Comprehensive analysis on phenotype and genetic basis of Chinese Fanconi anemia patients: dismal outcomes call for nationwide studies

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Abstract

Background Fanconi anemia (FA) is the most common inherited bone marrow failure (BMF) syndrome with 22 related genes identified. The ALDH2 rs671 variant has been proved related to accelerated progression of BMF in FA patients. The phenotype and genetic basis of Chinese FA patients have not been investigated yet.

Methods We analyzed the 22 FA-related genes of 63 BMF patients suspected to be FA. Clinical manifestations, morphological and cytogenetic feathers, ALDH2 genotypes, treatment, and outcomes of the definite cases were retrospectively studied.

Results 24 patients were confirmed the diagnosis of FA. The median age of BMF onset was 4.5-year old. The number of patients manifested as congenital malformations and growth retardation were 21/24 and 14/24, respectively. BM dysplasia and cytogenetic abnormalities were found in 15/23 and 10/22 patients. All the patients with abnormal karyotype also manifested as BM dysplasia or had evident blasts. Thirty-nine different variants were identified involving seven genes and including twenty-one novel variants. FANCA variants contributed to 58.33% of cases. Ten patients carried ALDH2-G/A genotype with a significantly younger age of BMF onset (p =0.024). Within the 22 patients adhering to continuous follow-up, 18 patients underwent hematopoietic stem cell transplantations (HSCTs). During the 33.5 months of follow-up, 8/22 patients died, seven of which were HSCT-related, and one patient who didn't receive HSCT died from severe infection.

Conclusion The phenotypic and genetic spectrum of Chinese FA patients is broad. Bone marrow dysplasia and cytogenetic abnormalities are prevalent and highly consistent. The overall outcome of HSCTs is disappointing. Nationwide multicenter studies are needed for the rarity and adverse outcome of this disease.

Introduction

Fanconi anemia (FA) is a rare genetic disease highly heterogeneous in clinical manifestations and genetics. Clinical manifestations primarily include congenital malformations, progressive bone marrow failure (BMF), and predisposition to hematopoietic and solid malignancies [1,2]. The most common congenital abnormalities include skin pigmentation, café au lait spots, short stature, and hypoplastic of radii and/or thumbs [2]. The time of BMF onset is variable but usually at pre-school age with the cumulative incidence of 90% by the age of 40 [3]. The malignancy risk in FA patients is mounting, especially the risks of myelodysplastic syndrome/acute myeloid leukemia (MDS/AML) and head and neck squamous cell carcinomas, which are several hundredfold higher than those of the general population [3–6].

Twenty-two genes (including three FA-like genes, e.g., RAD51C/FANCO, RAD51/FANCR, and BRCA1/FANCS, and the debatable FANCM) have been identified to be related to FA (Table S1), most of which are in autosomal recessive inheritance except FANCB and RAD51/FANCR, which are X-linked recessive and autosomal dominant, respectively. The 22 genes participate in the FA-BRCA pathway responsible for correcting interstrand crosslinks (ICLs) and other DNA damage events. Endogenous aldehyde is a genotoxic antigen and is detoxicated by aldehyde dehydrogenases (ALDHs) in vivo [7]. The mitochondrial ALDH2 isoform is the most efficient acetaldehyde-detoxifying enzyme in human [8]. Inactivating ALDH2 variant (rs671 G>A) is highly prevalent in East Asia and can abolish ALDH2 activity by a dominant negative effect [9]. ALDH2-A/A and ALDH2-G/A genotypes are related to accelerated progression of BMF and malignant transformation in FA patients [10].

Although the genetic basis, pathological mechanisms, and epidemiology of FA have been extensively studied, few researches focus on Chinese patients [11]. In the present study, we report 24 Chinese FA patients aiming to depict their genetic basis and clinical characteristics.

Subjects And Methods

Patient enrollment

We retrospectively analyzed 63 BMF patients who were suspected to be inherited BMF clinically in Hebei Yanda Lu Daopei Hospital from May 2012 to Dec. 2017. Detailed disease histories and examination files were retrieved from the electronic medical record system of our institute. All patients enrolled were confirmed BMF by morphology of BM aspiration and/or BM biopsy.
before chemotherapy or pre-hematopoietic stem cell transplantation (HSCT) conditioning regimen and should meet at least two following inclusive criteria: 1) growth retardation; 2) congenital physical malformations; 3) early onset of BMF (≤ 6 years old); 4) chronic onset of BMF with a progressive course (disease course > 6 months); 5) suggestive family history (consanguinity or family history of cancer or hematological disorders); 6) positive for chromosome breakage test. Other inherited syndromes manifested as BMF and malformations such as dyskeratosis congenita, Diamond-Blackfan anemia, and Neurofibromatosis-Noonan syndrome diagnosed based on syndromic presentations combined with genetic tests were excluded. The follow-up duration was defined as from the time of referral to the initiation of the study or from the time of referral to death.

Written informed consents were obtained from the patients or their statutory guardians and all tested family members in accordance with the Declaration of Helsinki. The study was approved by the ethics committee of the Hebei Yanda Lu Daopei hospital.

**Nucleic acid extraction**

Peripheral blood (PB), BM, or cryopreserved DNA samples of the patients and their parents were obtained. Genomic DNA was extracted from PB/BM nucleated cells using silica gel column method.

**High throughput sequencing, variant calling, and ALDH2 genotyping**

We carried out Sanger sequencing on the entire coding exons and flank regions of the three most common FA genes, FANCA, FANCC, and FANCG, in patients suspected to be inherited BMF from Apr. 2012 to May 2016. Targeted high-throughput sequencing (THS) has been applied since May 2016, and FANCD2 and BRCA2 were added in the panel. Whole genomic sequencing (WGS) was carried out using cryopreserved samples from patients who were highly suspicious of FA in clinic, and all the 22 FA and FA-like genes were analyzed (Table S1).

THS process has been described previously [12]. For the WGS, libraries were constructed with NEBNext Ultra II DNA Library Prep Kit for Illumina (New England Biolabs, US), followed by sequencing on Illumina HiSeq X Ten platform (Illumina, US) using HiSeq X Ten Reagent Kit v2.5 (Illumina, US) running on paired-end 150bp mode.

Reads yielded by the two kinds of sequencing were all aligned to the human reference genome (hg19) with the Burrow-Wheeler Aligner (BWA) mem. Variants were called according to Genome Analysis Toolkit (GATK) best practices using bam files. Final confident variants were annotated using annovar and oncotator. Variants with minimal allele frequency (MAF) ≥ 1% in general population were filtered out according to 1000 Genomes, EXAC, and gnomAD databases. The pathogenicity of the germline missense mutations was assessed by in silico prediction algorithms, and the putative causal variants were classified according to the standards and guidelines recommended by the American College of Medical Genetics and Genomics (ACMG) [13]. Only pathogenic, likely pathogenic or uncertain significance variants were considered causative in the present study. The process of CNV analysis based on WGS has been described elsewhere [12].

ALDH2 genotyping was based on WGS data or Sanger sequencing with primers 5’-TGCTATGATGTGTTTGGAGCC–3’ (forward) and 5’-ATTAGGGTCTCTGCTGGCCG–3’ (reverse).

**Validation by Sanger sequencing**

Polymerase chain reaction (PCR) and Sanger sequencing performed on the ABI 3500xL Genetic Analyzer (Thermo Fisher, US) were adopted to confirm all the variants reported in this study. Single nucleotide variants (SNVs) and small insertions/deletions (InDels) were validated by PCR and Sanger sequencing using the patients’ samples and their parents’ samples when accessible. For the validation of CNVs, the breakpoints were confirmed by Sanger sequencing using patients’ DNA, while the parental origins were only verified through PCR and agarose gel electrophoresis (AGE).
Results

Demography and clinical characteristics

A total of 24 patients (seven females and seventeen males) from non-related families were finally diagnosed as FA, including one who has already been reported (Case 9) [12]. The median referral age of this cohort was seven years old, and the median age of BMF onset was 4.5 years old (range, 1–15 years old). There were 23 Han Chinese and one Uyghur Chinese, and the geographical distribution spread nationwide though half of the patients came from the south or southwest of China. All patients were referred to our institute because of severe cytopenia except a thirty-year-old boy (Case 11) who was initially diagnosed as MDS for the myeloid dysplasia and increased myeloblasts indicated by BM smear. Five patients had an indicative family history, among which, two patients had family members died from anemia (Case 4, Case 17), two patients were from consanguineous families (Case 18, Case 24), and one patient was in vitro fertilized whose paternal grandmother died from pancreatic cancer (Table 1).

Fourteen (58.33%) patients were growth retarded, and 21 (85.7%) patients manifested as congenital malformations. Congenital abnormalities in our cohort included skin pigmentation (13/24), café au lait spots (6/24), spin and limbs deformation (12/24), craniofacial malformations (8/24), genitourinary system malformations (7/24), cardiovascular system defects (2/24), nervous system diseases (2/24), and endocrine system defects (2/24) (Table 1).

Thoroughly evaluation of the hematologic phenotype is crucial to FA patients since BM dysplasia or pathological cytogenetics relate to disease progression and adverse HSCT outcomes [5,14]. Twenty-three patients’ morphologic test results and 22 patients’ cytogenetics test results before pre-HSCT conditioning regimen and/or chemotherapy were available. BM dysplasia was found in 15/23 (65.22%) patients, including one AML with the myeloblast count of 41% (Case 6) and one myelodysplasia with the blast count of 6% (Case 11). Karyotypes were described according to the International System for Human Cytogenetic Nomenclature 2013 [15], at least 20 metaphases were analyzed for each assay. Cytogenetic abnormalities were found in 10/22 (45.45%) patients with clonality found in six patients, and half of the abnormal karyotypes involved chromosome 7 (–7, 7q-, or der(7)t(1;7)) (Table 2). The cytogenetic result of Case 6 who was diagnosed as AML was 46, XX, der(7)t(1;7)(q21;q36) [20], which was confirmed to be non-constitutional by matched peripheral blood, and the karyotype of patient Case 11 was highly complex (Table 2). All the patients with abnormal karyotypes also manifested as dysplasia on bone marrow smear or had evident blasts, suggesting the initiation of clonal evolution in the hematopoietic system.

Characteristics of variants

A total of 44 variants were identified involving seven different FA genes and composed by 16 missense mutations [16–19], ten large deletions [12], eight nonsense mutations [16], seven frameshift mutations [20–22], two splicing mutations, and one deep intron mutation [18] (Figure 1, Table 3). All the large deletions were found within the FANCA gene. The variant spectrum was broad with 21 (47.73%) novel variants identified in our cohort. The majority of variations were private except FANCA c.367C>T and FANCA c.3627–607_3765+268del which were shared by two patients separately. (Figure 2, Table 3). Furthermore, we did not find the most frequently occurred FANCA c.2546delC in our cohort, which accounts for over 30% FANCA mutations in Japanese and Korean patients [23,24].

Among the 24 patients, 17 patients carried compound heterozygous mutations, three patients carried homozygous mutations, three patients harbored monoallelic FANCB mutations who were all males, and one patient with a heterozygous FANCE mutation was identified. Biallelic FANCA variants caused 58.33% (14/24) of the cases, followed by monoallelic FANCB variants which constituted 12.5% (3/24) of the cases; both FANCD2 and FANCE mutations made up to 8.33% (2/24) of the cases, and FANCC, SLX4, and XRCC1 mutations caused one case each (Table 3). We did not find any case attributed to FANCG mutations, which is the second most prevalent responsible gene in eastern Asian in according to Japanese and Korean studies [23,24]. Despite the limited size of this cohort, we identified three FANCB variants, making it rank one of the most common causative genes in line with the Japanese study [23]. There were three homozygous mutations, FANCA c.1867C>T, FANCC c.545C>A, and XRCC2 c.1426G>A; the latter two mutations were carried by patients both came from consanguineous families, and the FANCC c.545C>A was carried by the only Uyghur patient in our cohort.
In the process of criminal variant identification, we adopted rigorous criteria and observed the canonical inheritance to the greatest extent. Majority of the patients were assigned with compelling variants with two exceptions. All variants were classified as pathogenic, likely pathogenic, or uncertain significance according to guidelines of ACMG but FANCD2 c.983G>A, which were harbored by Case 19. Albeit the ambiguous pathogenicity, it was rare in general population and was the only possible responsible variants found in this patient after the exhaustive search for variations in FA-related genes. Hence, we considered it to be deleterious and the causative mutation. Case 21 carried a heterozygous FANCE c.1111C>T mutation inherited from her father. This variant was regarded responsible for the patient’s disease for her characteristic clinical manifestations and no other variants identified in FA or FA-like genes while only FANCE c.1317–237C>G were found in the maternal allele with unclear clinical significance (not included in statistics) (Table 3).

**ADLH2 rs671 genotype**

14/24 (58.33%) patients in our cohort carried ALDH2-G/A genotype, and the other patients were all ALDH2-G/G genotype. There was no ALDH2-A/A genotype identified (Table 2). The age of BMF onset of ALDH2-G/A patients was significantly younger than that of the ALDH2-G/G patients ($p = 0.024$, t-test).

**Treatment and outcome**

Within the 24 patients, continuous medical records of 22 patients can be retrieved except Case 9 and Case 24, who only came to us once and were excluded in this section. All the 22 patients were eligible for HSCT for they were all transfusion-dependent, which was performed on 18 patients (81.81%). The numbers of patients accepted HSCT from HLA-matched unrelated donors (MUD), HLA-unmatched unrelated donors (UUD), HLA-haploidentical related (sibling or parental) donors (HRD), and HLA-matched related donors (MRD) were four, five, seven, and one, respectively. Another patient accepted HLA-unmatched unrelated cord blood (UUC) HSCT. The other four patients who did not undergo HSCT accepted androgen, cytokine, and/or intermittent transfusion support. All the patients with abnormal karyotype underwent HSCTs. In patients who underwent HSCTs, 11/18 (61.11%) were ALDH2-G/A genotype. The median follow-up duration was 33.5 months ranged from one month to 84 months. By the time of the study, eight patients (36.36%) have been dead. Seven of them were HSCT-related, mainly severe acute graft-versus-host disease (aGVHD) and infections, accounting for 38.89% HSCT patients. One patient did not receive HSCT died from severe infection (Table 3).

**Discussion**

The 24 patients displayed a wide range of clinical phenotype and genetic variation spectrum that all physiological systems besides hematopoietic system were involved (Table 1), and the responsible variants were detected in seven different genes (Figure 1, Table 3). In keeping with other studies, bone marrow dysplasia and abnormal karyotypes were prevailing (65.22% and 45.45%, respectively) and highly consistent [14,25], denoting the risk of hematologic malignant transformation, especially the ones with aberration in chromosome 7, which is the most prevalent cytogenetic abnormality in pediatric MDS and indicates an adverse long-term outcome even after HSCTs in MDS/AML patients [26]. ALDH2-G/A and ALDH-A/A genotypes are confirmed to be associated with more severe hematologic phenotype and more adverse outcomes of FA in Asian patients [10,14]. The same tendency was observed in our cohort, despite there was no patient carrying ALDH2-AA genotype.

All patients in our cohort presented with a more severe hematologic manifestation, and the proportion of patients who received HSCTs was higher than that of most studies [3–6,14,27]. Although BMF is the typical and most prevalent feature, our data may not reflect the actual behavior of FA since all the patients were referral to our institute seeking for HSCTs. Studies suggest the high HSCT-related mortality in FA patients, and of which infection and aGVHD were the two main causes [5,27]. In our cohort, 38.89% of HSCT patients died from HSCT-related acute complications. Furthermore, studies also suggest the overall dismal outcome that ten years cumulative risk of death was over 22% and the overall survival after 30 years of diagnosis dropped to below 40%; besides, the long-term survival of HSCT patients and non-HSCT patients were comparable [5,23,25,27], partly because the HSCT in the context of FA is specifically challenging. Therefore, even with the optimized pre-HSCT conditioning regimens that
is the reduced intensity and the introduction of fludarabine, meticulousness is needed in decision-making that whether HSCT is the most appropriate treatment strategy depends much on the severity of cytopenia and hematologic adverse events of a particular patient and the type of donor he/she could get.

The cumulative incidence of leukemia and solid tumors in the middle age of FA patients were reported to be ~20% and ~30%, respectively [4–6,28]. In our cohort, no patient developed hematologic or solid malignancies during the follow-up up to date except the ones initially diagnosed as AML (Case 6) and MDS (Case 11), but the longest duration of follow-up in our cohort was seven years, which may not be long enough for the malignant phenotype to emerge.

Although this study is limited by its cohort size, it is still informative and enriched the knowledge on Chinese FA patients which was nearly barren. By thoroughly investigating the clinical manifestations, morphologic and cytogenetic changes, genetic basis, and outcomes of 24 Chinese FA patients, this study displayed the broad phenotypic and genetic variant spectrum of Chinese FA patients and the current disappointing status of treatment including allogenic-HSCT which needs to be improved, and highlighted the urgency of nationwide multicenter studies for the rarity and adverse outcome of this entity so as to reveal the mask of Chinese FA patients and optimize the clinical managements.

Abbreviations

ACMG: American College of Medical Genetics and Genomics; AGE: agarose gel electrophoresis; aGVHD: acute graft-versus-host disease; ALDHs: aldehyde dehydrogenases; AML: acute myeloid leukemia; BMF: bone marrow failure; BWA: Burrow-Wheeler Aligner; CNVs: copy number variants; FA: Fanconi anemia; HLA: human leukocyte antigen; HRD: HLA-haploidentical related donors; HSCT: hematopoietic stem cell transplantations; ICLs: interstrand crosslinks; InDels: insertions/deletions; MAF: minimal allele frequency; MDS: myelodysplastic syndrome; MRD: HLA-matched related donors; MUD: HLA-matched unrelated donors; PB: peripheral blood; PCR: polymerase chain reaction; SNVs: single nucleotide variants; THS: targeted high-throughput sequencing; UUC: HLA-unmatched unrelated cord blood; UUD: HLA-unmatched unrelated donors; WGS: whole genomic sequencing

Declarations

Acknowledgments

The authors would like to thank the patients and their families for participating in the study.

Ethics approval and consent to participate

The study was approved by the ethics committee of the Hebei Yanda Lu Daopei hospital. The patients provided written informed consent for genetic analysis.

Consent for publication

Written informed consents were obtained from the patients or their statutory guardians and all tested family members for publication of clinical details.

Availability of data and materials

Sequencing data in fastq format generated in this study are available from the corresponding author on reasonable request.

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**Competing interests**

The authors declare that no competing interests in this study.

**Author's contributions**

DN reviewed medical history of the patients, analyzed sequencing data, and wrote the manuscript, JZ, FW, WZ, XM and LL performed the sequencing process, analyzed the data, and wrote the manuscript, XC and YZ analyzed the morphology and karyotype results, PC designed the bioinformatic analysis process. MX followed up the patients, TW and PW carried out the morphologic and cytogenetic study, WT, MW, and KC analyzed the clinical data and supervised the study, HL designed and supervised the study.

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### Tables

Table 1 Clinical features of the 24 FA patients

| Case No. | Gender | Age of referral (years) | Age of clinical BMF onset (years) | Congenital malformations | Growth retardation | Family history |
|----------|--------|------------------------|-------------------------------|-------------------------|------------------|----------------|
| 1        | F      | 16                     | 15                            | None                    | No               | Negative       |
| 2        | M      | 12                     | 2                             | S, C, M                 | Yes              | Negative       |
| 3        | M      | 7                      | 2                             | S, C, G                 | Yes              | Negative       |
| 4        | M      | 10                     | 5                             | S, G                    | Yes              | One sibling manifested as polydactyly and died from anemia |
| 5        | M      | 11                     | 5                             | C, G, M                 | Yes              | IVF and paternal grandmother died from pancreatic cancer |
| 6        | F      | 11                     | 10                            | C, H                    | Yes              | Negative       |
| 7        | M      | 7                      | 5                             | S, M                    | No               | Negative       |
| 8        | M      | 17                     | 10                            | S, M                    | No               | Negative       |
| 9        | M      | 7                      | 7                             | S, C, M, H              | No               | Negative       |
| 10       | M      | 5                      | 4                             | S, M                    | No               | Negative       |
| 11       | M      | 13                     | 13                            | E                       | Yes              | Negative       |
| 12       | M      | 9                      | 6                             | S                       | Yes              | Negative       |
| 13       | F      | 7                      | 7                             | M                       | Yes              | Negative       |
| 14       | M      | 7                      | 1                             | S, M                    | Yes              | Negative       |
| 15       | M      | 5                      | 2                             | S                       | No               | Negative       |
| 16       | M      | 4                      | 1                             | S, M                    | No               | Negative       |
| 17       | M      | 14                     | 5                             | C, G                    | Yes              | Two family members died from anemia |
| 18*      | M      | 6                      | 4                             | S, C, M                 | Yes              | 2nd degree consanguinity |
| 19       | M      | 4                      | 3                             | None                    | No               | Negative       |
| 20       | F      | 9                      | 4                             | S, M                    | No               | Negative       |
| 21       | F      | 7                      | 4                             | S, G, N                 | Yes              | Negative       |
| 22       | M      | 9                      | 7                             | None                    | No               | Negative       |
| 23       | F      | 6                      | 3                             | S, C, G                 | Yes              | Negative       |
| 24       | F      | 6                      | 4                             | S, M, H, E, G           | Yes              | 2nd degree consanguinity |

* Case 18 is a Uyghur Chinese.

F, female; M, male; S, skin and annex; C, craniofacial abnormalities; M, musculoskeletal system; G, genitourinary system; H, cardiovascular system; E, endocrine system; N, nervous system. IVF, in vitro fertilized.

Skin and annex abnormalities include skin pigmentation, café au lait spots, excess hair; craniofacial abnormalities include microcephalus, ptosis, hypertelorism, hypotelorism, flat nose bridge; malformations in musculoskeletal system include polydactyly, deformity of thumbs, absence of thumbs, hypoplasia of thenar eminence, and scoliosis; genitourinary system malformations include kidney malformation, hydronephrosis, indirect inguinal hernia, cryptorchidism, ovary absence, and uterine malformation/absence; cardiovascular system defects include patent ductus arteriosus and ventricular septal defect; nervous system abnormalities include encephalatrophy and moyamoya disease; endocrine system defects include hypothyroidism, primary adrenocortical insufficiency, and obesity.
Table 2 Bone marrow morphology, karyotype, chromosome breakage tests, and ALDH2 genotypes of the 24 FA patients.

| Case No. | BM morphology | BM karyotype | Chromosome breakage test | ALDH2 genotype |
|----------|---------------|--------------|--------------------------|----------------|
| 1        | Dysplasia     | 47,XX,+8[1]/46,XX[20] | Negative                | G/G            |
| 2        | Dysplasia     | NA           | Positive                 | G/A            |
| 3        | Dysplasia     | 47,XY,+15[1]/46,XY[20] | Positive                | G/A            |
| 4        | Hypoplasia    | normal       | Positive                 | G/A            |
| 5        | NA            | NA           | Positive                 | G/G            |
| 6*       | AML           | 46,XX,der(7)t(1;7)(q21;q36)[20] | Positive           | G/G            |
| 7        | Hypoplasia    | Normal       | Positive                 | G/A            |
| 8        | Dysplasia     | 46,XY,-7,+21[5]/46,XY[16] | Positive              | G/G            |
| 9        | Hypoplasia    | Normal       | Positive                 | G/G            |
| 10       | Hypoplasia    | Normal       | Positive                 | G/A            |
| 11**     | MDS           | Complex      | Positive                 | G/G            |
| 12       | Hypoplasia    | Normal       | Positive                 | G/A            |
| 13       | Dysplasia     | Normal       | Positive                 | G/G            |
| 14       | Dysplasia     | Normal       | Positive                 | G/G            |
| 15       | Dysplasia     | Normal       | Positive                 | G/G            |
| 16       | Hypoplasia    | Normal       | Positive                 | G/G            |
| 17       | Dysplasia     | 46,XY,del(7)(p13)[13]/46,XY[7] | Positive             | G/G            |
| 18       | Hypoplasia    | Normal       | Positive                 | G/A            |
| 19       | Dysplasia     | Normal       | Positive                 | G/G            |
| 20       | Dysplasia     | 46,XX,t(1;5)(p36.1;q13)[1]/46,XX[19] | Positive           | G/G            |
| 21       | Dysplasia     | 46,XX,del(14)(q24)[1]/46,XX[20] | Positive             | G/G            |
| 22       | Dysplasia     | 45,XY,-7[19]/46,XY[3] | Negative              | G/A            |
| 23       | Dysplasia     | 46,XX,del(7)(q22)[8]/46,XX,del(5)(p11)/46,XX[19] | Positive           | G/A            |
| 24       | Dysplasia     | Normal       | Positive                 | G/A            |

* Case 6 is diagnosed as acute myeloid leukemia. The myeloblasts count 41% of the nucleated cells according to morphologic test of bone marrow smears.

** Case 11 is diagnosed as myelodysplastic syndrome. His bone marrow morphology shows dysplasia was observed in his granulocytic lineage and megakaryocytic lineage with the myeloblasts count 6% of the nucleated cells. The result of his karyotype is: 46,XY,dup(1)(q21q23),add(2)(p11.2),add(3)(q27),der(5)t(1;5)(q21;q35),add(20)(p12)[17]/45,XY,der(1)(7::1q42->1q21::1p36.3->1q32::1q21->1q44::?),add(2)(p11.2),add(3)(q27),add(4)(p16),der(5)t(1;5)(q21;q35),-18,add(20)(p12),ace[2]/46,XY[1]

Chromosome breakage tests were induced by mitomycin C.

NA, not available; AML, acute myeloid leukemia; MDS, myelodysplastic syndrome.

Table 3 Variant details
| Case No. | Gene | Genomic location | Mutation 1 (maternal) | Ref./Com. | Genomic location | cDNA/Protein | Mutation 2 (paternal) | Ref./Com. |
|---------|------|------------------|----------------------|-----------|------------------|-------------|----------------------|-----------|
| 1       | FANCA | chr16:89831465   | c.2611C>G/p.L871V;c. | NA        | chr16:8980940-   | c.3627-607_3765+268del | 12        |
|         |      |                  |                      |           | 89809954         |             |                      |           |
| 2       | FANCA | chr16:89813593   | c.2557C>T/p.R853X    | 16        | chr16:89877396   | c.367C>T/p.Q123X | NA        |
| 3       | FANCA | chr16:89815145-  | c.3270_3271delCT/p.C1090RfsX25 | Novel    | chr16:89868906- | c.792+761_c.523-635del | Novel    |
|         | 89815146 |                  |                      |           | 89875410         |             |                      |           |
| 4       | FANCA | chr16:89877396   | c.367C>T/p.Q123X    | NA        | chr16:89818822   | c.2982-192A>G | 18        |
| 5*      | FANCA | chr16:89842183   | c.1867C>T/p.Q623X   | Novel     | chr16:89842183   | c.1867C>T/p.Q623X | Novel    |
| 6       | FANCA | chr16:89804935-  | c.3935-178_4368+74del | Novel    | chr16:89819567- | c.3627-607_3765+268del | Novel    |
|         | 89806139 |                  |                      |           | 89839134         |             |                      |           |
| 7       | FANCA | chr16:89811185-  | c.3239+397_3626+202del | Novel    | chr16:89858887   | c.1074_1075delGT/p.Y359PfsX49 | NA        |
|         | 89815741 |                  |                      |           | 12               |             |                      |           |
| 8       | FANCA | chr16:89826812-  | FANCA c.2852+1545_SPRE2 c.646-1671del | Novel    | chr16:89825071   | c.2894_2895delCT/p.P965RfsX9 | Novel    |
|         | 89919023 |                  |                      |           | 12               |             |                      |           |
| 9       | FANCA | chr16:89780001-  | VPS9D1 c.432-877_FANCA | 12        | chr16:89808940- | c.3627-607_3765+268del | 12        |
|         | 89822000 |                  | c.2981+2985del       |           | 89809954         |             |                      |           |
| 10      | FANCA | chr16:89823177-  | c.2853-333_2981+1808del | Novel    | chr16:89809270   | c.3703C>T/p.Q1235X | Novel    |
|         | 89825446 |                  |                      |           | 19               |             |                      |           |
| 11      | FANCA | chr16:89818619   | c.2990_2993delGTATA/p.S997MfsX28 | NA        | chr16:89862229   | c.987_990delTCAC/p.H330Af3X4 | 19,22     |
| 12      | FANCA | chr16:89816286   | c.3091C>T/p.Q1031X   | NA        | chr16:89792569- | c.3628C>G/p.R880G | NA        |
|         |                  |                  |                      |           | 89821767         |             |                      |           |
| 13      | FANCA | chr16:89806417   | c.3918dupT/p.Q1307SfsX6 | 20        | chr16:89831438   | c.2638C>G/p.R880G | NA        |
| 14      | FANCA | chr16:89858941   | c.1021C>T/p.Q341X    | Novel     | chr16:89811412   | c.3581C>T/p.P194L | 20        |
| 15      | FANCA | chr14:8466851    | c.1472T>A/p.V491E    | Novel     | —                 | —           | —                    | —         |
| 16      | FANCA | chr14:8479579    | c.1197+1insA         | Novel     | —                 | —           | —                    | —         |
| 17      | FANCA | chr14:8477390    | c.1018C>A/p.Q340K    | Novel     | —                 | —           | —                    | —         |
| 18*     | FANCC | chr9:97912346    | c.545C>A/p.S182Y     | Novel     | chr9:97912346    | c.545C>A/p.S182Y | Novel    |
| 19      | FANCD2 | chr3:10084828    | c.983G>A/p.R328Q     | NA        | chr3:1014634     | c.2574T>G/p.I858M | Novel    |
| 20      | FANCD2 | chr3:10132005    | c.3713T>A/p.M1238K   | NA        | chr3:10089999    | c.1279-2A>T | Novel    |
| 21      | FANCE | —                | —                    | NA        | chr6:35426215    | c.1111C>T/p.R371W | 17,18,19 |
| 22      | FANCE | chr6:35423547    | c.272C>T/p.S91L      | Novel     | chr6:35426215    | c.1111C>T/p.R371W | 17,18,19 |
| 23      | SLX4  | chr16:3645671    | c.1948C>T/p.L505F    | NA        | chr16:3633419    | c.4832A>G/p.E1611G | NA        |
| 24*     | ERCC4 | chr16:14015937   | c.257G>A/p.R86H      | NA        | chr16:14015937   | c.257G>A/p.R86H | NA        |

* Case 5, Case 18, and Case 24 carries homozygous variants.

NA, not available.

Table 4 Treatment and outcomes of the 24 FA patients.
| Case No. | Therapeutics | Donor type | Pre-HSCT conditioning regimen | Outcomes |
|----------|--------------|------------|-------------------------------|----------|
| 1        | HSCT         | UUD        | Bu+CTX+Flu+Alemtuzumab        | Alive    |
| 2        | HSCT         | UUD        | Bu+CTX+Flu+Alemtuzumab        | Dead (aGVHD, infections) |
| 3        | HSCT         | UUD        | Bu+CTX+Flu+ATG+Me-CCNU       | Alive    |
| 4        | HSCT         | MUD        | Bu+Flu+CTX+ATG               | Alive    |
| 5        | HSCT         | HRD        | Bu+CTX+Flu+ATG+Me-CCNU       | Alive    |
| 6        | HSCT         | HRD        | Decitabine+Ara-C+Bu+Flu+ATG+Me-CCNU | Dead (aGVHD, drug-induced encephalopathy) |
| 7        | Androgen and transfusion | — | — | Alive |
| 8        | HSCT         | HRD        | Decitabine+Ara-C+Bu+Flu+ATG+Me-CCNU | Dead (aGVHD, MODS) |
| 9        | Lost         | — | — | — |
| 10       | HSCT         | MRD        | Bu+Flu+CTX+ATG               | Dead (aGVHD, septic shock) |
| 11       | HSCT         | HRD        | Decitabine+Ara-C+Bu+Flu+ATG+Me-CCNU | Dead (aGVHD, septic shock) |
| 12       | HSCT         | UUC        | Bu+Flu+CTX+ATG               | Dead (aGVHD, pulmonary infection, CMV infection) |
| 13       | Androgen, cytokine, transfusion | — | — | Alive |
| 14       | Androgen and cytokine | — | — | Alive |
| 15       | HSCT         | HRD        | Bu+Flu+CTX+ATG               | Dead (aGVHD, TMA, pulmonary infection) |
| 16       | HSCT         | MUD        | Bu+Flu+CTX+ATG               | Alive    |
| 17       | HSCT         | HRD        | Bu+Flu+CTX+ATG               | Alive    |
| 18       | Androgen and transfusion | — | — | Dead (pulmonary infection, septic shock) |
| 19       | HSCT         | MUD        | Bu+Flu+CTX+ATG               | Alive    |
| 20       | HSCT         | UUD        | Bu+Flu+CTX+ATG               | Alive    |
| 21       | HSCT         | UUD        | TBI+CTX+Flu+ATG              | Alive    |
| 22       | HSCT         | HRD        | Bu+Flu+CTX+ATG               | Alive    |
| 23       | HSCT         | MUD        | Bu+Flu+CTX+ATG               | Alive    |
| 24       | Lost         | — | — | — |

HSCT, hematologic stem cell transplantation; UUD, HLA-unmatched unrelated donor; MUD, HLA-matched unrelated donor; HRD, HLA-haploidentical related donor; MRD, HLA-matched related donors; UUC, HLA-unmatched unrelated cord blood; Bu, Busulfan; CTX, cyclophosphamide; Flu, Fludarabine; ATG, antithymocyte globulin; Me-CCNU Semustine; TBI, total body irradiation; aGVHD, acute graft-versus-host disease; CMV, cytomegalovirus.

Figures
Figure 1

Variant distribution and composition of the 44 variants. a. variant distribution in our cohort. b. variant composition in our cohort.

Figure 2

Locations, frequencies, and types of variants in FANCA, FANCB, FANCC, FANCD2, FANCE, SLX4, and ERCC4 genes. The colored rectangles represent exons, different types of the variants are represented by different patterns with different colors denoted on the top of each gene except large deletions which are represented by the black horizontal bars denoted under FANCA gene.