A type 2 diabetes disease module with a high collective influence for Cdk2 and PTPLAD1 is localized in endosomes

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Abstract

Despite the identification of many susceptibility genes our knowledge of the underlying mechanisms responsible for complex disease remains limited. Here, we identified a type 2 diabetes disease module in endosomes, and validate it for functional relevance on selected nodes. Using hepatic Golgi/endosomes fractions, we established a proteome of insulin receptor-containing endosomes that allowed the study of physical protein interaction networks on a type 2 diabetes background. The resulting collated network is formed by 313 nodes and 1147 edges with a topology organized around a few major hubs with Cdk2 displaying the highest collective influence. Overall, 88% of the nodes are associated with the type 2 diabetes genetic risk, including 101 new candidates. The Type 2 diabetes module is enriched with cytoskeleton and luminal acidification–dependent processes that are shared with secretion-related mechanisms. We identified new signaling pathways driven by Cdk2 and PTPLAD1 whose expression affects the association of the insulin receptor with TUBA, TUBB, the actin component ACTB and the endosomal sorting markers Rab5c and Rab11a. Therefore, the interactome of internalized insulin receptors reveals the presence of a type 2 diabetes disease module enriched in new layers of feedback loops required for insulin signaling, clearance and islet biology.

Introduction

The insulin receptor (IR) belongs to the receptor tyrosine-kinase (RTK) family of cell-surface receptors [1, 2]. Early work on insulin and epidermal growth factor (EGF) revealed the presence of signaling molecules in hepatic endosomes fractions [3]. The concept of endosomal signaling is now well established [4], but the rules underlying IR trafficking and signaling compared with
those underlying the EGF receptor (EGFR) remain relatively unknown; this may be because proper insulin signaling and trafficking correlate with the maintenance of cell polarity [5].

Type 2 diabetes (T2D) is the result of a chronic energy surplus [6] coupled with a strong hereditary component. Estimates for the heritability of T2D range from 20 to 80% with a sibling relative risk of approximately 2, with obesity being an important driver in every population. The detailed genetic architecture of T2D was recently elucidated, and unlike type 1 diabetes (T1D) where the genetic risk is mostly concentrated in the HLA region, the genetic component explaining part of the heritability of T2D is primarily due to a combination of numerous common variants of small effect scattered across the genome [7–9]. T2D is characterized by both resistance to the action of insulin and defects in insulin secretion; the former has been an important motivating factor in the exploration of insulin signaling [1, 2]. Previous efforts to demonstrate that the genes mapping close to T2D risk loci are enriched for established insulin signaling pathways, however met with limited success; the most robust finding to date implicates seemingly unrelated cellular mechanisms, the majority of which affect insulin secretion and beta cell function [8–12].

An accumulation of proteins associated with T2D was previously observed in the interactome of the IRs endocytosed in an hepatic Golgi/endosomes fraction [13], suggesting the existence of a disease module at this intracellular locus that could help to further understand IR routing mechanisms, the primary mechanisms of the disease and drive the development of rational approaches for new therapies [14–16]. Here, starting from a proteome of IR-containing endosomes to narrow the space search, and the construction of a T2D-protomodule using validated genes, we reveal the presence of a T2D disease module with functional relevance both to insulin targets and insulin producing cells.

Materials and methods

Cell Fractions- Harlan Sprague-Dawley rats (female 120–140 g, b.w.) were purchased from Charles River Ltd. (St. Constant, Quebec, Canada) and were maintained under standard laboratory conditions with food and water available ad libitum, except that the food was removed 18 hours before the experiments. All animal procedures were approved by the Comité de Protection des animaux du Centre de Recherche du Centre Hospitalier de L’Université Laval (CPA-CRCHUQ, certificate 055–3). The G/E and the PM fractions were prepared and characterized in terms of enzyme markers, electron microscopy (EM) and ligand-mediated endocytosis, as originally described and used directly [3]. The G/E fraction was also characterized in terms of proteomic survey and construction of the protein interaction network (GEN) [13]. The compiled yield for the G/E fraction was 0.47 ± 0.04 mg protein/g liver weight (n = 57). A compiled yield of 2.4 ± 0.6 mg of protein/g of liver (n = 25) was obtained. IR-immuno-enriched endosomes were prepared as originally depicted [17] from the parent mixed hepatic Golgi/endosomal (G/E) fraction with only minor modifications [18]. Dynabeads (Dynal-A, Invitrogen, San Francisco, CA, USA) that were pretreated with 0.1% BSA and coated with the anti-IR β-subunit antibody (Sc-711, Santa Cruz Biotechnology, Santa Cruz, CA, USA), were incubated with freshly prepared G/E fractions (10 mg of protein) for 1 hour at 4˚C under gentle agitation. Beads were then rapidly rinsed before being subjected to EM, immunoblotting and mass spectrometry (MS) analysis. There was no major differences in the size and morphology of the vesicles immuno-isolated after 2 minutes or after 15 minutes of stimulation. They were relatively homogeneous with a diameter of 70–200 nm and some tubular elements. The IR was detected by Liquid chromatography-multiple reaction monitoring analysis (LC-MRM) using the peptide TIDSVTAQELR (P15127_IR, Q1 charge 660.3 (+2), Q3 charge 804.4 (2y7, +1) [13]. The amount of total protein bound to the beads was calculated by substracting the
nonbound from the starting material [17] [18]. A 16–23 range fold- purification over the parent fraction was measured.

Protein in-gel digestion- beads were washed 3 times with 50 mM ammonium bicarbonate buffer. They were suspended in 25 μl of 50 mM ammonium bicarbonate, following which trypsin (1 μg) was added. Proteolysis was done at 37˚C and stopped by acidification with 3% acetonitrile-1% TFA-0.5% acetic acid. Beads were removed by centrifugation, and peptides were purified from the supernatant by stage tip (C18) and vacuum dried before MS injection. Samples were solubilized into 10 μl of 0.1% formic acid and 5 μl was analyzed by mass spectrometry [19].

Mass spectrometry- peptide samples were separated by online reverse-phase (RP) nanoscale capillary liquid chromatography (nano LC) and analyzed by electrospray mass spectrometry (ES MS/MS). The experiments were performed with an Agilent 1200 nano pump connected to a 5600 mass spectrometer (AB Sciex, Framingham, MA, USA) equipped with a Nanoelectrospray ion source. Peptide separation occurred on a self-packed PicoFrit column (New Objective, Woburn, MA) packed with Jupiter (Phenomenex, Torrance, CA) 5 μl, 300A C18, 15 cm x 0.075 mm internal diameter. Peptides were eluted with a linear gradient from 2–30% solvent B (acetonitrile, 0.1% formic acid) in 30 minutes at 300 nl/min. Mass spectra were acquired using a data-dependent acquisition mode using Analyst software version 1.6. Each full scan mass spectrum (400 m/z to 1250 m/z) was followed by collision-induced dissociation of the twenty most intense ions. Dynamic exclusion was set for a period of 3 sec and a tolerance of 100 ppm. All MS/MS peak lists (MGF files) were generated using Protein Pilot (AB Sciex, Framingham, MA, USA, Version 4.5) with the paragon algorithm. MGF sample files were then analyzed using Mascot (Matrix Science, London, UK, version 2.4.0). MGF peak list files were created using Protein Pilot version 4.5 software (ABSciex) utilizing the Paragon and Progroup algorithms. (Shilov). MGF sample files were then analyzed using Mascot (Matrix Science, version 2.4.0) [20], and rodent databases (S1 Table). The number of newly identified proteins plateaued at approximately 10–20% of total for the second and third experiments, indicating that we were close to the completion point with this method [21] (S1A Fig).

Databases and network analyses- Conversion to human orthologs was performed using the InParanoid8 database (S2 Table). The PPIN was generated from a listing of protein-coding genes generated and named according to HUGO database nomenclature. Proteins found to be associated with IR in hepatic endosomes were included in the analysis: ATIC, PTPLAD1, SHP1, Cdk2, PLVAP1, CdkN1B and CCNE1. The interactions were curated using Y2H binary interactions of the CCSB human interactome, physical complexes and direct interactions from Intact, Database of Interacting Protein (DIP, UCLA), REACTOME, HITPREDICT and HINT databases, affinity complexes from BIOGRID and HPRD databases. Proteins having nonspecific interactions such as chaperones, ribosomal (RPL family) proteins, ubiquitylation and sumoylation processes (UBC, CUL), elongation factors were removed [22–24]. The Cytoscape platform (Version 3.2.0) was used for network visualization [25]. Self-loops and duplicated edges were removed prior the analyses. The cytoHubba algorithm was used to compute and rank nodes according to their centrality «Betweenness and Connectivity» scores in the network [22, 26, 27] (S3 Table). Cellular component grouping and functional analysis were performed after a gene ontology analysis with the Biological Networks Gene Ontology tool (BINGO version 2.44). Hypergeometric test was used for statistical analysis and the Benjamin & Hochberg False Discovery Rate correction was set as multiple testing correction when performing gene ontology analysis with BINGO. Kinase predictions were performed with GPS 3.0 [28], phosphosites [29] and NetworKIN [30] version 3.0 (KinomeXplorer) using the high-throughput workflow option. Data from GPS 3.0 were additionally filtered by a differential score (difference between Score and Cut of) higher or equal to 1.0. Networking associations
were considered if the Networkin score was observed to be higher than 2.0. Analyses were performed on November 10 2017. http://dx.doi.org/10.17504/protocols.io.

Candidate gene analysis and identification-

GO analysis-

We verified the probability of intra-cellular colocalization for candidates and seeds using the plugin BINGO adapted for the Cytoscape platform. We clustered the hybrid network (S2 Fig) based on enrichment in the same cellular compartment by GO. In IREP proteins coming from Golgi-endosomal fractions, 21 seeds were found to be enriched in the Golgi apparatus ($p < 5.6822 \times 10^{-14}$, after correction) and endosomes ($p < 1.1315 \times 10^{-16}$, after correction). Of the 126 IREP candidates identified by PPIN, 32 have at least three interactors among the 21 Golgi-endosomal seeds. The analysis was expanded to other compartments with 7 candidates interacting each with three seeds in the cytosol cluster ($p < 6.8728 \times 10^{-17}$, after correction), 10 candidates in the endoplasmic reticulum ($p < 1.6173 \times 10^{-12}$, after correction), 25 in the plasmamembrane ($p < 5.6570 \times 10^{-8}$, after correction), and 13 in the extracellular region ($p < 4.6024 \times 10^{-5}$, after correction). Taken together, 54 nonredundant IREP coding genes among the 126 identified by PPIN were found to be colocalized with validated seeds based on GO analysis (S3 Fig and S4 Table).

Fine-mapping approach-

We performed a linkage disequilibrium (LD) analysis and identified proximal SNPs correlated to diabetes GWAS signals ($p \leq 10^{-3}$) using replicated data as displayed in tables from the Wellcome Trust Case Control Consortium (WTCCC), GWAS Central portal, GWAS catalog or DIAGRAM GWAS-Metabochip or trans-ethnic data. This analysis provided a list of 130 IREP coding genes falling in genomic loci reliably associated with diabetes (S5 Table).

Genes expression analysis-

Most of the SNPs identified by GWAS are intergenic or fall in intronic regions of genes suggesting a regulatory role [7, 9]. Among the 130 candidates identified by fine-mapping, we verified which ones had SNPs experimentally shown to affect gene expression and to likely regulate some transcription factor binding as described in category-1 of high-confidence associations in the RegulomeDB database [31]. We identified 15 IREP coding genes fulfilling these criteria, consequently forming a first pool of IREP candidates based on gene expression regulation (S6 Table). A second pool was made-up of IREP genes showing or predicted to have similar patterns of expression with at least three of the 184 seeds by RNA-Seq analysis and simultaneously sharing regulatory binding motifs either for transcription factors or for miRNA. The candidates and seeds pairs were considered coexpressed if they were mutually ranked among the top 1% of coexpressed genes pairs by the Genefriends database [32]. The transcription factor targets (TFTs) or microRNA targets were analyzed using the top 10 grouping of the Gene Set Enrichment Analysis [33, 34] with ($p < 4.35 \times 10^{-16}$ after correction for TFTs and $p < 2.88 \times 10^{-6}$ for miRNA targets). In all, 296 IREP coding genes were found to share TFTs with at least three diabetes genes compared with 112 for miRNA targets and 109 for RNA-Seq. Only 80 genes from the RNA-Seq analysis were considered for the second pool of candidates because they simultaneously showed some shared binding targets with at least three DAGs for TFs (72 genes) and/or for miRNA (28 genes). Taken together, 94 non-redundant IREP coding genes from the first and second pools are considered candidates based on shared regulatory elements with validated DAGs (S6 Table).

IR endosomal autophosphorylation-

IR endosomal autophosphorylation was measured as previously reported [35] with minor modifications [13]. SiRNA in vivo: Rats were injected via the jugular vein with a scrambled or predesigned stabilized rat PTPLAD1 sequence (100 mg/100 g bw; IVORY in vivo siRNA GGGCAGUCUAAUUCGGUG UGCU, D-00203-0200-V; purified/desalted by RP/IEX-HPLC; Riboxx Life Sciences, Germany; Liver In vivo transfection reagent 5061, Altogen Biosystems, Las Vegas, CA) 48 and 24 hours before isolating the G/E fraction. The PTPLAD1 mRNA expression level was measured against GAPDH in liver sections using quantitative polymerase chain reaction (qPCR) and was decreased by 52 +/- 6.2%, n = 3.
Cell culture and analysis- HEK293 cells were maintained in DMEM high-glucose medium with 10% foetal bovine serum. PTPLAD1 siRNA knockdown was performed as previously described [13] using the predesigned human sequence as follows: GACCCAGAGGCAGGUAAACAUUACA NM_016395_STEALTH_367. Cells were transfected using Lipofectamine 2000™ (Life Technologies) for 48 hours and subjected to the described experiments. For overexpression experiments PTPLAD1 WT and Cdk2 WT were cloned into the pcDNA3 expression vector. Transfection was performed with Lipofectamine 2000™ and plasmid DNA (300 ng/ml). Cells were preincubated at 37°C without serum for 5 hours before insulin (35 nM) stimulation for the indicated times. Immunoprecipitation (IP) were done under solubilization conditions that preserve the integrity of insulin-dependent complexes (Empigen BB 0.3%, 2 hours, 4°C) [18].

Reagents and antibodies- Porcine insulin (I5523) was obtained from Sigma-Aldrich (St. Louis, MO, USA). The following antibodies were used: anti-phosphotyrosine (PY20, Sigma-Aldrich, St. Louis, MO, USA). The IR β-subunit (Sc-711), Rab5c (sc-365667) and Cdk2 (sc-163, sc-163AC) antibodies were obtained from Santa Cruz Biotechnology (Santa Cruz, CA, USA). The anti-PTPLAD1 was from Abcam (ab57143, Cambridge MA, USA). The anti-tubulin antibodies were obtained from Sigma-Aldrich (T5168, TUB 2.1, St. Louis, MO, USA). The anti-MAD2 was from Bethyl Laboratories (Montgomery, TX, USA). The RILP antibody was from Invitrogen (PA5-34357, Waltham, MA, USA). The generic anti-phosphothreonine was from Zymed (San Francisco, CA, USA). The anti-tubulin antibodies were obtained from Sigma-Aldrich (T5168, TUB 2.1, St. Louis, MO, USA). The antibody against Rab11a was from ThermoFisher Scientific (Rockford, IL, USA). Peroxidase-conjugated secondary antibodies were used (1:10,000, Jackson Immuno Research Laboratories, West Grove, PA, USA). Membranes (PVDF) were analyzed using a chemiluminescence kit (ECL, Perkin Elmer Life science, Boston, MA) or using an ImageQuant LAS 40 000 imager (GE Healthcare Biosciences, Baie d’Urfé, QC, CA). [γ-32P]-ATP (1000–3000 Ci/mmol) was from New England Nuclear Radiochemicals (Lachine, Québec). Other chemicals and reagents were of analytical grade and were purchased from Fisher Scientific (Sainte-Foy, Québec, CAN) or from Roche Laboratories (Laval, Québec, CAN).

Results and discussion
Rabs, V-ATPase subunits, tyrosine phosphatases, and cell cycle proteins shape IR-containing endosomes
To determine the proteomic environment of the internalized IRs, we performed a survey of IR-containing endosomes fractions. We started with a mixed Golgi/endosomes fraction (G/E) using a single dose of insulin (1.5 μg/100 g body weight [b.w.]) that resulted in 50% saturation of rat liver receptors. Fractions were prepared at the 2-minute time peak of IR accumulation and the 15-minute 50% decline time [13, 36] to collect a larger proteome. Freshly prepared fractions were then incubated with anti-IR (β-subunit)-coated magnetic beads, and endosomes were collected with a magnet [17, 18]. We identified a total of 620 proteins with high confidence (named IREP: IR Endosome Proteome, Fig 1A and S1 Table).

Gene ontology (GO) analysis revealed enrichment of proteins involved in trafficking and signaling (MGI database; Biological Network Gene Ontology (BINGO) tool). These were primarily represented by coat-forming elements, small GTPases, components of the actin cytoskeleton, microtubules and motor proteins of the microtubule cytoskeleton and regulators of the cell cycle (Fig 1A and 1B left panel). Immunoblotting analysis confirmed the peak of IR accumulation occurring at 2 minutes post-insulin injection (Fig 1B right panel). The protein PTPLAD1 (HACD3), previously observed to be associated with the IR in G/E fractions after insulin stimulation [13], was also detected here at 15 minutes post-insulin injection (Fig 1B,
right panel). Consistent with the presence of sets of Rabs [17, 37], thirteen Rabs were identified. They were shown to be involved in transport from early to recycling endosomes (Rab22a, 2 minutes post-insulin injection) or late recycling endosomes (Rab11a, Rab17) [38]. Rab8a, reported to act exclusively in the trans-Golgi network to plasma membrane transport, was identified at 15 minutes post-insulin injection (Fig 1A). Other Rabs identified at both times (Rab6a, Rab5c, Rab1a, Rab2b, Rab11b, Rab14, Rab1b, Rab7a and Rap1b) are all implicated in recycling, transcytotic or Golgi transport events [17, 38]. Among signaling proteins, the transmembrane protein tyrosine phosphatase (PTP) of the R subfamily [39], PTPRF (also named leukocytes antigen-related, LAR) was identified (Fig 1A). PTPRs are generally associated with IR tyrosine dephosphorylation [40–43], acting preferentially on the juxtamembrane sites Y960 and Y1146 located in the IR activation loop [41, 43]. The putative PTP Dnajc6 (also called auxilin) is a chaperone involved in clathrin-mediated endocytosis of EGFR [44, 45]. PTPN6 (SHP-1) is a known IR regulator in the liver [46]. The large representation of regulators of the cell cycle was less expected but is consistent with the attenuation of endocytosis during cell division [47]. The proton translocation machinery necessary to achieve optimal lumenal acidic pH is also particularly well represented (ATPv1a, ATPv1b2, ATPv1f, Atpv1e1, ATPv0a1; 2 minutes post-insulin injection) (Fig 1A and 1B left panel, S1 Table). Efficient acidification by V-ATPase is particularly important for the ligand dissociation-degradation sequence according to the law of mass action and is specific to insulin in contrast with EGF or prolactin complexes. This sequence is followed by a rapid recycling of the freed IR under the concerted action of endosomal protein tyrosine phosphatases (PTPs), thus supporting efficient circulating clearance [3, 48].

**Genes at risk for type-2 diabetes form a proto-module enriched for transport and oxygen species regulation**

Most of the established T2D genes are supported by low and high probability GWAS signals of their identified variants [8, 9]. To verify if the IREP is associated with T2D, we used complementary data sources (DIAGRAM consortium, SNPs provided in replicated GWAS from the NHGRI-EBI GWAS catalog and GWAS Central portal, source S7 Table) to compile a list of 452 T2D and associated trait genes on the basis of single-nucleotide polymorphisms (SNPs) identified in their genomic loci (diabetes-associated gene: DAG; p-value < 5 x 10^{-8}; S8 Table). This list also contains relevant genes associated with T2D Mendelian traits described in the OMIM database and tagged with the symbol (3) indicative of known molecular associations (S8 Table-sheet OMIM). To reduce false-positive associations, the 452 DAG products were validated in a physical protein interaction network (PPIN) [14, 24]. We gathered physical protein-protein interaction data from the Biological General Repository Interaction Datasets (BIOGRID), the human interactomes I and II generated with Y2H systems from the Center for System Biology (CCSB) interactome, Intact, Reactome, Database of Interacting Proteins (DIP, UCLA), HitPredict databases or from the Human Proteins Repository Database (HPRD). The network was visualized with Cytoscape [25]. The 452 DAG products formed a
PPIN of 184 proteins and 309 interactions we called the proto-T2D module (Fig 2A and S3 Table-sheet T2DN-protomodule).

The proto-T2D module is made up essentially of protein coding-genes from OMIM (26%), GWAS variants with a p-value < 1x10^-8; 69%, and GWAS variants with 5 x 10^-8 < p-value < 1 x 10^-8; 5%, (Fig 2B). It displays a scale-free topology relying on a few hubs of large size such as HNF4A surrounded by a majority of the peripheral nodes (more than 38% of the nodes have only one interactor) [14, 15] (Fig 2A and 2C and S3 Table-sheet ProtoT2Dmodule). Protein transport (p < 1.82 x 10^-34) and response to oxygen-containing compounds (p < 1.32 x 10^-32) are the most enriched cellular processes identified by a gene ontology (GO) analysis with the presence of trafficking proteins (Rab5b, RABEPP1, RABEPP2) and transcription factors from the HNF (HNF4A, HNF1A or HNF1B), FOX (FOXO3, FOXA2) and TCF families (TCF7L2, TCF4, and TCF19). Signaling modules associated with insulin sensitivity are also present (INSR, IRSs, GRB14, PTPN1) (Fig 2A and S3 Table-sheet GO analysis-T2D-protomodule).

The affinity for these biological processes is supported by the enriched subcellular component analysis which revealed an accumulation of the coding genes associated with risk for T2D in endosomes and endoplasmic reticulum among the major genes. Proteins from the histocompatibility complex are also among the most significant clusters in the T2D-protomodule (Fig 2A and S3 Table sheet GO analysis-T2D-protomodule).

Overall, 62% of the 452 selected DAGs are disconnected (267/452). Five factors likely contribute to this fragmentation as follows:

i) True lack of binary or indirect physical interaction.

ii) Interactome incompleteness [24].

iii) False positives (not all genes have a known mechanistic association with the disease), and genes associated with late complications of the disease.

iv) The T1D-T2D paradox and disease classification [7].

v) Missing heritability [8, 9].

A total of 101 high confidence candidate genes for type 2 diabetes risk are identified in IREP

Genes that fall within one of the known disease loci and whose protein products interact with a known risk factor are predicted to be 10-fold enriched in a true disease gene. By considering the cellular localization as well, the network information leads to a 1000-fold enrichment over random genes [14, 49]. We used a combination of approaches to identify candidate genes confidently associated with diabetes traits. The candidates were grouped and ranked according to i) their topological proximity with the 184 previously validated DAG “seeds” in the PPIN approach, ii) the probability of co-localization in the same subcellular locus (S4 Table), iii) the identification of proximal variants correlating with the diabetes GWAS signal by fine-mapping analysis (S5 Table) and iv) the similarity of gene expression regulation with the 184 seeds of the proto-T2D-module (S6 Table) (see Methods).

A total of 246 nonredundant IREP coding genes are associated with diabetes traits when considering each of the approaches individually. Of these, 38 were validated by at least three of the approaches mentioned previously. This list includes the Cdk2 gene which is located in the risk area composed of 4 blocks in strong LD around the T2D SNP rs2069408 (S4A Fig and S5 Table). ATIC, which was previously observed to be associated with the IR in endosomes
Fig 2. Diabetes-associated genes form a protomodule. (A) Overall, 452 diabetes-associated gene (DAGs; GWAS p value < 5 \times 10^{-8} and OMIM) products form a PPIN of 184 proteins and 309 interactions termed T2D-protomodule. (B) In total, 10% (11/102) of the high-confidence DAGs with a probability less than 5 \times 10^{-8}, 53% of the DAGs with a probability less than 1 \times 10^{-8} (141/266) and 49% of the OMIM genes (49/84) are recovered in the proto-T2D module, showing a tendency to select the highest level of reliability. (C) Nodes-degree distribution: More than 38% of nodes (70 nodes) in the proto-T2D module are peripheral with a minority of hubs from transcription factor families. The general topology of the protomodule is characteristic of a disease network with the presence of few central hubs of large size, surrounded by numerous peripheral hubs of smaller size (S3 Table).

https://doi.org/10.1371/journal.pone.0205180.g002
together with PTPLAD1 and AMPK [13]; PTPN6 (SHP-1); the small GTPases of Rab the family (Rab14, and Rab5c); and a series of coat components (i.e., AP complexes, CAV1, COPA, SEC23A and SEC24C) are also present (Table 1).

Sixty-three other IREP coding genes are shown to have reliable association with diabetes after validation with any two approaches. This list includes the HACD3 (PTPLAD1) gene, located in a risk area (S4B Fig and S5 Table), that was also previously associated with T2D in human islets [50]. PRKAA1 (AMPK); the Cdk2 regulators (CDKN1B, CCNE1); a V-ATPase subunit (ATP6VA1); the small GTPase Rab1a, Rab1b and Rab8a; several coat components; the putative tyrosine phosphatase DNAJC6; ACTB and TUBA also fall into this category (Table A in S4 Table). IREP is also enriched in genes associated with the T2D risk with 15 of the 184 validated DAGs from the human genome being identified indicating a nonrandom concentration of diabetes genes variants during IR endocytosis (p-value of 3.44 x 10^{-4}; hypergeometric test) (S2 Table- sheet IREP-HUGO).

Collectively, IREP consists of more than 20% (15 validated DAGs from the T2D-protomodule and 101 candidates) of gene products confidently associated with the T2D risk.

**The insulin receptor-containing endosome network (IREN) displays a type-2 diabetes disease module architecture**

A disease module can be defined as a connected subnetwork showing mechanistic evidence for a phenotype [14, 15]. To identify the molecular mechanisms associated with IREP, we constructed a PPIN of IR-containing endosomes. The cytoHubba algorithm was used to compute and to rank nodes in the network [26]. The resulting collated PPIN is formed by 313 nodes and 1147 edges (55% of IREP proteins; named IREN, Insulin Receptor Endosome Network). The general topology of IREN is based on few major hubs, with the kinase Cdk2 displaying the highest centrality. Large nodes represented by the IR itself, proteins of the actin cytoskeleton (ACTB), and those involved in vesicular trafficking (CAV1) were observed. More peripheral nodes were also present as follows: coats (GOLGA2, CLTC), V-ATPase subunits (ATP6V1A), and cargos (APOA1) (Fig 3 and S3 Table-sheet IREN).

From the 101 high-confidence candidates (Table 1 and Table A in S4 Table) and 15 of the 184 validated DAGs identified in IREP, 94 of the candidates and 10 of the DAGs are present in IREN. They form a single-connected subnetwork of 94 nodes with 330 interactions (Fig 3 and S3 Table). To test whether this module could arise by chance in the context of IR endocytosis, we made random reiterations of any 94 nodes of IREN. The results showed that the subnetwork is robust (p-value < 0.0049, S5 Fig; http://dx.doi.org/10.17504/protocols.io.[PROTOCOL DOI]). Its collective influence was analyzed by expanding it to the first adjacent nodes. This resulted in a connected network of 271 nodes (88% of nodes) and 1070 out of 1147 IREN interactions (Figs 3 and S5), coverage that is largely more than expected by chance (S5 Fig). GO analysis also revealed an enrichment for vesicle transport (p< 1.85 x 10^{-51}) and response to oxygen species (p < 8.52 x 10^{-20}), with the most enriched cell components being endosomes (p < 7.00 x 10^{-25}), Golgi apparatus (p < 1.77 x 10^{-27}) and endoplasmic reticulum (p < 2.13 x 10^{-23}), showing that the T2D-protomodule expansion coincides with a functional expansion (Fig 3 and S3 Table-sheet IREN, GO analysis). Of the 94 of the 101 high-confidence candidates in IREP, 94 have credible tyrosine phosphorylation motifs with the IREN kinases (Table 1 and S9 Table). Of the 87 present in IREN, 71 have at least one of their kinase-substrate interactions confirmed in IREN (Fig 3 and S9 Table), further emphasizing the mechanistic association. Taken together, these results indicate that IREN is a T2D-disease module.
**Table 1. List of candidates.** Thirty-eight IREP coding genes are validated for association with diabetes traits by at least three out of four distinct approaches. (PPIN) protein-protein interactions network, (GO) Gene Ontology, Subcellular co-localization, (GWAS) fine-mapping, (COEXPRESS) same expression pattern.

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https://doi.org/10.1371/journal.pone.0205180.t001
Cdk2 affects the association of IR with microtubules

A prerequisite in the description of interaction maps is the validation of hubs in terms of perturbation responses. We then tested examples here of hub complexes in terms of insulin response. We verified first whether Cdk2, which displays the highest centrality and is a high-confidence candidate (Table 1 and S4A Fig), is indeed associated with key elements. Microtubules, for instance, rely on dimerization of tubulin subunits alpha and beta for their assembly and Rab5-containing endosomes have a capacity to move along microtubules [51]. We noticed that the tubulin alpha subunit (TUBA), a reported substrate for the IR in vitro [52, 53], is preponderant within the IREN (Fig 3). We show that TUBA indeed readily associates with the IR after insulin stimulation in HEK293 cells, while TUBB has a different profile (Fig 4A). In addition, the association was nearly abolished upon Cdk2 overexpression, confirming the presence of complexes and indicating that Cdk2 levels can effect complexes organization within IREN (Fig 4A).

PTPLAD1 expression affects the IR autophosphorylation activity and association with Cdk2, Rab5c, Rab11a and actin

Compared with Cdk2, PTPLAD1 is an example of good centrality, but it is poorly studied and identified as a moderate candidate (Table A in S4 Table and Fig 3). Because the fatty acid elongation enzymatic activity was not confirmed, it was recently hypothesized that PTPLAD1 (HACD3) is involved in the elongation of specialized forms of 3-OH acyl-CoAs, such as those containing a short or branched alkylic chain [54]. An interaction with Rac1 was also reported [55]. PTPLAD1 has a well-positioned conserved cysteine C(X)5K motif in the soluble cytosolic loop, residues 257–279, and its partial deletion in cultured HEK293 cells coincided with IR tyrosine hyper-phosphorylation [13]. We verified whether PTPLAD1 acts on IR tyrosine phosphorylation outside a whole-cell context. We used an in vitro assay, whereby IR-loaded endosomes were incubated in the presence of ATP. We observed that a prior siRNA-mediated depletion in rat PTPLAD1 nearly abolished the PTPLAD1 presence from isolated endosomes, which coincided with a marked increase in IR autophosphorylation, demonstrating that the IR tyrosine-phosphorylated state is modified by PTPLAD1 (Fig 4B right panels). Low, but consistent, enzymatic activity towards the artificial substrate pNPP was measured (Fig 4B left panel) resembling the loss of PTP activity towards the IR observed previously observed after membrane solubilization [35]. In accordance with these results, and with prior PTPLAD1 depletion experiments [13], we observed that overexpression of PTPLAD1 in HEK293 cells markedly decreases the IR tyrosine-phosphorylated state (Fig 4C). Of further interest, the candidate Rab5c (Table 1) also associates with the IR and this association increases in an insulin-regulated (Fig 4C). Rab5a and b are well documented as playing a role in the early events of EGFR endocytosis but the role of Rab5c remains unclear [56]. Rab5c may therefore be particularly important for IR action as we noted the presence of an IR phosphorylation motifs located in the GTP binding site (Y83) (S9 Table) that resembles an inhibitory feedback loop described previously for Rab24 [57]. In support of this finding, we detected Rab5c and Cdk2, but not Rab11a, in anti-phosphotyrosine affinity complexes, and this association was markedly decreased after...
PTPLAD1 overexpression (Fig 4C). In addition, both IR and Rab5c were present in Cdk2 affinity complexes, and this association decreased following PTPLAD1 overexpression (Fig 4C). To further test the importance of PTPLAD1 on IR complexes, we verified and noticed an insulin-dependent association of IR with Rab11a, a known marker at the intersection between the endocytic and exocytic pathways [38, 58]. This supports a role for PTPLAD1 in cycling from endosomes to the plasmamembrane (PM). On another hand, we confirmed that under the same circumstances PTPLAD1 deletion, using siRNA, increases IR tyrosine phosphorylation and the presence of actin in IR immunoprecipitates (Fig 4D) [13]. We also observed an insulin-dependent recruitment of the Rab-interacting lysosomal protein (RILP) in IR immunoprecipitates that was nearly abolished by PTPLAD1 deletion (Fig 4D). RILP was demonstrated to be required for EGFR confinement and degradation in late endosome compartments [59] and is an inhibitor of V-ATPase activity [60]. This supports the importance of the PTPLAD1 node within IREN. Collectively, the data support the presence of dynamic insulin-dependent interactions for Cdk2 between the IR, PTPLAD1, Rab5c, Rab11a, tubulin and actin cytoskeletons whereby PTPLAD1 controls IR tyrosine phosphorylation and key interactions.

To verify the idea that cell cycle components have expanded their action on endocytic traffic, we verified whether the protein MAD2, which binds to the MAD2-interacting motif (MIM) located in the carboxyterminal domain of the IR β-subunit during clathrin-mediated endocytosis [61], is responsive to insulin at the cell surface. The results demonstrate that MAD2 readily disappears from the PM fractions following IR tyrosine kinase activation (Fig 4E), thus supporting the idea that cells use cell cycle regulators for both early [61] and later events of IR endocytosis. No MAD2 signals were detected in the G/E fraction, suggesting a rapid dissociation of the complexes together with the clathrin coat.

https://doi.org/10.1371/journal.pone.0205180.g004
Inhibition of V-ATPase shifts the IR accumulation rate in endosomes in vivo

V-ATPase subunits are well represented in IREP (Fig 1 and S1 Table), forming large, more peripheral nodes in IREN (Fig 3) with ATP6V1A being identified here as moderate candidate (Table A in S4 Table) for type 2 diabetes risk. We thus verified whether the kinetics of IR endocytosis are affected in vivo after treatments with two different potent V-ATPases inhibitors. We observed that the peak of IR accumulation in endosomes is markedly shifted towards later times of endocytosis following either concanamycin A or bafilomycin A1 pretreatments as demonstrated by immunoblotting and hexokinase activity measurements, indicating that V-ATPase reduces IR degradation and transport or decreases IR recycling to the plasma membrane [62] or both (Fig 5A). Concanamycin A does not affect IR on going IR autophosphorylation in vitro, showing that V-ATPase inhibitors do not inadvertently function through PTPs inhibition (Fig 5B left panel). We noted however a strong and consistent reconstituted threonine phosphorylation signal that was readily abolished by concanamycin A (Fig 5B right panel), suggesting the presence of additional feedback loop layers, which have yet to be characterized, informing the cell that the lumenal acidification process is optimized. We verified whether V-ATPases elements contains IR phosphorylation motifs. The kinase network analysis indicated that ATP6V1A (Table A in S4 Table) and ATP6V1E1 are indeed strong candidate substrates for the IR as well as for Cdk2 and AMPK (PRKAA1) (Fig 5C and S9 Table).

Conclusion

Using a combination of cell fractionation and computational approaches, we found a T2D disease module in IR-containing endosomes. The starting point of our analysis was a list of seed genes with established genetic T2D association and high GWAS p-values (1 x 10^{-8}) against the background of random variation. They carry enough information to build a robust T2D-protomodule (Fig 2). The functional specialization of the T2D-protomodule also found in IREN (Fig 3) is in accord with the connection of these processes (protein transport, transcriptional factors and response to oxygen) in insulin action [1, 2]. The topological features of a scale-free network, with the view that hubs with the highest influence represent important points in biological networks [14, 26], coupled with the large enrichment in T2D genetic risk is particularly well represented by Cdk2 (Fig 3 and Table 1). Cdk2 regulators were independently and repeatedly reported by GWAS and their role, with many other common variants, was interpreted more in terms of insulin production and secretion indicating that the beta-cell is a more appropriate place to find a T2D-disease module [9, 11, 63, 64]. Indeed, mice lacking Cdk2 are viable [65, 66] and targeted Cdk2 deletion in the pancreas induces glucose intolerance primarily by affecting glucose-stimulated insulin secretion [67]. Similar to endosomes, the secretory pathway consists of multiple dynamic compartments linked via anterograde and retrograde transport [68, 69]. The T2D-disease module thus can be co-functional in endosomes and insulin-secreting cells. In this regard, in the liver the presence of insulin-regulated Cdk2/cyclinE/p27kip1 complexes having a capacity to inhibit hybrid endosome formation in vitro has been previously reported [70].

In contrast with Cdk2, PTPLAD1 has less topological influence in IREN and is a less-studied protein. PTPLAD1 is, however, functionally well connected as the control of IR activity may be achieved at several endosomal targets by PTPLAD1 that, together with Cdk2, seems to have a considerable local influence on actin and microtubule networks, Rabs and V-ATPase. The finding that IR complexes are under the control of PTPLAD1 would also be particularly important because PTPLAD1 mobilization in response to insulin inputs has also interesting consequences by favoring tyrosine phosphorylated-IR quanta formation, which is considered as an emergent property of endosomes as signaling devices [4]. This PTP activity is yet to be
Fig 5. The pharmacological inhibition of V-ATPase affects the time peak of IR accumulation in endosomes. Rats that were treated with concanamycin A (Cona A, 4.0 μg/100 g, b.w.) or were left untreated, were then stimulated with insulin (1.5 μg/100 g, b.w.) for the indicated time and the G/E fractions were isolated. (A) Left panel, immunoblot of IR using the anti-IRβ subunit or αPY20 (95 kDa PY) antibodies (50 μg of protein). Right panel, rats were left untreated or treated with bafilomycin A1 (Baf A1, 0.5 μg/100 g, b.w.). IRs from G/E fractions prepared at the noted time following insulin administration (1.5 μg/100 g, b.w.) were partially purified by WGA-sepharose affinity chromatography and subjected to exogenous kinase assay. 

32P incorporation into poly Glu-Tyr (4:1) is expressed as pmol/μg protein. Values shown are means ± s.d. (P < 0.0001, 2 minutes and 15 minutes, n = 3). (B) G/E liver fractions were prepared at their IR concentration time peak (2 minutes after insulin injection; 1.5 μg/100 g, b.w.) and immediately suspended in the cell-free system for 0 and 2 minutes at 37 °C and in the presence of ATP and the absence or presence of fresh cytosol (diluted 1/10) and Cona A. After stopping the reaction (0 and 2 minutes), the fractions were immunoblotted (input 50 μg of protein; 12% resolving gels) with the anti-phosphotyrosine (anti-p-Tyr, left panel) or anti-phosphothreonine (anti-pThr, right panel) antibodies. (C) Subnetwork extracted from IREN (Fig 3) depicting the connectivity of V-ATPase subunits. The V-ATPase subunits ATP6V1A, ATP6V1E1, ATP6V1D1 and ATP6V1B2 containing high confidence IR-tyrosine kinase phosphorylation and Ser/Thr kinases Cdk2, PRKAA1 (AMPK) and Citron phosphorylation motifs (S9 Table) are marked according to the legend.

https://doi.org/10.1371/journal.pone.0205180.g005

fully characterized and can be supported elsewhere in the cell by the small fraction of the endoplasmic reticulum-associated PTP-1B with high specific activity that can reach the plasma membrane at specific points of cell-cell contact [71], by cytosolic PTPs (SHP1/2) (Fig 2 and Table 1) that couple to RTK phosphorylation in a negative feedback manner at the PM with longer delays [72], and by PTPRs that are thought to display low specific activity towards basal RTK autophosphorylation activities occurring at the cell surface [73]. Through a concerted action on microtubules and actin elements, IREN supports a model in which Cdk2 controls the microtubules-based traffic, and PTPLAD1 is an insulin-dependent switch deciding the choice of IR interaction with microtubules versus actin routing events (Fig 4). Interesting times are ahead for investigating insulin responses in the context of IREN. The question arises as to the extent of crosstalk between the IR-Tyr kinase and the predominantly Ser/Thr kinases (Cdk2, AMPK) that drive IR trafficking and signaling, and when and where this crosstalk occurs. Aprt from the presence of multiple high-confidence substrates for Cdk2 and AMPK in IREN, the current results strongly point to the internalized IR as a relatively pleiotropic writer in the disease module (Tables 1, S4 and S9. For example, ATP6V1E1: Y-464; AMPK: Y-247; ATIC: Y-151; Rab5c: Y-83) and PTPLAD1 as the insulin-dependent eraser with short delay. A related challenge will be systematically matching these phospho-sites to their cognate physiological readers [74].

Another connected example of the IR regulatory mechanism associated with the T2D genetic risk concerns the marked effect of the proton pumping activity on IR in vivo (Fig 5). A concrete problem for the cell concerns the energy sources, and it seems that an efficient solution was found to connect IR activity with intermediary metabolism and trafficking by linking V-ATPase subunits (continuous energy demand) with AMPK (energy sensor and action on IR trafficking) and the metabolic enzyme ATIC (ATP production) (Fig 5) further supporting the idea of the presence of an IR/ATIC/AMPK/PTPLAD1 circuit [13, 75]. The decreased presence of ATIC homodimers, using a small interface interactor, indeed activates AMPK and improve glucose intolerance in a mouse model [76]. We also noted the presence of related candidate enzyme, MTHFD1 (Fig 3 and Table 1: PPIN, GWAS, co-expression). The fact that V-ATPase controls the activity of AMPK [77] emphasizes the idea that all the conditions are present in IREN to auto-regulate this node and thus IR routing, signaling and hepatic clearance in relation to global cell energy status. The presence of the V-ATPase inhibitor RILP [60] in IR immunoprecipitates, which was nearly abolished by PTPLAD1 deletion (Fig 4D), further supports the idea that PTPLAD1 has a large capacity for action to decide IR routing towards early versus late compartments [59].

A facet of IR trafficking in endosomes that can affect indirectly insulin production and secretion is the insulin dissociation/degradation sequence occurring in endosomes, which supports
efficient hepatic insulin clearance [3, 48]. A reduction in hepatic insulin clearance is viewed as an adaptive mechanism that relieves the burden on pancreatic beta-cells [6, 78]. On the other hand, as shown by a mouse model, moderate chronic hyperinsulinemia can be the primary mechanism resulting in insulin resistance [79]. The idea that the complex genetic heterogeneity converges towards a single module co-functional in insulin-producing and target cells, implies a mechanistic promiscuity between insulin signaling, transport and production, that can explain the prevalence of insulin clearance in insulin sensitivity found in some animal models [80].

We acknowledge that some endosomal structures might not be accessible to the IR β-subunit antibody and the limitations inherent to the fractionation approaches such as true tubular connection between different organelles versus contaminants [17, 81]. Nonetheless, the present IREN helps us narrow the search space of the full organism interactome and focus a search in a well-localized network neighborhood. Quantitative proteomic approaches are needed to establish how changes in endosomes occur in space and time according to low (around 10% saturating) versus large saturating insulin doses. This will provide a more complete picture of IREN dynamics that takes the in vivo polarized situation into account [5].

In conclusion, our results establish that the endosomal apparatus contains a T2D disease module located in close proximity to the IR. It senses the state of IR activation and seems co-functional with insulin secretion and islets biology. It helps to explain disease heterogeneity and represents a valuable new resource to understand insulin action and to classify related metabolic traits [82]. Rewiring a network, distorted under the combined genetic and environmental pressures [83], with designed surface interactors [24], provides a mechanistic rationale for the exploration of personalized medicine and elaborate new necessary drugs [1, 2, 84].

Supporting information

S1 Fig. Controls IR containing endosomes and IPs. A) IREP, the number of newly identified proteins from one independent experiment to another (tryptic peptides, see Methods). B) upper panel: The G/E fraction was prepared 2 minutes after insulin injections and incubated with uncoated beads (C), with beads coated with an unrelated IgG (IgG) or beads coated with the anti-IR (2 min). The bound fraction was blotted and incubated with the antibody against IR β-subunit. Lower panel: HEK293 cells were preincubated in serum-free medium for 5 hours and then stimulated for the indicated times with insulin (35 nM). Left, IP with an unrelated IgG (IgG) or the anti-IR antibody (anti-IR). Right, IP with an unrelated IgG (IgG) or the anti-Cdk2 antibody. (TIF)

S2 Fig. The hybrid T2D-protomodule/IRE P module. In total, 126 IREP proteins were selected on the basis of having each at least three interactors among 112 of the 184 seeds of the T2D protomodule (S4 Table-sheet hybrid module). The 126 IREP coding genes make up the list of candidates based on the PPIN approach. The diabetes-associated genes (DAGs) are represented according to the colors indicated in the legend. Orange: Diabetes-associated traits. Blue: Insulin-associated traits. Dark green: Obesity-associated traits. Green: Glycemic traits (S3 Table-sheet validated DAGs seeds). White: IREP candidates. (TIF)

S3 Fig. The extracted IREN subnetwork of DAGs physically associated with candidates. The general topology of IREN is conserved and based on few major hubs, with the kinase Cdk2 displaying the highest centrality (S8 Table). Candidates (yellow and blue colors and black characters; Tables 1 and 2) and DAGs (pink color and black characters) form a single-connected disease module of 94 nodes (33% of IREN nodes) with 330 interactions (28.7% of IREN interactions). An expansion to the first level of adjacent nodes results in a connected subnetwork of 272 nodes.
(88% of nodes) covering 92% of interactions (1070 out of 1147 IREN interactions). The functional groups are represented according to the colors of borders indicated in the legends.

**S4 Fig. LD display (Haploview) of the CDK2 and HACD3 genes.** For all graphs, a reference track with chromosomal location, SNP position, and gene position runs along the top. (Red) indicates strong LD between markers; (white) no LD; (light blue) lack of information to evaluate LD. The block-like patterns of LD are evident in the triangles representing regions of high LD, divided by narrow areas where even adjacent markers are completely independent. The SNP in the red box are associated with T2D. A) Cdk2 genotyped in European population (CEU TSI FIN GBR) as part of the 1000 Genomes project (release 2013/05/03). The pop-up represents the haplotype block of the SNP rs2069408 implicated in T2D (S2 Table). B) The HACD3 genotype in an African population (YRI) as part of the 1000 Genomes project (release 2013/05/03). Three SNPs associate with T2D (rs3759852, rs3784448 and rs3743171, S2 Table) and SNPs in strong LD with these three SNPs in the gene HACD3 (PTPLAD1).

**S5 Fig. Random simulation of T2D subnetworks.** Subnetworks were constructed by 10 000 reiterations of 94 randomly selected IREN nodes. For each simulation, the number of interactions was computed protocols.iodx.doi.org/10.17504/protocols.io.sdqea5w). A) The distribution of the number of nodes for each subnetwork with the neighborhood of 94 selected nodes; the number of nodes observed with the T2D subnetwork is in red. B) The distribution of the number of interactions in the same subnetwork used in A; in red, the number of interaction observed with the T2D subnetwork. C) Distribution of the number of interactions; the number of interactions observed from the 94 T2D nodes is in red.

**S1 Table. Proteome: Proteins and spectra reports.**

**S2 Table. Listing of IREP proteins orthology and networks listing.**

**S3 Table. IREN and T2D-protomodule construction with Hubaa; GO analysis.**

**S4 Table. Gene ontology (GO) subcellular analysis.**

**S5 Table. Fine mapping analysis: LD analysis of IREP coding genes and DAGs variants.**

**S6 Table. TF motifs and coexpression analysis.**

**S7 Table. Source list of T2D and associated traits (glucose intolerance, obesity) genes.**

**S8 Table. Selected DAGs and validated seeds.**

**S9 Table. Kinase-substrate analysis based on Phosphositeplus, Networkin and GPS 3.0 databases.**
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References


