Novel and recurrent variants of ATP2C1 identified in patients with Hailey-Hailey disease

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Abstract

Hailey-Hailey disease (HHD) is a rare, late-onset autosomal dominant genodermatosis characterized by blisters, vesicular lesions, crusted erosions, and erythematous scaly plaques predominantly in intertriginous regions. HHD is caused by ATP2C1 mutations. About 180 distinct mutations have been identified so far; however, data of only few cases from Central Europe are available. The aim was to analyze the ATP2C1 gene in a cohort of Polish HHD patients. A group of 18 patients was enrolled in the study based on specific clinical symptoms. Mutations were detected using Sanger or next generation sequencing. In silico analysis was performed by prediction algorithms and dynamic structural modeling. In two cases, mRNA analysis was performed to confirm aberrant splicing. We detected 13 different mutations, including 8 novel, 2 recurrent (p.Gly850Ter and c.325-3 T > G), and 6 sporadic (c.423-1G > T, c.899 + 1G > A, p.Leu539Pro, p.Thr808TyrfsTer16, p.Gln855Arg and a complex allele: c.[1610C > G;1741 + 3A > G]). In silico analysis shows that all novel missense variants are pathogenic or likely pathogenic. We confirmed pathogenic status for two novel variants c.325-3 T > G and c.[1610C > G;1741 + 3A > G] by mRNA analysis. Our results broaden the knowledge about genetic heterogeneity in Central European patients with ATP2C1 mutations and also give further evidence that careful and multifactorial evaluation of variant pathogenicity status is essential.

Keywords Hailey-Hailey disease · ATP2C1 · Genodermatosis

Introduction

Hailey-Hailey disease (HHD, OMIM 16960, or Benign Chronic Pemphigus.) is a rare (incidence 1:50000) autosomal dominant genodermatosis. The symptoms, aggravating periodically, onset in third–fourth decade include blisters, vesicular lesions, crusted erosions, and erythematous scaly plaques, which occur mainly on groins, axillae, neck, and other intertriginous areas, and mucosa may also be involved. Lesions may be odorous and painful and lead to mobility affecting fissures (Li et al. 2016; Zamiri and Munro 2016). In histopathological findings, suprabasalar and intraepidermal keratinocyte acantholysis with a “dilapidated brick wall” appearance is due to abnormal epidermal Ca2+ distribution by secretory pathway Ca(2+) ATPase 1 (hSPCA1) caused by mutation in its gene: calcium-transporting ATPase type 2C member 1 (ATP2C1) (Cheng et al. 2010; Micaroni et al. 2016; Cialfi et al. 2016). Importantly, ATP2C1 is expressed in all tissues, although HHD clinical symptoms are solely isolated to the skin. Four isoforms differing by alternative processing of the C-terminus are produced, but only few of ATP2C1 mutations localized beyond the core of 26 exons are present in each transcript (Nellen et al. 2017). The majority of ATP2C1 mutations lead to a premature termination codon (PTC); thus the dominant inheritance pattern of HHD seems to result from haploinsufficiency. Nevertheless, as around 1/3
of mutations lead to missenses or in-frame rearrangements, other mechanisms may be involved (Dobson-Stone et al. 2002; Kitajima 2002). Thus, to understand the pathophysiological molecular mechanism of HHD, further investigation is required. Worldwide, only about 300 individuals have been described so far with 179 distinct ATP2C1 variants (Nellen et al. 2017). The majority of them are Asians, and only few cases from Central Europe were published, including a not genotyped case report from Poland (Rácz et al. 2005; Sudbrak et al. 2000; Chlebicka et al. 2012).

Herein, we report the results of the first genetic investigation in 18 Polish HHD patients together with characterization of splicing mutations and in silico structural dynamic modeling of novel missense mutations.

**Patients and methods**

Eighteen probands of Polish descent (Table 1) with clinical HHD manifestation (according to Matsuda et al. (2014)) have been enrolled in the study, together with their relatives, if available. The average age at diagnosis was 29 years old (range: 15–40). All patients gave informed consent for participation.

All coding exons of ATP2C1 were analyzed using Sanger sequencing (primers and PCR conditions available on request) or panel next generation sequencing (customized KAPA Library Preparation Kit - Roche) using MiSeq (Illumina). The variants were annotated against NCBI RefSeq: NM_014382.3 and checked for presence in the GnomAD, ClinVar, HGMD Professional and ATP2C1 LOVD v.3.0 databases.

Novel missense mutations were analyzed using in silico algorithms: DANN, MutationTaster, FATHMM, FATHMM-MKL, GERP, MutationAssessor, SIFT, Provean, and PolyPhen2, classified according to ACMG guidelines (Richards et al. 2015) and visualized using dynamic structural modeling (Yasara Structure Package v.15.7.12). Briefly, isoform 1a of hSPCA1 (NP_055197.2) was modeled by homology, using eight closest templates identified in RCSB Protein Data Bank (PDB) records, IDs: 3N5K, 4BEW, 1WPG, 2YN9, 2YFY, 2ZXE, 4RET, and 4HYT. For each template, up to five alternative sequential alignments have been tested. Finally, the best scored models were built on the basis of 1WPG (37.2% of sequence identity and 56.7% of sequence similarity within 802 residues of 919 being aligned) and 3N5K (36.8%, 56.2%, and 810, respectively) PDB records. However, the final hybrid model, which combines the optimal parts of the top models, was scored substantially higher than the latter and thus was further used.

Novel intronic mutations were evaluated with the use of three splice site prediction algorithms: MaxEnt, NNSPLICE, and HSF. In order to confirm the putative cryptic splicing of mutations c.325-3 T > G and c.1741 + 3A > G, RNA was isolated from peripheral blood leukocytes, reverse transcribed, and PCR amplified and analyzed using Sanger sequencing (Fig. 1). As negative and positive controls, we included RNA isolated from a healthy person and from a HHD patient with the already known mutation c.1308 + 1G > T.

**Results**

ATP2C1 variants were detected in 17/18 probands, resulting in a detection frequency of 94%. Overall, we detected 13 different heterozygous ATP2C1 variants, (6 missense or nonsense, 3 splice site, 1 complex allele (missense and intronic in cis), and 3 deletions or duplications). Eight of them (8/13, 61%, Table 1) are novel, i.e., c.2548G > T, c.325-3 T > G, c.423-1G > T, c.899 + 1G > A, c.1616 T > C, c.2408 2420dup, c.2564A > G, and a complex allele: c.[1610C>G;1741 + 3A > G]. Identified mutations localize in the following exons: 26 (2/13), 18 (3/13), and (single variant each) in 7, 8, 12, 21, 23, and 25 and introns 4, 6, and 11. The molecular dynamic modeling or/and in silico prediction analysis (Table 2) together with mRNA analysis of putative splicing mutations (Fig. 1) enabled us to confirm the likely pathogenic status of these novel variants. Precisely, novel c.325-3 T > G, c.1741 + 3A > G, and recurrent c.1308 + 1G > T mutations cause in-frame skipping of exons 5, 18, and 15, respectively, which seems to severely affect the protein structure.

**Discussion**

The majority of mutations (61%) identified in this study have never been reported before, including two recurrent novel splice site c.325-3 T > G (intron 4) and nonsense c.2548G > T (exon 26) mutations, identified in 2/17 (12%) and 4/17 (24%) in different Polish families, respectively. This could suggest specific founder mutations in this ethnic population. All ATP2C1 missense mutations are localized in exons 8, 12, 18, and 26, which is partially in concordance with previous observations clustering in exons 12, 13, 18, 21, and 23 (Micaroni et al. 2016).

Half (4/8) of the novel mutations could easily be classified as pathogenic due to introduction of a premature stop codon (p.Gly850Ter, p.Thr808TyrfsTer16) or change in the conserved consensus sequence of the canonical splice sites (c.423-1G > T, c.899 + 1G > A). The novel missense mutations, p.Gln855Arg, p.Leu539Pro and p.Thr537Arg detected in cis with 1741+3A>G, were analysed using molecular dynamic modeling and standard in silico tools.

Molecular dynamic modeling showed that conversion of Gln855 into Arg would distort the transmembrane helical structure and form a positively charged region located in the...
Table 1  Results of genotyping

| Fam. No. |Chr3(GRCh37):| HGVS ver.15.11 NM_014382.3:| HGVS ver.15.11 NP_055197.2:| Localization| Putative protein domain| No of probands (and family data)| Classification| Prediction algorithms| Additional data |
|----------|----------------|------------------------|------------------------|---------------|------------------------|---------------------------|----------------|------------------------|----------------|
| Novel mutations |
| 1, 2 | g.130656269 T > G | c.325-3 T > G | p.Ala109_Gln120del† | Intron 4 | M2 | 2: detected in two distinct families (family 1: the first case in the family, the young-adult son of the patient have some slight clinical symptoms, but did not agree for clinical evaluation and genetic test, family 2 - no data) | Likely | Pathogenic (PM4,PM2, PP3, PP4) | mRNA analysis performed |
| 3 | g.130660434 G > T | c.423-1 G > T | p.? | Intron 6 | S2 or A | 1 (DNA of affected son of the proband not analyzed) | Pathogenic (PVS1, PM2, PP3) | MaxEnt: −100.0% | Canonical splice site |
| 4 | g.130678186 G > A | c.899 + 1 G > A | p.? | Intron 11 | M4 | 1 (mutation present in proband and in his affected father) | Pathogenic (PVS1, PM2, PP3) | MaxEnt: −100.0% NNSPLICE: −99.1% HSF: −3.1% | In silico modeling performed |
| 5 | g.130698132 C > G | c.1610 C > G | p.Thr537Arg | Exon 18 | N | 1 (mutations in cis present in patient and his affected mother) | Likely | Pathogenic (PM1, PM2, PP2, PP3) | PolyPhen-2: Probably damaging (score: Hum Div 0.985/1, Hum Var: 0.936/1) SIFT (v6.2.0): Tolerated (score: 0.05, median: 3.58) MutationTaster (v2013): disease causing (p value: 1) | PolyPhen-2: Probably damaging (score: Hum Div 0.985/1, Hum Var: 0.936/1) SIFT (v6.2.0): Tolerated (score: 0.05, median: 3.58) MutationTaster (v2013): disease causing (p value: 1) | mRNA analysis performed |
| 5 | g.130698266 A > G | c.1741 + 3A > G | p.Val524_Ile580del† | Intron 18 | N | Likely | Pathogenic (PM4,PM2, PP3, PP4) | MaxEnt: −100.0% | In silico modeling performed |
| 6 | g.130698138 T > C | c.1616 T > C | p.Leu539Pro | Exon 18 | N | 1 (no family data available) | Likely | Pathogenic (PM1, PM2, PP2, PP3) | PolyPhen-2: Probably damaging (score: Hum Div 0.992/1, Hum Var: 0.936/1) SIFT (v6.2.0): Tolerated (score: 0.05, median: 3.58) MutationTaster (v2013): disease causing (p value: 1) | The new reading frame ends in a STOP codon 16 positions downstream from Thr808 |
| 7 | g.130717154_130717166dup | c.2408_2420del | p.Thr808TyrfsTer16 | Exon 25 | L4 | 1 (no family data available) | Pathogenic (PVS1, PM2, PP3) | PTC | | |
| Fam. No. | Chr3(GRCh37): HGVS ver.15.11 | HGVS ver.15.11 NM_014382.3: | Localization | Putative protein domain | No of probands (and family data) | Classification ACMG | Prediction algorithms | Additional data |
|----------|-----------------------------|-----------------------------|--------------|------------------------|-------------------------------|---------------------|---------------------|-------------------|
| 8,9,10,11 | g.130718422G > T c.2548G > T | p.Gly850Ter Exon 26 | M9 | 4 (in 3 cases mutations were present in at least two affected members of the family, in one case the DNA of several affected relatives of the proband were not analyzed) | Pathogenic (PVS1, PM2, PP3) | PTC | The mRNA produced might be targeted for nonsense mediated decay (NMD) |
| 12 | g.130718438A > G c.2564A > G | p.Gln855Arg Exon 26 | M9 | 1 (no family data available) | Likely Pathogenic (PM2, PP2, PP3) | PolyPhen2:Probably damaging (score: Hum Div 1/1, Hum Var: 1/1) | In silico modeling performed |
| Recurrent mutations | | | | | Reference | | | |
| 13 | g.130660532dupA c.519dup | p.Arg174ThrfsTer4 Exon 7 | A | 1 (mutation present in proband and his child, in whom symptoms appear) | Dobson-Stone et al. 2002 |
| 14 | g.130672792G > A c.659G > A | p.Gly220Glu Exon 8 | A | 1 (no familial data available) | Nellen et al. 2017 |
| 15 | g.130,682,919 T > C | p.Leu335Pro Exon 12 | S4 | 1 (mutation found in proband and 2 affected 1st degree relatives) | Ma et al. 2008 |
| 16 | g.130698260A > G c.1738A > G | p.Ile580Val Exon 18 | N | 1 de novo (mutation not detected in parental DNA) | Dobson-Stone et al. 2002 |
| 17 | g.130715627dupC c.2234dup | p.Ala746SerfsTer11 Exon 23 | M6 | 1 (no familial data available) | Meng et al. 2015 |

† according to mRNA analysis

A: Actuator domain; N: Nucleotide-binding domain; S: (1–5) stalk helices in the cytoplasm; M: (1–10) transmembrane helices
proximity of the ion channel, which seemingly would affect protein location and Ca\(^{2+}\) transport. The effect of p.Leu539Pro is less clear; however it is possible that this substitution would destabilize hydrophobic core formed between β-sheet structures and hence influence ATPase alpha subunit interactions, which in turn could affect ATP binding, its hydrolysis, and finally ion transportation. Unfortunately, no family data were available for probands with p.Leu539Pro and p.Gln855Arg; thus the genotype-phenotype segregation could not be performed.

The clinical significance of another missense, the p.Thr537Arg in exon 18 is more difficult to evaluate. In GnomAD, no records for p.Thr537Arg can be found. The prediction algorithms (PolyPhen, SIFT, MutationTaster) indicated possible pathogenic effect of p.Thr537Arg. Contradictory to them, dynamic structural modeling showed that this solvent exposed substitution most probably does not result in significant conformational change. Furthermore, the p.Thr537Arg was found in cis with a novel, mutation c.1741 + 3A > G in intron 18, which leads to exon 18 in-
frame skipping as we have shown by mRNA analysis. Thus, the protein, if at all synthetized, lacks 57 codons including codon 537. This example of a complex allele containing two variants is not reported before, and c.[1610C > G;1741 + 3A > G], which both were assigned as potentially pathogenic by common prediction algorithms, draws attention on an important issue of careful pathogenicity status evaluation, especially when only selected exons are investigated. Importantly, when p.Thr537Arg status was evaluated alone, it was assigned as “likely pathogenic” using ACMG classification (Richards et al. 2015), which later changed into “uncertain significance” when we detected c.1741 + 3A > G and proved its impact on splicing.

Novel c.325-3 T > G and recurrent c.1308 + 1G > T mutations also lead to in-frame exons skipping (of exons 5 and 15, respectively). Moreover, given that skipping of exons 5 and 15 due to other mutations have been described before (Kitajima 2002; Matsuda et al. 2014; Xiao et al. 2019), our observation indicates that despite distinct molecular lesions, the functional effect of mutations may be similar, which could be significant with regard to the purposes of personalized treatment.

In summary, this is the first report of genetic analysis in Polish HHD patients. Thirteen variants were identified and characterized, including eight unreported before and two recurrent. The results further show heterogeneity in the ATP2C1 mutational spectrum, with possible ethnic-specificity. Last but not the least, by showing a case of complex allele c.[1610C > G;1741 + 3A > G], we also point that careful in silico and extended molecular analysis is essential with respect to proper interpretation of mutation pathogenicity.

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Compliance with ethical standards

Conflict of interest  The authors declare that they have no conflict of interest.

Ethical approval  All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Statement of informed consent  Informed consent was obtained from all individual participants included in the study.

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