mutTCPdb: a comprehensive database for genomic variants of a tropical country neglected disease—tropical calcific pancreatitis

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Abstract

Tropical calcific pancreatitis (TCP) is a juvenile, non-alcoholic form of chronic pancreatitis with its exclusive presence in tropical regions associated with the low economic status. TCP initiates in the childhood itself and then proliferates silently. mutTCPdb is a manually curated and comprehensive disease specific single nucleotide variant (SNV) database. Extensive search strategies were employed to create a repository while SNV information was collected from published articles. Several existing databases such as the dbSNP, Uniprot, miRTarBase2.0, HGNC, PFAM, KEGG, PROSITE, MINT, BIOGRID 3.4 and Ensemble Genome Browser 87 were queried to collect information specific to the gene. mutTCPdb is running on the XAMPP web server with MYSQL database in the backend for data storage and management. Currently, the mutTCPdb enlists 100 variants of all 11 genes identified in TCP, out of which 45 are non-synonymous (missense, nonsense, deletions and insertions), 46 are present in non-coding regions (UTRs, promoter region and introns) and 9 are synonymous variants. The database is highly curated for disease-specific gene variants and provides complete information on function, transcript information, pathways, interactions, miRNAs and PubMed references along with remarks. It is an informative portal for clinicians and researchers for a better understanding of the disease, as it may help in identifying novel targets and diagnostic markers, hence, can be a source to improve the strategies for TCP management.

Database URL: http://lms.snu.edu.in/mutTCPDB/index.php
Introduction

Tropical Calcific Pancreatitis (TCP) is a juvenile, non-alcoholic idiopathic chronic pancreatitis, prevalent in the tropical regions with unknown aetiology and is defined under the idiopathic category of TIGAR-O (i) toxic-metabolic, (ii) idiopathic, (iii) genetic, (iv) autoimmune, (v) recurrent and severe acute pancreatitis or (vi) obstructive) classification system (1). In 1980, juvenile tropical pancreatitis was described by the authors in the journal Lancet (2). Then, in the same year, Tan et al. described specifically ‘TCP’, for the first time (3). TCP is associated with clinical manifestations that include severe abdominal pain, pancreatic calculi, steatorrhea, bilateral parotid enlargement, cyanotic hue of lips and fibrocalculus pancreatic diabetes (FCPD). FCPD is the characteristic feature of TCP and a distinct form of diabetes (Ketosis resistant) as described by World Health Organization (4). There are also evidences of head mass or tumors observed in 30–75% of TCP patients as reported by the authors (5). Hence, TCP can lead to malignancy in late stage. Morphologically, TCP is characterized by pancreatic ductal dilation, large dense calculi (6), pancreatic atrophy and fibrosis (7–9). Zuidema, in 1959, first reported pancreatic calculi and symptoms of malnutrition in many patients (10). Down the line, most studies were reported from Indian subcontinent. However, reports were also recorded from other tropical countries in Asia (11) (Malaysia, China, Japan, Bangladesh (12)), Africa (13) (Uganda, Nigeria (14)), South America (Brazil) and Mexico (15). A survey on chronic pancreatitis (CP) in Asia-Pacific region found that 70% of CP patients in India and China fall in the criteria of TCP (16). The symptoms of TCP overlap with other types of pancreatitis (alcoholic, hereditary and drug induced), therefore there is still a lack of strategic clinical management of this disease.

Extensive calcification of exocrine pancreas, large stones and dilated pancreatic duct are some of the phenotypic diagnostic markers for TCP in non-alcoholic patients. But there are no potential molecular/genetic biomarkers identified till now, because of the lack of objective knowledge about the pathophysiology involved in TCP. The treatment for TCP is mostly surgical in the form of removal of all intraductal stones. Chromogenic techniques (17) such as endoscopic retrograde cholangiopancreatography, computed tomography scan, trans-abdominal ultrasonography, and so on can clinically diagnose the pervasive calcification in the exocrine pancreas.

Our database, ‘mutTCPdb’ contains single nucleotide variants (SNVs) present in genes, so far identified in TCP patients (patients with non-alcoholic chronic pancreatitis). The genes defined in the database are; extracellular calcium sensing receptor (CASR), cystic fibrosis transmembrane conductance regulator (CFTR), chymotrypsin-C, cathepsin B (CTSB), pancreatic secretory granule major glycoprotein GP2 (GP2), protease, serine, 1 (trypsin 1) (PRSS1), protease, serine, 2 (trypsin 2) (PRSS2), regenerating islet-derived 1 alpha (REG 1 A), serine peptidase inhibitor, Kazal type 1 (SPINK1), transcription factor 7-like 2 (TCF7L2) and carboxypeptidase A (CPA1). These genes have variants that are associated with non-alcoholic chronic pancreatitis in the tropical regions. There is a research article, which negates the association of angiotensin converting enzyme (ACE) gene to TCP, based on direct DNA sequencing studies of ACE gene (18). Recently, an association between CLDN2-MORC4 and PRSS1-PRSS2 locus variants have been established with TCP (19). A variant C677T in 5, 10-methylenetetrahydrofolate reductase was found to be associated with tropical chronic pancreatitis patients from North Indian population (20).

In general, ‘mutTCPdb’ was created with an objective to organize the sparsely distributed data on genetic variants studied so far in TCP. This manually curated comprehensive database is an integrated and updatable mutations resource for clinicians and scientists working on TCP.

Materials and methods

Data collection

TCP is a neglected tropical disease, therefore limited information is available. Initially, the PubMed (http://www.ncbi.nlm.nih.gov/pubmed) was searched with terms ‘TCP’ [Title/Abstract/Text Word], ‘Tropical Pancreatitis’ [Title/Abstract/Text Word], ‘Tropical Chronic Pancreatitis’ [Title/Abstract/Text Word] and ‘Chronic calcific non-alcoholic pancreatitis’ [Title/Abstract/Text Word]. Consequently, around 500 published articles since 1980 were enlisted for the above queries till July 2017. Additionally, the NCBI portal (https://www.ncbi.nlm.nih.gov/), Ensemble Genome Browser 87 (http://asia.ensembl.org/index.html) and UNIPROT (http://www.uniprot.org/) sources were also used for supplementary information.

Database organization

After retrospectively screening the literature, SNVs were extracted from published articles and the data were organized in various levels. Each variant in a specific gene was annotated with an ID (e.g. tcp8461). The initial query starts with Gene_symbol (e.g. CASR, CFTR), Entrez_ID (e.g. 846) or Gene_ID (e.g. ENSG0000036828) and then divided into the following headings (Figure 1).

(A) Summary
This section provides the information of every gene associated with TCP and studied for variants. The significance of
the respective gene to be involved in TCP was curated from publications on PubMed (https://www.ncbi.nlm.nih.gov/pubmed).

(B) SNV information
The information for SNVs in respective genes is subdivided in the following descriptors:

1. **Variants.** The variant information of each single nucleotide variant (SNV) in a gene is divided in two categories: (i) databases-derived information and (ii) literature-derived information.

   ‘Literature-derived information’ segment has the curated information (genomic data) for each variant available in the published article. If the data for a specific variable (e.g. Chromosome Refseq_ID, Nucleotide Refseq_ID etc.) were not available in the literature, the column was left with a hyphen (-).

   For each variant, the ‘database-derived information’ segment has complete transcript information available on dbSNP database build 150 (http://www.ncbi.nlm.nih.gov/SNP/) with respect to human genome assembly, GRCh38.p7. Every variant is defined and attached with a ‘remark’ column to simplify the user query. Besides, cross reference (Cross_Ref) column list other diseases, if associated with that particular variant.

2. **Evidence information.** This section contains all the relevant information from the published literature about variants identified in that particular gene.
3. Transcript information. The data in this table include, gene_name, Alias, Entrez_ID, Locus, Strand, Gene_ID, Transcript_ID, Protein RefSeq_ID, Length(aa), mRNA RefSeq_ID, Length(bp), Exons, Coding exons, Protein RefSeq_ID, Type, CCDS, Uniprot_ID. The information respective to each variable is extracted from NCBI portal and Ensemble Genome Browser 87.

(C) MicroRNA predictions
This tab includes validated MicroRNAs (miRNAs) targeting each TCP-related genes. The miRNA data were obtained from miRWalk2.0 algorithm and miRTarBase database (21).

(D) Cross-reference
The miscellaneous identifiers for each gene are listed in this section for user reference.

Web interface and data browser
The web interface for mutTCPdb is uncomplicated, researcher friendly and interactive. mutTCPdb focuses on providing better navigation through individual sections to increase data discoverability. There are six tabs provided at the top of the interface (‘Home’, ‘About’, ‘Browse’, ‘Stats and FAQs’, ‘Submit Data’ and ‘Contact Us’) through which users can navigate and explore the required information. mutTCPdb is running on XAMPP web server with MYSQL database in the backend for data storage and management. Text query box is provided at the top of each page to search by Gene name, Entrez ID or Transcript ID related to every gene. The data can be accessed either by browsing through the predefined lists (provided at ‘Browse’ tab) or through search box. The snapshots of the database are illustrated in Figures 2 and 3.

Query processing scripts are written in PHP and Perl. The database is tested and works well with commonly available web browsers, such as Mozilla Firefox, Google Chrome, safari and Microsoft Internet Explorer.

There is a provision of submitting new data to mutTCPdb under the ‘Submit Data’ tab. The user can submit the data using an interactive form and then upload the file in any format. Once the data are submitted, the admin will verify the information from the references cited and also check for the duplicate entry in mutTCPdb. If the admin finds no discrepancy, the information will be uploaded and made available to the users. Additionally, mutTCPdb 1.0 will be updated regularly, every 24 months, with new data and novel features.

Statistics and results
The database (mutTCPdb) consists of 100 variants found within 11 genes studied so far associated with TCP patients (complete statistics is described in Figure 4). These variants (Figure 5) were further classified into missense (n = 37), nonsense (n = 1), deletions–insertion variants (n = 7) and synonymous variants (n = 9). Non-coding variants were subdivided as, 5’UTR (Untranslated regions) variants (n = 15), 3’UTR variants (n = 5) and intronic variants (n = 26). Deletion–insertion variants were positioned in introns, exons or in UTRs. We used GeneHancer to check whether the variants present in the non coding regions fall in gene enhancer regions or not (22). Out of all the genes in the
database, we could only find seven variants in 5'UTR regions present in CTB gene and two variants in TCF7L2 gene, to fall in enhancer regions (Table 1). Each variant was specifically extracted from a published article and linked with its population data. The ‘Description’ and ‘Function’ heading gives the information about the gene location, activity and role. The ‘Significance in TCP’ heading explains the rationale behind a gene to be considered as pathogenic in TCP. For example, the CPA1 gene has been included in database based on the research article by Witt et al., which describes the variants in CPA1 gene to be associated with TCP (23). Also, the article describes that the reduced activity of mutant CPA1 gene during experimental study might be due to reduced secretion of CPA1 as a result of misfolding in ER as stated in ‘Significance in TCP’ tab. Hence, the selected genes have been curated from published articles on the basis of variants present in that gene, respectively, identified in TCP patients.

**Conclusion**

mutTCPdb provides information regarding SNVs associated with TCP. This is first effort to manually resource all the variants present in the literature and make a comprehensive repository for TCP related SNV. Till now, neither the diagnosis nor the medications are specific for TCP because of overlapping symptoms with other forms of pancreatitis and lack of defined etiopathogenesis. ‘mutTCPdb’ will help in
characterizing the disease further by studying TCP at the molecular level and unfold the enigma hovering over the pathogenesis of TCP, which can rather be interpreted as an inflammation without a cause'. In future, mutTCPdb will be used to correlate present data with the next generation sequencing results like Exome and RNA sequencing, which can definitely enrich the current database. The first update of mutTCPdb will be released next year and further updates will be available after every 2 years.

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**Conflict of interest** None declared.

**References**


**Table 1.** The table indicates the 5'UTR variants in CTSB and TCF7L2 genes with there corresponding Gene Enhancer IDs in the second column

<table>
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<th>TCP ID</th>
<th>Gene Enhancer ID</th>
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<td>tcp69342</td>
<td>GH10G0113047</td>
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</table>

The first column has TCP variant ID’s for CTSB and TCF7L2 genes.


