Novel mutations of epidermolysis bullosa identified using whole-exome sequencing in Indonesian Javanese patients

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SUMMARY

Epidermolysis bullosa (EB) is a group of inherited blistering skin diseases known to have heterogeneity of phenotypes and genotypes. There are four main types of EB: simplex, junctional, dystrophic, and Kindler syndrome, which are further classified into 34 distinct subtypes. Twenty different gene mutations are responsible for the loss of function and integrity of the basal membrane zone. In limited-resource settings such as Indonesia, diagnoses of hereditary skin disease often rely on clinical features. This limitation was managed by using the Clinical Diagnostic Matrix EB for clinical diagnosis support and whole-exome sequencing for genetic analysis. This study is the first whole-exome sequencing analysis of Javanese Indonesian patients with EB. The genetic analysis from four patients with EB identified all novel mutations unreported in the dbSNP database. There are Kindler syndrome with \textit{FERMT1} frameshift mutation in exon 4, at c.388A (p.I130fs), which causes truncated protein; junctional EB generalized intermediate (JEB-GI) subtype with missense mutation at \textit{LAMB3} gene position c.A962C (p.H321P); and recessive dystrophic EB (RDEB) a missense mutation at \textit{COL7A1} gene position c.G5000T (p.G1667V). The whole-exome sequencing was further verified by Sanger sequencing. The new mutations' finding is possibly due to the limited genetic database in the Malayo-Polynesian ethnic group. Indonesia has hundreds of ethnic groups, and the Javanese is the largest ethnic group that populates Indonesia. Genetic data of these ethnic groups is important to be established in the international genetic database. This combination of clinical diagnostic and genetic analysis tools with whole-exome sequencing confirmed the challenging diagnosis of epidermolysis bullosa.

Keywords

clinical diagnostic matrix, Kindler syndrome, junctional epidermolysis bullosa, dystrophic epidermolysis bullosa, Malayo-Polynesian ethnic group

1. Introduction

Epidermolysis bullosa (EB) is an inherited skin disorder characterized by flaky skin and blisters after light trauma or friction. The wounds tend to occur repeatedly and may become chronic, which causes high morbidity. This disease's clinical manifestations vary, ranging from local wounds on the hands and feet to generalized blisters or erosion and failure to thrive (1).

Epidermolysis bullosa comprises four major subtypes: EB simplex, junctional EB (JEB), dystrophic EB (DEB), and Kindler syndrome (KS). Each subtype is classified based on the molecular structure mutation in the dermis-epidermis (2). Before high-throughput next-generation sequencing (NGS) was developed, diagnosis of EB required an invasive procedure such as skin biopsy followed by microscopic examination and/or direct immunofluorescence to determine the mutation locations and finally, the results were confirmed by Sanger sequencing (SS) as the gold standard of diagnosis (3).

Caused by mutations in 20 different genes, EB involves the loss of function and integrity of the basal membrane zone structure. These genes code structured proteins in the intermediate filaments (keratin 5 and 14), focal adhesions, and desmosome complexes (desmoplakin, plakophilin, and plakoglobin). They are also vital to form intraepidermal adhesions and the
dermo-epidermal anchoring complex in the basement membrane zone of the mucous membrane and skin (integrin 64, collagen type XVII, laminin 332, collagen type VII, integrin three subunits, and kindlin-1) (4). Unfortunately, SS can only test single or a few genes at a time; thus, it is time-consuming and costly to do SS in cases when multiple genes need to be tested. Recently, direct procedures with NGS or targeted NGS have been performed to overcome the previous procedure's limitations. NGS screens all the suspected genes involved in EB using a single sample (3).

The NGS technology can simultaneously sequence the peptides/proteins coded by a known gene in the genome called whole-exome sequencing (WES). Even though WES can only sequence 1.5% of the entire human genome, 85% of the pathogenic mutations occur in regions encoded by protein, where WES is preferable. In recent years, the decreased cost of WES has developed a potential preference to use WES as a clinical practice (5).

In a limited-resource setting such as Indonesia, EB and EB subtypes’ diagnosis depends on clinical findings and hematoxylin-eosin staining biopsy results. Electron microscopic and immunofluorescence examinations are not routinely done because they are rarely available in many centers. Thus, this condition implies a holistic approach to care for patients with EB. WES sequencing for analyzing all genes simultaneously to find the pathogenic variants will have a tremendous impact on both the novel search for disease gene discoveries and the efficient diagnosis of known genetic diseases in Indonesia.

Novel mutations or novel variants can be identified when they are not present in unrelated healthy control samples that were obtained from the exome sequence databases Exome Aggregation Consortium (ExAC) browser (https://gnomad.broadinstitute.org) and other databases such as dbSNP, 1000 Genomes Project (http://www.1000genomes.org), and Genome-Wide Association Studies (GWAS).

Javanese comprises the most prominent traits in Indonesian people who populate Java Island, and genetic testing with WES for the inherited disorder to the researchers’ knowledge has never been done before in Javanese Indonesians. The prevalence of EB in Indonesia remains unknown. Based on case reports between 2013-2020, there were nine cases in Javanese traits, which is less than expected. Diana et al. reported around 31 EB cases in Indonesia in 2018 based on the Epidermolysis Bullosa Community Indonesia database (6). Even though genetic variants have been reported extensively in other populations, a study in the Javanese Indonesian population is lacking.

This study explores the use of WES to diagnose EB cases in Javanese Indonesians. This study highlights the first-tier use of WES to analyze genetic variation in Javanese EB cases. It will hopefully become a standard diagnostic tool for EB and other genetic disorders in Indonesia.

2. Materials and Methods

The clinical subtype diagnosis of EB was performed using a clinical matrix diagnostic tool (7), consisting of nine of the most often identified subtypes of EB. From these, the clinicians chose the most appropriate clinical findings that matched the available matrix.

DNA extraction from patients, parents, and healthy controls were obtained after informed consent for genetic analyses and medical information disclosure. Ethical clearance was completed following the Declaration of Helsinki with approval from the Universitas Gadjah Mada Medical and Health Research Ethics Committee. Four patients with EB, five unaffected parents, and ten healthy controls were included in this study.

Genomic DNA was purified using the AllPrep DNA/RNA Mini Kit (QIAGEN, Hilden, Germany). The concentration and quality of genomic DNA (gDNA) extracted from peripheral blood were examined with NanoDrop ND-100 (Thermo Scientific) and agarose gel electrophoresis.

Whole-exome sequencing was performed at Novogene (Hongkong, China). DNA was captured using the Agilent SureSelect Human All Exon kit and sequenced on an Illumina HiSeq platform. The sequence was aligned to the human genome reference sequence (hg38) using the Burrows-Wheeler Aligner (v0.7.8-r455). Duplicate reads were removed using Picard (v1.111), single nucleotide polymorphisms (SNPs) and Indels (insertions and deletions) were identified (SAMtools v1.0). The detected variants were then annotated using ANNOVAR (8). We defined database annotation variants with reported allele frequencies of 0.5% or greater in one of the following databases: dbSNP, 1000 Genomes Project (http://www.1000genomes.org), Genome-Wide Association Studies (GWAS), and Exome Aggregation Consortium (ExAC). We focused on variants of known causal genes in EB diseases, including KRT5, KRT14, DST, EXPH5, DSP, PLEC, LAM3A3, LAMB3, ITGA3, LAMC3, ITGB4, ITGA6, JUP, COL7A1, COL17A1, EXPH5, FERMT1, PKP1, and TGM5. From those variants, we filtered deleterious predicted variants according to the following criteria: frameshift, in-frame insertion and deletions, stop codon changes, missense unless predicted innocuous by SIFT, polyphen-2, Mutation Taster, likelihood ratio test (LRT), Mutation Assessor, and Functional Analysis Through Hidden Markov Models (FATHMM).

The SS was performed to confirm the pathogenic variants in the patient, parents, and the control group using an ABI PRISM® BigDye Terminator Cycle Sequencing Kit v3.1 (Applied Biosystems; Thermo Fisher Scientific, Inc.). The primer sequences of each of the suspected pathogenic variant genes
FERMT1 and LAMB3 were chosen based on previous studies (9,10), the COL7A1 exon 53-54 were modified from Cristianto et al. in 1997 with forward 5’CCTTGAGAACTGCTTGCTTC3’ and reverse 5’GAAGATGGAGGTTCAGC3’ (11). The sequence reads using 4peaks application (https://nucleobytes.com/4peaks).

To identify the novelty of EB variants in Indonesia, the literature search was conducted using PubMed and Google Scholar with the term "epidermolysis bullosa" "southeast Asia" "Indonesia" "Javanese" and the full texts of retrieved literature were read carefully. Personal communication with clinicians was also established to find EB cases in Central Java and Yogyakarta. Novel variants are considered when the variant annotation is not found in the databases: dbSNP, 1000 Genomes Project (http://www.1000genomes.org), Genome-Wide Association Studies (GWAS), and the Exome Aggregation Consortium (ExAC).

3. Results and Discussion

Four patients with clinically confirmed EB were included in this study. The clinical diagnosis was made with support from clinical matrix diagnostic tools, and all patients underwent genetic analysis. Two patients are not related, but Cases 2 and 3 were siblings. Two patients are male, and two are female with an age range of 10 to 22 years. Three from 4 subjects were missense mutations; one subject was a frameshift mutation. The subtypes of EB were varied from Kindler syndrome, JEB-GI, and RDEB-GS. All patients are genetically of Javanese ethnicity.

Each of the pathogenic EB variants was found to be deleterious by its rare or low frequency. We documented novel mutations in all subjects with WES to be deleterious by its rare or low frequency. We

<table>
<thead>
<tr>
<th>Characteristic variant</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Annotation</strong></td>
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<td>SNP</td>
<td>SNP</td>
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<tr>
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<td>NM_000094</td>
<td>COL7A1</td>
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<td>NM_001017402</td>
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<td></td>
<td>exon 4</td>
<td>exon 9</td>
<td>exon 9</td>
<td>exon 54</td>
</tr>
<tr>
<td><strong>cDNA</strong></td>
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<td>c.A962C</td>
<td>c.A962C</td>
<td>c.G5000T</td>
</tr>
<tr>
<td><strong>Known variant</strong></td>
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<td>Novel'</td>
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<tr>
<td><strong>SIFT</strong></td>
<td>Deleterious</td>
<td>Deleterious</td>
<td>Deleterious</td>
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<td></td>
<td>(0.002)</td>
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<td>(0.002)</td>
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<tr>
<td><strong>Polyphen2</strong></td>
<td>Probably damaging (0.999)</td>
<td>Probably damaging (0.999)</td>
<td>Probably damaging (1.0)</td>
<td></td>
</tr>
<tr>
<td><strong>Mutation taster</strong></td>
<td>-</td>
<td>Disease-causing</td>
<td>Disease-causing</td>
<td>Disease-causing</td>
</tr>
<tr>
<td><strong>LRT</strong></td>
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<td>Deleterious</td>
<td>Deleterious</td>
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<tr>
<td></td>
<td>-</td>
<td>Tolerated (-0.2)</td>
<td>Tolerated (-0.2)</td>
<td>Tolerated (-4.62)</td>
</tr>
<tr>
<td><strong>FATHMM</strong></td>
<td>-</td>
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</tbody>
</table>

Variant not found in dbSNP, 1000 Genomes Project, GWAS, ExAC, nor ESP database.

Clinical WES is a valuable diagnostic tool to pinpoint rare genetic alterations in an unbiased and efficient way to confirm a diagnosis despite extensive testing and workup. The use of WES as a first-tier test would substantially reduce the time to diagnosis, and it is only a half to one-quarter the cost of traditional testing (12). WES has a consistently higher diagnostic yield than routine clinical SS. However, the Sanger method is still considered the "gold standard" for clinical DNA sequencing. Nevertheless, SS has limitations because it can only test single or a few genes at a time and becomes costly when multiple genes are tested before reaching a diagnosis (13). Hamilton et al. in 2016 demonstrated that WES has high concordance with SS with 97.3% of variants concordant (13).

However, WES has some limitations in poor targeting by the exome capture kit, high guanine-cytosine (GC) content, and the presence of repetitive sequences, leading to inadequate coverage or sequencing inaccuracy. The exome capture kit used to define allele frequencies in our study was using the 1000 Genomes Project, GWAS, ExAC, and dbSNP. Unfortunately, none of the commercial arrays for SNPs databases were created from polymorphism data generated in Southeast Asian Malays. In the future, it is essential to develop population-specific databases as a country-based molecular variation database since different genetic backgrounds may have different susceptibility to the haploid in the sufficiency of variants or mutations. SNPs may contribute to the genetic disorder, or they may be in linkage disequilibrium with other causal variants and mutations (14).

Indonesia, located in Southeast Asia, comprises more than 17,000 islands, including the five biggest islands, which are Sumatra, Java, Borneo (Kalimantan),...
Sulawesi, and Papua. Indonesia is the world’s 4th most populous country with over 267 million people, and Java is the most populous island. Indonesia consists of hundreds of distinct native ethnic and linguistic groups, with the largest being Javanese (https://en.wikipedia.org/wiki/Indonesia). Our study is located in Central Java and Yogyakarta, which is mostly populated by people with Javanese ancestry.

Case 1, a 12-year-old boy with clinical manifestation of skin brittle, poikilodermia, meatal stenosis, aphthous ulcer, and history of hernia inguinal (Figure 1). On genetic analysis, we identified a homozygous mutation in the location of 389 FERMT1, an insertion of A-base, that converts an isoleucine codon (AAT) at codon 130 to asparagine codon (AAT) and leads to a premature termination codon (PTC) at codon 132 (p.I130fs). The parents were heterozygous for the mutation and were considered obligate carriers. The location of the mutation is at the F1 domain, which is considered highly conserved. We confirmed the diagnosis of Kindler syndrome. The mutation of c.388dupA p.I130fs is the first reported in the literature. The duplication of FERMT1 in exon 4 has been reported previously (15), but the correlation between duplication and severity of the clinical findings is still unclear. The mutation in our study is in the F1 segment of FERM, which is predicted to cause loss of the FERM domains of the protein, thus resulting in the detached adhesin bond and membrane matrix (16). Furthermore, the isoleucine protein in the F1 domain is a highly conserved domain (17). This mutation appears to contribute to the pathogenesis of several organs, not only manifesting in skin atrophy and poikilodermia but also including dysfunction of the oral, intestinal, and urinary systems.

Cases 2 and 3 are siblings with various clinical manifestations (Figure 2). The chronic insole ulcers...
and finger ulcers were seen in the 10-year-old younger brother, but not in his sister, instead, a chronic ulcer at the philtrum was seen. Both of them have ulcers behind their ear, and unsurprisingly they have the same homozygous mutation in the location of 962 LAMB3 A>C. The histidine codon (CAC) converts to a proline residue (CCC). The mother also has the same homozygous variant, but interestingly she did not have any clinical manifestations. The father has a heterozygous variant in the same allele. The diagnosis confirmed junctional EB intermediate. Mutation of the LAMB3 gene is the hallmark of JEB, the critical encoding component of the hemidesmosome-anchoring filament complex that links the keratin cytoskeleton to the lamina densa of the dermo-epidermal junction (18).

In the majority of the JEB cases, 80% of mutations predominantly affect the LAMB3 gene. However, the clinical spectrum is highly variable depending on the mutation's type and position (19). We identified siblings with the same mutation of LAMB3 at the same loci c.A962C (p.H321P), but they demonstrated different phenotypes. Other polymorphisms in each person may correlate with various clinical manifestations. However, this possible explanation needs to be addressed in future studies.

In Case 4, the result of WES and SS had high concordance. The WES presented a mutation in the G>T exon 54 COL7A1, and SS reported a heterozygous mutation at codon 5000 in the affected child (Figure 3). However, we did not find the variant in the mother nor wildtype control. Unfortunately, paternal DNA cannot be defined. Thus, we diagnosed the patient with recessive dystrophic epidermolysis bullosa. COL7A1 is responsible for dystrophic EB with dominant and recessive genetic patterns, though genetic analyses of the COL7A1 gene in affected individuals revealed that most mutations detected are recessive DEB (RDEB) (20). The majority of severe RDEB cases are caused by PTC mutations resulting from nonsense, frameshift, or splice-site mutations on both COL7A1 alleles, which result in either nonsense-mediated decay of the mRNA or truncated polypeptides, and the result is unable to assemble the functional protein and the mRNA degraded within the cell. Milder cases of RDEB are often caused by compound heterozygous mutations: one PTC mutation and a missense mutation (21). Our patient had a heterozygous missense mutation of COL7A1 at exon 54 substitution 5000G>T that substitutes glycine for valine. Mallipedi et al. in 2003 pointed out the need to search for further mutations when they found glycine substitution in sporadic cases (22). However, our case did not reveal other mutations on COL7A1.

The heterogeneous mutations in our case caused disease phenotypes and led to moderate-to-severe DEB forms, unlike in other articles, which need to be further examined (23).

Murata et al. in 2003 demonstrated that the presence of recurrent mutations could be classified into specific ethnic groups and worldwide mutations (20). Our four cases revealed novel mutations that were not recorded in any public SNP database (dbSNP). Our novel finding is probably because of the limited number of EB Malay traits in reported articles, resulting in a lack of identified recurrent specific ethnic group mutations in Malay ancestry. Abu Sa'd et al. in 2006 also stated that diagnostic strategies based upon molecular epidemiological features that are determined in Western origin populations might not apply to Middle Eastern populations (24). However, two studies reported two Sundanese Indonesian families with dystrophic EB. They reported the mutations in COL7A1 at c.6218G>A; p.Gly2073Asp, c.5945G>T; p.Gly1982Val and c.6253G>T; p.Gly2085Trp (25,26). We did not find other articles associated with EB genetic examination in Indonesian traits.

As a limitation, the number of subjects was too small to analyze a high diagnostic yield in WES compared to standard genetic testing. The low incidence of cases may hamper subject recruitment.

In conclusion, we demonstrated a novel mutation in Kindler syndrome on c.388A (p.I130fs), junctional epidermolysis bullosa intermediate on c.A962C (p.H321P), and recessive dystrophic epidermolysis bullosa on 5000G>T (p.G1667V) as Javanese traits. In Figure 3. Clinical manifestations of RDEB and electropherogram of COL7A1 mutation. (A) skin erosion and right point finger contracture in Case 4; and (B) heterozygous missense mutation at c.5000G>T, which was not shown in wildtype control nor the mother.
these cases, practical genetic analysis was best performed using whole-exome sequencing, a less invasive genetic diagnostic tool, with a reduced time to diagnosis.

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