



Article

The Spectrum of FANCM Protein Truncating Variants in European Breast Cancer Cases

Gisella Figlioli ¹, Anders Kvist ², Emma Tham ³, Jana Soukupova ⁴, Petra Kleiblova ⁵ Taru A Muranen ⁶, Nadine Andrieu ^{7,8}, Jacopo Azzollini ⁹, Judith Balmaña ^{10,11}, Alicia Barroso 12, Javier Benítez 12,13,14, Birgitte Bertelsen 15, Ana Blanco 16,17,18, Bernardo Bonanni 19, Åke Borg 2, Joan Brunet 20, Daniele Calistri 21, Mariarosaria Calvello 19, Mariarosaria Calvello 19, Stepan Chvojka ²², Laura Cortesi ²³, Esther Darder ²⁰, Jesús Del Valle ²⁰, Orland Diez ^{10,24}, ENIGMA Consortium ²⁵, Séverine Eon-Marchais ^{7,8}, Florentia Fostira ²⁶, GENESIS Study Collaborators 7, Francesca Gensini 27, Claude Houdayer 28, Marketa Janatova 4, Johanna I Kiiski ⁶, Irene Konstantopoulou ²⁶, Katerina Kubelka-Sabit ²⁹, Conxi Lázaro ²⁰, Fabienne Lesueur ^{7,8}, Siranoush Manoukian ⁹, Ruta Marcinkute ³⁰, Ugnius Mickys ³¹, Virginie Moncoutier 32, SWE-BRCA Group 33, Aleksander Myszka 34, Tu Nguyen-Dumont 35,36, Finn Cilius Nielsen 15, Rimvydas Norvilas 30,37, Edith Olah 38, Ana Osorio 12,13, Laura Papi 27, Bernard Peissel⁹, Ana Peixoto³⁹, Dijana Plaseska-Karanfilska⁴⁰, Timea Pócza³⁸, Maria Rossing 15, Vilius Rudaitis 41, Marta Santamariña 16,17,18, Catarina Santos 39, Snezhana Smichkoska 420, Melissa C Southey 35,36, Dominique Stoppa-Lyonnet 32, Manuel Teixeira ^{39,43}, Therese Törngren ², Angela Toss ²³, Miguel Urioste ⁴⁴, Ana Vega ^{16,17,18}, Zdenka Vlckova 45, Drakoulis Yannoukakos 26, Valentina Zampiga 21, Zdenek Kleibl 40, Paolo Radice 46, Heli Nevanlinna 60, Hans Ehrencrona 33,470, Ramunas Janavicius 30,37 and Paolo Peterlongo 1,*

- Genome Diagnostics Program, IFOM the FIRC Institute for Molecular Oncology, Milan 20139, Italy; gisella.figlioli@ifom.eu
- Division of Oncology and Pathology, Department of Clinical Sciences Lund, Lund University, Lund SE-22381, Sweden; Anders.Kvist@med.lu.se (A.K.); ake.borg@med.lu.se (Å.B.); Therese.Torngren@med.lu.se (T.T.)
- Department of Clinical Genetics, Karolinska University Hospital and Department of Molecular Medicine, Karolinska Institutet, Stockholm 17176, Sweden; Emma. Tham@ki.se
- Institute of Biochemistry and Experimental Oncology, First Faculty of Medicine, Charles University, Prague 12853, Czech Republic; jana.soukupova@lf1.cuni.cz (J.S.); mjana@lf1.cuni.cz (M.J.); zdekleje@lf1.cuni.cz (Z.K.)
- Institute of Biology and Medical Genetics, General University Hospital and First Faculty of Medicine, Charles University, Prague 12800, Czech Republic; pekleje@lf1.cuni.cz
- Department of Obstetrics and Gynecology, Helsinki University Hospital and University of Helsinki, HUS, Helsinki 00029, Finland; taru.a.muranen@helsinki.fi (T.A.M.); johanna.kiiski@helsinki.fi (J.I.K.); Heli.Nevanlinna@hus.fi (H.N.)
- Inserm, U900, Institut Curie, PSL University, Paris F-75005, France; nadine.andrieu@curie.fr (N.A.); Severine. Eon-Marchais@curie.fr (S.E.-M.); fabienne.lesueur@inserm.fr (G.S.C.); fabienne.lesueur@curie.fr (F.L.)
- Mines ParisTech, Fontainebleau F-77300, France
- Department of Medical Oncology and Hematology, Unit of Medical Genetics Fondazione, IRCCS Istituto Nazionale dei Tumori, Milan 20133, Italy; Jacopo. Azzollini@istitutotumori.mi.it (J.A.); Siranoush.Manoukian@istitutotumori.mi.it (S.M.); bernard.peissel@istitutotumori.mi.it (B.P.)
- Hereditary Cancer Group, Vall d'Hebron Institute of Oncology (VHIO), Barcelona 08035, Spain; Jbalmana@vhebron.net (J.B.); odiez@vhio.net (O.D.)
- Department of Medical Oncology, University Hospital Vall d'Hebron, Barcelona 08035, Spain
- Human Genetics Group, Human Cancer Genetics Programme, Spanish National Cancer Research Centre, Madrid 28029, Spain; abarroso@cnio.es (A.B.); jbenitez@cnio.es (J.B.); aosorio@cnio.es (A.O.)
- Spanish Network on Rare Diseases (CIBERER), Madrid 28029, Spain

Cancers 2020, 12, 292 2 of 10

Genotyping Unit, CEGEN, Human Cancer Genetics Programme, Spanish National Cancer Research Centre, Madrid 28029, Spain

- Center for Genomic Medicine, Copenhagen University Hospital, Rigshospitalet, Copenhagen 2100, Denmark; Birgitte.Bertelsen@regionh.dk (B.B.); finn.cilius.nielsen@regionh.dk (F.C.N.); caroline.maria.rossing@regionh.dk (M.R.)
- ¹⁶ Fundación Pública Galega Medicina Xenómica-SERGAS, Santiago de Compostela 15706, Spain; ana.blanco@usc.es (A.B.); santamarinapena@gmail.com (M.S.); ana.vega@usc.es (A.V.)
- ¹⁷ Instituto de Investigación Sanitaria de Santiago de Compostela (IDIS), Santiago de Compostela 15706, Spain
- ¹⁸ Centro de Investigación en Red de Enfermedades Raras (CIBERER), Madrid 28029, Spain
- Division of Cancer Prevention and Genetics, IEO, European Institute of Oncology IRCCS, Milan 20141, Italy; bernardo.bonanni@ieo.it (B.B.); mariarosaria.calvello@ieo.it (M.C.)
- Hereditary Cancer Program, Catalan Institute of Oncology, ONCOBELL-IDIBELL-IDIBGI-IGTP, CIBERONC, Barcelona 08908, Spain; Jbrunet@iconcologia.net (J.B.); edarder@iconcologia.net (E.D.); jdelvalle@iconcologia.net (J.D.V.); conxi.lazaro@gmail.com (C.L.)
- Biosciences Laboratory, Istituto Scientifico Romagnolo per lo Studio e la Cura dei Tumori (IRST) IRCCS, Meldola 47014, Italy; daniele.calistri@irst.emr.it (D.C.); valentina.zampiga@irst.emr.it (V.Z.)
- Centre for Medical Genetics and Reproductive Medicine, Gennet, Prague 17000, Czech Republic; stepan.chvojka@gennet.cz
- ²³ University Modena Hospital, Modena 41124, Italy; hbc@unimore.it (L.C.); angela.toss@unimore.it (A.T.)
- Area of Molecular and Clinical Genetics, University Hospital Vall d'Hebron, Barcelona 08035, Spain
- ²⁵ QIMR Berghofer Medical Research Institute, Brisbane, QLD, Australia; contact@enigmaconsortium.org
- InRASTES, Molecular Diagnostics Laboratory, National Centre for Scientific Research "Demokritos", Athens 15310, Greece; florentia_fostira@hotmail.com (F.F.); reenakon@gmail.com (I.K.); yannouka@gmail.com (D.Y.)
- Department of Experimental and Clinical Biomedical Sciences, University of Florence, Florence 50134, Italy; francesca.gensini@unifi.it (F.G.); laura.papi@unifi.it (L.P.)
- Genetics Department, F76000 and Normandy University, UNIROUEN, Inserm U1245, Normandy Centre for Genomic and Personalized Medicine, Rouen University Hospital, Rouen, France; claude.houdayer@chu-rouen.fr
- Department of Histopathology and Cytology, Clinical Hospital Acibadem Sistina, Skopje 1000, Republic of North Macedonia; katerina.kubelka@acibademsistina.mk
- Hereditary Cancer Center, Hematology, Oncology and Transfusion Medicine Center, Vilnius University Hospital Santaros Klinikos, Vilnius 08410, Lithuania; ruta.marcinkute@gmail.com (R.M.); rimvydas.norvilas@santa.lt (R.N.); Ramunas.Janavicius@santa.lt (R.J.)
- National Center of Pathology, Vilnius University Hospital Santaros Klinikos, Vilnius 08410, Lithuania; ugnius.mickys@vpc.lt
- Service de Génétique, Institut Curie, Inserm, U830, Paris Descartes University, Paris F-75005, France; virginie.moncoutier@curie.fr (V.M.); Dominique.StoppaLyonnet@curie.fr (D.S.-L.)
- Division of Clinical Genetics, Department of Laboratory Medicine, Lund University, Lund SE-22100, Sweden; hans.ehrencrona@skane.se
- ³⁴ Institute of Medical Sciences, University of Rzeszow, Rzeszow 35-310, Poland; amyszka@univ.rzeszow.pl
- Precision Medicine, School of Clinical Sciences at Monash Health, Monash University, Clayton 3168, Australia; tu.nguyen-dumont@monash.edu (T.N.-D.); melissa.southey@monash.edu (M.C.S.)
- 36 Department of Clinical Pathology, The University of Melbourne, Melbourne 3010, Australia
- Department of experimental, preventive and clinical medicine, State Research Institute Centre for Innovative Medicine, Vilnius 08410, Lithuania
- Department of Molecular Genetics, National Institute of Oncology, Budapest 1122, Hungary; e.olah@oncol.hu (E.O.); timipocza@oncol.hu (T.P.)
- Department of Genetics, Portuguese Oncology Institute of Porto (IPO Porto), Porto 4200-072, Portugal; analuisamoura@ipoporto.min-saude.pt (A.P.); catarinasantos@ipoporto.min-saude.pt (C.S.); manuelteixeira@ipoporto.min-saude.pt (M.T.)
- Research Centre for Genetic Engineering and Biotechnology 'Georgi D. Efremov', Macedonian Academy of Sciences and Arts, Skopje 1000, Republic of North Macedonia; dijana@manu.edu.mk
- Department of Gynaecology, Center of Obsterics and Gynaecology, Vilnius University Hospital Santaros Klinikos, Vilnius 08410, Lithuania; Vilius.Rudaitis@santa.lt
- Medical Faculty, University Clinic of Radiotherapy and Oncology, Ss. Cyril and Methodius University in Skopje, Skopje 1000, Republic of North Macedonia; smicko@t.mk

Cancers 2020, 12, 292 3 of 10

- Biomedical Sciences Institute, University of Porto, Porto 4050-313, Portugal
- Familial Cancer Clinical Unit, Human Cancer Genetics Programme, Spanish National Cancer Research Centre, Madrid 28029, Spain; murioste@cnio.es
- 45 Department of Medical Genetics, GHC Genetics, Prague 11000, Czech Republic; Vlckova@ghc.cz
- Department of Research, Unit of Molecular Bases of Genetic Risk and Genetic Testing, Fondazione IRCCS Istituto Nazionale dei Tumori, Milan 20133, Italy; Paolo.Radice@istitutotumori.mi.it
- Office for Medical Services, Region Skåne, Department of Clinical Genetics and Pathology, Laboratory Medicine, Lund SE-22100, Sweden; hans.ehrencrona@med.lu.se
- * Correspondence: paolo.peterlongo@ifom.eu; Tel.: +39-02-57430-3867

Received: 13 December 2019; Accepted: 21 January 2020; Published: 26 January 2020



Abstract: Germline protein truncating variants (PTVs) in the *FANCM* gene have been associated with a 2–4-fold increased breast cancer risk in case-control studies conducted in different European populations. However, the distribution and the frequency of *FANCM* PTVs in Europe have never been investigated. In the present study, we collected the data of 114 European female breast cancer cases with *FANCM* PTVs ascertained in 20 centers from 13 European countries. We identified 27 different *FANCM* PTVs. The p.Gln1701* PTV is the most common PTV in Northern Europe with a maximum frequency in Finland and a lower relative frequency in Southern Europe. On the contrary, p.Arg1931* seems to be the most common PTV in Southern Europe. We also showed that p.Arg658*, the third most common PTV, is more frequent in Central Europe, and p.Gln498Thrfs*7 is probably a founder variant from Lithuania. Of the 23 rare or unique *FANCM* PTVs, 15 have not been previously reported. We provide here the initial spectrum of *FANCM* PTVs in European breast cancer cases.

Keywords: breast cancer predisposition; breast cancer risk factors; *FANCM* truncating variants; mutation spectrum; PTVs

1. Introduction

FANCM, the Fanconi anemia (FA) complementation group M gene (OMIM609644), was originally described as one of the members of the FA molecular pathway [1] that is primarily responsible for the repair of the DNA inter-strand crosslinks through homologous recombination. FA is a recessive DNA repair disease characterized by bone marrow failure, congenital malformations, chromosome fragility, and cancer. Although FANCM is part of the core complex in the FA pathway, accumulating evidence indicates that protein truncating variants (PTVs) in this gene are not causative of FA. Recently, eight individuals who were not diagnosed with FA were found to harbor bi-allelic FANCM PTVs. Three individuals developed early-onset cancers including lymphoblastic leukemia and squamous cancers [2]. In addition, five carriers of homozygous FANCM PTVs were identified among females diagnosed with breast cancer, two of which were diagnosed with early onset disease [3]. Hence, it seems that bi-allelic FANCM PTVs cause cancer predisposition with a greater risk for early onset development.

In the last years, results from case-control studies have indicated that mono-allelic *FANCM* PTVs are breast cancer risk factors. The c.5101C > T (p.Gln1701*, rs147021911) is frequent in the Finnish population with carrier frequency = 0.0162 versus 0.00168 in non-Finnish Europeans (NFE), as reported in gnomAD v2.1.1 (https://gnomad.broadinstitute.org/; [4]). In the Finnish population, p.Gln1701* was found to be associated with breast cancer risk with an odds ratio (OR) of 1.86 and with greater effects in familial cases, and for estrogen receptor-negative (ER-negative) and triple-negative breast cancer (TNBC) subtypes [5]. This variant was later found to be also associated with breast cancer survival and treatment outcome [6]. A second *FANCM* PTV, the c.5791C > T (p.Gly1906Alafs12*, rs144567652), which is annotated and hereafter referred to as p.Arg1931*, showed association with breast cancer risk in familial cases with OR = 3.93 [7]. This PTV was later found to also be associated with risk for TNBC in Finnish cases with OR = 5.14 [8]. A sequencing analysis of the entire *FANCM* coding region showed

Cancers 2020, 12, 292 4 of 10

an excess of a third common PTV, the c.1972C > T (p.Arg658*, rs368728266), in German breast cancer cases versus controls, and confirmed that FANCM PTVs have a particularly high risk (OR = 3.75) for TNBC [9]. We recently tested p.Arg658*, p.Gln1701*, and p.Arg1931* in 67,112 European breast cancer cases and 53,766 controls. In these analyses, we observed that p.Arg658* was associated with increased risk of ER-negative disease and TNBC with ORs of 2.44 and 3.79, respectively, and that p.Arg1931* was associated with the risk of ER-negative breast cancer with an OR of 1.96 [10].

BRCA1, BRCA2, and PALB2 are established breast cancer predisposition genes conferring high risk especially for ER-negative disease and TNBC. Similarly to other genes such as BARD1, RAD51D, BRIP1, and RAD51C [11], FANCM is emerging as a breast cancer predisposing factor conferring a greater risk specifically for these breast cancer subtypes. However, the spectrum of FANCM PTVs in Europe has never been investigated. Through a project call addressed to ENIGMA consortium collaborators [12], we collected data from European female breast cancer probands who were subjected to the sequencing of the FANCM coding region, resulting as carriers of a FANCM PTV. This study describes the spectrum of FANCM PTVs in this cohort of European breast cancer cases.

2. Results and Discussion

The 114 European breast cancer probands included in this study carried 27 different *FANCM* PTVs. Four, namely c.1491dupA (p.Gln498Thrfs*7, rs797045117), p.Arg658*, p.Gln1701*, and p.Arg1931*, were each identified in at least 8 probands and were classified as "common *FANCM* PTVs". The remaining 23 variants were unique or were found in not more than three probands and were classified as "rare *FANCM* PTVs". Additional data on common and rare *FANCM* PTVs were derived from previously published studies based on *FANCM* sequencing of German breast cancer probands [9,13]. Altogether, these and our data allowed us to show the geographic distribution of the four common PTVs and the rare PTVs combined in a cohort of breast cancer probands (Figure 1).

Of the 114 breast cancer probands, 86 (75.4%) carried one of the four common *FANCM* PTVs (Table 1). Of these 86 probands, 8 (9.3%), carried the p.Gln498Thrfs*7, 13 (15.1%) the p.Arg658*, 38 (44.2%) the p.Gln1701*, and 27 (31.4%) carried the p.Arg1931*. We observed that the relative frequency of the two most common PTVs, p.Gln1701* and p.Arg1931*, differ with respect to the country of origin of carriers. The p.Gln1701* is the most common PTV in Northern Europe while it has a lower relative frequency in Southern Europe. An opposite frequency gradient along the North–South axis appeared to exist for p.Arg1931*, which is the most common PTV in Southern Europe but more rare in Northern Europe. Consistently, in Central Europe, p.Gln1701* and p.Arg1931* are both common with similar relative frequencies. The relative frequencies of the *FANCM*:p.Arg658*, the third most common *FANCM* PTV, suggest that this variant has the highest frequency in Central Europe, is present in Southern Europe, and seems to be absent or very rare in Scandinavia. The fourth common *FANCM* PTV p.Gln498Thrfs*7 was identified in 6 of the 7 probands from Lithuania, suggesting it is a founder variant from this specific geographic area (Table 1, Figure 1).

Cancers **2020**, 12, 292 5 of 10

Table 1. Eighty-six breast cancer probands who carry one of the four common *FANCM* PTVs.

Center/Study	Country of Origin	PTV	Age at BC Diagnosis	ER-Status	BC/OC Famil History
CZECANCA-Charles University, Prague	Czech Republic	p.Arg658*	53	ER-pos	yes
CZECANCA-Charles University, Prague	Czech Republic	p.Arg658*	85	NA	no
CZECANCA-Charles University, Prague CZECANCA-Gennet, Prague	Czech Republic	p.Arg658*	35	NA ER nos	yes
CZECANCA-Gennet, Frague	Czech Republic Czech Republic	p.Arg658* p.Arg658* (#)	66 28	ER-pos ER-pos	yes
CZECANCA-Gric, Frague	Czech Republic	p.Gln1701*	51	ER-pos ER-neg	yes yes
CZECANCA-Charles University, Prague	Czech Republic	p.Gln1701*	36	ER-pos	no
CZECANCA-Gennet, Prague	Czech Republic	p.Gln1701*	40	ER-neg	no
CZECANCA-Gennet, Prague	Czech Republic	p.Gln1701*	43	NA	no
CZECANCA-GHC, Prague	Czech Republic	p.Gln1701*	45	ER-pos	yes
CZECANCA-GHC, Prague	Czech Republic	p.Gln1701*	44	ER-pos	no
CZECANCA-Charles University, Prague	Czech Republic	p.Arg1931*	41	ER-neg	yes
ZECANCA-Charles University, Prague	Czech Republic	p.Arg1931*	46	ER-pos	no
CZECANCA-Charles University, Prague	Czech Republic	p.Arg1931*	44	ER-pos	yes
CZECANCA-GHC, Prague	Czech Republic	p.Arg1931*	63	NA ED	no
Copenhagen Breast Cancer Study Copenhagen Breast Cancer Study	Denmark Denmark	p.Gln1701* p.Gln1701*	58 50	ER-neg ER-pos	yes
Copenhagen Breast Cancer Study	Denmark	p.Arg1931*	27	ER-pos	no yes
Helsinki Breast Cancer Study	Finland	p.Gln1701*	46	ER-pos	yes
Ielsinki Breast Cancer Study	Finland	p.Gln1701*	43	ER-pos	yes
Ielsinki Breast Cancer Study	Finland	p.Arg1931*	64	ER-pos	yes
GENE SISters	France	p.Gln1701*	44	ER-pos	yes
nstitut Curie, Paris	France	p.Gln1701*	63	NA	yes
GENE SISters	France	p.Arg1931*	54	ER-pos	yes
GENE SISters	France	p.Arg1931*	39	ER-pos	yes
GENE SISters	France	p.Arg1931*	65	ER-pos	yes
GENE SISters	France	p.Arg1931*	45	ER-pos	yes
GENE SISters	France	p.Arg1931*	43	NA	yes
nstitut Curie, Paris	France	p.Arg1931*	48	NA	yes
National Center for Scientific Research-Demokritos, Athens	Greece	p.Arg658*	33	ER-neg	no
National Center For Scientific Research-Demokritos, Athens	Greece	p.Gln1701*	61	ER-pos	yes
National Center For Scientific Research-Demokritos, Athens	Greece	p.Arg1931*	63	ER-neg	yes
National Center For Scientific Research-Demokritos, Athens	Greece	p.Arg1931*	42, 45	ER-pos	no
Hungarian Breast and Ovarian Cancer Study	Hungary	p.Arg658*	44	ER-pos	yes
Hungarian Breast and Ovarian Cancer Study	Hungary	p.Arg658*	35	ER-pos	yes
ondazione IRCCS-Istituto Nazionale dei Tumori, Milan	Italy Italy	p.Arg1931*	49 32	NA EP nos	yes
stituto Scientifico Romagnolo per lo Studio e la Cura dei umori, Meldola	Italy	p.Arg1931*	32	ER-pos	no
'ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	31	ER-neg	no
Ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	35	ER-neg	no
ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	46	ER-pos	no
ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	39	ER-pos	no
/ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	58	ER-pos	yes
/ilnius University Hospital Santaros Klinikos	Lithuania	p.Gln498Thrfs*7	75	ER-pos	no
/ilnius University Hospital Santaros Klinikos	Lithuania	p.Arg658*	39	ER-pos	no
Macedonian Breast Cancer Study	Macedonia	p.Arg1931*	65	ER-neg	no
Vroclaw Medical University, Wrocław	Poland	p.Gln498Thrfs*7	63	ER-neg	no
Vroclaw Medical University, Wrocław	Poland	p.Gln498Thrfs*7	52	NA	yes
Vroclaw Medical University, Wrocław	Poland	p.Arg658*	62	ER-pos	yes
Portuguese Oncology Institute-Porto Breast Cancer Study	Portugal	p.Arg658*	29	NA	yes
fundación Pública Galega Medicina Xenómica, Santiago de	Spain	p.Arg658*	40	ER-neg	no
Compostela					
panish National Cancer Research Centre, Madrid	Spain	p.Arg658*	38	ER-pos	yes
Catalan Institute of Oncology, Barcelona	Spain	p.Arg1931*	48	ER-pos	yes
Catalan Institute of Oncology, Barcelona panish National Cancer Research Centre, Madrid	Spain Spain	p.Arg1931*	55 47	ER-pos NA	yes
wedish Breast Cancer Study	Sweden	p.Arg1931* p.Gln1701*	34	ER-neg	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Gln1701*	58	ER-neg ER-neg	yes yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	21	ER-neg	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Gln1701*	71	ER-neg	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	71	ER-neg	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	57	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	37	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	68	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	43	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	54	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	56	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	48	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	27	ER-pos	no
wedish Breast Cancer Study	Sweden	p.Gln1701*	42	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	60	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	44	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	57	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	32	ER-pos	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	46	NA NA	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	49	NA NA	yes
wedish Breast Cancer Study	Sweden	p.Gln1701*	36	NA NA	no
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden Sweden	p.Gln1701*	46 39	NA NA	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden Sweden	p.Gln1701* p.Gln1701*	28	NA NA	no
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Gln1701* p.Gln1701*	47	NA NA	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden Sweden	p.Gin1701* p.Arg1931*	36	ER-neg	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Arg1931*	47	ER-neg ER-neg	yes no
wedish Breast Cancer Study	Sweden	p.Arg1931*	52	ER-pos	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Arg1931*	55 55	ER-pos ER-pos	yes
wedish Breast Cancer Study wedish Breast Cancer Study	Sweden	p.Arg1931*	36	ER-pos ER-pos	no
		p.Arg1931*	39	NA	yes
wedish Breast Cancer Study	Sweden				

PTV, protein truncating variant; BC, breast cancer; OC, ovarian cancer; ER, estrogen receptor; pos, positive; neg, negative; NA, not available. (#) This individual also carries the rare *FANCM*: c.3088C > T (p.Arg1030*) PTV and is also included in Table 2.

Cancers 2020, 12, 292 6 of 10

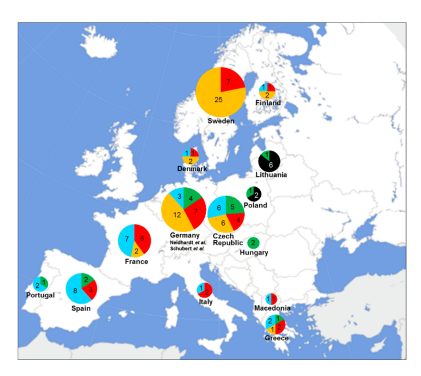


Figure 1. Distribution of the common and rare *FANCM* PTVs found in the 114 European female breast cancer probands. The common PTVs p.Gln498Thrfs*7, p.Arg658*, p.Gln1701*, and p.Arg1931* are indicated in black, green, orange, and red, respectively; the rare PTVs are indicated in light blue. Numbers of PTV carriers are reported according to their country of origin. The 26 PTV carriers from Germany are derived from previously published studies (Neidhardt et al., 21 probands [9]; Schubert et al., 5 probands [13]).

Table 2. Twenty-nine breast cancer probands who carry one of the 23 rare FANCM PTVs.

Center/Study	Country of Origin	PTV	Annotated in Databases	Age at BC Diagnosis	ER-Status	BC/OC Family History
CZECANCA-AGEL, Novy Jicin	Czech Republic	c.1798C > T; p.Gln600*	no	37	ER-pos	yes
CZECANCA-Charles University, Prague	Czech Republic	c.3898G > T; p.Glu1300*	no	37	ER-pos	no
CZECANCA-Charles University, Prague	Czech Republic	c.3979_3980delCA;	LOVD, gnomAD	50	ER-pos	yes
, ,	•	p.Gln1327Valfs*16			•	*
CZECANCA-Gennet, Prague	Czech Republic	c.2260C > T; p.Arg754*	gnomAD	51	ER-neg	no
CZECANCA-Gennet, Prague	Czech Republic	c.2260C > T; p.Arg754*	gnomAD	34	ER-neg	no
CZECANCA-GHC, Prague	Czech Republic	c.3088C > T; p.Arg1030* (#)	no	28	ER-pos	ves
Copenhagen Breast Cancer Study	Denmark	c.3745_3748del;	no	50	ER-pos	NA
		p.Thr1249Glnfs*7			•	
Helsinki Breast Cancer Study	Finland	c.4021 4022del	no	39	ER-pos	yes
GENE SISters	France	c.1432C > T; p.Arg478*	no	48	NA	yes
GENE SISters	France	c.3745dupA; p.His1248fs	no	63	NA	yes
Institut Curie, Paris	France	c.1868delA; p.Arg624Glufs*46	no	56	ER-neg	ves
Institut Curie, Paris	France	c.2260C > T; p.Arg754*	gnomAD	43	ER-neg	yes
Institut Curie, Paris	France	c.2590delG; p.Asp864Ilefs*12	no	64	ER-neg	yes
Institut Curie, Paris	France	c.4930C > T; p.Arg1644*	gnomAD	46	NA	yes
Institut Curie, Paris	France	c.5363delCTAG;	no	43	ER-neg	no
		p.Ser1788Leufs*15				
National Center For Scientific	Greece	c.4843A > T; p.Lys1615*	LOVD	33	NA	ves
Research-Demokritos, Athens						,
National Center For Scientific	Greece	c.5314_5315delTG; p.Cys1772*	LOVD, gnomAD	47	ER-pos	ves
Research-Demokritos, Athens	Office C	c.oo11_0010dc110, p.c.y01712	DO VD, BROHLI ID	.,	ER POS	yes
Fondazione IRCCS-Istituto Nazionale dei	Italy	c.2354T > G; p.Leu785*	no	35	NA	no
Tumori, Milan	itary	C.25541 > G, p.Ecuros	110	55	1421	110
Macedonian Breast Cancer Study	Macedonia	c.5048 5052delAAAGA;	gnomAD	32	ER-neg	no
Macedonian Dieast Cancer Study	Maccaona	p.Lys1683Argfs*3	grionia i D	32	Living	110
Portuguese Oncology Institute-Porto Breast	Portugal	c.2320G > T; p.Glu774*	no	42	ER-neg	yes
Cancer Study	1 ortugui	C.2320G > 1, p.Glu774	110	144	Living	yes
Portuguese Oncology Institute-Porto Breast	Portugal	c.5766 5769del;	no	43	ER-neg	Troc
Cancer Study	1 Ortugai	p.Thr1923Profs*2	110	4.5	ER-fieg	yes
Catalan Institute of Oncology, Barcelona	Spain	c.624_625delAA; p.Ile208fs	no	23	NA	no
Catalan Institute of Oncology, Barcelona	Spain	c.1303C > T; p.Gln435*	no	39	ER-pos	yes
Catalan Institute of Oncology, Barcelona	Spain	c.5766_5769del;		31	ER-pos ER-pos	•
Catalan institute of Oncology, barcelona	эраш	p.Thr1923Profs*2	no	31	EK-pos	no
Fundación Pública Galega Medicina	Spain	c.2320G > T; p.Glu774*	no	57	ER-neg	****
	эраш	C.2520G > 1; p.Glu774	110	37	EK-fleg	yes
Xenómica, Santiago de Compostela	Country	- 250/ 2500 I-14 A A A	Clinivan LOVID	49	NTA	
Fundación Pública Galega Medicina	Spain	c.2586_2589delAAAA;	ClinVar, LOVD,	49	NA	yes
Xenómica, Santiago de Compostela	Country	p.Lys863Ilefs*12	gnomAD		NTA	
Spanish National Cancer Research Centre, Madrid	Spain	c.1196C > G; p.Ser399*	ClinVar	66	NA	yes
	Constant	- 2220C - T - Cl. 774*		40.75	FD	
Spanish National Cancer Research Centre,	Spain	c.2320G > T; p.Glu774*	no	40, 65	ER-pos	yes
Madrid	Constant	- 4042 A - T - I 1615*	LOVID	41	NTA	
Spanish National Cancer Research Centre, Madrid	Spain	c.4843A > T; p.Lys1615*	LOVD	41	NA	yes

PTV, protein truncating variant; BC, breast cancer; OC, ovarian cancer; ER, estrogen receptor; pos, positive; neg, negative; NA, not available. (#) This individual also carries the common PTV, p.Arg658*, and is also included in Table 1.

Cancers 2020, 12, 292 7 of 10

The high relative frequency of *FANCM*:p.Gln498Thrfs*7 observed in Lithuania could indicate a founder effect in this relatively homogenous Baltic population. Similarly, haplotype analyses previously conducted in this population found a strong founder effect for several recurrent pathogenic variants in the high-risk predisposition genes *BRCA1* and *BRCA2* [14,15]. Calculations from haplotype analysis and paleo-demographic data indicate that a bottleneck phenomenon occurring approximately in the 4th–5th century AD in the Lithuanian region could have contributed to decreased genetic diversity and subsequent expansion of specific alleles [14].

We compared the geographic distribution of the four common PTVs observed in our study cohort with that reported in the gnomAD database v2.1.1 [4]. In gnomAD, 630 European individuals carried one of the p.Gln498Thrfs*7, p.Arg658*, p.Gln1701* or p.Arg1931* PTVs with relative proportions of 3.2% (20/630), 3.3% (21/630), 54.0% (340/630), and 39.5% (249/630), respectively. About half of the p.Gln498Thrfs*7 gnomAD carriers (9/20) were from Estonia, confirming that this variant has a specific origin in the Baltic countries. In gnomAD, p.Arg658* was prevalently found in NFE with the higher frequency of the carries (8/21) found in North-western Europeans and only one carrier found among Southern Europeans, which is similar to what we observed. Having being found in 340 gnomAD individuals, FANCM:p.Gln1701* is by far the most common PTV in Europe. As mentioned before, p.Gln1701* was particularly frequent in Finland and was observed more frequently in North-western than Southern Europeans, which is consistent with our data. FANCM:p.Arg1931* was the second most common FANCM PTV in Europe and the most common FANCM PTV in Southern Europeans in the gnomAD dataset. However, this variant was reported to be the most common also in North-western Europeans, not confirming the South–North frequency gradient observed in our cohort.

Of the 114 breast cancer probands, 29 (25.4%) carried one of the 23 rare FANCM PTVs (Table 2). Interestingly, a proband from the Czech Republic was a bi-allelic carrier of FANCM PTVs having inherited the rare p.Arg1030* variant from the mother, and the common p.Arg658* variant from the paternal side of the family (Tables 1 and 2). This carrier was diagnosed with breast cancer at a young age (28 years), which is consistent with published data showing that females with bi-allelic FANCM PTV may develop early onset disease [3]. Eight of the 23 rare PTVs were annotated in the public ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/), LOVD (https://www.lovd.nl/), or gnomAD databases and 15 were novel. Of the 23 rare PTVs, 19 were unique; two, p.Arg754* and p.Glu774*, were found in three probands; and two, p.Lys1615* and p.Thr1923Profs*2, in two probands. Given these low frequencies, we could not speculate on the origins or the geographic distribution of any of the rare FANCM PTVs. Considering the numbers of carriers of common PTVs and rare PTVs in each country, we observed that in Spain and France, there was an apparent excess of carriers of rare PTVs. On the contrary, in Sweden and Lithuania, no carriers of rare PTVs were found (Figure 1).

3. Materials and Methods

The individuals included in our analysis were contributed by ENIGMA consortium collaborators [12] interested in studying the effect/role of *FANCM* on breast cancer predisposition. They were originally ascertained at several Centers or National Studies and considered eligible for mutation sequencing analysis of breast and/or ovarian cancer predisposing genes in the frame of diagnostic or research purposes. In general, the ascertainment criteria for mutation testing were based on the presence of family history for breast or ovarian cancer, and on the early onset or specific clinical subtype of breast cancer. In the present study, we included 114 probands who received (i) a diagnosis of breast cancer, (ii) tested negative for *BRCA1* or *BRCA2* pathogenic variants, (iii) underwent complete sequencing of the *FANCM* coding region, resulting as a carrier of a PTV. Unfortunately, it was not possible to retrieve the number of probands that were originally subjected to mutation sequencing analysis at each participating Center or National Study. The 114 carriers of *FANCM* PTVs were ascertained in 20 Centers or National Studies from 13 European countries (Table S1) including Czech Republic (20, 17.5%), Denmark (4, 3.5%), Finland (4, 3.5%), France (15, 13.2%), Greece (6, 5.3%), Hungary (2, 1.8%), Italy (3, 2.6%), Lithuania (7, 6.1%), Macedonia (2, 1.8%), Poland (3, 2.6%), Portugal (3, 2.6%),

Cancers 2020, 12, 292 8 of 10

Spain (15, 11.4%), and Sweden (32, 28.1%). Some of these FANCM PTVs carriers were described in studies previously published, as indicated (Table S1). Of the 114 probands, 58 (50.9%) were ER-positive, 27 (23.7%) were ER-negative, and 80 (70.2%) had a positive family history of breast or ovarian cancer. The mean age at breast cancer diagnosis was 46.7 years (min = 21, max = 85). All the 114 probands provided written informed consent for their data to be used for research purposes or specifically agreed to be tested for new variants in potential breast cancer associated genes. The participation of all the Centers or National Studies in the present work (Table S1) was approved by their ethic committees. Specifically, these are the Ethics Committee of the General University Hospital in Prague (Approval code 87/14); Ethics Committee of Carlos III Institute of Health in Madrid (CEI PI 06_2013-v2); by the appropriate Advisory Committee on the Treatment of Health Research Information (Comité Consultatif de Protection des Personnes dans la Recherche Biomédicale (CCPPRB) Ile-de-France III) and by the National Data Protection authority; Ethics Committee of Fondazione IRCCS Istituto Nazionale dei Tumori in Milan (INT 7/15); Ethics Committee of the Capital Region of Denmark (H-4-2010-050); Ethics Committee of Hospital District of Helsinki and Uusimaa (HUS/1597/2016); Ethical Sub-Committee on Medicine, Pharmacy, Veterinary and Dental Medicine, Macedonian Academy of Sciences and Arts (09-1047/1 from 20/04/2016); IPO (Instituto Português de Oncologia) Porto institutional review board (nr. 219-019); Comité Territorial de Ética de la Investigación de Santiago -Lugo, Conselleria de Sanidade, Xunta de Galicia (2018/200); Ethics Committee of the Area Vasta Romagna e I.R.S.T (CEAV, 1727/2012 I.5/43); Provincial Ethics Committee (209/16) and the Modena Hospital (3387); Ethics Committee of University of Rzeszow, Poland (resolution no. 9/06/2015), University of Melbourne Human Research Ethics Committee, Melbourne, Australia; National Scientific and Ethics Committee (ETT-TUKEB), 20998-0/2010-1018EKU (845/PI/010); IDIBELL (Institut d'Investigacio Biomedica de Bellvitge, Barcelona) Ethics Committee PR278/19; Lithuanian Bioethics Committee (LT-HCCA-003 protocol), Vilnius Regional Ethics Committee (Nr. 58200-12-27(1-61); Ethics Committee of Lund, Sweden (Dnr 2011/349 and 2011/652); Bioethics Committee of NSCR (National Centre of Scientific Research) Demokritos (BcncSrD-240/eH Δ /11.3, updated on 29 June 2015).

4. Conclusions

We described the geographic distribution and relative prevalence of 27 different *FANCM* PTVs detected in 114 female European breast cancer probands with no pathogenic variants in *BRCA1* or *BRCA2* genes. As far as we know, this is the largest available collection of breast cancer probands carrying *FANCM* PTVs. In our study-cohort, we found that the *FANCM*: p.Gln1701* had high relative frequency in Northern Europe and low relative frequency in Southern Europe, whereas p.Arg1931* was the most common PTV in Southern Europe and less common in Northern Europe. The p.Arg658* had higher frequency in Central Europe, while p.Gln498Thrfs*7 is probably a founder variant from Lithuania. Fifteen of the 27 PTVs described are novel, including one found in three probands, and one found in two probands. Further data are warranted to provide an extensive spectrum of *FANCM* PTVs in Europe.

Supplementary Materials: The following are available online at http://www.mdpi.com/2072-6694/12/2/292/s1, Table S1: Center or Study contributing the 114 *FANCM* PTV probands. Some of these probands were previously described as indicated; sequencing details are provided.

Author Contributions: Conceptualization, G.F., R.J. and P.P.; Formal analysis, G.F. and P.P; Data curation (providing samples and clinical data), G.F., A.K., E.T., J.S., P.K., T.A.M., N.A., J.A., J.B., A.B., J.B., B.B., A.B., B.B., Å.B., J.B., D.C., M.C., S.C., L.C., E.D., J.D.V., O.D., S.E.M., F.F., F.G., C.H., M.J., J.I.K., I.K., K.K.S., C.L., F.L., S.M., R.M., U.M., V.M., A.M., T.N.-D., F.C.N., R.N., E.O., A.O., L.P., B.P., A.P., D.P.K., T.P., P.R., M.R., V.R., M.S., C.S., S.S., Melissa C Southey, D.S.L., M.T., T.T., A.T., M.U., A.V., Z.V., D.Y., V.Z., Z.K., H.N., H.E., R.J. and P.P.; Writing—original draft preparation, G.F., R.J. and P.P.; Writing—review and editing, G.F., A.K., E.T., J.S., P.K., T.A.M., F.L., D.P.K., P.R., M.R., Z.K., H.N., H.E., R.J. and P.P.; Supervision, P.P. All authors have read and agreed to the published version of the manuscript.

Cancers 2020, 12, 292 9 of 10

Funding: This research was partially funded by Associazione Italiana Ricerca sul Cancro (AIRC; IG2015 no.16732) to P. Peterlongo, a fellowship from Fondazione Umberto Veronesi to G. Figlioli and by the Italian Ministry of Health with Ricerca Corrente and 5x1000 funds. E. Tham is supported by Region Stockholm (ALF). The Czech study was supported by a grant of the Ministry of Health of the Czech Republic NV16-29959A. CNIO study was partially supported by projects PI16/00440 and PI19/00640, supported by the Instituto de Salud Carlos III, cofunded by European Regional Development Fund (ERDF), the Spanish Network on Rare Diseases (CIBERER) and BRIDGES project H2020. Financial support for GENESIS resource and genotyping was provided by the Ligue Nationale contre le Cancer (grants PRÉ05/DSL, PRE07/DSL, PRE11/NA), the French National Institute of Cancer (INCa grant No b2008-029/LL-LC) and the comprehensive cancer center SiRIC, (Site de Recherche Intégrée sur le Cancer: Grant INCa-DGOS-4654). HEBCS was funded by Helsinki University Hospital Research Fund, Sigrid Juselius Foundation, The cancer Society of Finland. A.Vega is supported by the Spanish Health Research Foundation, Instituto de Salud Carlos III (ISCIII) through Research Activity Intensification Program (contract grant numbers: INT15/00070, INT16/00154, INT17/00133), and through Centro de Investigación Biomédica en Red de Enferemdades Raras CIBERER (ACCI 2016: ER17P1AC7112/2018); Autonomous Government of Galicia (Consolidation and structuring program: IN607B), and by the Fundación Mutua Madrileña (call 2018). This work was supported by the Australian National Health and Medical Research Council (APP1029974 and APP1074383) and by a Victorian Life Sciences Computation Initiative grant (number VR0182) on its Peak Computing Facility, an initiative of the Victorian Government. T.N-D is a Career Development Fellow of the National Breast Cancer Foundation (Australia, ECF-17-001). M.C.S. is a National Health and Medical Research Council (Australia) Senior Research Fellow. The Hungarian Breast and Ovarian Cancer Study was supported by Hungarian Research Grants KTIA-OTKA CK-80745 and NKFI OTKA K-112228 to E. Olah. Lithuanian study was supported by The Research Council of Lithuania grant SEN18/2015 and P-MIP-20-25 to R. Janavicius. The ICO was supported by the Carlos III National Health Institute funded by FEDER funds—a way to build Europe—[PI16/00563, PI19/00553 and CIBERONC]; the Government of Catalonia [Pla estratègic de recerca i innovació en salut (PERIS) Project MedPerCan, 2017SGR1282 and 2017SGR496]; and CERCA program.

Acknowledgments: The authors are particularly grateful to individuals participating in this study and their families. We would like to thank the personnel of the Genetic Testing for Hereditary Tumors Laboratory -Cogentech, Spiros Tavandzis from the AGEL Laboratories in Novy Jicin for his valuable contribution to the study, and Daniela Zaffaroni (Istituto Nazionale dei Tumori, Milan). GENESIS (GENE SISters) is a French national study sponsored by UNICANCER (Sinilnikova et al. BMC Cancer 2016). We wish to thank the genetic epidemiology platform (the PIGE, Plateforme d'Investigation en Génétique et Epidemiologie: Séverine Eon-Marchais, Marie-Gabrielle Dondon, M. Marcou, D. Le Gal, L. Toulemonde, J. Beauvallet, N. Mebirouk, E. Cavaciuti), the biological resource centre (S. Mazoyer, F. Damiola, L. Barjhoux, C. Verny-Pierre, V. Sornin) and all the GENESIS collaborating cancer clinics.. HEBCS thanks Drs. Kristiina Aittomäki and Carl Blomqvist and research nurse Outi Malkavaara for their help with patient data and samples. FPGMX: members of the Cancer Genetics group (IDIS): Belinda Rodríguez, Olivia Fuentes and Miguel Aguado. ICO wants to thank all the members of the ICO Hereditary Cancer Program team led by Dr. Gabriel Capella. SWE-BRCA (The Swedish BRCA1 & BRCA2 Study Collaborators): Gothenburg, Sahlgrenska University Hospital: Zakaria Einbeigi, Anna Öfverholm. Linköping University Hospital: Marie Stenmark-Askmalm, Ekaterina Kuchinskaya. Lund University Hospital: Hans Ehrencrona, Therese Törngren, Anders Kvist, Åke Borg. Stockholm, Karolinska University Hospital: Brita Arver, Annika Lindblom, Emma Tham. Umeå University Hospital: Beatrice Melin. Uppsala University Hospital: Ylva Paulsson-Karlsson.

Conflicts of Interest: The authors declare no conflict of interest.

References

- 1. Meetei, A.R.; Medhurst, A.L.; Ling, C.; Xue, Y.; Singh, T.R.; Bier, P.; Steltenpool, J.; Stone, S.; Dokal, I.; Mathew, C.G.; et al. A human ortholog of archaeal DNA repair protein Hef is defective in Fanconi anemia complementation group M. *Nat. Genet.* **2005**, *37*, 958–963. [CrossRef] [PubMed]
- 2. Bogliolo, M.; Bluteau, D.; Lespinasse, J.; Pujol, R.; Vasquez, N.; d'Enghien, C.D.; Stoppa-Lyonnet, D.; Leblanc, T.; Soulier, J.; Surralles, J. Biallelic truncating FANCM mutations cause early-onset cancer but not Fanconi anemia. *Genet. Med.* **2018**, *20*, 458–463. [CrossRef] [PubMed]
- 3. Catucci, I.; Osorio, A.; Arver, B.; Neidhardt, G.; Bogliolo, M.; Zanardi, F.; Riboni, M.; Minardi, S.; Pujol, R.; Azzollini, J.; et al. Individuals with FANCM biallelic mutations do not develop Fanconi anemia, but show risk for breast cancer, chemotherapy toxicity and may display chromosome fragility. *Genet. Med.* **2018**, 20, 452–457. [CrossRef] [PubMed]
- 4. Lek, M.; Karczewski, K.J.; Minikel, E.V.; Samocha, K.E.; Banks, E.; Fennell, T.; O'Donnell-Luria, A.H.; Ware, J.S.; Hill, A.J.; Cummings, B.B.; et al. Analysis of protein-coding genetic variation in 60,706 humans. *Nature* 2016, 536, 285–291. [CrossRef] [PubMed]

Cancers 2020, 12, 292 10 of 10

5. Kiiski, J.I.; Pelttari, L.M.; Khan, S.; Freysteinsdottir, E.S.; Reynisdottir, I.; Hart, S.N.; Shimelis, H.; Vilske, S.; Kallioniemi, A.; Schleutker, J.; et al. Exome sequencing identifies FANCM as a susceptibility gene for triple-negative breast cancer. *Proc. Natl. Acad. Sci. USA* **2014**, *111*, 15172–15177. [CrossRef] [PubMed]

- 6. Kiiski, J.I.; Fagerholm, R.; Tervasmaki, A.; Pelttari, L.M.; Khan, S.; Jamshidi, M.; Mantere, T.; Pylkas, K.; Bartek, J.; Bartkova, J.; et al. FANCM c.5101C>T mutation associates with breast cancer survival and treatment outcome. *Int. J. Cancer* **2016**, *139*, 2760–2770. [CrossRef] [PubMed]
- 7. Peterlongo, P.; Catucci, I.; Colombo, M.; Caleca, L.; Mucaki, E.; Bogliolo, M.; Marin, M.; Damiola, F.; Bernard, L.; Pensotti, V.; et al. FANCM c.5791C>T nonsense mutation (rs144567652) induces exon skipping, affects DNA repair activity and is a familial breast cancer risk factor. *Hum. Mol. Genet.* **2015**, *24*, 5345–5355. [CrossRef] [PubMed]
- 8. Kiiski, J.I.; Tervasmaki, A.; Pelttari, L.M.; Khan, S.; Mantere, T.; Pylkas, K.; Mannermaa, A.; Tengstrom, M.; Kvist, A.; Borg, A.; et al. FANCM mutation c.5791C>T is a risk factor for triple-negative breast cancer in the Finnish population. *Breast Cancer Res. Treat.* **2017**, *166*, 217–226. [CrossRef] [PubMed]
- 9. Neidhardt, G.; Hauke, J.; Ramser, J.; Gross, E.; Gehrig, A.; Muller, C.R.; Kahlert, A.K.; Hackmann, K.; Honisch, E.; Niederacher, D.; et al. Association Between Loss-of-Function Mutations Within the FANCM Gene and Early-Onset Familial Breast Cancer. *JAMA Oncol.* 2017, 3, 1245–1248. [CrossRef]
- 10. Figlioli, G.; Bogliolo, M.; Catucci, I.; Caleca, L.; Lasheras, S.V.; Pujol, R.; Kiiski, J.I.; Muranen, T.A.; Barnes, D.R.; Dennis, J.; et al. The FANCM:p.Arg658* truncating variant is associated with risk of triple-negative breast cancer. *npj Breast Cancer* **2019**, *5*. [CrossRef]
- 11. Shimelis, H.; LaDuca, H.; Hu, C.; Hart, S.N.; Na, J.; Thomas, A.; Akinhanmi, M.; Moore, R.M.; Brauch, H.; Cox, A.; et al. Triple-Negative Breast Cancer Risk Genes Identified by Multigene Hereditary Cancer Panel Testing. *J. Natl. Cancer Inst.* **2018**, *110*, 855–862. [CrossRef]
- 12. Spurdle, A.B.; Healey, S.; Devereau, A.; Hogervorst, F.B.; Monteiro, A.N.; Nathanson, K.L.; Radice, P.; Stoppa-Lyonnet, D.; Tavtigian, S.; Wappenschmidt, B.; et al. ENIGMA—Evidence-based network for the interpretation of germline mutant alleles: An international initiative to evaluate risk and clinical significance associated with sequence variation in BRCA1 and BRCA2 genes. *Hum. Mutat.* **2012**, *33*, 2–7. [CrossRef]
- 13. Schubert, S.; van Luttikhuizen, J.L.; Auber, B.; Schmidt, G.; Hofmann, W.; Penkert, J.; Davenport, C.F.; Hille-Betz, U.; Wendeburg, L.; Bublitz, J.; et al. The identification of pathogenic variants in BRCA1/2 negative, high risk, hereditary breast and/or ovarian cancer patients: High frequency of FANCM pathogenic variants. *Int. J. Cancer* 2019, 144, 2683–2694. [CrossRef] [PubMed]
- 14. Janavicius, R.; Rudaitis, V.; Feng, B.J.; Ozolina, S.; Griskevicius, L.; Goldgar, D.; Tihomirova, L. Haplotype analysis and ancient origin of the BRCA1 c.4035delA Baltic founder mutation. *Eur. J. Med. Genet.* **2013**, *56*, 125–130. [CrossRef] [PubMed]
- 15. Janavicius, R.; Rudaitis, V.; Mickys, U.; Elsakov, P.; Griskevicius, L. Comprehensive BRCA1 and BRCA2 mutational profile in Lithuania. *Cancer Genet.* **2014**, 207, 195–205. [CrossRef] [PubMed]



© 2020 by the authors. Licensee MDPI, Basel, Switzerland. This article is an open access article distributed under the terms and conditions of the Creative Commons Attribution (CC BY) license (http://creativecommons.org/licenses/by/4.0/).