Whole Exome Sequencing for Mutation Screening in Hemophagocytic Lymphohistiocytosis

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Abstract

Background: Hemophagocytic lymphohistiocytosis (HLH) is an immune system disorder characterized by uncontrolled hyper-inflammation owing to hypercytokinemia from the activated but ineffective cytotoxic cells. Establishing a correct diagnosis for HLH patients due to the similarity of this disease with other conditions like malignant lymphoma and leukemia and similarity among its two forms is difficult and not always a successful procedure. Besides, the molecular characterization of HLH due to the locus and allelic heterogeneity is a challenging issue.

Materials and Methods: In this experimental study, whole exome sequencing (WES) was used for mutation detection in a four-member Iranian family with children suffering from signs and symptoms of HLH disease. Data analysis was performed by using a multi-step in-house WES approach on Linux OS.

Result: In this study, a homozygous nucleotide substitution mutation (c.551G>A:p.W184*) was detected in exon number six of the *UNC13D* gene. W184* drives to a premature stop codon, so produce a truncated protein. This mutation inherited from parents to a four-month female infant with an autosomal recessive pattern. Parents were carrying out the heterozygous form of W184* without any symptoms. The patient showed clinical signs such as fever, diarrhea, hepatosplenomegaly, high level of ferritin, and a positive family history of HLH disease. W184* has a damaging effect on cytotoxic T lymphocytes, and natural killer cells. These two types of immune system cells without a healthy product of the *UNC13D* gene will be unable to discharge toxic granules into the synaptic space, so the inflammation in the immune response does not disappear.

Conclusion: According to this study, WES can be a reliable, fast, and cost-effective approach for the molecular characterization of HLH patients. Plus, WES specific data analysis platform introduced by this study potentially offers a high-speed analysis step. This cost-free platform doesn't require online data submission.

Keywords: Hemophagocytic Lymphohistiocytosis, Sequencing, UNC13D

Introduction

Progress in controlling communicable diseases increases the relative importance ofnon-infectious diseases, including genetic disorders Hemophagocytic lymphohistiocytosis (HL H) is a type of immunodeficiency syndromes characterized by uncontrolled hyper-inflammation due to the high secretion of cytokines from the

activated but ineffective cytotoxic T lymphocytes (CTLs) and natural killer (NK) cells (2). Clinical symptoms and laboratory findings associated with HLH include prolonged fever, hepatosplenomegaly, cytopenia, hemophagocytosis, decreasing level of fibrinogen, and increasing level of ferritin and triglycerides (3). The etiology of HLH is now considered as a genetic defect and

cause failures in the immune response agents. So against infectious far, distinct genetic defects in responsible genes for the killing ability of CTLs and NK cells have been identified, which are divided into familial HLH (FHL) and syndromes associated with **HLH** (4). FHL is a rare autosomal recessive disease with an incidence rate of 1.2 births per 100,000, which is more prevalent in children's infancy period (5). Four out of five types of FHL (FHL 2-5) have been determined as a responsible associated FHL. FHL-1 has with no specific gene candidate except for a cytogenetic location (9q21.3-

22). Perforin deficiency which described by Stepp et al. as the first genetic defect involving in familial HLH, later classified as type 2 of FHL (FHL-2) (6). Table I shows that deficiencies in some other are responsible for syndromes genes associated with HLH, including GS2, HPS2. Except CHS, and for the all other perforin gene, genes are accountable for vesicle priming, fusion, or trafficking of cytotoxic docking, granules in CTLs and NK cells. Other genetic diseases that predispose patients to HLH, mainly induced by Epstein bar virus (EBV) infection, including X-linked proliferative syndromes 1 and 2, CD27 deficiency, Interleukin-2-Inducible T-Cell Kinase (ITK) deficiency, and XMEN 7). Infectious syndrome (4,agents. including EBV, protozoa, bacteria, and induce HLH symptoms. fungi. can However, identifying these agents in HLH patients does rule not genetic HLH out (8).

Clinical signs of genetic and acquired form of HLH are indistinguishable. In other words, because of phenotype similarity between two types and another condition like malignant lymphoma and leukemia (7), the diagnosis of HLH is not always an easy-going procedure (9). Therefore, all of the patients with HLH symptoms are receiving a recommendation for performing a genetic test (10). However,

allelic and locus heterogeneity doubled difficulty. Generally, diagnosis application of conventional direct DNA sequencing methods such as Sanger sequencing (11)for molecular characterization of affected patients, which involves many genes, is a time-consuming and expensive method. Hence, many patients may remain undiagnosed or died due to the delay in initiating a suitable especially developing treatment, in countries. The mean life expectancy for patients who are not receiving proper therapy is about two months, whereas, more than 55% of patients have a positive outcome by suitable treatment (10). Nextgeneration sequencing (NGS) technologies, especially whole exome sequencing (WES), provide a reliable, fast, and cost-effective approach for clinicians and researchers through the re-sequencing of all coding regions (exome) in one test instead of the gene by gene approach (12, 13). For example, in a study by Mukda et al., evaluation of 25 HLH samples led to the identifying disease-causing mutations in genes, including PFR1, UNC13D, STXBP2, LYST, and XIAP (14). In another study, WES has implemented the identification 16 HLH samples to of harboring mutations in *PFR1* and *LYST* genes (15). This study use WES to underlie aimed to mutational landscape of a patient from a four-member family manifesting clinical symptoms of HLH.

Materials and Methods Patient information

A four-member Iranian family with two children, a four years old boy and a four-month infant girl, participated in this study. Parents were the first cousins. The infant, due to some complications, was hospitalized in the Children Medical Center, Pediatrics Center of Excellence, Tehran, Iran. Specialized clinicians reported clinical symptoms such as fever, diarrhea, and hepatosplenomegaly. Laboratory investigation had revealed

anemia (hemoglobin 5.9 mg/l) and cytopenia (White blood cell (WBC)= $2.12 \ 10^3/\mu l$, Platelet (PLT)=25000Lymphocyte (Lymph)=71.3%, Neutrophil (Neut)=15.7%, Monocyte (Mono)=8.5%, and Eosinophil (Eos)=4.4%). investigation The infectious diseases throughout the culture of blood, stool, urine, and bone marrow aspiration was negative. The result of prothrombin time (PT), partial thromboplastin time (PTT), and international normalized ratio (INR) 50 was 17 secs. secs. and 1.6. respectively. Besides, by first blood evaluation, the C reactive protein (CRP) and erythrocyte sedimentation rate (ESR) levels were 67.2 mg/l and 5 mm/h, respectively. Alanine aminotransferase (ALT) and aspartate aminotransferase (AST) were 205 and 124 respectively. Serology for viral infection, including human immunodeficiency virus (HIV), EBV, and viral hepatitis (A, B, C), was negative. Serum immunoglobulin G (IgG) against Cytomegalovirus (CMV) was positive; whereas, immunoglobulin M (IgM) was negative, which means there is no recent viral infection in the patient. Based on the explanation above, probably IgG antibodies of the mother, which transferred into the child placental, caused this incident. Further lab findings showed a decreased level of fibrinogen (58 mg/dl), increased level of ferritin (6200 ng/ml), and an increased level of triglyceride (404 mg/dl). Analysis of bone marrow by flow cytometry showed a lymphocyte population of 84% of total cells that was normal for T and B cells (CD 3= 58%, CD 4= 31%, CD 5= 20%, CD 7= 22%, CD 8= 28%, CD 19= 2%, CD 20=5%, and CD 22= 2%). The expert clinicians confirmed the HLH diagnosis based on existing symptoms and diagnostic criteria (HLH-2004) (16). Additionally, this family lost their previous infant with similar symptoms that was suspected HLH disease. After initial treatment, WBC

and ESR increased to 6700 and 65, respectively, and serum ferritin, ALT, and AST decreased gradually to 450, 76, and 48, respectively.

Ethical Consideration

This study was performed under the ethical permission code of No: IR.IUMS.REC.1394.27436 from Ethical Committees of Iran University of Medical Sciences and earlier, informed consent signed by infant's parent.

DNA extraction and quality control

Peripheral blood collected into a 5 ml EDTA coated vacuum, and then DNA extraction from leukocytes performed with QIAamp DNA Blood Mini kit (Qiagen, Germany, Cat. No. 51104) according to the manufacturer's instructions. The purity of DNA was assessed by spectrophotometry using NanoDrop 2000 (Thermo Fisher Scientific, Waltham, MA, USA). The integrity of DNA was evaluated by agarose gel electrophoresis.

Whole exome sequencing

Sequencing was performed with Illumina HiSeq 2000/2500 sequencer (Macrogen, Seoul, South Korea) by subjecting two µg high-quality gDNA to Agilent SureSelect Target Enrichment Kit with an average coverage of 100-fold and desired readlength of 101 bp. For increasing sensitivity and reliability detection of duplicate and possible fusions regions, paired-end sequencing employed. was After sequencing, raw data were converted into the Fastq file format by running the (support.illumina.com/downloads/bcl2fast q-conversion-software-v2-20.html).

Bioinformatics analysis

All steps for NGS data analysis was performed on Linux OS. Accordingly, quality control was performed on Fastq files by using the **FastQC** tool (www.bioinformatics.babraham.ac.uk), then further preparation, such as trimming issues, was performed by the Fastx-toolkit (hannonlab.cshl.edu/fastx_toolkit/). The alignment of reads against the human reference genome (GRch38) was

processed by BWA-MEM (17) and result SAM (Sequence stored in the Alignment/Map) file format. Some postalignment steps, such as converting SAM file to BAM (Binary Alignment/Map), deduplicating, and sorting the BAM file, were processed by the Picard tools (broadinstitute.github.io/picard/). coverage percentage was performed by Bed tools 2.26) (bedtools.readthedocs.io/en/latest/) and integrative genomics viewer (IGV) (v 3) (18), the latter has the capability of direct bam file visualization. GATK pipeline (19, 20) was settled for the base quality score SNP (Single nucleotide recalibration, polymorphism), and INDEL (Insertion and Deletion) calling, INDEL, **SNP** and realignment, and variants score recalibration by using Genome Analysis Toolkit (GATK v 3.7) (21). Raw variants were stored in a text-based file format, namely VCF (variant call format). A biological knowledge-based platform from Kggseq (v 1) tool was used to identify genetic loci responsible for human diseases/traits. According to this multi-step platform, quality assessment, filtration, annotation, and pathogenicity prediction were applied to the variants. For example, allele frequency-based filtering was performed against the 1000 genome project, dbsnp 141, Exome Aggregation (ExAC Consortium r 0.3.1). and Exome ESP6500AA (NHLBI GO Sequencing Project). Then, the Mouse phenotype (22),Deciphering Developmental Disorders (DDD) study (23), OMIM and PubMed database were used for annotation of the remained variants. In addition, dbSNFP (24) was used for functional prediction and all potential annotation of single-nucleotide synonymous variants (nsSNVs) by compiling prediction scores from 26 prediction algorithms such as SIFT (25), PolyPhen-2 (26), and CADD. Additionally, the DAVID database was used for further comprehensive functional annotation (27). Pathway and proteinprotein interaction analysis were performed by using the KEGG pathway database (28) and STRING database (https://string-db.org), respectively. Plus, Interpro Scan was used for detecting possible domains in the proteins (29).

Sanger sequencing validation

All variants were identified by the WES confirmed using approach Sanger sequencing, which was performed on corresponding regions. PCR primers were designed by Primer (http://bioinfo.ut.ee/primer3-0.4.0/) checked by primer-blast webtool (Table II). PCR condition was done as follows: 95°C/4min, followed by 30 cycles with 95°C/30 s, 66°C/30 s, and 72°C/45 s. Finally, the result of sequencing was analyzed using the Chromas software v 2.6 (http://technelysium.com.au).

The limitation of this study includes bioinformatics required sophisticated and systems extensive data storage capabilities, which can be costly. Moreover, the amount of produced data, heterozygosity, high repeat content, short DNA read lengths, and the high error rate can impact the outcome.

Results

four-member Iranian family with children suffering signs from and symptoms of HLH disease was investigated in this study for genetics diagnosis by WES, which is crucial for them. The family pedigree was made after genetic consulting (Figure 1). performing exome sequencing, quality assessment of raw Fastq files showed that total reads were 66,582,362, with GC constancy of 51%. Besides, more than 99% of reads were above 20 Phred quality (>O2O). Moreover. score alignment statistics showed that 79% of all reads mapped to the human reference genome (GRch38) with a mean depth of 70 at the regions. target The total number of SNPs and INDELs fond in discovery section was 52456. Prioritization step resulted in 110 potentials diseasecausing variants (Figure 2). Among these damaging variants and according to the clinical symptoms, known genes, locus responsible for HLH disease, homozygous nucleotide substitution in the exon six of *UNC13* homolog D gene (munc13-4 protein) was found that caused a premature stop codon at Tryptophan 184 position (NM 199242: c.551G>A:p.W184*). The minor allele frequency (MAF) for this variant based on the ExAC database was 0.000008. The CADD score was 30, and the LRT prediction score was a deleterious effect. The corresponding region was amplified set of primers specific PCR conditions. Sanger sequencing result showed that candidate from WES analysis homozygous substitution for the patient, and her parent was carrying heterozygous form (Figure 3). According to the DAVID server and KEGG database, the gene products of the UNC13 family (Pathway ID: hsa04721) the transmembrane protein that functions as the regulators of docking, priming, and fusion of vesicles to the plasma membrane. In the immunologic synapse, munc13-4 motives secretion of lytic granules, and the occurrence of FHL-3 is due to the defect in this protein. The result of the InterPro Scan showed that this protein had conserved Munc13 homology (MHD 1 & 2) flanked by two calciumbinding domains (C2) (Figure Additionally, protein-protein interaction revealed analysis that the UNC13D participated in product the leucocyte mediated toxicity process by interacting STXBP2. STX11. and (Figure 4B). The finding of this study highlighted a particular platform in whole exome sequencing experiments speculated on both clinical and research applications of these sequencing platforms for rare complex disorders in the future. study indicated **WES** This that experiments were able to identify some rare variants of diseases, including HLH, which consider a life-threating illness.

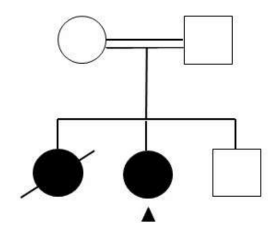


Figure 1. Family Pedigree. Squares and circles indicate males and females, respectively; open symbols indicate unaffected individuals; whereas, filled symbols indicate affected individuals; Proband (triangle) has lost her sister due to the HLH.

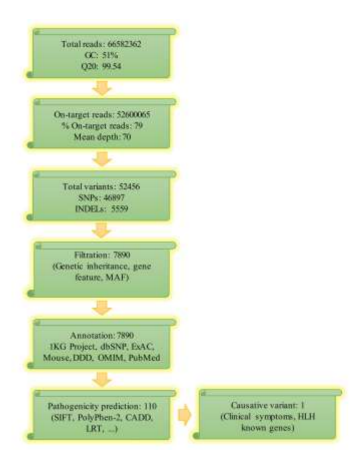


Figure 2. WES data analysis statistics from raw data to the causative variant

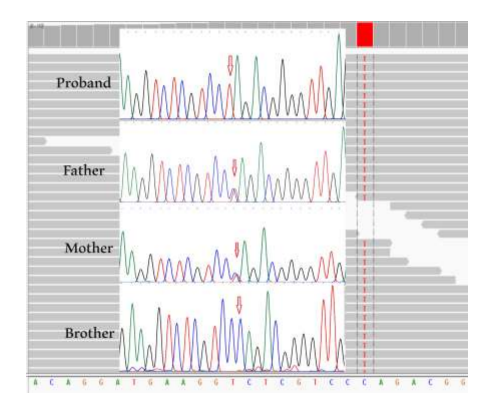


Figure 3. Mutational analysis (WES and Sanger sequencing) of the *UNC13D* gene in the proband and her family. This illustration shows an IGV screenshot of a homozygous substitution (c.551C>T) in the proband and further Sanger sequencing confirmation for the patient (arrow) and her parents (as heterozygous); whereas, her brother is in a healthy state (chromatogram is a result of the reverse strand).

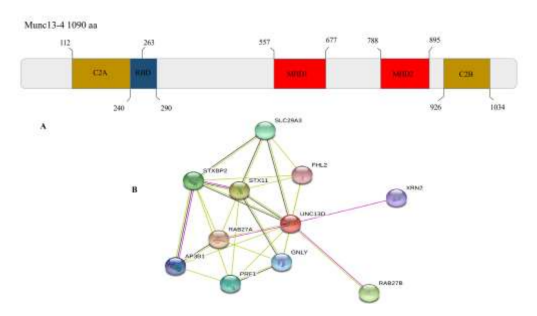


Figure 4. Schematic illustration of munc13-4 domains and specified residue number (A, Based on InterproScan result). The unc13d network revealed that this protein had interaction with ten other gene products and participated in the leucocyte mediated toxicity process (B, Based on string database).

Table I. The responsible genes for HLH and syndromes associated HLH.

Syndromes associated HLH				
n				
-q42.2				
EBV associated syndromes				
31				

Table II. Forward (F) and reverse (R) PCR primers designed by Primer 3.

Primer	Tm (°C)	Length	Product size (bp)
F 5'-TGACCCTCCTTGCCCCATCAG-3'	63	21	431
R 5'- TGCCTCGTCTTCCCCACTCCT-3'	64	21	

Discussion

The molecular characterization of HLH patients owing to the heterogenic nature of HLH is a challenging issue. Patients also need an early and correct molecular diagnosis. In this study, to understand the possible genetic damage related to the patient, whole exome sequencing on the Illumina HiSeq2500 machine was performed using Agilent SureSelect Human All Exon kit. The familial HLH type 3 (FHL-3, OMIM # 608898) was confirmed in the patient according to the clinical manifestation, pedigree, WES, and Sanger sequencing result. FHL-3 is a genetic defect with autosomal recessive inheritance and responsible for 10%-32% of genetic HLH cases. FHL-3 considered the second most common familial HLH subtype (the first one is FHL-2). This lifethreating disease requires an early and correct diagnosis to prevent compliance and mortality (9, 30). Genetic defects in

the *UNC13D* gene (OMIM# 608897) cause the FHL-3. This gene is located on chromosome 17, comprises of 32 exons, which translated to a 120 KD protein, namely munc-13 homolog D or munc13-4 (UniProtID: Q70J99), and functions in the vesicle maturation, docking, priming, and fusion during secretary process (30). It is demonstrated that MHDs region (MHD1 and MHD2) are present in a variety of proteins from human, C.elegans, Drosophila melanogaster, mouse, rat, some which function membrane's as trafficking regulators. Munc13A munc13C are brain-specific, while other family members express ubiquitously (31). Human munc13-4 has an expression at a high level in thymus, spleen, peripheral leukocytes, and mast cells (30, 32). According to a model proposed by Willenbring and Johnson (2017), this protein is involved in the regulation of the cytolytic granules secretion

cytotoxic cells, such as CTLs and NK ones. In other words, munc13-4 plays a vital role in intracellular trafficking. In the immunologic synapses, exocytosis perforin and granzyme is a responsibility of munc13-4. This event trigger apoptosis in the target cells (33). The hypothetical mechanism molecular behind munc13-4 works introduced by Elstak et al. in 2011. They explained that secretory event induces by triggering effector cells (such as T lymphocytes) and followed by the formation of a complex consist of small GTPase rab27 and munc13d in the phagocytic cells. Co-localization of these two proteins recruit other effectors and facilitates the tethering of secretory lysosome to the plasma membrane. Tethering can occur when munc13 binds to syntaxin11 or $Doc2\alpha$, which causes interaction with SNAP-23. Then, the secretory lysosome can fuse to the plasma membrane (34). Genetic defects in the munc13-4 and other effectors engaged in exocytosis machinery dramatically eliminate the killing efficacy of cytotoxic cells, so it provides an environment for infections, such insurgent as influenza virus, EBV, Cytomegalovirus (CMV), and Streptococcus pneumonia (35, 36). Human Gene Mutation Database (HGMD) contains 113 mutations related to the *UNC13D* gene (www.hgmd.cf.ac.uk), and it confirmed that mutations in this gene lead to FHL-3 (30). In addition, some studies have described an association between variants of the UNC13D gene and idiopathic systemic juvenile arthritis (sJIA) (37, 38). The *UNC13D* gene belongs to the patient has a premature stop codon (W184*) producing a truncated munc13-4, which is incompatible with functional protein and finally subjected to cell degradation machinery. Therefore, in the immunologic synapse, stimulated, but ineffective cytotoxic cells secrets high amount cytokines, including Interleukin-1 (IL-1), IL-6, and Tumor Necrosis Factor α (TNFα). These released cytokines cause prolonged fever. Furthermore, the TNF α ,

along with Interferon γ (INF γ), prevents the proliferation of blood cells and the of triglycerides. hydrolysis Activated macrophages cause an increased level of ferritin, whereas the level of fibrinogen decreased by them. Finally, the clinical picture of familial HLH disease present in the patient (9). To reduce the mortality rate associated with HLH treatment should initiate the elimination of possible infectious agents and activated CTLs. administrate drugs such Then Dexamethasone, Etoposide, Cyclosporine A. In familial forms, the restoration of the immune system requires Hematopoietic stem cell (HSCs) transplantation (3, 39). Reports on the implementation of NGS especially technologies, WES. detecting the harmful mutation in the HLH patients showed the accuracy effectiveness of these methods. Ferarri et al. (2017) revealed causative mutations in the UNC13D, STX11, and STXBP2 genes in 11 out of 12 patients by using WES (40). Moreover, Qian et al., (2014), by evaluating 1709 HLH samples, reported a 253 Kilo base pair inversion and deep intronic mutations in the UNC13D gene. They also discovered the W184* mutation harbored in the two Caucasian patients (41).

Conclusion

In conclusion, one of the more significant findings of this study was that the use of WES for determining the molecular diagnosis of a complex genetic disease like HLH could be a fast, reliable, and costeffective approach for clinicians and families. Additionally, designed a local multi-step, WES specific data analysis platform, and cost-free introduced by this study potentially offer a high-speed analysis step. No need for online data submission. Users capable are replicating analysis in all stages.

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Conflict of interest

All authors contributed equally to this work and declare that there is no conflict of interest regarding the publication of this article.

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