Enrichment of rare variants in population isolates: single \textit{AICDA} mutation responsible for hyper-IgM syndrome type 2 in Finland

Luca Trotta1, Timo Hautala2, Sari Hämäläinen3, Jaana Syrjänen4, Hanna Viskari4,5, Henrikki Almusa1, Maija Lepisto1, Meri Kaustio1, Kimmo Porkka6, Aarno Palotie1,7, Mikko Seppänen4,8,9,10 and Janna Saarela*,1,10

Antibody class-switch recombination and somatic hypermutation critically depend on the function of activation-induced cytidine deaminase (AID). Rare variants in its gene \textit{AICDA} have been reported to cause autosomal recessive AID deficiency (autosomal recessive hyper-IgM syndrome type 2 (HIGM2)). Exome sequencing of a multicase Finnish family with an HIGM2 phenotype identified a rare, homozygous, variant (c.416T>C, p.(Met139Thr)) in the \textit{AICDA} gene, found to be significantly enriched in the Finnish population compared with other populations of European origin (38.56-fold, \textit{P}<0.001). The population history of Finland, characterized by a restricted number of founders, isolation and several population bottlenecks, has caused enrichment of certain rare disease-causing variants and losses of others, as part of a phenomenon called the Finnish Disease Heritage. Accordingly, rare founder mutations cause the majority of observed Finnish cases in these mostly autosomal recessive disorders that consequently are more frequent in Finland than elsewhere. Screening of all currently known Finnish patients with a HIGM2 phenotype showed them to be homozygous for p.(Met139Thr). All the Finnish p.(Met139Thr) carriers with available data on their geographic descent originated from the eastern and northeastern parts of Finland. They were observed to share more of their genome identity by descent (IBD) than Finns in general (\textit{P}<0.001), and they all carried a 207.5-kb ancestral haplotype containing the variant. In conclusion, the identified p.(Met139Thr) variant is significantly enriched in Finns and explains all thus far found AID deficiencies in Finland.


\section*{INTRODUCTION}

Primary immunodeficiency disorders (PIDDs) encompass a wide range of genetically determined inborn errors of immunity. Presently, >5000 variants affecting the function have been reported in over 250 genes as causative of 265 PIDDs, and novel defects continue to be discovered.\textsuperscript{1}

Hyper-IgM syndromes (HIGMs) include a genetically heterogeneous group of PIDDs defined by early-onset recurrent infections and autoimmunity, absence or very low levels of IgG, IgA and IgE, but elevated or normal serum IgM levels.\textsuperscript{2} This phenotype typically results from inherited defects in proteins involved in class-switch recombination (CSR) and somatic hypermutation (SHM). Classical HIGM-causing genes include \textit{CD40LG}, \textit{AICDA}, \textit{CD40} and \textit{UNG}.	extsuperscript{3} CSR, SHM and central B-cell tolerance critically depend on normal activation-induced cytidine deaminase (AID) function.\textsuperscript{4–6} AID also participates in removal of epigenetic memory by active demethylation.\textsuperscript{7} The immunologic HIGM phenotypes of AID and uracil DNA glycosylase (UNG) deficiencies closely resemble each other and are relatively easy to screen. In these, no CD19+CD27+IgD–IgM– switched memory B (smB) cells can be found in blood, whereas marginal zone CD19+CD27+IgD–IgM+ B (MZB) cells are normal or high. AID deficiency is estimated to affect \textless 2/10\textsuperscript{6} individuals.\textsuperscript{8}

The population history of Finland is characterized by a restricted number of founders, isolation, several population bottlenecks and recent expansion of the population. This has led to the enrichment of some deleterious variants and loss of others, creating a phenomenon called the Finnish Disease Heritage (FDH). By definition, FDH disorders are more frequent in Finland than elsewhere, and a majority of the Finnish patients share the same founder mutation.\textsuperscript{9}

Here, we have identified the Finnish founder allele causing HIGM2 and assessed its prevalence in Finns compared with other populations.

\section*{MATERIALS AND METHODS}

This study was conducted in accordance to the principles of the Helsinki Declaration and was approved by the Coordinating Ethics Committee of

1\textsuperscript{Institute for Molecular Medicine Finland, University of Helsinki, Helsinki, Finland; 2Department of Internal Medicine, Oulu University Hospital, Oulu, Finland; 3Department of Internal Medicine, Kuopio University Hospital, Kuopio, Finland; 4Department of Internal Medicine, Tampere University Hospital, Tampere, Finland; 5Department of Virology, University of Tampere, School of Medicine, Tampere, Finland; 6Institute for Molecular Medicine Finland, University of Helsinki, Helsinki, Finland; 7Hematology Research Unit, Biomedicum Helsinki and Hematology Division, Comprehensive Cancer Center, University of Helsinki and Helsinki University Hospital, Helsinki, Finland; 8Program in Medical and Population Genetics, Broad Institute of MIT and Harvard, Cambridge, MA, USA; 9Rare Disease Center, Children's Hospital, University of Helsinki and Helsinki University Hospital, Helsinki, Finland; 10Adult Immunodeficiency Unit, Inflammation Center, University of Helsinki and Helsinki University Hospital, Helsinki, Finland}

\textsuperscript{10}These authors contributed equally to this work.

*Correspondence: Dr M Seppänen, Rare Disease Center, Children's Hospital, University of Helsinki and Helsinki University Hospital, PO Box 280, FI-00029 HUS, Helsinki, Finland. Tel: +358 9 47180201; Fax: +358 2 941 25737; E-mail: miko.seppanen@helsinki.fi or Dr J Saarela, Institute for Molecular Medicine Finland, University of Helsinki, PO Box 20, FI-00014, Helsinki, Finland. Tel: +358 40 5123 801; Fax: +358 2 941 25737; E-mail: janna.saarela@helsinki.fi

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Abbreviations: F, female; M, male; Ig, immunoglobulin; NA, not assessed; ND, not detectable (values under the detection threshold of the analysis method). The p.(Met139Thr) variant in the AICDA gene refers to the following reference sequences: RefSeq NG_011588.1 and NM_020661.2 (GRCh37.p13).

*For more detailed history see Supplementary Information.*
Patients

The index case of the family I (I-I) was immunologically characterized in Helsinki University Hospital. She has undetectable level of smB cells and a typical clinical picture of HIGM2 (Table 1). Next, patient cohorts of the pediatric and adult immunodeficiency units of all five Finnish university hospitals were screened for patients with a phenotype compatible with either AID or UNG deficiency. All subsequent patients (families II–IV) with (1) low or absent IgA, IgG and IgE levels but normal or high IgM levels according to the laboratory reference values, together with (2) missing smB cells but normal or high levels of MZB in B-cell phenotyping (for methods see Haapaniemi et al.) were included in the study. Study subjects underwent clinical and immunological evaluations at Helsinki, Kuopio, Oulu and Tampere University Hospitals. All available patient records since June 1959 were reviewed and patients interviewed. Altogether, four families were identified (Table 1 and Figure 1). Patient histories are described in detail in the Supplementary Information.

Molecular genetics

Genomic DNA of the studied individuals was isolated using standard salt precipitation protocols. Exome sequencing was performed in the two index patients of family I and in two of their healthy relatives to investigate the genetic basis of their familial disease presentation. A Nextera Rapid Capture Exome kit (Illumina, San Diego, CA, USA) was used for library preparation and exome enrichment and sequencing was performed on a HiSeq 1500 platform (Illumina). The data were analyzed using a version 2.7 of the in-house developed analysis pipeline for quality control and variant identification (VCP). Detailed sequencing statistics and procedures for read alignment and variant calling are provided in the Supplementary Information. Additional patients were screened for AICDA allelic variants by Sanger sequencing. The analysis of the AICDA gene (GRCh37.p13:12:8754762-8765463) and the primer design were performed using the following genomic and transcript sequences: RefSeq NG_011588.1 and NM_020661.2. The identified variants and other patient data were deposited in the LOVD database (http://databases.lovd.nl/shared/genes/AICDA) (variant ID: AICDA_000004; individual IDs: 00058568, 00058569, 00058570, 00058572, 00058573, 00058574 and 00058575).

DNA variants were verified by restriction endonuclease digestion. See Supplementary Information for the sequence of primers and further description of the procedures.

Population analysis

We performed a population-based analysis of the identified sequence variant frequency by using data of 60,786 individuals from Exome Aggregation Consortium including 34,699 individuals of European origin, of whom 3013 were Finnish.

The geographic distribution in Finland of the p.(Met139Thr) alleles (RefSeq NM_020661.2; c.416T>C; rs200858797) was illustrated based on the information obtained from the study subjects and from the carriers included in the SISu project (https://sisu.fimmm.fi/)12 for which such data were available and in three Finnish sample collections (the Finnish Twin Cohort study, the National Finrisk Study and the Migraine Family Study; Supplementary Information, Supplementary Table 1 and Figure 2). The analysis of pairwise segmental sharing was conducted on a set of 6755 Finns included in epidemiological and clinical Finnish sample collections, of whom 20 were p.(Met139Thr) carriers, using 113 common markers genotyped using HumanCoreExome-24 BeadChips (Illumina; 1000 Genomes;
indicated by a purple spot, listing the number of the family (from I to IV).

AIDCA biallelic AICDA previously been tested in a reference laboratory to have wild-type from Eastern Finland (Figure 1). The index of the family had surrounding the p.(Met139Thr) were screened for a putative shared allele, (v. 1.07, http://pngu.mgh.harvard.edu/purcell/plink/). Next, according to the National Finrisk Study and the Migraine Family Study) (Supplementary clinical Finnish sample collections (the Finnish Twin Cohort study, the SISu). The p.(Met139Thr) variant included in

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Figure 2 Distribution of the AICDA p.(Met139Thr) carriers in Finland. Blue triangles point to the geographical origin of the Finnish carriers (n = 27) of the p.(Met139Thr) variant included in SISu and in epidemiological and clinical Finnish sample collections (the Finnish Twin Cohort study, the National Finrisk Study and the Migraine Family Study) (Supplementary Table 1). Yellow symbols indicate the birthplaces of carriers’ parents, if discordant. The birthplaces of the patients identified in this study are indicated by a purple spot, listing the number of the family (from I to IV). For families III and IV, the mother corresponds to ‘a’ and the father to ‘b’. The black dots mark the main municipal areas.

www.1000genomes.org; accessed 18 April 2014). A cryptic relatedness analysis was performed by using the identity by descent (IBD) estimation on the above-mentioned set of unaffected individuals (Supplementary Information).

In parallel, a segregation analysis of polymorphisms in a 2-Mb region encompassing the p.(Met139Thr) was executed in the 32 carriers included in the above mentioned clinical and epidemiological sample collections in addition to the SISu data set (Supplementary Table 2) by utilizing PLINK (v. 1.07, http://pngu.mgh.harvard.edu/purcell/plink/). Next, according to the observed haplotype blocks, a total of 80 markers located in a 1.1-Mb region surrounding the p.(Met139Thr) were screened for a putative shared allele, carried on in the comprehensive group of 32 carriers (Supplementary Table 3).

Statistical analysis
Pearson’s χ² test (10⁸ simulations) was used to evaluate the different p.(Met139Thr) allelic distributions among Finns and the more heterogeneous European populations (Supplementary Table 4), and the load of hidden relatedness among the carriers and the general population was weighed using Welch’s two-sample t-test (10⁸ simulations).

RESULTS
Genetic analysis
We first studied a family with four affected individuals originating from Eastern Finland (Figure 1). The index of the family had previously been tested in a reference laboratory to have wild-type AICDA and UNG. However, exome sequencing revealed a known biallelic AIDCA variant in the living affected members (p.(Met139Thr)) that has previously been shown to cause HIGM2. The two healthy relatives carried one copy of the variant. Targeted Sanger sequencing of an archived sample from the index verified the presence of the same biallelic sequence change (I-I, Figure 1). Thereafter, all remaining Finnish patients with a compatible phenotype (n = 4) were screened and found homozygotes for the p.(Met139Thr) variant (Figure 1 and Table 1).

Population analysis
Overall, we found the HIGM2 causing p.(Met139Thr) alteration to have a frequency of 0.012% in a total of 57 391 exomes provided by the Exome Aggregation Consortium (ExAC). More detailed analysis of the data revealed an allelic frequency of 0.0047% in 31 686 individuals of European ancestry (non-Finns) and the absence of the variant in non-European populations (22 692 individuals). Compared with other populations of European origin, a statistically significant 38.56-fold allelic frequency was observed in Finns with 11 uniallelic carriers in 3013 exomes (0.18%, P < 0.001; Supplementary Table 4), resulting in the calculated theoretical frequency of AID deficiency of 0.81/10⁶ in those of Finnish ancestry. Other AICDA variants showed no substantial differences in frequencies between the populations (data not shown).

Because of the enrichment of the p.(Met139Thr) variant in Finland, we studied its geographical distribution based on the information on birthplace retrieved from the studied subjects, and from those 27 out of 31 carriers within the SISu cohort and other Finnish sample collections with such data available. Interestingly, all of the AID deficiency patients and 24 of the 27 carriers originated from the late settlement regions of Eastern and Northeastern Finland, suggesting shared origin for the p.(Met139Thr) alleles in all these individuals (Figure 2). The remaining three carriers were born in Helsinki area that has experienced substantial immigration from the rest of the country during recent centuries. Thus, we searched for possible shared haplotype in the region surrounding AICDA by using the exome data for 3325 individuals of the SISu cohort, including 11 p.(Met139Thr) carriers. We first retrieved the haplotype structure of the 2 Mb genomic region encompassing the p.(Met139Thr) and observed clear haplotype blocks 90 kb upstream and 51 kb downstream of the variant (Supplementary Table 5). Further examination of the genomic region flanking AICDA using the UCSC Genome Browser revealed the presence of a 10-kb recombination hot spot encompassing the gene that likely weakens the possibility of tracking a conserved ancestral allele. Nonetheless, by combining the genetic data of all the 31 carriers of the two different population-based data sets (exome data of the SISu cohort and genotyping data of the Finnish epidemiological and clinical cohorts) and the two exome sequenced familial carriers, and by monitoring the alleles seen in each haplotype block, we identified a 207.5-kb core haplotype including the p.(Met139Thr) variant shared by all the carriers (Figure 3). The minimal shared region was restricted by recombination in five individuals, whereas the core haplotype extended significantly further in the others (Figure 3). Further comparison of the pairwise genome-wide IBD showed higher values in the group of p.(Met139Thr) carriers (average piHat = 0.007 ± 0.0027) than in the general population (piHat = 0.003 ± 0.005), displaying significant increased relatedness within the carriers (P = 1.59E−12).

DISCUSSION
In the current study, we identified a Finnish founder mutation for AID deficiency. The rare recessive p.(Met139Thr) allelic variant in the AICDA gene causes the disease in all known Finnish patients. The variant, previously confirmed to affect the AID function in a
single HIGM2 patient of unknown origin,\textsuperscript{14} exhibits a significant (38.56-fold) enrichment in Finns compared with the data from other European populations.

There are at least 117 previously published cases of AID deficiency and, currently, at least 43 autosomal recessive or dominant negative causative \textit{AICDA} variants have been reported.\textsuperscript{16–19} The observed p.(Met139Thr) change affects an evolutionarily conserved amino acid residue in the APOBEC-like domain, and \textit{in silico} analyses are consistent with a deleterious effect, resulting in severely impaired CSR.\textsuperscript{15} Interestingly, a different causative missense substitution affecting the same amino acid has been found in three Turkish patients with HIGM2 (RefSeq NM_020661.2; c.415A\textsuperscript{+}G, p.(Met139Val); \textit{UNG} c.1352G\textsuperscript{+}A, p.Asp451Glu).\textsuperscript{19} This disrupts the AID activity \textit{in vitro}.\textsuperscript{22}

Given the known frequency of uniallelic p.(Met139Thr) in 3013 Finns, the predicted prevalence for homozygous individuals was \(\sim 0.81/10^6\). However, the presently known prevalence of AID deficiency in Finland is 1.5/10\(^6\), greatly exceeding the estimate in literature, although this was partly based on the incidence in French-Canadians, the other known population with an \textit{AICDA} founder mutation.\textsuperscript{8} Currently, there are relatively few Northern and Northeastern Finns in the SiSu cohort, potentially explaining the observed difference between theoretical and known prevalence. To evaluate the contribution of other variants causing \textit{HIGM}s in the Finnish population, we performed a similar population-based comparison of allele frequencies for all the variants in genes affecting CSR (\textit{AICDA}, \textit{UNG}, \textit{CD40} and \textit{CD40LG}). None of the other allelic variants were significantly enriched in Finns compared with other populations. No reliable data are available for the CHH-associated variants.

As almost all p.(Met139Thr) carriers originated from the late settlement areas of Eastern and Northeastern Finland, the geographic distribution of the variant fits well with the known inhabitation patterns of the country and suggest a single origin.\textsuperscript{29,30} Consequently, we made an effort to validate this hypothesis by analyzing the genomic region encompassing the \textit{AICDA} gene in the individuals of the \textit{SiSu} cohort. We identified linkage disequilibrium blocks upstream and downstream the p.(Met139Thr), suggesting the actual architecture of the region as reflecting the remnants of a wider previous haplotype that has potentially included the variant. The small number of the carriers and, mostly, the presence of a recombination hot spot of 10 kb surrounding the \textit{AICDA} gene\textsuperscript{31} could have limited our ability to identify the ancestral allele. In order to overcome these limitations, we further studied the haplotype structure of the surrounding areas in all the p.(Met139Thr) carriers included in the \textit{SiSu} project and three epidemiological and clinical Finnish sample collections. A segregation analysis revealed a shared haplotype of 207.57 kb including the variant, with no apparent recombination events in the 31 carriers. This was surrounded by a partially conserved genomic region of 901 kb where a limited amount of recombination events had taken place. The outlined genetic structure comprised the likely ancestral founder allele. Our hypothesis of a single mutation event and shared ancestry was also further strengthened by the finding that all p.(Met139Thr) carriers shared more of their genome than the general population (2.25-fold increased IBD).

AID deficiency is clinically characterized by severe antibody deficiency, lymphatic nongranulomatous hyperplasia with hyperplastic germinal centers and inflammatory complications like hemolytic
autoimmunity, chronic hepatitis, diarrhea and aseptic arthritis.16,17 Our patients display a uniform matching phenotype. The patients of family I have a longer follow-up than most patients in the current literature, lending insight into the long-term consequences of the disease. To the best of our knowledge, the patients have developed several previously unreported systemic, renal and gastrointestinal autoimmune complications (Table 1 and Supplementary Information). However, aggressively substituted younger patients in families II–IV seem to have few autoimmune problems. Unlike in common variable immunodeficiency, granulomatous lymphadenitis is not a previously described feature of AID deficiency. A pronounced and difficult to treat granulomatous lymphadenopathy was noted in family I and confirmed by biopsies. Unfortunately, no archived tissue samples were available. As this occurred during a familial tuberculoid mini-epidemic, it suggests that infectious causes of granulomas should always be excluded in AID deficiency. Opportunistic lethal infections in I–II were likely caused by secondary immunosuppression and are also not a feature of AID deficiency. Whether AID deficiency is able to cause spontaneously terminated pregnancies should be further studied (cf. Supplementary Information).

In summary, we identified a single variant affecting the function of the protein accounting for all diagnosed AID deficiencies in Finns. In all likelihood, p.(Met139Thr) is a Finmajor founder mutation and AID deficiency belongs to the FDH. This phenomenon closely resembles the known p.Arg112Cys founder allele in French Canadians, but 3p.(Met139Thr) is even more prevalent in Finns.8 Taken together, these findings underline the correlation between the genetic structure of the population and the distribution of genetic disorders, and emphasize the benefits of researching population isolates with systematized health records available.

CONFLICT OF INTEREST
The authors declare no conflict of interest.

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