ion transmembrane transport423.8E-14GO:0005205 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007269 BPNeurotransmitter secretion173.6E-13GO:0007269 BPNeurotransmitter secretion387.1E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.	GO:0009887 BP	Organ morphogenesis	12	2.9E-4
GO:0008285 BPNegative regulation of cell proliferation253.1E.4GO:0048709 BPOligodendrocyte differentiation73.6E.4GO:000381 BPRegulation of alternative mRNA splicing, via spliceosome77.8E.4GO:000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E-3GO:0030054 CCExtracellular matrix191.7E-3T2DM71.4E-26GO:0030054 CCCell junction1471.4E-26GO:0030054 CCCell junction468.7E-21GO:0030054 CCTerminal bouton468.7E-21GO:0005216 MFIon transport1125.1E-18GO:00021734 CCPresynaptic membrane312.4E-14GO:0005216 MFIon channel activity481.8E-14GO:0005216 MFIon channel activity405.0E-13GO:0005216 MFIon transport223.8E-14GO:0005216 MFIon transport405.0E-13GO:0005217 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005586 CCPlasma membrane503.5E-12GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007269 BPNeurotransmitter secretion173.6E-13GO:0007269 BPNeurotransmitter secretion387.1E-11GO:0007267 BPNeurotransmitter	GO:0035925 MF	mRNA 3'-UTR AU-rich region binding	5	3.0E-4
GO:0048709 BPOligodendrocyte differentiation73.6E-4GO:0000381 BPRegulation of alternative mRNA splicing, via spliceosome77.8E-4GO:000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E-3GO:0031012 CCExtracellular matrix191.7E-3TZDMT1.4E-2660:003054 CCCell junction1471.4E-26GO:0030054 CCCell junction468.7E-2160:0043195 CCTerminal bouton468.7E-21GO:0005216 MFIon channel activity1125.1E-1860:0042734 CCPresynaptic membrane312.4E-14GO:0005216 MFIon channel activity405.0E-133.6E-133.6E-133.6E-13GO:0005216 CCSynaptic vesicle393.6E-133.6E-133.6E-133.6E-13GO:0005214 MFVoltage-gated ion channel activity405.0E-133.6E-13GO:0005205 MFCalcium ion binding1112.9E-123.5E-12GO:0005209 MFCalcium ion binding1112.9E-123.5E-12GO:000756 BPChemical synaptic transmission441.4E-143.6E-13GO:0007269 BPChemical synaptic transmission272.2E-10GO:000756 CVoltage-gated potassium channel complex272.2E-10GO:000756 PNeurotransmitter secretion173.6E-13GO:000756 PBNeurotransmiter secretion173.6E-14GO:000756 CVoltage-gated potassium channel complex27 </td <td>GO:0008285 BP</td> <td>Negative regulation of cell proliferation</td> <td>25</td> <td>3.1E-4</td>	GO:0008285 BP	Negative regulation of cell proliferation	25	3.1E-4
GO:0000381 BPRegulation of alternative mRNA splicing, via spliceosome77.8E-4GO:000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E-3GO:0031012 CCExtracellular matrix191.7E-3T2DM </td <td>GO:0048709 BP</td> <td>Oligodendrocyte differentiation</td> <td>7</td> <td>3.6E-4</td>	GO:0048709 BP	Oligodendrocyte differentiation	7	3.6E-4
GO:0000122 BPNegative regulation of transcription from RNA polymerase II promoter371.1E-3GO:0031012 CCExtracellular matrix191.7E-3T2DMGO:0030054 CCCell junction1471.4E-26GO:0030054 CCAxon981.9E-26GO:0043195 CCTerminal bouton468.7E-21GO:0006811 BPIon transport1125.1E-18GO:0005216 MFIon channel activity481.8E-14GO:00042734 CCPresynaptic membrane312.4E-14GO:0005216 MFIon channel activity423.8E-14GO:0008021 CCSynaptic vesicle393.6E-13GO:0005204 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:000586 CCPlasma membrane5093.5E-12GO:000576 BPChemical synaptic transmission2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007765 CCChromosome, centromeric region387.1E-11GO:0007767 CCChromosome, centromeric region387.1E-11GO:0006813 BPPotassium ion transport341.1E-9GO:0005267 MFPotassium ion transport341.1E-9GO:0005767 MFPotassium ion transport341.1E-9GO:0005767 MFPotassium ion transport341.1E-9GO:0005267 MFPotassium ion transport341.1E-9 <t< td=""><td>GO:0000381 BP</td><td>Regulation of alternative mRNA splicing, via spliceosome</td><td>7</td><td>7.8E-4</td></t<>	GO:0000381 BP	Regulation of alternative mRNA splicing, via spliceosome	7	7.8E-4
GO:0031012 CC Extracellular matrix 19 1.7E-3 T2DM <	GO:0000122 BP	Negative regulation of transcription from RNA polymerase II promoter	37	1.1E-3
T2DM G0:0030054 CC Cell junction 147 1.4E-26 G0:0030424 CC Axon 98 1.9E-26 G0:0043195 CC Terminal bouton 46 8.7E-21 G0:0006811 BP Ion transport 112 5.1E-18 G0:0005216 MF Ion channel activity 48 1.8E-14 G0:0042734 CC Presynaptic membrane 31 2.4E-14 G0:0042734 CC Synaptic vesicle 39 3.6E-13 G0:0008021 CC Synaptic vesicle 39 3.6E-13 G0:0005244 MF Voltage-gated ion channel activity 40 5.0E-13 G0:0005509 MF Calcium ion binding 111 2.9E-12 G0:0005509 MF Calcium ion binding 111 2.9E-12 G0:0005866 CC Plasma membrane 509 3.5E-12 G0:0007268 BP Chemical synaptic transmission 44 1.4E-11 G0:0007269 BP Neurotransmitter secretion 17 3.6E-11 G0:000757 CC Chromosome, centromeric region 38 7.1E-11 G0:0006813 BP Potassium ion transport 26 1.4E-9	GO:0031012 CC	Extracellular matrix	19	1.7E-3
GO:0030054 CC Cell junction 147 1.4E-26 GO:0030424 CC Axon 98 1.9E-26 GO:0043195 CC Terminal bouton 46 8.7E-21 GO:0005216 MF Ion transport 112 5.1E-18 GO:0005216 MF Ion channel activity 48 1.8E-14 GO:00042734 CC Presynaptic membrane 31 2.4E-14 GO:000301 CC Synaptic vesicle 39 3.6E-13 GO:0005204 MF Voltage-gated ion channel activity 40 5.0E-13 GO:0005209 MF Calcium ion binding 111 2.9E-12 GO:0005509 MF Calcium ion binding 111 2.9E-12 GO:0005886 CC Plasma membrane 509 3.5E-12 GO:0007268 BP Chemical synaptic transmission 44 1.4E-11 GO:0007269 BP Neurotransmitt	T2DM			
GO:0030424 CC Axon 98 1.9E-26 GO:0043195 CC Terminal bouton 46 8.7E-21 GO:0006811 BP Ion transport 112 5.1E-18 GO:0005216 MF Ion channel activity 48 1.8E-14 GO:0042734 CC Presynaptic membrane 31 2.4E-14 GO:0034765 BP Regulation of ion transmembrane transport 42 3.8E-14 GO:0008021 CC Synaptic vesicle 39 3.6E-13 GO:0005244 MF Voltage-gated ion channel activity 40 5.0E-13 GO:00050509 MF Calcium ion binding 111 2.9E-12 GO:0005244 MF Voltage-gated ion channel activity 40 5.0E-13 GO:0005509 MF Calcium ion binding 111 2.9E-12 GO:0005886 CC Plasma membrane 54 3.6E-13 GO:0007268 BP Chemical synaptic transmission 44 1.4E-11 GO:00007269 BP Neurotransmitter secretion 17 3.6E-11 GO:0000775 CC Chromosome, centromeric region 38 7.1E-11	GO:0030054 CC	Cell junction	147	1.4E-26
GO:0043195 CC Terminal bouton 46 8.7E-21 GO:0006811 BP Ion transport 112 5.1E-18 GO:0005216 MF Ion channel activity 48 1.8E-14 GO:0042734 CC Presynaptic membrane 31 2.4E-14 GO:0034765 BP Regulation of ion transmembrane transport 42 3.8E-14 GO:0008021 CC Synaptic vesicle 39 3.6E-13 GO:0005244 MF Voltage-gated ion channel activity 40 5.0E-13 GO:0005509 MF Calcium ion binding 111 2.9E-12 GO:0005509 MF Calcium ion binding 111 2.9E-12 GO:0005886 CC Plasma membrane 509 3.5E-12 GO:00007268 BP Chemical synaptic transmission 44 1.4E-11 GO:0007269 BP Neurotransmitter secretion 17 3.6E-13 GO:00007269 BP Neurotransmitter secretion 38 7.1E-11 GO:0000757 CC Chromosome, centromeric region 38 7.1E-11 GO:00006813 BP Potassium ion transport 34 1.1E-9 </td <td>GO:0030424 CC</td> <td>Axon</td> <td>98</td> <td>1.9E-26</td>	GO:0030424 CC	Axon	98	1.9E-26
GO:0006811 BPIon transport1125.1E-18GO:0005216 MFIon channel activity481.8E-14GO:0005216 MFIon channel activity312.4E-14GO:0042734 CCPresynaptic membrane312.4E-14GO:0034765 BPRegulation of ion transmembrane transport423.8E-14GO:0008021 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding1112.9E-12GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007269 BPNeurotransmitter secretion173.6E-13GO:0007269 BPNeurotransmitter secretion387.1E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0006813 BPPotassium ion transport341.1E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0043195 CC	Terminal bouton	46	8.7E-21
GO:0005216 MFIon channel activity481.8E-14GO:0042734 CCPresynaptic membrane312.4E-14GO:0034765 BPRegulation of ion transmembrane transport423.8E-14GO:0008021 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding1112.9E-12GO:0005886 CCPlasma membrane5093.5E-12GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007268 BPChemical synaptic transmission173.6E-11GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005887 FCPotassium ion transport261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E 10	GO:0006811 BP	Ion transport	112	5.1E-18
GO:0042734 CCPresynaptic membrane312.4E-14GO:0034765 BPRegulation of ion transmembrane transport423.8E-14GO:0008021 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding1112.9E-12GO:0005509 MFCalcium ion binding543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007269 BPNeurotransmitter secretion173.6E-11GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005887 CCIntersport261.4E-9GO:0005887 CCIntersport of plasma membrane15057	GO:0005216 MF	Ion channel activity	48	1.8E-14
GO:0034765 BPRegulation of ion transmembrane transport423.8E-14GO:0008021 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0045211 CCPostsynaptic membrane543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0042734 CC	Presynaptic membrane	31	2.4E-14
GO:0008021 CCSynaptic vesicle393.6E-13GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0045211 CCPostsynaptic membrane543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:000587 CCIntegral component of plasma membrane1505.7E-10	GO:0034765 BP	Regulation of ion transmembrane transport	42	3.8E-14
GO:0005244 MFVoltage-gated ion channel activity405.0E-13GO:0005509 MFCalcium ion binding1112.9E-12GO:0045211 CCPostsynaptic membrane543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0008021 CC	Synaptic vesicle	39	3.6E-13
GO:0005509 MFCalcium ion binding1112.9E-12GO:0045211 CCPostsynaptic membrane543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0005244 MF	Voltage-gated ion channel activity	40	5.0E-13
GO:0045211 CCPostsynaptic membrane543.6E-13GO:0005886 CCPlasma membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0005509 MF	Calcium ion binding	111	2.9E-12
GO:0005886 CCPlasm membrane5093.5E-12GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:000587 CCIntegral component of plasma membrane1505.7E-10	GO:0045211 CC	Postsynaptic membrane	54	3.6E-13
GO:0006810 BPTransport2271.1E-11GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0005886 CC	Plasma membrane	509	3.5E-12
GO:0007268 BPChemical synaptic transmission441.4E-11GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0005267 MFPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0006810 BP	Transport	227	1.1E-11
GO:0008076 CCVoltage-gated potassium channel complex272.2E-10GO:0007269 BPNeurotransmitter secretion173.6E-11GO:000775 CCChromosome, centromeric region387.1E-11GO:0006813 BPPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0007268 BP	Chemical synaptic transmission	44	1.4E-11
GO:0007269 BPNeurotransmitter secretion173.6E-11GO:0000775 CCChromosome, centromeric region387.1E-11GO:0006813 BPPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0008076 CC	Voltage-gated potassium channel complex	27	2.2E-10
GO:0000775 CCChromosome, centromeric region387.1E-11GO:0006813 BPPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505.7E-10	GO:0007269 BP	Neurotransmitter secretion	17	3.6E-11
GO:0006813 BPPotassium ion transport341.1E-9GO:0005267 MFPotassium channel activity261.4E-9GO:0005887 CCIntegral component of plasma membrane1505 7E 10	GO:0000775 CC	Chromosome, centromeric region	38	7.1E-11
GO:0005267 MF Potassium channel activity 26 1.4E-9 GO:0005887 CC Integral component of plasma membrane 150 5.7E-10	GO:0006813 BP	Potassium ion transport	34	1.1E-9
GO-0005887 CC Integral component of plasma membrane 150 5 7E 10	GO:0005267 MF	Potassium channel activity	26	1.4E-9
1.10 5.7E-10	GO:0005887 CC	Integral component of plasma membrane	150	5.7E-10

Table II. Top 20 enriched GO terms for differentially	expressed genes in the sciatic nerve of T1- and T2DM mode	ls, respectively.
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GO, gene ontology; BP, biological process; MF, molecular function; CC, cellular component; T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Table III. Coincident enriched Kyoto Encyclopedia of Genes and Genomes pathways in T1- and T2DM.

	T1D	М	T2DM	
Term	Gene count	P-value	Gene count	P-value
mmu04010:MAPK signaling pathway	19	7.5E-4	29	3.2E-2
mmu04512:ECM-receptor interaction	10	1.4E-3	15	6.9E-3
mmu04390:Hippo signaling pathway	11	1.8E-2	24	1.2E-3
mmu04360:Axon guidance	9	4.5E-2	17	3.8E-2

T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

C

Table IV. Enriched Kyoto Encyclopedia of Genes and Genomes pathways in T1- and T2DM, respectively.

Table V. Top 10 proteins with relatively high connectivity degrees in the protein-protein interaction networks for T1- and T2DM.

Term	count	P-value
T1DM	10	
mmu04010:MAPK signaling pathway	19	7.5E-4
mmu04512:ECM-receptor interaction	10	1.4E-3
mmu05202:Transcriptionalmisregulation in cancer	14	1.7E-3
mmu05200:Pathways in cancer	24	2.5E-3
mmu05205:Proteoglycans in cancer	15	3.8E-3
mmu04310:Wnt signaling pathway	11	1.1E-2
mmu05214:Glioma	7	1.4E-2
mmu05222:Small cell lung cancer	8	1.4E-2
mmu04015:Rap1 signaling pathway	14	1.4E-2
mmu04330:Notch signaling pathway	6	1.7E-2
mmu04914:Progesterone-mediated	8	1.7E-2
oocyte maturation		
mmu04390:Hippo signaling pathway	11	1.8E-2
mmu00100:Steroid biosynthesis	4	1.9E-2
mmu05218:Melanoma	7	2.1E-2
mmu04614:Renin-angiotensin system	5	2.2E-2
mmu04520:Adherens junction	7	2.3E-2
mmu04510:Focal adhesion	13	2.5E-2
mmu04913:Ovarian steroidogenesis	6	3.0E-2
mmu04360: Axon guidance	9	4.5E-2
mmu05210:Colorectal cancer	6	4.6E-2
mmu05212:Pancreatic cancer	6	4 9E-2
	0	1.52.2
12DM	10	4 (5 40
mmu05033:Nicotine addiction	19	1.6E-10
mmu04080:Neuroactive ligand-receptor interaction	49	1.4E-7
mmu04723:Retrograde endocannabinoid signaling	23	8.5E-6
mmu04721:Synaptic vesicle cycle	17	1.2E-5
mmu05032:Morphine addiction	20	6.7E-5
mmu04514:Cell adhesion molecules (CAMs)	27	2.4E-4
mmu04724:Glutamatergic synapse	21	4.3E-4
mmu04024:cAMP signaling pathway	30	4.9E-4
mmu04923:Regulation of lipolysis in adipocytes	13	1.1E-3
mmu04390:Hippo signaling pathway	24	1.2E-3
mmu04020:Calcium signaling pathway	26	2.7E-3
mmu04972:Pancreatic secretion	17	3.8E-3
mmu00561:Glycerolipid metabolism	12	4.2E-3
mmu04911:Insulin secretion	15	5.6E-3
mmu04727:GABAergic synapse	15	6.2E-3
mmu04512:ECM-receptor interaction	15	6.9E-3
mmu04261:Adrenergic signaling in cardiomyocytes	20	2.1E-2
mmu04022:cGMP-PKG signaling pathway	22	2.2E-2
mmu04810:Regulation of actin cytoskeleton	26	2.3E-2
mmu04726:Serotonergic synapse	18	2.4E-2
mmu04725:Cholinergic synapse	16	2.6E-2
mmu04728:Dopaminergic synapse	18	2.8E-2
mmu04610:Complement and coagulation cascades	12	3.0E-2
mmu04010:MAPK signaling pathway	29	3.2E-2
mmu04970:Salivary secretion	12	3.3E-2
mmu04360:Axon guidance	17	3.8E-2
5		

T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

23

4.6E-2

mmu04062:Chemokine signaling pathway

T1DM			T2DM		
Rank	Protein	Degree	Rank	Protein	Degree
1	Lrrk1	76	1	Top2a	171
2	Trp53	65	2	Cdk1	144
3	Actb	60	3	Plk1	126
4	Actr1b	42	4	Aurkb	120
5	Fn1	40	5	Plk4	116
6	Yes1	38	6	Cenb1	115
7	Mapk12	38	7	Aurka	114
8	Prkacb	33	8	Mki67	113
9	Pten	32	9	Kif11	112
10	Ywhaz	28	10	Cdc20	111

T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Regarding DPN, it has been suggested that perturbation of insulin receptor signaling due to insulin resistance may cause neurons to become more vulnerable to metabolic insults and contribute to the pathogenesis of neuropathy (31). However, how gene expression is altered under the insulin-resistant state in nerve tissue remains unclear for T2DM. In the present study, a number of unique GO terms and signaling pathways were determined for the sciatic nerve profile of T2DM mice, which may result from an insulin-resistant state. For instance, a majority of the GO terms among the top 20 enriched terms unique to T2DM were related to the biological processes associated with cell junctions, ion activity and membrane activity. Previous studies indicated that the insulin-resistant state induced phosphorylation and downregulated of the expression of certain connexin (Cx) proteins including Cx43, which may constitute a potential mechanism underlying the pathogenesis of insulin resistance and its complications (32,33). Furthermore, it was demonstrated that Ca2+ overload in the mitochondria caused production of superoxide and functional impairment of multiple tissues, which may result in β -cell failure and insulin resistance in target tissues, further aggravating the complications of diabetes (34). Therefore, it may be speculated that the impairment of nerve tissues under the insulin-resistant state is attributable to these DEGs in the aforementioned biological processes in T2DM.

In addition, it was apparent that the efficacy of different treatment strategies for the two types of DM may be attributed to certain DEGs. For example, treatment of obesity appeared as critical as glycemic control for preventing the development of DPN in T2DM (5). In the present study, C-C chemokine receptor 7 (Ccr7), which was among the proteins with the highest degrees of connectivity in the PPI network for T2DM, could interact with the chemokine ligand 19 (Ccl19). The Ccl19-Ccr7 pathway may serve an important role in development of high-fat-induced obesity and subsequent insulin resistance (35). Therefore, it may be suggested that a potential treatment strategy is through Ccr7 targeting to alleviate insulin resistance and neuropathy in T2DM.

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J	υ	J

pathway

Table VI. Information on the modules of the protein-protein interaction networks in T1- and T2DM.					
Module	Node number	P-value	Protein with highest connectivity degree (degree)	GO term number	KEGG path number
T1DM					
1	15	9.4E-5	Srsf1 (22)	18	0
T2DM					
1	125	4.2E-14	Top2a (171)	151	8
2	28	2.9E-10	Frrs11 (28)	2	0
3	24	8.3E-8	Ccr7 (46)	79	5
4	41	1.0E-4	Mapk4 (72)	40	4
5	11	2.8E-4	Gabrg2 (25)	37	5
6	32	3.4E-4	Scn8a (52)	74	4
7	20	6.0E-4	Aldh1a2 (30)	34	9

T1DM, type 1 diabetes mellitus; T2DM, type 2 diabetes mellitus.

Limitation of the current study included the data being obtained from online databases, meaning the genetic backgrounds of the mice could vary substantially. Further genetic analyses are therefore warranted to identify genes and determine the molecular differences in neuropathy between the two types of DM based on different strains of mice.

In conclusion, the present study revealed the gene expression profiles and signaling pathways associated with the sciatic nerve in T1- and T2DM mouse models. The DEGs and signaling pathways may indicate unique biological processes and treatment strategies for the two types of DM. Further molecular biological experiments are required to validate the function of the DEGs and signaling pathways in DPN.

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Availability of data and materials

All data used and/or analyzed during this study are included in this published article.

Authors' contributions

XL designed the study and aided in drafting of the manuscript. YG and ZLQ performed the bioinformatics analysis and drafted the manuscript. DZL and GLS analyzed the data. ZQH and YCG interpreted the results and prepared the figures. XL and ZQH edited and revised manuscript. All authors approved the final version of the manuscript to be published.

Ethics approval and consent to participate

Not applicable.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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