ATP7A Clinical Genetics Resource – A comprehensive clinically annotated database and resource for genetic variants in ATP7A gene

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ATP7A is a critical copper transporter involved in Menkes Disease, Occipital horn Syndrome and X-linked distal spinal muscular atrophy type 3 which are X linked genetic disorders. These are rare diseases and their genetic epidemiology of the diseases is unknown. A number of genetic variants in the genes have been reported in published literature as well as databases, however, understanding the pathogenicity of variants and genetic epidemiology requires the data to be compiled in a unified format. To this end, we systematically compiled genetic variants from published literature and datasets. Each of the variants were systematically evaluated for evidences with respect to their pathogenicity and classified as per the American College of Medical Genetics and the Association of Molecular Pathologists (ACMG-AMP) guidelines into Pathogenic, Likely Pathogenic, Benign, Likely Benign and Variants of Uncertain Significance. Additional integrative analysis of population genomic datasets provides insights into the genetic epidemiology of the disease through estimation of carrier frequencies in global populations. To deliver a mechanistic explanation for the pathogenicity of selected variants, we also performed molecular modeling studies. Our modeling studies concluded that the small structural distortions observed in the local structures of the protein may lead to the destabilization of the global structure. To the best of our knowledge, ATP7A Clinical Genetics Resource is one of the most comprehensive compendium of variants in the gene providing clinically relevant annotations in gene.

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1. Introduction

Copper (Cu) is an essential trace element vital to the health of all living organisms and is crucial for the normal development of the nervous system. It is required for a variety of biological processes in the body including myelin formation and connective tissue synthesis. It is required by cuproenzymes like ceruloplasmin, a major circulating transport protein for copper, lysyl oxidase, an enzyme essential for initiating the crosslinking of connective tissues by oxidizing peptidyl lysine in elastin and collagen [1]. Human body requires copper for its cellular metabolism. The copper transporter 1, CTR1, plays a major role for cellular Cu uptake in the intestinal epithelium and in hepatic cells. Various chaperones such as Antioxidant 1, Copper Chaperone (ATOX1) and, Human Cytochrome c oxidase copper chaperone (hCOX17) distribute the Cu to specific proteins or organelles. Cu is transported from enterocytes into blood circulation by ATP7A whereas ATP7B incorporates Cu in liver ceruloplasmin and also mediates biliary excretion of Cu [2].

Various genes are involved in Cu metabolism, such as CTR1 which is a high affinity Cu transporter mediating copper uptake, DMT1 another mediator of Cu uptake. Another important gene involved in Cu transport is ATOX1 which transports Cu from the cytosol to transporters ATP7A and ATP7B. ATP7A and ATP7B are located in the trans golgi network, ATP7A pumps Cu into blood and ATP7B is responsible for maturation of ceruloplasmin [3]. ATP7A located at chromosome Xq21.1 spans about 150 kb of genomic DNA and contains 23 exons. The size of individual coding exons varies between 77 and 726 bp and introns vary in size between 196 bp and approximately 60 kb [4]. As described by [5] ATP7A is a member of the P1B-subfamily of the P-type ATPases.
The gene encodes a transmembrane copper-transporting P-type ATPase. Copper-transporting ATPases (Cu-ATPases) mediate the copper transport, these ATPases are required for the growth, development and also various physiological processes. Cu-ATPases transport copper from the cytosol across cellular membranes, this process reduces the intracellular copper concentration and also controls the copper homeostasis in the body.

Genetic anomalies in the Cu-ATPase encoding ATP7A are associated with a fatal neurological disorder such as Menkes disease (MD), Occipital Horn Syndrome (OHS) which is a mild form of MD and X-linked distal spinal muscular atrophy. ATP7A is ubiquitously expressed in extracellular cells and tissues, therefore, systemic defects are caused due to its absence or inactivation, in MD or OHS patients. MD is an X-linked recessive disorder which results in copper deficiency. Two forms of MD have been described: classic MD and mild MD, based on the symptoms mild MD is a less severe form. Classical MD is characterized by neurological defects and peculiar “kinky” hair. MD is caused by mutations in the ATP7A gene. One-third of the MD cases are due to de novo mutations in ATP7A [6]. Copper absorption is impaired in MD and intracellular copper trafficking is disrupted by mutations in the ATP7A gene [4]. Similar to MD, OHS is also a rare disorder occurring due to impaired copper absorption. OHS patients are characterized by “occipital horns” which are downward pointing exostosis arising from the occipital bone and certain connective tissue deformities comprising cutis laxa, hernias, joint laxity and bladder diverticula [7]. X-linked distal spinal muscular atrophy type 3 is a rare distal hereditary motor neuropathy. This disease is characterized by weakness of distal muscles of hands and feet, distal motor weakness particularly of the lower limbs [8].

In the present study, we have developed a comprehensive clinically relevant resource for genetic variants in the ATP7A gene by systematic curation and annotation of the genetic variants, with appropriate classification according to ACMG-AMP guidelines. We have also estimated the carrier frequencies of the variants in global populations thus explicating the genetic epidemiology of the disease. In this manuscript, we also present our molecular modelling studies for six variants, to elucidate the stability of these variants and the mutation-structure relationship. To the best of our knowledge, this resource is one of the most comprehensive compendium of variants in the ATP7A gene, with systematic annotation and classification according to the ACMG-AMP guidelines.

2. Materials and methods

2.1. Data curation and contents

The genetic variants in ATP7A associated with diseases were curated from publications indexed in PubMed and PubMed Central databases using formatted queries including the gene name and disease names. The data was systematically entered into a preformatted spreadsheet. Additional variants listed in ClinVar and Locus Specific Databases were also checked and linked to the data curation. The preformatted spreadsheet also collected a number of fields including the genome build, variant loci including the chromosome, start position, end position, Reference base (Ref), alternative base (Alt); gene and the amino acid change was collected. In addition, the technique used for identification, ethnicity, geographical origin and population from where the variant was identified were collected. The nomenclature of the variants as per the Human Genome Variation Society guidelines were also generated. In addition, the variant types as defined by nonsense, frameshift, splicing, and deletion/insertion, missense, UTR3', UTR5' were also annotated.

2.2. Validation of curated data

All the variants which were curated from the various platforms were rechecked in order to ensure the correctness of the variant positions as well as the variant notations as per the Human Genome Variation Society (HGVS) nomenclature guidelines. Two web based utilities – LUMC Mutalyzer and Variant Validator were utilized.

Mutalyzer (https://mutalyzer.nl/) is a Web interface used for constructing, validating, and transforming sequence variant descriptions. Mutalyzer evaluates sequence variant nomenclature according to the guidelines of the Human Genome Variation Society (HGVS) nomenclature. The DNA tool Position converter was used for finding the missing pieces of data obtained originally. The Position Converter depends on mapping information from the NCBI. The Position Converter converts the variant description positions from the chromosomal position for a specific human genome build i.e GRCh37 / hg19 to a position relative to RefSeq transcript reference sequences present in the local mapping database or vice versa.

Variant Validator (https://variantvalidator.org/) is a web-based variant validation tool which provides an interface which allows the validation of genomic variations published in scientific literature or databases. It is a Web interface which is used to describe sequence variants. We validated all the curated variants through Variant Validator, in order to correct the erroneous variants published in the literature sources if any [9].

2.3. Functional annotation of the variants

All the curated variants post-validation were systematically annotated using the ANNOVAR package. The gene based annotations were retrieved for the variants. In addition, Computational annotations of the variants such as deleteriousness-prediction scores such as SIFT, Polyphen, CADD etc for predicting the deleterious nature of the variants were obtained. In addition, the allele frequencies across ExAC, gnomAD, ESP6500, 1000 genome datasets were retrieved.

2.4. Interpretation of pathogenicity of sequence variants

The variants were systematically analysed and classified as per the guidelines for interpretation of sequence variants as put forward by the ACMG & AMP.

The following attributes were decided on the basis of population data i.e. allele frequencies of the variant from the three databases 1000 Genomes Project, ESP6500 exome-sequencing project, ExAC 65,000 exomes data set. BA1 (Allele frequency is >5%); BS1 (Allele frequency in between 0.01 and 0.05); PM2 (Allele frequency absent or <0.0005). The following attributes were decided on the bases of computational data like SIFT, Polyphen and CADD score generated through annovar run. PP3 (If two of the three computational data shows deleterious effect); BP4 (If two of the three computational data shows tolerated/benign effect). The following attributes were decided on the basis of functional data; PS3 (If in vitro or in vivo functional assay support the damaging effect of the variant on the gene or gene product); PM1 (Variant must be present in functionally important protein domain) this was evaluated from the pfam protein families database. The following attributes were decided on the basis of Clinvar significance; PP5 (If Clinvar reports the variant as pathogenic or likely pathogenic); BP6 (If Clinvar reports the variant as Benign or likely benign). The following attributes were decided on the bases of segregation data; PP1 (Cosegregation with disease in multiple affected family members); BS5
(Lack of segregation in affected members of a family); PS2 (De novo confirmed); PM6 (De novo assumed); PP4 (Large family segregation); The following attributes were decided on the bases of variants; BP7 (Synonymous variants); PVS1 (Frameshift, Non-sense, Splice-site variants); PP2 (Missense variant in a gene that has a low rate of benign missense variation); BP1 (Missense in gene where only truncating cause disease); Other attributes; PS1 (Same amino acid change, previously described as pathogenic); PM5 (Missense change at a position where a different amino acid change described as pathogenic before [10]. These criteria were used to annotate each variant to the five classes as “pathogenic,” “likely pathogenic,” “uncertain significance,” “likely benign,” and “benign” as per the algorithm.

2.5. Comparison of allele frequencies in different global datasets

The variants that were classified as Pathogenic and Likely pathogenic according to the ACMG guidelines, were mapped to different global datasets of 1000 Genomes, ExAC and gnomAD version 2. The allele frequencies in the different subpopulations of the variants mapping to different datasets were compared to the global allele frequency in their respective dataset using Fisher’s Exact test. The p values thus obtained were corrected using bonferroni correction method. The allele frequencies in the subpopulations having corrected p value less than 0.05 were defined to be significantly different as compared to the global allele frequency of that variant in the respective dataset.

2.6. Molecular modeling studies

The variants that were classified as “pathogenic and likely pathogenic” according to ACMG guidelines, were subjected to detailed molecular modeling studies to provide a mechanistic explanation for their pathogenicity. Although we have multiple entries with pathogenic and likely pathogenic labels, we restricted our studies only to those that have tertiary structures. To select the variants that have tertiary structures, we systematically matched the position of variants against the structure of Copper-transporting ATPase 1 (ATP7A). Though there are 6 isoforms of ATP7A reported, we chose isoform-4 (contains 1500 amino acids) for our studies since it was reported as a canonical sequence.

2.7. Assessment of structural stability upon mutations

After systematically matching the respective positions of mutations onto the structure, we selected several mutations for our modeling studies. All those mutations underwent structural stability calculations using multiple tools. We assumed a consensus-based ranking would provide a better understanding of the deleterious nature of the mutations via their structural distortions. Hence, we used 9 tools which consist of standalone molecular modeling packages such as FoldX [11], Schrödinger with two differ (ent force fields, OPLS2005 [12] and OPLS3 [13], MOE [14] and web servers such as CUPSAT [15], mCSM [16], SDM [17], iMutant 2.0 [18] and POPMUSIC [19]. The outputs of these tools were categorized as ‘destabilizing,’ ‘stabilizing’ and ‘neutral’ based on the reference values of individual tools and a heat map was plotted.

2.8. Molecular dynamics simulations

In order to corroborate the results obtained from the previously used tools, we performed molecular dynamics simulations for 50 ns. We found that the mutation A629P lies in the solution structure of the apo form of the sixth soluble domain of ATP7A (PDB ID: 2KMV) [22]. To perform MD simulations, we selected the first structure from the ensemble of solution structures and mutations were introduced using COOT [23]. Later, all these mutants along with the wild types (WT) were subjected to MD simulations with Desmond molecular dynamics package by preparing the proteins using the protein preparation wizard of Schrödinger Maestro, where the protein preparation was executed by adding hydrogen atoms, specifying the bond orders, incorporating protonation states for protein residues, optimizing hydrogen bond network and running a short energy minimization with a RMSD cutoff of 0.30 Å. The prepared proteins (both WT and mutants) were soaked into an orthohemibic water box that contained TIP3P water molecules. The size of the box was set in such a way to include the entire protein. Also, the system was neutralized by applying the respective number of Cl- atoms. All MD simulations were performed using NPT ensemble where the pressure and temperature was set to default i.e. 1.01325 bar and 300 K respectively. We recorded the trajectory and energy at every 50 ps and 1.2 ps respectively.

2.9. Database and web server

The database search tool was designed to allow users to easily explore variants in gene ATP7A from the database. The variant data is stored in MongoDB v3.4.10. The data can be accessed through a web interface running on Apache HTTP server using PHP 7.0. The user-friendly web interface for querying the database is coded in PHP 7.0, AngularJS, HTML, Bootstrap 4 and CSS. MongoDB v3.4.10 was used to keep track of data processing through the web interface.

3. Results

3.1. Compendium and classification of genetic variants

The compendium of curated variants encompasses a total of 602 variant entries in the ATP7A gene. These variants were derived from a total of 64 publications [6, 8 and 24–82] and encompassed variants reported from 17 countries. Of the total compendium of variants, a total of 404 variants were unique and a large majority of variants mapped to exons 18,18% (328/404), while a small number were intronic 12% (49/404), splicing 6.18% (25/404) and UTR 0.4% (2/404) variants (Fig. 1A). The variants were also classified by their potential functional implication. A majority of variants were classified as non-synonymous (226/404), while (14/404) were classified as synonymous. (52/404) variants caused a stop-gain, while (31/404) variants caused a frameshift, (3/404) were non-frameshift and (2/404) were insertions. The variant classes are summarised in Fig. 1B.

3.2. Classification of variants as per the ACMG & AMP guidelines

The variants were further reclassified as per the ACMG & AMP guidelines for interpretation of the pathogenicity of variants. All variants were classified into one of the five categories - Pathogenic, Likely Pathogenic, Benign, Likely Benign and Variant of Uncertain Significance (VUS). The classification revealed that 87 variants could be classified as Pathogenic (21.53%), while 56/404 could be classified as Likely pathogenic (13.86%), 22/404 could be classified as likely benign 5.44% and 10/404 as benign (2.47%). A vast majority of variants, 229/404 could only be classified as a VUS (56.68%) for lack of evidence to classify them as Pathogenic or Benign (Fig. 1C). Of the Pathogenic / Likely Pathogenic variants, 136 are
associated with Menkes disease, while 11 are associated with X-linked distal spinal muscular atrophy type 3 and 6 are associated with OHS (Fig. 1D).

3.3. Distribution of likely pathogenic and pathogenic variants in the ATP7A gene

Our analysis revealed a total of 143 variants which could be classified as pathogenic / likely pathogenic as per the ACMG & AMP guidelines. The variants were systematically mapped to the ATP7A protein structure and domains. A total of four functional protein domains were annotated as per Pfam database using maf-tools package in R-programming. These included six heavy metal associated domains (HMA), E1-E2 ATPase, HAD like hydrolase and ATPase IB-1. With an exception of a few variants, all the pathogenic/likely pathogenic variants mapped to functional domains of the protein (Fig. 2).

3.4. Global allele frequencies

All the variants annotated as Pathogenic and Likely Pathogenic were further checked in global population genome datasets for their allele frequencies.

The datasets considered include the 1000 Genomes, ExAC and the gnomAD (version 2). Only a total of 5 likely pathogenic variants, of the 143 likely pathogenic and Pathogenic variants, mapped to any of the global datasets. The variant rs782237314 shows frequency of 0.00002 in the gnomAD dataset and ExAC dataset while it was absent in the 1000 Genomes dataset. This variant was only observed in the non-Finnish european subpopulation of both the datasets. The variant rs367775730 shows frequency of 0.0002 in the gnomAD dataset and ExAC dataset while it was absent in the 1000 Genomes dataset. This variant had a significantly high allele frequency in the Ashkenazi jewish subpopulation as compared to the global allele frequency of gnomAD dataset. The variant
rs374162669 shows a frequency of 0.0003 in 1000 Genomes and 0.0001 in gnomAD and ExAC datasets. The variant rs138958687 shows frequency of 0.0006 in the 1000 Genomes and 0.0007 in the gnomAD and ExAC datasets. This variant has significantly different allele frequency in the South Asian, Non-Finnish european and Finnish subpopulations as compared to the global allele frequency in the gnomad dataset, while in the ExAC dataset, it showed a significant difference in the Finnish subpopulation as
compared to the global allele frequency. The variant rs781995242 shows frequency of 0.00001 in the gnomAD and ExAC datasets while it was absent in the 1000 Genome dataset. Fig. 3 shows the distribution of pathogenic variants in the population datasets.

3.5. Molecular modeling studies

After systematically matching the respective positions of mutations, we selected 6 variants for our studies; A629P, R844C, R844H, P852L, G876R and V1180D. These variants lie in three different domains of the ATP7A structure. In our stability predictions, we found FoldX predicted all of them as ‘neutral’. Similarly, when applied to two different force fields, i.e., OPLS2005 and OPLS3, Schrödinger predicted as ‘stabilizing’ for variants R844C, R844H, P852L, G876R. In the same way, SDM predicted A629P and R844H as ‘stabilizing’. Also, POPMUSIC predicted R844C, R844H as ‘stabilizing’. The iMutant 2.0 was unable to predict any effect for V1180D. However, most of the tools in our consensus-based stability prediction predicted the variants as ‘destabilizing’. The stability prediction heatmap is shown in Fig. 4A.

Since the majority of the tools predicted these variants as destabilizing, we assumed that our MD simulations would provide more insights into the reasons for the destability. To elucidate the mutation-structure relationships we mainly computed and compared the root mean square deviation (RMSD) of the $C_a$ and root mean square fluctuations (RMSF) of individual residues of mutant and WT protein over a 50 ns simulation time. The potential energy of mutant and WT was also calculated and compared between the mutants and WT.

In the case of A629P (Fig. 4B), the RMSD of the mutant does not deviate much when compared to the WT (Fig. 4C). There was only
0.10 KJ difference between potential energies of mutant (4.25 x 10^-4 KJ) and WT (4.15 x 10^-4 KJ) throughout the 50 ns MD simulations (Fig. 4D). We found that the RMSF of mutant has deviated significantly at one of the flanking regions when compared to the WT (Fig. 4E and F). Our analysis revealed that the side chains of lysine and glutamic acid at the flanking regions were fluctuating drastically. When comparing the ensembles of the experimental structures, the RMSF of lysine and glutamic acid at the flanking region are also highly distorted as we observed in our MD simulations. From our simulation studies, we could conclude that the A629P mutation didn’t impart any structural changes when compared to the WT. Our results are in good agreement with the previously reported experimental studies [20].

In the case of R844C, R844H, P852L and G876R (Fig. 5A), the RMSD of all the mutants agree with that of WT (Fig. 5B). RMSDs of all the proteins including WT were stabilized between 2 and 4 Å. Similarly, the potential energies (Fig. 5C) of the mutants and WT were between 6.35 x 10^-4 KJ to 6.45 x 10^-4 KJ which indicate the energies of the systems are well maintained throughout the MD simulations irrespective of the mutation. We then computed the RMSF of the mutants and compared with that of WT and found that the RMSF of both mutants and WT are stable (Fig. 5D–H). These results lead us to think that any of these mutations do not impose detectable structural changes when compared to the WT.

Finally, for V1180D (Fig. 6A), we found that the RMSD does not deviate much for the mutant when compared to the WT. For both mutant and WT, the RMSD was stabilized between 10 and 12 Å (Fig. 6B). To understand the drastic jump in RMSD, we critically analyzed the distortions in the structures throughout the MD simulations and found that most of the structural distortions happened in the region that connects β3 and β4. High flexibility for this region was previously reported in the experimental studies [22]. The potential energy and RMSF of both mutant and WT were almost the same throughout the MD simulations (Fig. 6C and D). However, when we compared the overall secondary structural elements, we found that there was a 5% reduction in the beta strands for the mutant. The mutation V1180D was located at the β4 strand. When valine is replaced with aspartic acid, there is not sufficient space to accommodate the side chain of the aspartic acid and due to that there is structural rearrangement which is reflected in the reduction in the secondary structural elements.

3.6. Database interface and features

The ATP7A genetic variant resource features a user friendly web based interface which allows query and browsing the compendium of variants. The search interface allows query of the compendium using variant, ACMG, AAchange and dbSNP ID. Variants can be queried by formatted queries containing the chromosome, variant position, reference and alternate alleles. The query retrieves the list of all the matching entries from the database in a tabular form with an option to open the variant details. The variant details are compiled in 4 broad sections The first section includes basic information of the variant i.e. Gene name, Chromosome, Genome Build, HGVS Nomenclature, genomic locus, reference and alternate bases and the amino acid change. The second panel provides more information on the variant with respect to the functional effect of the mutation, and the classification as per the ACMG & AMP guidelines. This section also provides information on the disease along with the inheritance patterns, technique used for identification of the variant, ethnicity from which the variant has been reported and links to the variant in dbSNP database. The last sections provide information on the geographical / population information from where the variant was reported and the publication from which the variant information was retrieved.
4. Conclusions

The ATP7A Genetic Variant Resource is a comprehensive compendium of genetic variants in the ATP7A gene. The variant compendium indexes 404 variants in the ATP7A gene. The database also provides the annotations as per the ACMG & AMP guidelines putting together the disease associations and evidence to qualify the pathogenicity as derived from integrative analysis of datasets and published literature to aid clinicians and researchers in the clinical interpretation of variants.

Our MD studies revealed no large changes in the local structures of the protein. However, it may be possible that even the small structural distortions that we observed in our studies may affect the stability of the global structure. Also changes in the microenvironment of the mutations that we could not detect could play an important role in protein destabilization. We also admit that longer-timescale simulation may reveal more significant structural changes which could lead to the destabilization of the protein. We agree that our MD simulation studies have limitations in terms of time duration. Due to the incomplete structure of ATP7A, the predictions may not be fully indicative of the global stability of ATP7A. Additionally, the lack of experimental support to corroborate the simulation results (or vice versa) means further studies are necessary.

4.1. Genotype to phenotype correlation

Molecular diagnosis of MD can play a vital role in better prognosis and thus resulting in better disease management. In order to overcome the problem of clinical heterogeneity in MD patients, the defects should be properly characterized. Mutations like nonsense, frameshift-insertion, deletions which are truncating can cause complete disruption of the protein function, such mutations can lead to severe clinical phenotype. A young patient harboring ATP7A Q1168X mutation showed severe manifestation of MD [83]. Patients showing classical MD phenotype have shown to harbor ATP7A R201X, Q303X, R980X, R986X but exceptionally K1408X was found in a patient with milder phenotype of MD. ATP7A N1304S variant was observed in patient with mild form of MD i.e. OHS. It is observed that MD with milder form are characterized by mutations showing some residual activity while mutations with no residual activity manifested classical form of MD [41].

To the best of our knowledge, this is the most comprehensive collection of genetic variants in ATP7A gene and central point of information for the genetic epidemiology, carrier screening and functional genomics in MD and OHS.

Author contribution

VS and BK conceived and designed the study. AM, and MK, collected the data and ACMG classification. DKV and KYJZ performed molecular dynamics simulations. AM, MP and MK designed the figures, KP contributed to the search engine. All authors reviewed and approved the final manuscript as submitted and agree to be accountable for all aspects of the work.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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