Data-Driven Modeling of Knowledge Assemblies in Understanding Comorbidity Between Type 2 Diabetes Mellitus and Alzheimer’s Disease

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Abstract.

Background: Recent studies have suggested comorbid association between Alzheimer’s disease (AD) and type 2 diabetes mellitus (T2DM) through identification of shared molecular mechanisms. However, the inference is predominantly literature-based and lacks interpretation of pre-disposed genomic variants and transcriptomic measurables.

Objective: In this study, we aim to identify shared genetic variants and dysregulated genes in AD and T2DM and explore their functional roles in the comorbidity between the diseases.

Methods: The genetic variants for AD and T2DM were retrieved from GWAS catalog, GWAS central, dbSNP, and DisGeNet and subjected to linkage disequilibrium analysis. Next, shared variants were prioritized using RegulomeDB and Polyphen-2. Afterwards, a knowledge assembly embedding prioritized variants and their corresponding genes was created by mining relevant literature using Biological Expression Language. Finally, coherently perturbed genes from gene expression meta-analysis were mapped to the knowledge assembly to pinpoint biological entities and processes and depict a mechanistic link between AD and T2DM.

Results: Our analysis identified four genes (i.e., ABCG1, COMT, MMP9, and SOD2) that could have dual roles in both AD and T2DM. Using cartoon representation, we have illustrated a set of causal events surrounding these genes which are associated to biological processes such as oxidative stress, insulin resistance, apoptosis and cognition.

Conclusion: Our approach of using data as the driving force for unraveling disease etiologies eliminates literature bias and enables identification of novel entities that serve as the bridge between comorbid conditions.

Keywords: Alzheimer’s disease, comorbidity, systems biology, type 2 diabetes mellitus

INTRODUCTION

In recent years, comorbidities are inspected with a different perspective. The new route in understanding possible comorbidities has changed from classical approaches that use magnitude, severity, patterns, and burden to comparing disease associated events, pathways, and maps [1, 2]. By establishing comorbid...
associations from assessment scores of these aspects, classical approaches fail to explain the biology underlying diseases. Hence, biological entities such as genes, proteins, and miRNAs and their involvement in biological processes and pathways have been studied to unravel insights about comorbidity.

The possible association between type 2 diabetes mellitus (T2DM) and Alzheimer’s disease (AD) has enticed significant interest from the scientific community since the identification of typical events of T2DM in AD and vice-versa. For instance, the brains of AD patients are reported to exhibit T2DM-related mechanisms including impaired insulin signaling, insulin resistance, and impaired glucose metabolism [3]. Moreover, hyperphosphorylated microtubule associated protein tau (MAPT) leading to formation of neurofibrillary tangles (NFTs), one of the hallmarks of AD, is a consequence of abnormal glycogen synthase kinase 3 beta activity in the insulin signaling pathway [4]. On the other hand, the presence of abnormally processed islet amyloid polypeptide (Aβ) in the pancreas of T2DM patients mimics amyloid-β protein precursor (AβPP)-derived deleterious amyloid-β (Aβ) in AD brains [5]. In addition to these, the comorbid link between T2DM and AD has been established through several studies reporting shared biological processes such as oxidative stress, mitochondrial dysfunction, inflammation, and advanced glycation end products [6].

While most of the speculations are based on individual experiments, studies, or review articles, the putative mechanisms explaining the comorbidity are still unknown. To address this issue, disease-specific knowledge assemblies are created by systematic retrieval of biological information from literature and compared for identifying shared pathophysiological mechanisms. In this regard, Kodamullil et al. (2015) have undertaken a systems biology approach to create cause-and-effect models and proposed single nucleotide polymorphism (SNP)-based mechanisms as the link between the diseases. This is one of the first and few studies that mechanistically depicts and compares disease etiologies [7]. A broader scenario representing mechanistic crosstalk between several pathways such as insulin signaling, neurotrophin signaling, inflammatory regulators, and MTOR signaling in AD and T2DM was demonstrated in our previous work [8]. Interestingly, we have also suggested that metformin, an FDA approved T2DM drug, could be one of the risk factors for developing AD in old age of the diabetic patients. Through this study, the consideration of metformin in drug repositioning in AD has been questioned by depicting the role of metformin in contributing to augment characteristic features of AD such as neuroinflammation, formation of Aβ, and NFTs. Therefore, the hypothesis of drug-induced comorbidity cannot be ruled out. In this context, prolonged use of anti-psychotic drugs has been previously reported to induce symptoms of Parkinson’s disease (PD) [9–11]. The authors have rationalized this assumption by identifying blocked dopamine receptors and calcium channels by the drugs, both of which are impaired in PD. However, the postulation about this aspect of drugs in inducing a disease as a side-effect is still at its infancy.

The prevalence of study bias, which eventually leads to literature bias, is due to the fact that proteins with known biomedical functions and associated signaling pathways are studied recurrently [12, 13]. And because knowledge assemblies massively depend on literature resource, they inherit pre-existing bias. Therefore, chances are higher that literature aided inferences could represent biased knowledge. Taking this into consideration, Naz et al. (2017) have analyzed genomic data and performed functional assessment of prioritized SNPs using literature to depict stress-induced comorbid association in AD and PD [14]. This approach not only eliminates biasedness of over-representation of well-known biological entities and processes, but also identifies new genes and associated events which can serve as putative drug targets and drugable mechanisms. In this study, we have implemented a similar strategy in deciphering the comorbid link between T2DM and AD. The genomic data (i.e., SNPs) for AD and T2DM were fetched from curated public databases and subjected to linkage disequilibrium (LD) analysis. After filtering for shared SNPs in both diseases and prioritizing them based on their relevance to the diseases, we constructed cause-and-effect computable, network models using Biological Expression Language (BEL) [15]. The language enables conversion of unstructured textual information from literature into structured computer-readable triples (i.e., subject-predicate-object). The parsing and compilation of several triples after syntactic and semantic validation generates network models, which are also known as knowledge assemblies. Next, we added the dimension of high-throughput data as the driving force of our analysis by mapping differentially expressed genes to our knowledge assemblies. Finally, a mechanistic graph tailored by analysis of genomic and transcriptomic data was created from the knowledge assembly to explain the comorbid link between T2DM and AD.
Materials and Methods

Firstly, a knowledge assembly embedding prioritized SNP was created from the literature. To this knowledge assembly, we mapped expression profiles from our gene expression analysis. Finally, we filtered the knowledge assembly with those genes which were consistently perturbed. The overall methodology implemented in this study can be divided into 1) data collection, 2) data analysis, and 3) interpretation. Firstly, we collected SNPs and gene expression (GE) datasets for AD and T2DM from freely accessible public databases (i.e., GWAS Catalog [16], GWAS Central [17], dbSNP [18], and DisGeNET [19]). Secondly, for SNP data, we conducted LD analysis followed by identification of shared SNPs and their prioritization using Polyphen-2 [20] and RegulomeDB [21]. Likewise, for GE datasets, we performed differential GE analysis followed by meta-analysis of AD and T2DM datasets. Lastly, we built a literature-derived knowledge assembly representing the results of the SNP analysis and mapped expression profiles of genes from the meta-analysis. A schematic diagram illustrating the methodology is shown in Fig. 1 and described in detail in the following sections.

Retrieval of AD and T2DM SNPs from curated SNP databases

We retrieved a total of 1,130, 1,516, 1,420, and 1,062 SNPs associated to AD from GWAS Catalog, GWAS Central, dbSNP, and DisGeNET, respectively. Similarly, we extracted 1,791, 1,069, 1,865, and 1,522 SNPs associated to T2DM from GWAS Catalog, GWAS Central, dbSNP, and DisGeNET, respectively. To ensure the accuracy of our search, we queried dbSNP, GWAS central, and DisGeNET with corresponding Medical Subject Headings (MeSH) identifiers of AD (i.e., D000544) and T2DM (i.e., D003924). Likewise, we used Experimental Ontology Factor (EFO) identifiers of AD (i.e., EFO_0000249) and T2DM (i.e., EFO_0001360) for GWAS Catalog.

Linkage disequilibrium analysis and SNP prioritization

Using a total of 5,128 and 6,247 SNPs associated to AD and T2DM, respectively, we performed a LD analysis using the R-package haploR [22]. The function queryHaploReg was used with the default $r^2$ threshold of 0.8 to perform the analysis. This yielded 77,486 SNPs in AD and 130,807 SNPs in T2DM. Out
of these, 3,572 SNPs were shared between the diseases. Next, depending on whether the SNPs occur in coding or non-coding region of the gene, we used two databases to functionally annotate these shared SNPs. The impact of the SNPs located in the coding region and the resulting amino acid mutation along with the prediction, either benign or possibly damaging, was assessed using Polyphen-2. Likewise, the assessment of the SNPs in the regulatory region (non-coding) was performed using the function queryRegulome from R-package haploR. Subsequently, we prioritized them using RegulomeDB scores based on current ENCODE releases, Chromatin States from the Roadmap Epigenome Consortium as well as updates to DNase foot printing, Position Weight Matrix for TF binding, and DNA Methylation, and ENSEMBL SNP’s functional consequences [21, 23].

**Literature corpus and cause-and-effect model using Biological Expression Language**

The functional annotation of SNPs using Polyphen-2 and RegulomeDB helps in prioritization of SNPs. Nonetheless, these databases lack their putative roles in a disease context. In this study, we aimed at depicting mechanistic causal graphs embedding prioritized SNPs and their corresponding genes. This was achieved by building a comprehensive knowledge assembly using MEDLINE as the source of literature. The MeSH terms “Alzheimer Disease” and “Diabetes Mellitus, Type 2” were used to query PubMed (Date:02-12-2019) to create separate literature corpus of both diseases. The total number of articles for AD and T2DM were 90,215 and 127,020, respectively. Furthermore, through text mining, we created literature corpora that only contained shared SNPs and genes from LD analysis. The new corpus corresponding to AD and T2DM had a total of 14,293 and 9,032 articles, respectively. Next, we used BEL to capture causal and correlative relationships between the entities from the corpora. The language serves as an efficient platform to create computable knowledge assemblies by compiling relationships which are formulated in the form of triples. The conversion of regular text to BEL was assisted by BELIEF, a semi-automatic workflow to systematically extract BEL relationships from the corpus [24]. The outputs of the BELIEF workflow were manually curated to ensure high quality of the BEL relationships and then compiled using PyBEL for visualization [25].

**Meta-analysis of gene expression datasets**

In this study, our objective is to perform functional assessment of shared SNPs between AD and T2DM with the help of literature derived knowledge assemblies. In the Introduction section, we have already mentioned the possible bias that results from a purely literature-based construction of knowledge assemblies. Therefore, in order to tackle this issue, we mapped and investigated genes with consistent patterns of perturbed expressions to the knowledge assembly as such genes are more likely to be important in disease pathophysiology. A total of 14 GE datasets, 7 each for AD and T2DM, were selected from GEO (Gene Expression Omnibus). The selection of the datasets was done based on the criterion that the samples must be from humans (i.e., patients) diagnosed with AD or T2DM. Moreover, we did not consider datasets that used cell lines, induced medical conditions, animal models and modified genes or environments for expression analysis. The datasets were analyzed with GEO2R tool to identify differentially expressed genes in both diseases [26]. However, because expression patterns of the same disease are inconsistent and non-reproducible [27, 28], we performed a meta-analysis of the AD and T2DM GE datasets independently. This was achieved by using MetaVolcanoR, an R package with an algorithm based on voting approach and p-values of differentially expressed genes [29]. This allowed us to identify consistent patterns of perturbed gene expression across all the datasets. A brief description of each of the datasets is provided in Supplementary File 1.

**RESULTS**

**Linkage disequilibrium analysis**

The distribution analysis of 3,572 shared SNPs revealed that chromosome 1 had the highest number of SNPs, i.e., 495, followed by chromosome 17 (295 SNPs) and chromosome 8 (289 SNPs). The distribution of SNPs over all the chromosomes is shown in Supplementary File 2. The shared SNPs were mapped to 236 genes and the top 5 genes with the highest number of SNPs were lipoprotein lipase (LPL) (CHR 8, 153 SNPs), ubiquitin conjugating enzyme E2 D3 (UBE2D3) (CHR 4, 128 SNPs), leptin receptor (LEPR) (CHR 1, 116 SNPs), FTO alpha-ketoglutarate dependent dioxygenase (FTO) (CHR 16, 94 SNPs), and EF-hand calcium binding domain
Assessment of SNPs with Polyphen-2 and RegulomeDB

A total of 64 SNPs, mapped to 50 genes, were identified by Polyphen-2 to be responsible for amino acid substitutions in their corresponding proteins. Out of these, 50 mutations were predicted to be benign while the remaining 14 mutations were predicted to be possibly damaging. Interestingly, mutations in the few well-characterized genes in AD and T2DM such as apolipoprotein E (APOE), brain derived neurotrophic factor (BDNF), and insulin receptor substrate 1 (IRS1) were classified as “possibly damaging”. The full list of Polyphen-2 output is provided in Supplementary File 4. Likewise, a total of 127 SNPs, mapped to 52 genes, were identified by RegulomeDB to be located in the functional region of their corresponding genes. This was indicated by the scores ranging between 1a and 1f. Genes such as APOE, translocase of outer mitochondrial membrane 40 (TOMM40), and interleukin 6 (IL6) were among the examples for the genes that were mapped to the 127 SNPs. The full list of RegulomeDB output is provided in Supplementary File 5.

Results from meta-analysis of GE datasets

The meta-analysis of AD datasets showed 206 genes to exhibit consistent patterns of perturbed expression, where 49 genes were underexpressed and 157 genes were overexpressed. Similarly, in T2DM, a total of 142 genes regulated persistently, with 13 genes showing downregulation and 129 genes that were consistently upregulated. Out of these, 3 genes, i.e., interferon gamma inducible protein 16 (IFI16), syntrophin beta 2 (SNTB2), and laminin subunit alpha 4 (LAMA4) were found to be overexpressed in meta-analyses of AD and T2DM. The full list of differentially expressed genes and plots showing expression patterns of each datasets are provided in Supplementary Files 6 and 7, respectively. The implementation of GE meta-analysis after differential GE analysis is justified by our findings that the number of coherently perturbed genes reduced with increasing number of GE datasets. This implies the ability of GE meta-analysis to yield robustness and convergence of expression patterns.

Comorbidity in AD and T2DM explained by mechanistic BEL graphs

The knowledge assemblies representing AD and T2DM were combined to investigate the role of shared SNPs and their corresponding genes along with consistently perturbed genes from our meta-analyses. The merged network had a total of 692 nodes and 1,793 edges. The top 5 biological processes based on highest degree of node centrality were insulin resistance, inflammatory response, aggregation of Aβ, apoptotic process, and oxidative stress. Interestingly, cystatin C (CST3), BDNF, peroxisome proliferator activated receptor gamma (PPARG), MAPT, and LEPR were the top 5 genes in the network. We mapped persistently perturbed genes from the meta-analyses to the network and used them as driving force of our comorbid analysis. The rationale supporting this implementation are 1) abnormal expression of genes and their activities mutilate biological processes and pathways and thus are responsible for manifesting disease characteristics, and 2) it overcomes the risk of representing biased knowledge. A mechanistic graph embedding corresponding genes of our SNP analysis and abnormally expressed genes is shown in Fig. 2 It had a total of 41 nodes and 45 edges and comprised of 4 genes (i.e., cholinergic receptor nicotinic alpha 3 subunit (CHRNA3) (CHR 15, 6 SNPs), catechol-O-methyltransferase (COMT) (CHR 22, 4 SNPs), nuclear receptor subfamily 1 group H member 3 (NR1H3) (CHR 11, 13 SNPs), and transforming growth factor beta 1 (TGFβ1) (CHR 19, 4 SNPs) sharing 27 SNPs between AD and T2DM.

As shown in Fig. 2, COMT is known to influence synaptic plasticity and dopamine metabolism, both of which are associated with cognition. In this context, two point mutations (i.e., Val108Met and rs4680 → Val158Met) in this gene were identified to be predictors of cognition scores in AD patients through independent studies [30, 31]. Interestingly, the latter mutation along with rs4646312 in COMT has been associated with T2DM [32]. Also, the 900delC variant form in COMT correlates to chronic renal insufficiency in T2DM [33]. Along the same lines, C47T variant in superoxide dismutase 2 (SOD2) is associated with cognition [34]. Likewise, ATP binding cassette subfamily G member 1 (ABCG1), which is upregulated by NR1H3 [35], has been linked with T2DM because of its involvement in obesity and lipid metabolism [36, 37]. In AD, ABCG1 is reported to inhibit the process of formation of Aβ through inhibition of
Fig. 2. Mechanistic comorbid association: The cartoon diagram represents causal interactions between entities and events led by prioritized SNPs, their corresponding genes and consistently dysregulated genes. Genes such as COMT, MMP9, ABCG1, and SOD2 were identified to contribute to biological processes such as oxidative stress, apoptosis and cognition, all of which are associated with T2DM and AD. Since the onset of T2DM takes place earlier than AD, this diagram is inclined toward AD-related phenomena (i.e., NFT aggregation, synaptic plasticity, etc.) which are influenced by downstream T2DM-related genes.

presenilin 1 (PSEN1) [38]. The amyloid hypothesis in AD considers PSEN1 as one of the two important enzymes that is responsible for abnormal cleaving of AβPP [39], the other being beta-secretase 1 (BACE1) [40]. The accumulated Aβ leading to production of reactive oxygen species (ROS) which further increases oxidative stress to consequently trigger apoptosis is well understood through several studies in AD [41, 42]. Similarly, transcription factor 7 like 2 (TCF7L2)-activated Wnt signaling has been reported to generate ROS and impair glucose homeostasis [43]. The excess of ROS thus produced is detrimental as it results in insulin resistance [44]. Moreover, another study has identified dysfunctional cathepsin D (CTSD) to increase both insulin resistance and apoptosis [45, 46]. In addition, SOD2 induced oxidative stress [47], TNFRSF1A associated via death domain (TRADD) activated Tnf signaling, and cyclin dependent kinase inhibitor 1A (CDKN1A) activated mitogen-activated protein kinase 9 (MAPK9) [48], mitogen-activated protein kinase kinase kinase 5 (MAP3K5), and Fas cell surface death receptor (FAS) also lead to apoptosis [49]. In contrast, the suppression of apoptosis takes place through CHRNA3 [50] and TGFBI activated BCL2 apoptosis regulator (BCL2) [51]. The identification of two SNPs (i.e., rs1801270 and rs1059234) in CDKN1A positively correlated with NFT aggregation in AD patients [49]. The binding of protein tyrosine phosphatase non-receptor type 11 (PTPN11) and MAPT promotes neurogenesis by activating nerve growth factor [52]. Besides, matrix metallopeptidase 9 (MMP9) is also suggested to play a part in neurogenesis and other biological processes such as hypertension, demyelination, and inflammatory response [53, 54]. Lastly, triggering receptor expressed on myeloid cells 2 (TREM2) activity is found to influence inflammatory and immune response [55].

DISCUSSION

In this study, we formulated an integrative approach of combining data and knowledge to unravel new insights about the possible association between AD and T2DM. Our data-driven modeling of knowledge assemblies represents highly
specialized knowledge on comorbidities. While most of the data analytics workflow end up in gene set enrichment analysis, our approach has opened up a new avenue of mechanism-centric interpretation of data. We have used two different data modalities to guide the extraction process of relevant literature knowledge. Firstly, we identified shared SNPs between AD and T2DM from curated resources and built a knowledge assembly around prioritized SNPs and their corresponding genes. Although literature can also be used as a source of SNP information, it is important to note that we have considered only curated databases for retrieval of SNPs. This decision can be explained by the fact that curated databases ensure association between a SNP and a disease with a given statistical significance (i.e., p-value). Unlike curated databases, some SNPs mentioned in the literature might not have any association with a disease because the statistical power of association is below par [56–59]. Therefore, by including such SNPs, we would be adding possible false positives in our analysis and, thus, diminishing the quality of the results. Secondly, as we are aware of the literature bias in knowledge assemblies, we identified consistently perturbed genes by conducting a GE meta-analysis and used these signals to mechanistically link AD and T2DM. Our results illustrate that genes such as COMT, MMP9, SOD2, and ABCG1, which do not belong to the realm of well-known genes in AD and T2DM, are involved in important biological processes of both diseases. This suggests dysfunctional activities of these genes could be the bridge between these diseases. Moreover, our findings endorse and strengthen the proposition of AD and T2DM comorbidity suggested by epidemiological, preclinical, and pathophysiology studies by identifying novel genes.

The genetic variants of AD and T2DM amassed in our study are readily explorable and bear the potential to yield new insights. For instance, genomic loci dependent SNPs can be functionally assessed to uncover their roles in the underlying comorbid mechanisms. This would enable identification of "genomic hotspots" that are closely associated to AD and T2DM. However, this study does not address this aspect due to time constraints and it is out-of-scope of our objectives. Also, we have not considered the role of epigenetic modifications in the comorbid association between AD and T2DM. Nonetheless, our knowledge assemblies can be used as the starting point for assimilating epigenetic modifications concerning AD and T2DM.

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SUPPLEMENTARY MATERIAL

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