TITLE PAGE

TITLE: Evaluation of rare variants in the new Fanconi Anemia gene ERCC4 (FANCQ) as familial breast/ovarian cancer susceptibility alleles.

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Abstract:

It has been recently reported that bi-allelic mutations in the ERCC4 (FANCQ) gene cause Fanconi Anemia (FA) subtype FA-Q. To investigate the possible role of ERCC4 in breast and ovarian cancer susceptibility, as occurs with other FA genes, we screened the 11 coding exons and exon-intron boundaries of ERCC4 in 1573 index cases from high risk Spanish familial breast and ovarian cancer pedigrees that had been tested negative for BRCA1 and BRCA2 mutations and 854 controls. The frequency of ERCC4 mutation carriers does not differ between cases and controls, suggesting that ERCC4 is not a cancer susceptibility gene. Interestingly, the prevalence of ERCC4 mutation carriers (1 in 288) is similar to that reported for FANCA while there are approximately 100 fold more FA-A than FA-Q patients, indicating that most bi-allelic combinations of ERCC4 mutations are embryo lethal. Finally, we identified additional bone fide FA ERCC4 mutations specifically disrupting interstrand crosslink repair.
The *ERCC4* (*FANCQ*) gene (MIM 133520) encodes for a DNA repair endonuclease (XPF) that plays essential roles in nucleotide excision repair (NER) and interstrand crosslink repair (ICLR) (Gregg, et al., 2011). Bi-allelic mutations in *ERCC4* have been linked to Xeroderma Pigmentosum (XP [MIM 278700; 610651; 278720; 278730; 278740; 278760; 278780; 278750]) (Sijbers, et al., 1996) and progeria (XFE [MIM 610965]) (Niedernhofer, et al., 2006) diseases. Very recently, it has been published that bi-allelic mutations in the gene are also causative of disease in previously unclassified Fanconi Anemia (FA) patients subtype FA-Q and, therefore, the use of *FANCQ* as an alias for *ERCC4* was approved by the HUGO Gene Nomenclature Committee (Bogliolo, et al., 2013). Individuals harbouring mutations in *ERCC4* show high variability on their clinical manifestations, ranging from mild XP symptoms (sun sensitivity, freckling of the skin and basal or squamous cell carcinomas) to the dramatic accelerated aging symptoms of a progeroid syndrome, and finally, as recently shown (Bogliolo, et al., 2013) in FA patients, to malformations and progressive bone marrow failure. It has been also demonstrated that a balance between NER and ICLR activities due to different effects caused by distinct mutations in *ERCC4* determined the final clinical outcome (Bogliolo, et al., 2013). On the other hand, given that FA-Q patients are normal in FANCD2 monoubiquitination, it has been suggested that mono-allelic mutations in *ERCC4* could be linked to the breast and ovarian hereditary cancer syndrome as it occurs with other FA genes such as *BRCA2/FAND1* (Howlett, et al., 2002), *BRIPI/FANCIJ* (Seal, et al., 2006), *PALB2/FANCN* (Rahman, et al., 2007) and *RAD51C/FANCO* (Meindl, et al., 2010; Osorio, et al., 2012) (MIM 300514, 610832, 614082, and 614083, respectively) (Levy-Lahad, 2010).
To investigate the possible role of *ERCC4* in breast and ovarian cancer susceptibility, we screened by DHPLC (Denaturing High Performance Liquid Chromatography) and direct sequencing the 11 coding exons and exon-intron boundaries of the *ERCC4* gene in 1573 index cases from high risk Spanish familial breast and ovarian cancer pedigrees that had been tested negative for mutations in *BRCA1* and *BRCA2* and 854 controls without personal or familial antecedents of cancer. Criteria for inclusion of cases and controls, and methods of screening for mutations in *BRCA1/2* have been previously published (Romero, et al., 2011; Fernandez-Rodriguez, et al., 2012; Osorio, et al., 2012; Bonache, et al., 2013). We identified five and four unique variants among cases and controls among which three and four respectively were considered as putatively deleterious. Missense mutations were evaluated using the program CONDEL that predicts the pathogenicity of non synonymous variants using a consensus deleteriousness score that combines various tools such as SIFT, Polyphen2 and MutationAssessor (http://bg.upf.edu/condel/home). Those missense variants predicted to be deleterious by CONDEL were considered as predicted to affect protein function and this was studied later by functional analysis; those predicted to be neutral were considered as such and not further evaluated (Table 1).

Regarding cases, one of the mutations found, c.584+1G>A in intron 3, was confirmed to cause skipping of exon 3 of the gene and a premature stop codon (PTC) (data not shown). The other two mutations, p.Arg150Cys in exon 3 and p.Ser786Phe in exon 11 were later functionally investigated. Regarding controls, we identified two different frameshift mutations c.540_541delAG in exon 3 (found in two independent controls) and c.2291delG in exon 11, both predicted to cause PTCs. Even though this later PTC is very C-terminal and could potentially result in a shorter but partially functional protein, this mutation results in a truncated XPF protein that lacks the double helix-hairpin-helix
(HhH2) domain involved in heterodimerization with ERCC1 and DNA binding (de Laat, et al., 1998), very similar to a pathogenic ERCC4 mutation (c.2371_2398dup28; p.Ile800Thrfs*24) that functionally disrupts NER and ICLR activities (Bogliolo, et al., 2013). The last deleterious variant found in controls was the missense p.Arg689Ser in exon 11 previously found in a FA patient, and demonstrated to cause abnormal nuclease activity and to specifically disrupt ICLR (Bogliolo, et al., 2013). No differences were found regarding localization of mutations in the gene among cases and controls (Table 1).

All variants reported have been submitted to the Leiden Open Variation Database (LOVD).

To evaluate the functional impact of the missense variants, we cloned a- HA-tagged WT ERCC4 cDNA in a pBABE-puro retroviral vector (Addgene plasmid 14430, kindly shared by Dr. LM Martins) in IRES with the GFP cDNA, and the c.448C>T, c.2065C>A and c.2357C>T variants were introduced by site directed mutagenesis (Bogliolo, et al., 2013). The resulting constructs were transduced in NER and ICLR deficient Ercc4 KO mouse embryonic fibroblasts (MEFs) and, after puromycin selection, the green cells were sorted to achieve a purity of over 98% by FACS (Bogliolo, et al., 2013). Due to the bicistronic nature of the IRES construct, we were able to assess the stability of the mutant XPF proteins using GFP as a reference, since both proteins are encoded by the same mRNA (Pelletier and Sonenberg, 1988). The p.Arg689Ser variant reduced by 40% the stability of XPF, while the p.Arg150Cys and p.Ser786Phe variants had no impact on protein stability (Fig. 1A and 1B). UVC sensitivity of Xpf KO MEFs was complemented with the expression of both p.Arg150Cys- and p.Ser786Phe-XPF (Fig. 1C), and only the Xpf KO MEFs expressing p.Ser786Phe-XPF or p.Arg689Ser showed a FA phenotype in terms of mitomycin C (MMC) sensitivity (Fig. 1D), MMC-induced cell cycle arrest at the G2/M phase (Fig. 1E) and DEB-induced chromosome fragility (Fig. 1F). These data
confirms that, resembling p.Arg689Ser, p.Ser786Phe specifically disrupts ICLR and, therefore, is a bone-fide FA mutation. Interestingly, both mutations are located in the nuclease domain of XPF. Despite a mild MMC sensitivity (Fig. 1D and 1E), Ercc4 KO MEFs expressing p.Arg150Cys-XPF did not show DEB-induced chromosome fragility (Fig 1F). These data, together with the protein stability and UVC sensitivity data, indicate a null impact of the c.448C>T variant on XPF NER functions and a mild effect on ICLR activity.

In conclusion, the frequency of Spanish individuals heterozygous for pathogenic mutations in the ERCC4 gene is approximately 0.3% and it does not differ between familial breast/ovarian cancer patients and healthy controls (p=0.251), suggesting that mono-allelic mutations in ERCC4 are not linked to cancer susceptibility in the general population. Similar results were found with SLX4 that, like ERCC4, acts downstream FANCD2 monoubiquitination but upstream the homologous recombination step of ICL (Fernandez-Rodriguez, et al., 2012). The prevalence of ERCC4 mutation carriers (1 in 288) is similar to that reported for FANCA. However, there are approximately 100 fold more FA-A than FA-Q individuals, suggesting that over 90% of bi-allelic combinations of ERCC4 mutations are embryo lethal in humans. All reported XP patients subtype XPF world-wide have at least one missense mutation disrupting NER while all missense mutations found in 2,427 Spanish individuals have substantial NER activity explaining why there are no reported XPF families in Spain.
References:


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The authors declare no competing interests of any kind.

Web resources section


URL for Leiden Open Variation Database: http://databases.lovd.nl/shared/login

Legend to figure

Figure 1: Funcional studies of ERCC4 missense variants. (A) Western blot analysis showing HA-XPF and GFP expressions in the transduced MEFs as a measure of XPF protein stability. XPF levels were quantified with ImageJ software and are expressed as a ratio relative to the GFP levels. (B) XPF levels were quantified as in (A) and expressed as a percentage of the WT. Mean and SEM of 6 experiments are shown (* p=0.005 t-student test). (C) UVC sensitivity of Ercc4 KO MEFs expressing the different alleles of ERCC4. Cells were analysed 72 h post irradiation. Data represent means and SD of three independent experiments. (D) MMC sensitivity in the same cells as in C. Data represent means and SD of two independent experiments. (E) MMC-induced G2/M cell cycle arrest in the same cells as in (C). (F) DEB-induced chromosome fragility test arrest in the same cells as in (C). Chromatid breaks were scored in 25 metaphase per point. Methods for western blot, site-directed mutagenesis, cDNA transduction, antibodies used and the
experiments presented in D, E and F were performed as reported earlier (Trujillo, et al., 2012; Bogliolo, et al., 2013).
<table>
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<th>Rank</th>
<th>Nucleotide change</th>
<th>Protein change</th>
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<th>MAF reported</th>
<th>Phenotype</th>
<th>Predicted to affect protein function</th>
<th>Functional assay</th>
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<td>3</td>
<td>c.446C&gt;T&lt;sup&gt;a&lt;/sup&gt;</td>
<td>p.Arg150Lys</td>
<td>m.1454G&gt;A in 3p21.3</td>
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<td>3</td>
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<td></td>
<td></td>
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<td></td>
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<td>p.Thr62Ala</td>
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<td>p.Arg686Ser</td>
<td>m.1483C&gt;T in 2p16.1</td>
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<td>Yes</td>
<td>disrupt ICAR</td>
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<tr>
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<td>p.Leu754fsfs06</td>
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<td></td>
<td>Case</td>
<td>Yes</td>
<td>disrupt ICAR</td>
</tr>
</tbody>
</table>

<sup>a</sup> Mutation nomenclature listed uses GenBank reference sequences NM_000258.2 with numbering starting at the A of the first ATG, following the HGVS guidelines (www.hgvs.org/mutnomenclature).

<sup>b</sup> Apart from the 854 controls in which XPR1/BCDF was fully sequenced, these variants were specifically analyzed in 300 additional controls. The mutation gave rise to skipping of exon 3 (c.289-594del) that was confirmed at the cDNA level (data not shown).

<sup>c</sup> MAF (Minor Allele Frequency) in European American population as described in the NHLBI Exome Sequencing Project (ESP) (http://evs.gs.washington.edu/ES/).

<sup>d</sup> Mutation found in two independent controls.