Next-generation sequencing reveals novel variants and large deletion in FANCA gene in Polish family with Fanconi anemia

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Abstract

Background: Fanconi anemia (FA) is the most common inherited bone marrow failure syndrome. However, establishing its molecular diagnosis remains challenging. Chromosomal breakage analysis is the gold standard diagnostic test for this disease. Nevertheless, molecular analysis is always required for the identification of pathogenic alterations in the FA genes.

Results: We report here on a family with FA diagnosis in two siblings. Mitomycin C (MMC) test revealed high level of chromosome breaks and radial figures. In both children, array—Comparative Genomic Hybridization (aCGH) showed maternally inherited 16q24.3 deletion, including FANCA gene, and next generation sequencing (NGS) disclosed paternally inherited novel variants in the FANCA gene—Asn1113Tyr and Ser890Asn. A third sibling was shown to be a carrier of FANCA deletion only.

Conclusions: Although genetic testing in FA patients often requires a multi-method approach including chromosome breakage test, aCGH, and NGS, every effort should be made to make it available for whole FA families. This is not only to confirm the clinical diagnosis of FA in affected individuals, but also to enable identification of carriers of FA gene(s) alterations, as it has implications for diagnostic and genetic counselling process.

Keywords: Fanconi anemia, Thrombocytopenia, Chromosome breakage test, aCGH, FANCA gene

Background

Fanconi anemia (FA) is the most common form of inherited bone marrow failure syndromes (IBMFS) related to developmental abnormalities. Clinical symptoms, present in about 75% of patients, most often include short stature, microcephaly, thumb and radial side limb defects, abnormal skin pigmentation, and genitourinary defects. Progressive bone marrow failure occurs in the first decade of life, initially often expressed by leukopenia or thrombocytopenia. The most common malignancies occurring in patients with FA are acute myeloid leukemia and solid tumors of the head and neck, skin, and gastrointestinal as well as genitourinary systems [1, 2]. At the cellular level, FA is characterized by sensitivity to interstrand crosslinking agents (ICL), such as diepoxybutane (DEB) and mitomycin C (MMC) [3, 4]. FA is mostly caused by biallelic pathogenic variants in any of the 21 genes: FANCA, FANCC, FANCD1(BRCA2), FANCD2, FANCE, FANCC, FANCG, FANCI, FANCJ(BRIP1), FANCL, FANCM, FANCP(PALB2), FANCO(RAD51C), FANCP(SLX4), FANCQ(ERCC4), FANCS(BRCA1), FANCS(BRCA1).
FANCT (UBE2T), FANCU (XRXX2), FANCV (MAD2L2/REV7), FANCW (RFWD3), FANCY (FAP100). Variants in FANCB (presenting X-linked recessive inheritance pattern) and FANCR/RAD51 (presenting autosomal dominant inheritance pattern) have also been reported as disease-causing, therefore a total of 23 genes have been implicated in the pathogenicity of FA so far [5, 6]. In two-thirds of FA patients pathogenic variants occur in FANCA, making it the most frequently mutated FA gene. Approximately 20% of patients carry mutations in FANCC or FANCG, and mutations in the other 20 FA genes account for 0.1–4% cases per gene. FANCA variants include single nucleotide variants, small insertions, and deletions which may be detected by Next Generation Sequencing (NGS). However, 20–40% of FANCA mutations are large deletions, which size span a wide range from ~1 to 545 kb, identified by array—Comparative Genome Hybridization (aCGH) or multiplex ligation-dependent probe amplification (MLPA) [7, 8].

The wider study we performed under the research grant WL.131 aimed to provide a detailed and accurate genetic diagnosis for all Fanconi anemia individuals and their families by establishing a comprehensive diagnostic approach for this rare condition. In this paper, we focus on clinical presentation and the results of genetic analyses in one selected family showing that a broad approach comprising chromosome breakage test followed by chromosomal microarray analysis and NGS is not only sufficient to provide the genetic diagnosis in FA patients, but also proves beneficial when extended to the whole family.

**Methods**

**Clinical and hematological report**

A 5-year-old male (Patient 1—IV.3) was born with a birth weight of 2520 g, and Apgar score of 10. He was the first child of unrelated healthy parents. There was a history of three recurrent miscarriages in his mother, as well as a presence of congenital defects and early infant death on the maternal side of the pedigree (Fig. 1a). An extra thumb, a heart defect with patent ductus arteriosus (PDA), and abnormal localization of the right kidney were noted at newborn period. PDA was closed with coil implementation at the age of 6 months. During the following years short stature, low weight, café-au-lait spots, and conductive hearing impairment became apparent. At the age of five he was referred to the Clinical Genetic Department where slight epicanthic folds, large and short neck, kyphoscoliosis, pes plano-valgi were also noted (Fig. 1b). The boy met the criteria of congenital abnormality score (CABS) 3 (1 point each for heart anomaly, kidney defect, and hearing impairment) [9].

For the first 2 years of life complete blood count was normal except for the episode of acute hemolytic anemia due to the PDA (this presented with elevated unconjugated bilirubin levels, reticulocyte count, and decreased haptoglobin). Complete recovery was achieved after additional coil deployment and infusion of 7 units of red blood cell concentrate.

The next episode of cytopenia occurred at the age of 2 years with decreased platelet count on pneumonia. In the third year of life the median platelet and neutrocyte counts were $110 \times 10^3/\mu l$ (91–129) and $1 \times 10^3/\mu l$ (0.98–2.8), respectively. In the fourth year of life these results amounted to $78 \times 10^3/\mu l$ (35–103) and $1 \times 10^3/\mu l$ (0.67–1.5), in the fifth one to $58 \times 10^3/\mu l$ (35–99) and $0.95 \times 10^3/\mu l$ (0.53–1.9), respectively. For the first time hemoglobin reduction to 9.8 g/dl occurred at 6 years of age, and at the same time platelet count dropped below $20 \times 10^3/\mu l$. The boy started to demonstrate hemorrhagic symptoms (bruising, epistaxis) and became dependent on platelet transfusions. Myelogram showed low cellularity (20–30%) represented nearly exclusively by the erythroid lineage cells.

After the diagnosis of FA was confirmed by genetic testing, an allogeneic hematopoietic stem cell transplantation (allo-HSCT) was performed from the matched unrelated donor (MUD). Following conditioning with busulfan (0.8 mg/kg/day, 4 days), cyclophosphamide (10 mg/kg/day, 4 days), fludarabine (30 mg/m²/day, 4 days), and antithymocyte globulin (40 mg/kg/day, 4 days), peripheral blood cells of $4.5 \times 10^8$ CD34+ /kg and $0.87 \times 10^8 CD3+ /kg$ were infused. Cyclosporine (days $-3$ to $+100$) with mycophenolate mofetil (45 mg/kg/day days +1) for graft-versus-host disease (GvHD).

![Fig. 1](image)

*Summary of clinical and genetic findings in the presented family. a Pedigree of FA family. Squares—males, circles—females, open—unaffected individuals, filled—affected individuals, half filled—carriers of deletion or mutation. MMC+/- = presence or absence of heterozygous deletion encompassing FANCA gene detected by aCGH, FANCA+/- = presence or absence of mutation in FANCA gene detected by NGS, Ca/Ca? = presence or possible presence of cancer. Black arrow indicates the proband. b–c Congenital anomalies in siblings with FA. B—short stature, posture defect, extra thumb and pes plano-valgi in patient 1 (IV.3). C—large short neck, corrected extra thumb, and cafe-au-lait spots in patient 2 (IV.4). d–e Metaphase spreads in patients 1 (d) and 2 (e)—MMC test. Green arrows show gaps, chromatid breaks (chb) andacentric fragments (ace). Red arrows show radial figures. Blue arrow shows chromatid interchange figure. f Partial array-CGH result showing the 16q24.3 deletion in the proband (patient IV.3). The same heterozygous deletion was detected in his siblings and mother. g Visualisation of FANCA sequencing reads showing c.3378TA>T; p.Asn1113Tyr variant (g1), and Sanger sequencing electrophoregram plots confirming the variant (g2—wild type found in the mother III.3, and younger daughter IV.6, g3—mutation in the heterozygous father III.2, g4—mutation in the only copy of the gene in IV.3 and IV.4 siblings).*
Fig. 1 (See legend on previous page.)
prophylaxis was administered. The time to neutrophil (>0.5 × 10⁹/l), platelet (>20 × 10⁹/l) engraftment was 14 and 9 days, respectively. The first reticulocytes emerged on day +10, and maximal reticulocyte count (41%) was achieved on day +17, with complete donor chimerism. Only minor feverish episodes complicated the post-transplant course. There were neither symptoms of viral infections reactivation nor graft versus host disease. The boy receives regular follow-up care, and up to date presents with normal ranges of a complete blood cell counts. Timeline summary of clinical and diagnostic information for this patient is shown in Fig. 2.

His 1 year younger sister (Patient 2—IV.4) was born with a birth weight of 2325 g and Appgar scores of 10. An extra thumb, PDA, and ectopic pelvic left kidney were noted at the newborn period. Due to suspicion of Fanconi anemia in the family, the girl was referred for genetic evaluation. On physical examination a large philtrum, large short neck, and large chest were noted (Fig. 1c). The girl fulfilled the criteria of CABS2 score. She shows normal growth parameters according to standard growth curves, her PDA is stable and does not require surgical intervention. The complete blood count has been monitored since the girl was five. In her sixth and seventh years of life the median neutrocyte count was 1.4 × 10³/µl, while the median platelet count was 128 × 10³/µl and 97 × 10³/µl, respectively.

The youngest sister (Patient 3—IV.6) and the parents have not presented symptoms similar to those of older siblings.

**G-banding analysis and chromosome breakage test**

Heparinized venous blood (≥ 5 ml) was used to prepare whole-blood cultures as usual for routine cytogenetic analysis. Four cultures per individual and healthy control were prepared by adding 0.5 ml blood to 4.5 ml complete RPMI medium (Gibco) supplemented with 15% fetal bovine serum (Gibco), gentamicin and phytohemagglutinin (PHA) (both Gibco), as prescribed (Joenje and co-workers). The cultures were incubated for 72 h with 0, 50, 150, or 300 nM mitomycin C (MMC; Sigma Aldrich). After treatment with colcemid (0.1 µg/ml) for 50 min, cells were harvested, incubated with 0.075 M KCl for 20 min at 37 °C, and fixed with 75% methanol and 25% acetic acid. Cytogenetic preparations were stained for 2 min in a 5% Giemsa solution (Merck); banding technique was applied only to slides from cultures without MMC to analyze the karyotype. The microscope analysis was performed with a Nikon E600 microscope and later analyzed by computer assisted metaphase system (ASI, Israel).

Chromosome instability data were analyzed and calculated (number of metaphases with breakage, mean chromosome break events per aberrant metaphase, and tri-,
tetra-, and multi-radials frequency) [10]. A healthy control was assured.

Nucleic acid extraction
Blood samples were collected from the proband, his sisters and parents. Whole genomic DNA was extracted from nucleated cells by silica gel column methods according to the manufacturer’s protocol (Qiagen) for aCGH assay and by using Prepito TM for NGS assay.

Array—Comparative Genomic Hybridization (aCGH)
aCGH was performed using the commercially available platform (Sure Print G3 Human CGH 8 × 60 Microarray Kit, Agilent Technologies, Santa Clara, CA, USA), according to the manufacturer’s protocol. Copy number analysis was performed using Agilent CytoGenomics Software 5.0.

Next generation sequencing (NGS)
A targeted next-generation sequencing (MiSeq, Illumina) was performed using a targeted multigene custom panel including ATM, BLM, BRIP1, FANCA, FANCC, FANCG, NBN, and PALB2 genes. The sequencing library preparation was performed using SeqCap EZ Choice NimbleGen (Roche). The sample was sequenced with Illumina technology on Miseq sequencer with 2 × 75 bp reads. Demultiplexing of the sequencing reads was performed with Illumina’s bcl2fastq2 v2.19.0. Adapters were trimmed with Skewer version 0.2.9 [11]. The reads were aligned to GRCh37/hg19 reference sequence using BWA-MEM [12]. Read duplicates were removed using Picard 2.18.2 (http://broadinstitute.github.io/picard/). Variant call was performed with GATK v4.0.3.0 HaplotypeCaller [13, 14] and FreeBayes (v1.2.0-2-g29c4002) [15]. Variants have been annotated with databases (i) VEP97 [16]: annotations Sift, Polyphen2, (ii) dbNSFPv4.0 [17] annotations: MutationAssessor, MutationTaster, DANN, FATHMM, (iii) ESP6500, (iv) GnomAD, (v) dbSNP, (vi) ClinVar and (vii) 1000 Genomes [18].

For the purpose of the study all the genetic analyses were performed in all siblings and their parents.

Results

G-banding analysis and chromosome breakage test
Cytogenetic GTG analysis showed normal karyotypes in all family members.

A standard chromosome breakage test with mitomycin C revealed chromosomal hypersensitivity to crosslinking agent in patients 1 (IV.3) and 2 (IV.4) (Fig. 1d–e). The number of metaphases with breaks, the mean number of breaks per aberrant metaphase, and radials frequency were higher than the normal range for non-FA cells (Table 1). The cytogenetic findings were compatible with Fanconi anemia. The results of patient 3 (IV.6) and the parents were normal.

Array—Comparative Genomic Hybridization (aCGH)
In patients 1 (IV.3), 2 (IV.4), and 3 (IV.6), aCGH analysis showed a heterozygous deletion of ~93 kb in chromosome 16q24.3 (Fig. 1f), encompassing the whole FANCA gene (arr[hg19] 16q24.3(89804014_89897040)×1). Parental array analysis showed the same heterozygous deletion in the mother (III.3). No other copy number imbalances of potential clinical significance have been detected.

Next generation sequencing (NGS)
Next-generation sequencing analysis revealed in patients IV.3 and IV.4 the presence of three rare variants (with minor allele frequency <1%), all localized in cis, in the only allele of FANCA gene: (i) a novel variant c.3337A > T; p.(Asn1113Tyr) found in exon 33 [NM_000135.2; NP_000126.2] (Fig. 1g1); (ii) c.2669G > A; p.(Ser890Asn), rs1055448646 in exon 28, and (iii) c.1614G > A; p.(Gly538=), rs747421581 in exon 17. According to ACMG classification [19], the p.Asn1113Tyr is a likely pathogenic variant—it meets the criteria of PM1 (is located in a mutational hotspot and/or critical and well-established functional domain), PM2 (is absent in mutation databases such as gnomAD), PP2 (missense variants are a common cause of the disease), PP3 (bioinformatic algorithms predict its

| Patient | Total cells | % Cells with breaks | Breaks/cell | Radial forms |
|---------|-------------|---------------------|-------------|--------------|
| 1       | 22 cells    | 14.00%              | 0.14 br/cell| None         |
| 2       | 21 cells    | 0.00%               | 0.00 br/cell| None         |
| Control | 20 cells    | 0.00%               | 0.00 br/cell| None         |

| Patient | Total cells | % Cells with breaks | Breaks/cell | Radial forms |
|---------|-------------|---------------------|-------------|--------------|
| 1       | 50 cells    | 96.00%              | 8.48 br/cell| 65           |
| 2       | 50 cells    | 100.00%             | 10.84 br/cell| 107          |
| Control | 50 cells    | 4.00%               | 0.05 br/cell| None         |

Table 1 Cytogenetic findings of spontaneous and MMC-induced chromosome aberrations (breaks, radial forms) of examined patients 1 (IV.3) and 2 (IV.4) compared to healthy control group
The FANCA gene (OMIM: 607139) is located on chromosome 16q24.3, contains 43 exons along a coding sequence of 4.3 kb, spans approximately 80 kb [23, 24] and is by far the most commonly mutated gene in patients with FA worldwide, accounting for the disease in the 60–70% of FA families. However, the percentages may differ for certain ethnic populations [8, 22, 25–29]. Therefore, molecular testing usually starts with this particular gene [8, 29]. Extensive allelic and non-allelic heterogeneity of FANCA gene mutations causes major problems in molecular diagnostics. If no mutation in FANCA is detected, the screening is extended to other FA genes [31]. The next most frequently mutated genes that are responsible for ~10–15% of cases are FANCC and FANCG [8, 22, 27, 28]. Causative mutations in other FA genes are rare and account for 0.1–3% of cases per gene [8, 22].

Several diagnostic approaches for FA have been described in the literature. The diagnosis of FA is still classically based on clinical evaluation followed by exposing patients’ lymphocytes to MMC or DEB, and performing subsequent quantification of all types of chromosomal breakages and radial forms [29, 30]. Although a positive chromosome breakage test is highly indicative for FA, it is not 100% specific for FA and can produce false-negative and false-positive results [31, 32]. Therefore, a molecular investigation is needed for definitive and accurate diagnosis, prognosis assessment and genetic counseling for FA families [28]. This, however, can be a daunting, complex, and time-consuming task [8, 20, 23, 29] due to genetic heterogeneity, large (and constantly growing) number of FA genes, together with a wide spectrum of private mutations [33].

Complete molecular diagnosis of the disease-causing specific pathogenic variants is fundamental for confirmation of the diagnosis, management of patients, genetic counseling, carrier testing, and prenatal diagnosis. Molecular diagnosis is also critical for the identification of appropriate donors for bone marrow transplantation, and for exploring novel therapeutic approaches [7, 8, 29, 31].

FA-related genes present a remarkable mutational heterogeneity [26]. The ever-growing portfolio of mutations includes point mutations, small insertions/deletions, splicing defects, and large deletions [8, 34]. The latter appears to be a frequent subtype of variation, especially in FANCA, accounting for 20–40% of all pathogenic variants of this gene [7, 8, 22, 35, 36].

Significant proportion of large deletions results from a high content of Alu repeats in this region. Alu repeats can facilitate intrachromosomal recombinations and generate intragenic copy number variants through unequal crossing-over and their presence is directly correlated with deletion breakpoints [28, 29, 36–38]. Alu elements can also induce exon skipping with the effect of both upstream and downstream alternative exons [23, 35].

Several methods have been used in screening protocols for FANCA. However, it became clear that they must incorporate methods that can detect point

![Table 2 Clinically relevant molecular findings in FANCA gene in examined family](https://example.com/table2)

| Patient         | Asn1113Tyr | Ser890Asn | Large deletion | Genotype |
|-----------------|------------|-----------|----------------|----------|
| IV.3/brother    | +          | +         | +              | m/del    |
| IV.4/older sister | +         | +         | +              | m/del    |
| IV.6/younger sister | −       | −         | +              | −/del    |
| III.3/mother    | −          | −         | +              | −/del    |
| III.2/father    | +          | +         | −              | m/−      |

m mutation, del deletion

Discussion

Fanconi anemia is a complex disorder, therefore understanding its underlying molecular mechanisms is critical not only for the diagnosis but also for the clinical management of the affected individuals and their families [20].

Clinical presentation of FA patients is highly variable even among individuals within the same family or among patients within the same complementation group, therefore the diagnosis should be confirmed by a positive chromosomal breakage test and identification of a causal pathogenic mutation in one of the FA-related genes [21, 22].

Table 2 shows summary of detected variants.

To confirm the pathogenic character of FANCA gene variants identified in affected children and their father, functional studies should be performed. Our classification of mutations done in accordance with mutation databases has only presumable character.

### Table 2 Clinically relevant molecular findings in FANCA gene in examined family

| Patient         | Asn1113Tyr | Ser890Asn | Large deletion | Genotype |
|-----------------|------------|-----------|----------------|----------|
| IV.3/brother    | +          | +         | +              | m/del    |
| IV.4/older sister | +         | +         | +              | m/del    |
| IV.6/younger sister | −       | −         | +              | −/del    |
| III.3/mother    | −          | −         | +              | −/del    |
| III.2/father    | +          | +         | −              | m/−      |

m mutation, del deletion
mutations as well as small and large deletions [26, 36]. In the past, after clinical diagnosis and a positive chromosome breakage test, FA patients’ molecular characterization was achieved by Sanger sequencing of candidate genes. This approach for FA is time-consuming, costly and, the most importantly, may not detect all types of disease-causing mutations [21]. NGS seems to facilitate a more comprehensive diagnostic strategy for FA patients’ characterization, especially with the development of targeted panels for FA genes [22, 26].

Large deletions can be detected by multiplex ligation-dependent probe amplification assay [8]. However, MLPA is a targeted assay and can identify only indicated DNA sequences. Moreover, it identifies only deleted exons, but not the precise breakpoints [7]. Until now, there has been very little effort made to expand the screening for all FA genes at once and to define the precise molecular nature of particular deletions. FANCA deletion analysis by MLPA was previously proposed as an initial step in a comprehensive mutation screening strategy [8, 21, 35]. Moreover, 90% of all FA gene deletions occur in FANCA, and approximately half of them extend beyond the FANCA gene region, affecting the integrity of neighboring genes and influencing patient’s phenotype [7, 8, 35]. Thus, the larger insight in the phenotypic (including clinical) picture of the patients is needed.

Currently, aCGH can be considered as an integral component of a comprehensive strategy for identifying disease-causing variants in FA genes. Efforts should be made to correlate these variants with associated phenotypic changes. The identification of precise breakpoints possible by aCGH in contrast to MLPA, allows for quick screening of deletions in family members and provides insight into the deletion driving mechanisms [7, 8].

In our study, we present a family with compound alterations in FANCA gene.

In all children, a heterozygous deletion of chromosome region 16q24.3 was detected, including the entire FANCA gene. The same deletion was present in the mother. As this heterozygous deletion did not explain clinical features in patients IV.3 and IV.4, mutational analysis using next generation sequencing was performed, which identified the presence of a novel paternally inherited c.3337A>T; p.(Asn1113Tyr) pathogenic variant in exon 33 of FANCA gene accompanied by c.2669G>A; p.(Ser890Asn) variant, classified as VUS (variant of uncertain significance). To finally determine the meaning of each variant, it is necessary to perform functional tests.

This study has its limitations. Functional studies, together with protein modelling, were outside the initial scope of this study and its funding. These will, however, be performed in a follow-up project, and the results will be made available separately.

In the era of precision medicine, detailed diagnostic work-up and genetic testing are of great importance for accurate diagnosis, prognostic estimation, and personalized treatment, especially for rare and life-threatening disorders.

Conclusions

Our study adds to observations made by other authors, that the application of evolving new technologies can help mutation detection in genetically heterogeneous diseases become more economical, affordable, and efficient. Mutation identification in a FA patient, and establishing a carrier status in relatives, is an important part of the diagnostic and genetic counselling process for each FA family [7, 20, 25].

Genetic testing is a valuable tool for families with FA suspicion. There are several reasons for it. First, it confirms the diagnosis of FA among patients with inconclusive chromosome breakage tests or those with clinical features overlapping with other genetic syndrome features. Second, the identification of the family mutations could be useful for prenatal screening in future pregnancies. Third, heterozygous carriers could be identified within the family with FA and the genetic counseling could also be offered to these individuals.

In conclusion, application of NGS-based gene panel testing for all known FA genes in combination with chromosomal microarrays for FANCA or, as a new option—copy number variant (CNV) testing with NGS, provides a comprehensive approach for mutational FANCA screening, including identification of large deletions, and should be offered to all clinically diagnosed FA patients. It proves to be a valuable tool for clinical management, including reproductive counseling and prenatal diagnosis for whole families. Identification of novel FANCA mutations provides a better understanding of molecular mechanisms and further genotype–phenotype correlation searching of the condition.

Abbreviations

FA: Fanconi anemia; ICL: Interstrand Crosslinking Agents; DEB: Diepoxybutane; MMC: Mitomycin C; aCGH: Array-Comparative Genomic Hybridisation; micro-arrays; NGS: Next Generation Sequencing; MLPA: Multiplex Ligation-dependant Probe Amplification; PDA: Patent Ductus Arteriosus; CABS: Congenital Abnormality Score; allo-HSCT: Allogeneic Hematopoietic Stem Cell Transplant; MUD: Matched Unrelated Donor; GvHD: Graft-versus-Host Disease; CVN: Copy Number Variation; VUS: Variant of Uncertain Significance.

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Author contributions
AR and OH made a substantial contribution to the conception and design of the study, analysis, interpretation of data, and manuscript preparation. OH and JIW made a substantial contribution to data acquisition and supervised the study. JS-S made a substantial contribution to acquisition of clinical data. KJ, EL, ABK, JS-S, DL and KC made a substantial contribution to analysis and interpretation of data. MMK made a substantial contribution to analysis and interpretation of data and manuscript preparation. MD contributed to the interpretation of data and manuscript preparation. All the authors were involved in revising the manuscript critically for important intellectual content. All the authors read and accepted the final manuscript.

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Availability of data and materials
All data generated or analysed during this study are included in this published article.

Declarations

Ethics approval and consent to participate
The study was conducted in compliance with all ethical research standards. Ethical approval was provided by the Ethics Committee of the Nicolaus Copernicus University. Consent to participate was obtained from all participants of the study and their parents (in case of children under 16 years old).

Consent for publication
Signed informed consent was obtained from all participants (and parents of children) of this study for publishing the clinical data, including the photographs.

Competing interests
The authors declare that they have no competing interests.

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