A never-ending story: the steadily growing family of the FA and FA-like genes

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Abstract

Among the chromosome fragility-associated human syndromes that present cancer predisposition, Fanconi anemia (FA) is unique due to its large genetic heterogeneity. To date, mutations in 21 genes have been associated with an FA or an FA-like clinical and cellular phenotype, whose hallmarks are bone marrow failure, predisposition to acute myeloid leukemia and a cellular and chromosomal hypersensitivity to DNA crosslinking agents exposure. The goal of this review is to trace the history of the identification of FA genes, a history that started in the eighties and is not yet over, as indicated by the cloning of a twenty-first FA gene in 2016.

Keywords: DNA repair, leukemia, Fanconi anemia, chromosomal abnormalities.

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Introduction

Fanconi anemia (FA) is a rare human genetic syndrome associated with bone marrow failure (BMF), myelodysplasia (MDS) and a predisposition to acute myeloid leukemia (AML) and head and neck cancer. FA was described in 1927 by the Swiss pediatrician Giuseppe Fanconi, who reported a family with three affected siblings exhibiting anemia and developmental defects (Lobitz and Velleuer, 2006).

The clinical phenotype of FA patients is extremely heterogeneous. Beyond their hematological problems, which constitute the major hallmark of the disease, approximately 70% of these patients present developmental abnormalities, including abnormal radius, absent or supernumerary thumbs, microcephaly, microphthalmia, slow growth rate, café-au-lait spots, skin hyper- and hypo-pigmentation, kidney and urogenital defects, and hypoplasia of the testes. The estimated frequency of the syndrome is 1 in 250,000 - 350,000 live births, with a carrier frequency of approximately 1 in 200 (Rosenberg et al., 2011; Fanconi Anemia Research Fund Inc, 2014).

During the seventies, several groups around the world contributed to the definition of the two major cellular characteristics of the pathology: its particular chromosome fragility and its hypersensitivity to DNA interstrand crosslink (ICL)-inducing agents (Fujiwara and Tatsumi, 1975; Latt et al., 1975; Fornace et al., 1979; Novotna et al., 1979; Ishida and Buchwald, 1982). Indeed, FA cells appear exquisitely sensitive at both the cellular (survival) and chromosomal levels to the exposure to chemicals such as mitomycin C (MMC), diepoxybutane, cis-platinum and photoactivated psoralens. Since it is difficult to distinguish FA patients from individuals suffering from other inherited or idiopathic BMF syndromes on their clinical characteristics alone, the diagnosis of FA is based on the chromosomal response to ICL-inducing agents. Indeed, cytogeneticists score both the basal and induced frequency of chromosome aberrations as well as their subtypes, i.e., tri- and quadriradials, whose presence is quite specific for FA cells (Pinto et al., 2009; Fanconi Anemia Research Fund Inc, 2014).

Based on both the chromosome fragility and the hypersensitivity to exposure to DNA damaging agents, it was quickly suspected that the proteins whose loss of function caused FA must be involved in the DNA damage response and, more specifically, in a DNA repair mechanism. Indeed, although alternative functions associated with each individual or subgroup of FANC proteins exist (Joenje et al., 1981; Rosselli et al., 1994; Fagerlie et al., 2001; Pang et al., 2001; Briot et al., 2008; Pagano et al., 2003, 2012; Zanier et al., 2004; Myers et al., 2011; Justo et al., 2014; Parodi et al., 2015; Sumpter et al., 2016), the well-established “canonical” function of the proteins is to work along a “linear” pathway that addresses replication stresses, assuring the transmission of a stable genome from one cell to the daughters and acting both during DNA replication to cope with stalled replication forks and in G2 and M phases.
to resolve underreplicated regions before cell division (Ceccaldi et al., 2016; Lopez-Martinez et al., 2016; Michl et al., 2016). How the other, noncanonical functions of the FANC proteins that are involved in cytokine production/response, inflammation (Rosselli et al., 1994; Fagerlie et al., 2001; Pang et al., 2001; Zanier et al., 2004; Briot et al., 2008), mitophagy Sumpter et al., 2016), and oxygen free radical metabolism (Joenje et al., 1981; Pagano et al., 2003; Pagano et al., 2012) as well as the subtle defects in immunity (Myers et al., 2011; Justo et al., 2014; Nguyen et al., 2014; Parodi et al., 2015) impact the clinical and cellular phenotypes of the patients remains a challenge for the future understanding of the pathology.

The 21 currently identified FANC and FANC-like genes (Table 1) encode proteins assembled into three biochemically and functionally defined groups (Wang, 2007). The first group contains several FA and FA-associated proteins (some named FAAPs) that coinmunoprecipitate in the same supramolecular complex. This group is formally called “FANCcore complex” and exhibits the E3 ubiquitin ligase activity responsible for the monoubiquitination of two downstream FANC proteins, FANCD2 and FANCI, which constitute group II. The third group consists of proteins directly involved in DNA metabolism, including structure-specific endonucleases (XP-F and SLX4) and several proteins involved in homologous recombination. Therefore, cells that are defective in genes coding for proteins from group III have normal levels of FANCD2 and FANCI monoubiquitination. The majority of patients (not less than 85%) harbor mutations in genes encoding proteins of the first group. Figure 1 schematically demonstrates the subcellular localization and the assembling pattern of the FANC proteins inside the nucleus in the presence of DNA damage and replication stress, where some of them form subnuclear foci, which can be observed by immunofluorescence microscopic analysis. Briefly, the strongest evidence in the literature supports the presence of three main FANCcore subcomplexes in the cytoplasm and/or in the nucleus, representing triads of proteins - FANCA, FANCG and FAAP20; FANCC, FANCE and FANCF; and FANCB, FANCL and FAAP100. In the nucleus, they assemble into a unique complex on the “cargo” FANCm, which is also associated with the FAAP24 and MIF 1/2 proteins as well as the Bloom-Associated Proteins (BLAPs) (Meetei et al., 2003b; Guo et al., 2009), which are prevented from sliding onto DNA by lesions or a stalled fork. Following an interaction with UBE2T, the complex locally monoubiquitinates FANCD2 and FANCI, a process that requires ATM- and/or ATR- and CHK1-mediated phosphorylation events on both the FANCcore complex proteins and FANCD2 and

| Complementation group | Estimated frequency | Gene name | Gene alias | Chromosomal position | Protein M.W. (kDa) | Core component | Cloning date | Bona fide FANC gene |
|-----------------------|---------------------|-----------|-----------|----------------------|-------------------|----------------|-------------|-------------------|
| A                     | 60-70               | FANCA     |           | 16q24.3              | 162.7             | Yes            | 1996        | Yes               |
| B                     | rare                | FANCB     | FAAP95    | Xp22.2               | 97.7              | Yes            | 2004        | Yes               |
| C                     | 10-15               | FANCC     | BRCA2     | 9q22.3               | 63.4              | Yes            | 1992        | Yes               |
| D1                    | 1-5                 | FANCD1    | BRCA2     | 3q12.3               | 384.2             | Yes            | 2002        |                  |
| D2                    | 1-5                 | FANCD2    | BRCA2     | 3p25.3               | 164.1             | Yes            | 2001        | Yes               |
| E                     | rare                | FANC      |           | 6p21.3               | 58.7              | Yes            | 2000        | Yes               |
| F                     | rare                | FANCF     |           | 11p15                | 42.2              | Yes            | 2000        | Yes               |
| G                     | 10-15               | FANCG     | XRCC9     | 9p13.3               | 68.5              | Yes            | 1998        | Yes               |
| I                     | rare                | FANCI     |           | 15q26.1              | 149.3             | Yes            | 2007        | Yes               |
| J                     | rare                | FANCJ     | BACH1; BRIP1| 17q22.2             | 140.9             | ?              | 2005        |                   |
| L                     | rare                | FANCL     |           | 2p16.1               | 42.9              | Yes            | 2003        | Yes               |
| M                     | rare                | FANCM     |           | 14q21.2              | 232.2             | Yes            | 2005        |                   |
| N                     | rare                | FANCN     | PALB2     | 16p12.12             | 131.3             | 2007           |            |                   |
| O                     | rare                | FANCO     | RAD51C    | 17q22                | 42.2              | 2001           |            |                   |
| P                     | rare                | FANCP     | SLX4      | 16p13.3              | 200               | Yes            | 2011        |                   |
| Q                     | rare                | FANCQ     | ERCC4; XPF | 16p13.12             | 104.5             | Yes            | 2013        |                   |
| R                     | rare                | FANC     | RAD51     | 15q15.1              | 37                | 2015           |            |                   |
| S                     | rare                | FANCS     | BRCA1     | 17q21                | 207.7             | 2015           |            |                   |
| T                     | rare                | FANCT     | UBE2T     | 1q32.1               | 22.5              | Yes            | 2016        |                   |
| U                     | rare                | FANCU     | XRCC2     | 7q36.1               | 31.9              | 2016           |            |                   |
| V                     | rare                | FANCY     | REV7      | 1p36.22              | 24.3              | 2016           |            | Yes               |
FANCI. The FANCcore complex monoubiquitinates FANCD2 and FANCI on lysine 561 and 523, respectively. Together, monoubiquitinated FANCD2 and FANCI represent the central link between the upstream FANC proteins (group I or the FANCcore complex) devoid of direct “DNA repair function” and the downstream proteins of group III that are involved in DNA metabolism. Indeed, following their monoubiquitination, FANCD2 and FANCI assemble in chromatin-associated foci, where they colocalize with several other proteins involved in homologous recombination, including other FANC and FANC-like representatives. The USP1:UAF1 dimer deubiquitinates both FANCD2 and FANCI.

It is important to note that in recent years, the criteria to be considered as a “bona fide FA gene” have become more stringent and are now based on the clinical phenotype. Indeed, whereas the loss of function of all the identified genes leads to the primary FA cellular characteristics, including the ICL hypersensitivity and chromosome fragility, the clinical traits of some patients fail to reach the canonical features of the FA syndrome, namely BMF and the MDS. The genes mutated in those patients are now excluded from the “bona fide FA gene” group and are considered as “FA-like genes” (Bofliolo and Surralles, 2015). Nevertheless, the number of bona fide FANC and FANC-like genes continues to grow and it is unlikely to stop anytime soon.

**History of the identification of FANC genes**

**One disease, many genes.**

The existence of genetic heterogeneity in FA was demonstrated at the beginning of the eighties by the pioneering work of the groups of Manuel Buchwald and Karl Sperling, which used a cell fusion approach (Zakrzewski and Sperling, 1980; Duckworth-Rysiecki et al., 1985). In particular, Duckworth-Risiecky and collaborators reported the existence of two FA complementation groups: A and B.
non-A. The complementation analysis was based on the rescue of the cellular and chromosomal sensitivity to MMC exposure in the hybrid cells compared to the two cell lines fused to obtain the hybrid (Duckworth-Rysiecki et al., 1985).

The nineties.

Seven years later, in 1992, the Buchwald group published two seminal manuscripts, with the first recognizing that the non-A group was heterogeneous, assembling three complementation groups: B, C and D (Strathdee et al., 1992a). The second manuscript reported the cloning of the first FANC gene, FANCC (Strathdee et al., 1992b). To identify the mutated gene, Strathdee and collaborators followed a functional complementation strategy. They transfected an EBV-based cDNA expression library into the HSC-536 B-lymphoblastoid cell line (previously assigned to the FA-C complementation group) and isolated three overlapping cDNA sequences able to independently complement the huge cellular and chromosomal hypersensitivity to MMC of the transfected cells. The proband carried a mutation of maternal origin that changed leucine 544 to a proline (L544P), a modification predicted to disrupt an alpha helical secondary structure of the protein (Strathdee et al., 1992b). Initially elusive, the mutation affecting the allele of paternal origin, a deletion of 327 bp resulting in the removal of exons 1 and 2, was identified some years later (Parker et al., 1998).

A new complementation group was added to the list in 1995. By analyzing 13 unrelated FA patients, Joenje et al. (1995) identified a fifth FA complementation group, named FA-E. The following year, two groups cloned FANCA, the most frequently mutated FANC gene, using two alternative strategies. One group identified the gene by the same functional complementation approach as used by the Buchwald group (Lo Ten Foe et al., 1996). Alternatively a consortium of several laboratories working to identify breast cancer susceptibility genes adopted the chromosome walking strategy (Fanconi anaemia/Breast Cancer, 1996) after the localization of a putative FANCA gene in the q24.3 region of the chromosome 16 (Pronk et al., 1995).

In 1997, the number of complementation groups established by the original cell fusion approach grew again, totaling 8: A to G (Joenje et al., 1997). In 1998, de Winter et al. (1998) used functional complementation to isolate a cDNA able to rescue the MMC hypersensitivity of a standard FA-G cell line. The identified sequence, renamed FANCG, was similar to that of a gene cloned one year before on the basis of its capability to complement the MMC hypersensitivity of a CHO UV40 mutant clone called XRCC9 (Liu et al., 1997).
The 2000s.

The third millennium opened with the cloning of two new FANC genes. Still using the functional complementation cloning approach, de Winter and collaborators identified the genes whose loss of function was associated with the FA complementation groups E and F, i.e., *FANCE*, (de Winter et al., 2000a), previously mapped on chromosome 6 (Waisfisz et al., 1999), and *FANCF* (de Winter et al., 2000b).

In 2001, Timmers and collaborators reported that the cell lines originally categorized in the FA complementation group D (FA-D) could be separated into two groups, named D1 and D2, and they identified the gene mutated in the D2 group, *FANCD2*, by positional cloning and chromosome transfer, owing to the previous identification of the gene on chromosome 3p (Whitney et al., 1995; Hejna et al., 2000; Timmers et al., 2001).

The next year, D’Andrea and colleagues decided to test the hypothesis that the inactivation of *BRCA1* and *BRCA2*, the most famous and frequently mutated genes in familial predisposition to breast cancer and whose loss of function results in a cellular phenotype similar to that described for FA, could also be involved in FA. A systematic sequencing of *BRCA1* and *BRCA2* was performed in several FA cell lines that belonged to complementation groups without an assigned gene. This “target gene” approach allowed the identification of biallelic variations in *BRCA2* in the FA-D1 standard cell line HSC-62. The variants were successively validated as inactivating mutations (Howlett et al., 2002), assigning *BRCA2* to the list of FANC genes as *FANCD1*. Howlett and collaborators also identified some variants of *BRCA1* by examining the HSC230 cell line, the FA-B standard cell line. However, these variants failed to be confirmed as bona fide inactivating mutations.

In 2003, a big step forward in the genetics of FA was achieved thanks to the work of Meetee et al. (2003b) in the Weidong Wang laboratory. They purified a BLM-associated supramolecular complex containing two salt-concentration-dependent separable groups of proteins as follows: the BLM-associated proteins (BLMAPs) and Fanconi anemia-associated polypeptides (FAAPs). Mass spectrometry analysis of the isolated FAAPs identified some of the known FANC proteins (FANCA, FANCC, FANCE, FANCF, and FANCN), and the unknown components were identified as FAAP43, 90/95, 100, and 250/300 on the basis of their molecular mass. The biochemical approach of the Wang group allowed the cloning of new FANC genes by a “protein to gene” walking route. Indeed, after the identification of the amino acid composition of the FAAPs, the authors were able to match this to the sequence of each corresponding gene, to look for mutations in cells from FA patients and/or to determine the gene function(s) by analyzing the phenotypic consequences of the engineered inactivation of these genes in model cells or mice. Moreover, the work of Meetee and collaborators also provided the first indication that at least some of the FANC proteins work together inside a molecular complex.

The same year, Meetee et al. (2003a) identified FAAP43 as the PHD finger protein 9 coding gene, or *PHF9*. It was known that the inactivation of the mouse homolog of *PHF9*, *Pog* (for proliferation of germ cells), resulted in infertility and the MMC hypersensitivity of bone marrow cells, the two more consistent features presented by the already obtained FA mouse models. To definitively validate that PHF9 belonged to the FANC gene family, Meetee and coworkers identified inactivating PHF9 mutations in a cell line, EUFA868, isolated from a patient not previously assigned to a complementation group. The EUFA868 cell line used to clone *PHF9* was assigned to FA-L and *PHF9* was also named *FANCL* (Meetee et al., 2003a). *PHF9/FANCL* encodes the ubiquitin E3 ligase of the FANC core complex, which mediates the FANCD2 and FANCI monoubiquitination (Meetee et al., 2004b). A second FA patient bearing a mutation in *FANCL* was identified six years later (Ali et al., 2009).

In 2004, Meetee et al. (2004a) identified the coding sequence of FAAP95 as being similar to the one named FLJ34064, a sequence localized on the X chromosome. Mutations in FLJ34064 were found in several FA cell lines, including the HSC-230 cell line, the standard for FA complementation group B. Thus, FAAP95 was renamed *FANCB*. *FANCB* is the only known FANC gene localized on the X chromosome. Consequently, its inactivation affects only males. *FANCB* is silenced via the methylation of its promoter on the X chromosome that undergoes inactivation during embryogenesis. Since the X-inactivation is stochastic, i.e., it affects either the paternal or the maternal X chromosome randomly, it is expected that the expression of a gene subjected to inactivation will exhibit mosaicism. In the case of FANCB+/- female carriers, the large majority of the lymphocytes and fibroblasts of the body express the WT gene, suggesting that the cells that express the mutated gene are counterselected, probably due to their growth difficulties, and are rapidly lost (Meetee et al., 2004a).

In 2005, Meetee et al. (2005) identified FAAP250 as KIAA1596, a human protein with sequence similarities to DNA repair proteins, including the yeast MPH1 and the human ERCC4/XPF. siRNA-mediated depletion of KIAA1596 in cellular models affected FANCD2 monoubiquitination and increased MMC sensitivity, arguing for the assignment of FAAP250 to the FANC gene family. Mutations were then identified in an FA patient who was not assigned to a known complementation group. The gene was called *FANCM*, and the new complementation group FA-M. Nevertheless, the attempts to complement the MMC hypersensitivity of cell lines derived from the patient by transfection of the wild-type *FANCM* cDNA failed. Surprisingly, it was demonstrated that he also carried biallelic mutations in *FANCA*. Therefore, even if the loss of function of FANCM by targeted mutagenesis in mice or by siRNA-
mediated depletion in human cells results in an FA-like cellular phenotype and in spite of its interaction with several other FANC proteins, the lack of patients with the major features of FA and only the loss of FANCM function impedes the assignment of the protein to the group of the "bona fide" FANC genes (Meetei et al., 2005; Singh et al., 2009).

In the same year, 2005, three groups identified the gene mutated in FA-J cells and named it FANCJ (Levitus et al., 2005; Levrán et al., 2005; Litman et al., 2005). Following unsuccessful attempts to identify the mutated gene by a complementation cloning strategy, Levitus et al. (2005) opted for a positional cloning strategy and identified in eight FA-J cell lines several pathogenic mutations in the gene encoding the DEAH-box DNA helicase and binding partner of BRCA1 BRIP1/BACH1 (BRCA1-Immunoprecipitated Protein 1/BRCA1-associated C-terminal helicase-1), which was previously cloned by Cantor et al. (2001). Using a genome-wide scan, Levrán et al. (2005) identified a homozygous region on chromosome 17q23 in which there were two interesting candidate genes: RAD51C and BRIP1. Inactivating mutations were found only in BRIP1/BACH1. Finally, Litman et al. (2005) identified BRCA1/BACH1 mutations in two families that were associated with an early onset of breast cancer and found the same recurrent nonsense mutation, the R798X mutation in exon 17, in both the breast cancer and FA-J families. This mutation affected the helicase domain of the protein, and since it was identified in people of different ethnic origins, it likely represents a hot spot of mutation or an inactivating event that remains compatible with survival (Levitus et al., 2005). Thus, BRIP1/BACH1 is also called FANCJ.

In 2007, following alternative approaches, three independent groups cloned the 13th FANC gene, FANCJ, which is the paralog of FANCN2 (Dorsman et al., 2007; Meijer, 2007; Sims et al., 2007; Smogorzewska et al., 2007). FANCJ was identified by a linkage analysis approach (Dorsman et al., 2007), by a bioinformatical screening for FANCD2 homologs (Sims et al., 2007) and by a proteomic search for ATM and ATR targets (Smogorzewska et al., 2007). It was described as the gene mutated in cells belonging to FA complementation group I.

Also in 2007, the 14th FANC gene was cloned and assigned to FA-N, a previously unrecognized FA complementation group (Reid et al., 2007; Xia et al., 2007). The identified gene, FANCN, was known to encode PALB2, isolated by immunoprecipitation one year prior as the Partner and Localizer of BRCA2 (Xia et al., 2006). Reid et al. (2007) followed a candidate gene approach, sequencing 82 FA patients with unelucidated genetic causes and identified mutations inactivating PALB2 in 7 individuals belonging to independent families. By Western blot analysis, Xia and collaborators noticed the lack of a full-length PALB2 in an unassigned FA cell line. Subsequent DNA sequencing allowed the identification of the inactivating mutations in PALB2 (Xia et al., 2007).

In 2010, aiming to identify the gene responsible for the pathology in a Pakistani family with FA by a genome-wide mapping approach, Vaz et al. (2010) identified a homozygous mutation in the RAD51C gene. Successively, in vitro functional studies showed that the identified mutation resulted in the loss of RAD51 focus formation in response to DNA damage, a defect that could be rescued by the ectopic expression of wild-type RAD51C. On this basis, the authors proposed to assign the acronym FANCO to RAD51C. RAD51C is also recognized as a gene associated with breast and ovarian cancer predisposition (Somayaji et al., 2012).

In 2011, several groups focused their work on SLX4, a gene previously identified in yeast and flies as well as in humans and involved in the cellular response to DNA ICLs (Mullen et al., 2001; Wu et al., 2004; Lee et al., 2005; Fekairi et al., 2009; Munoz et al., 2009; Svendsen et al., 2009). Kim et al. (2011) and Stoepker et al. (2011) decided to sequence the SLX4 gene in several patients with an FA-like phenotype, who until that time had not been assigned to any of the sixteen known complementation FA groups. They successfully identified some patients with mutations in the SLX4 coding sequence. On the other hand, Crossan et al. (2011) found that Slx4-null mice recapitulated the features of FA. Thus, SLX4 was also named FANCP. It codes for a structure-specific endonuclease that can be found in a complex with XP-F/ERCC1 and MUS81/EME1, proteins involved in protecting the genome during S and M phases.

Surprisingly, in 2013, inactivating mutations in ERCC4/XP-F, whose loss of function was previously associated with the skin cancer predisposition syndrome Xeroderma pigmentosum complementation group F, were also identified to be associated with an FA-like phenotype by whole-exome and Sanger sequencing of the DNA of unclassified FA individuals (Bogliolo et al., 2013). ERCC4/XP-F was thus renamed FANCQ. Analyses of the consequences of the identified mutations clearly demonstrated that compared to the NER-associated mutations, these mutations altered different regions, affecting an alternative function of the protein (Bogliolo et al., 2013). The association of XP-F mutations with an FA phenotype was successfully validated by an independent analysis (Kashiyama et al., 2013). Therefore, genetic defects in the structure-specific endonuclease XP-F/ERCC1 can result in xeroderma pigmentosum, Cockayne syndrome, Fanconi anemia, XFE progeria and cerebro-oculo-facio-skeletal syndrome (Manandhar et al., 2015).

Three important articles were published in 2015. Using whole-genome sequencing, Ameziane et al. (2015) identified a heterozygous dominant negative de novo mutation in the RAD51-encoding gene in an atypical FA patient and suggested adding the acronym FANCR to the major ho-
mologous recombination player known to play a role in both the resistance (if overexpressed) or sensitivity (when mutated or underexpressed) of cancer cells to radio- and chemotherapies and whose haploinsufficiency is involved in the congenital mirror movement neurological disorder (Depienne et al., 2012).

The second 2015 article was published by Sawyer et al. (2015). They described a patient with a complex FA-like phenotype carrying hereditary biallelic mutations in BRCA1. Indeed, the patient was identified in 2013 (Domechek et al., 2013), but the clinical phenotype of the patient was ascertained definitively only two years later. Therefore, thirteen years after the discovery that the gene for FA-D1 is BRCA2, BRCA1 has likewise obtained the acronym FANC.

Likewise, Hira et al. (2015) described one individual harboring the cellular features and symptoms of FA and bearing biallelic mutations in the gene coding for an E2 ubiquitin-conjugating enzyme, UBE2T. UBE2T was originally identified by Zhang et al. (2000) and recognized as the principal ubiquitin E2 ligase of the FANC core complex by Machida et al. (2006). UBE2T was, indeed, renamed FANCT.

In 2016, XRCC2 was identified as FANCU (Park et al., 2016). However, the patient who carried the XRCC2 mutations failed to show bone marrow failure. XRCC2 belongs to a group of RAD51 paralogs, which includes RAD51B, C and D. Therefore, with XRCC2/FANCU, the list of genes of which inactivating mutations could be associated with an FA-like phenotype and with breast cancer predisposition and/or homologous recombination now includes BRCA2, BRCA1, PALB2, BRIP1/BACH1, RAD51 and RAD51C.

Also in 2016, the 21st and last (but probably not for long) FA or FA-like gene was cloned (Bluteau et al., 2016). These authors identified a child with severe BMF harboring biallelic inactivating mutations in the gene encoding the translesion DNA synthesis (TLS) protein subunit REV7 (also known as MAD2L2), which was named FANCV. FANCV plays a central role in the bypass of the unhooked ICL downstream FANCD2/FANCI, allowing the progression of the process that leads to the HR-mediated rescue of the DNA replication impeded by the stall and collapse of an ongoing replication fork at the DNA lesion.

What next?

While 21 FANC genes have been identified and although the alphabet is near its end, the story is probably far from being finished. Obviously, patients bearing mutations in new genes are expected to be extremely rare. However, just looking at FANCN and the FANC core complex partners, no fewer than five genes could claim the title of “FANC or FANC-like gene”: FAAP20, FAAP24 and FAAP100, MHF1 and MHF2. The depletion or deletion of these proteins results in an FA-like cellular phenotype and mouse mutants, when derived, present a phenotype similar to that of the majority of the FANC-KO mouse models. However, because patients with these mutations are rare, the formal attribution of a FANC acronym to previous genes is currently impossible. Loss-of-function mutations of USP1, the FANCD2/FANCI deubiquitinase, result in an FA-like phenotype in a mouse model and in human cells. Indeed, to be unable to monoubiquitinate FANCD2 or to have FANCD2 constitutively monoubiquitinated represents a similarly poor fate for a cell. However, again, no patient bearing USP1 mutations and presenting the FA clinical symptoms have yet been identified. Moreover, the loss of function of the other components of the structure-specific endonuclease heterodimers MUS81-EME1 (and possibly also EME2), XPF-ERCC1 and SLX4-SLX1 could also be associated with patients with an FA-like phenotype, although, for the moment, their potential mutations are associated with either lethal or extremely strong clinical phenotypes that probably preclude the possibility of their assignment to FA. Additionally, mutations in several other known HR-associated proteins could also result in an FA or an FA-like phenotype in some rare families.

In conclusion, the story of the identification of the FANC genes allows to appreciate the evolution of the genetic and molecular techniques to identify disease-associated genes and to better define their links with the pathological traits. Also, considering the divergent clinical phenotypes associated with the loss of function of the gene products involved in the resistance to DNA crosslinking agents, it seems important to stress again that not all the ICL-repair proteins can nowadays claim to be members of the FANC gene group, even if they are involved in the FANC pathway. In the future, the upper part of the FANC pathway, consisting of the FANC core complex-encoding genes FANCD2/FANCI and some of the proteins of the third group (FANCQ, FANCV), will probably be considered separately from the bottom part, whose associated gene products are involved in homologous recombination biochemistry and in breast and ovarian cancer predisposition, for which biallelic germline inactivation results in strong clinical phenotypes.

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