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Genetic landscape of a large cohort of Primary Ovarian Insufficiency: New genes and pathways and implications for personalized medicine

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Summary

Background Primary Ovarian Insufficiency (POI), a public health problem, affects 1-3.7% of women under 40 yielding infertility and a shorter lifespan. Most causes are unknown. Recently, genetic causes were identified, mostly in single families. We studied an unprecedented large cohort of POI to unravel its molecular pathophysiology.

Methods 375 patients with 70 families were studied using targeted (88 genes) or whole exome sequencing with pathogenic/likely-pathogenic variant selection. Mitomycin-induced chromosome breakages were studied in patients' lymphocytes if necessary.

Findings A high-yield of 29.3% supports a clinical genetic diagnosis of POI. In addition, we found strong evidence of pathogenicity for nine genes not previously related to a Mendelian phenotype or POI: *ELAVL2*, *NLRP11*, *CENPE*, *SPATA33*, *CCDC150*, *CCDC185*, including DNA repair genes: *C17orf53* (*HROB*), *HELQ*, *SWI5* yielding high chromosomal fragility. We confirmed the causal role of *BRCA2*, *FANCM*, *BNC1*, *ERCC6*, *MSH4*, *BMPR1A*, *BMPR1B*, *BMPR2*, *ESR2*, *CAV1*, *SPIDR*, *RCBTB1* and *ATG7* previously reported in isolated patients/families. In 8.5% of cases, POI is the only symptom of a multi-organ genetic disease. New pathways were identified: NF-κB, post-translational regulation, and mitophagy (mitochondrial autophagy), providing future therapeutic targets. Three new genes have been shown to affect the age of natural menopause supporting a genetic link.

Interpretation We have developed high-performance genetic diagnostic of POI, dissecting the molecular pathogenesis of POI and enabling personalized medicine to i) prevent/cure comorbidities for tumour/cancer susceptibility genes that could affect life-expectancy (37.4% of cases), or for genetically-revealed syndromic POI (8.5% of cases), ii) predict residual ovarian reserve (60.5% of cases). Genetic diagnosis could help to identify patients who may benefit from the promising in vitro activation-IVA technique in the near future, greatly improving its success in treating infertility.

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Keywords: Primary ovarian insufficiency; Personalized medicine; NF-KB; Post - translational regulation; Meiosis/ DNA repair genes; Mitophagy

Research in context

Evidence before this study

Infertility is a major public health problem, affecting 10% of couples worldwide. Primary Ovarian Insufficiency (POI) is one of the most common syndromes implicated in female infertility, affecting ~1-3.7% of women under 40 years. POI most often leads to permanent sterility and severe complications: osteoporosis, cardiovascular disease and even neurologic degenerative diseases. Remarkably, a decrease in longevity was also observed which could be due to specific causes. But most causes are currently unknown. Despite the technological leap in genetics, the clinical impact on the diagnosis and management of POI remains unknown, as well as the molecular mechanisms disturbed in POI.

Added value of this study

To get insight into this disorder, we studied the feasibility of a first line genetic diagnosis in a large cohort of 375 patients including 70 families and looked for new responsible genes and altered signalling pathways. We have sequenced either the whole coding part of the genome or 88 known causing genes. A high-yield diagnosis of 29.3% was obtained supporting the use of genetics routinely to diagnose all unexplained POI. Interestingly, we identified 9 genes not previously related to POI or Mendelian disease and confirm 13 others previously reported in isolated patients or families. We showed how the molecular dissection of the pathways involved leads to personalized management of patients.

The main family is the DNA repair/meiosis/mitosis gene family (37.4% of cases), but it is also a tumour/cancer susceptibility gene family. Lifelong monitoring is therefore necessary to prevent or treat the appearance of these tumours. The second major family involved is that of follicular growth genes (35.4%). Strikingly, in 8.5% of cases, POI is the only single visible expression of a complex multi-organ genetic disease. A full patient assessment is required.

Other novel pathways identified could yield novel therapeutic targets: i.e. immunity, gene regulation, mitochondrial autophagy.

Three genes had been implicated in the large variance in the age of natural menopause, confirming a genetic link and a continuum between the two conditions, the difference may be related to the severity of the genetic variants involved, major in POI.

Implications of all the available evidence

We describe here a high-performance genetic diagnosis of POI, critical to understand the pathogenesis of POI

and leading to personalized medicine. These results have major implications for preventing/curing comorbidities related to tumour/cancer susceptibility genes that could affect lifespan or multi-organ disease revealed by genetics. Apart from the infertility of patients, it is therefore their entire state of health that must be assessed and treated as soon as the patient consults for POI. This should be done according to the underlying cause which must imperatively be identified beforehand.

Genetics is also critical to establish a fertility prognosis for the promising technique of in vitro follicular activation in the future by predicting a residual ovarian reserve (60.5% of cases). The selection of patients who could benefit from this technique, the genetic cause highlighting existing follicles in the ovaries blocked in their growth, could clearly improve its success in the treatment of infertility of POI. The pathways identified could also provide future therapeutic targets.

Introduction

Infertility affects 15% of couples worldwide and is a public health problem. Primary Ovarian Insufficiency (POI) affects ~1-3.7% of the women before 40 years^{1,2} with cessation of ovarian functions.¹ About 60–70% of cases remain idiopathic.¹ Women with POI are usually definitively infertile but spontaneous pregnancies have been described in 3.5% of patients with secondary amenorrhea or spaniomenorrhea.³ Moreover, various epidemiological studies have shown a decreased lifespan or an increased risk of all-cause mortality in infertile women.^{4,5} It is thus a priority to identify the causes involved. Next generation sequencing (NGS) especially whole exome sequencing (WES) has recently identified several genetic causes of POI, although in single or rare families, mostly consanguineous.¹ Due to this high genetic heterogeneity, the clinical impact of the etiological diagnosis of POI remains unknown. In clinical practice, only karyotype and FMR1 studies are routinely performed with respective diagnosis yields of 7–10% and 3–5%.¹

In this study, we describe the genetic landscape of POI in a large cohort of 375 patients, using WES or targeted NGS comprising all human POI genes known to date, and with a classification of variants using the American College of Medical Genetics and Genomics (ACMG) criteria.⁶ Our study revealed that a powerful etiological diagnosis is possible in a high proportion of patients, namely: 29.3%. We show here that

personalized genetic counselling and management of patients and family is possible based on the identification of the genetic cause. Our study also highlights genes not previously related to Mendelian phenotype, pathways and mechanisms in POI, potentially allowing identification of new therapeutic targets.

Methods

Ethics statement

The study was approved by all the institutions involved and by the Agence de Biomedecine (reference number PFS12-002). Written informed consent was received from participants prior to inclusion in the study.

Patients

Between 2017 and 2022, 375 patients were referred to our university hospital laboratory by different hospitals in Europe, Turkey, Africa, and Asia. We performed NGS analysis in patients who lacked the FMR1 premutation and had a normal karyotype. POI was diagnosed in the presence of primary amenorrhoea (PA), secondary amenorrhoea (SA) or spinoamenorrhoea (SP) for more than 4 months associated with follicle-stimulating hormone (FSH) plasma level ≥ 25 IU/L before the age of 40 years. Patients with known etiology of POI, such as chemotherapy, radiotherapy, or extensive ovarian surgery, were excluded from the study. The following clinical data were requested before the NGS analysis: menstrual cycle (PA, SA, SP), pubertal development, ethnicity, spontaneous pregnancies or pregnancies after in vitro fertilization (IVF), and familial history. Information on autoimmune disease or the presence of extra-ovarian symptoms—such as intellectual disability, hyperandrogenism, neurological and ophthalmic disorders, deafness, tumour/cancer, in the patient and/or family, family history of male or female infertility, was also requested. We also requested ultrasonography to assess ovarian and follicular sizes, the number of follicles, as well as a complete hormonal assay in particular the serum level of follicle stimulating hormone (FSH), luteinizing hormone (LH), estradiol, anti-Mullerian Hormone (AMH), thyroid stimulating hormone (TSH), and autoantibodies (anti-TPO, anti-TG, and anti-21-hydroxylase antibodies).

Molecular genetics studies

- a) A custom-made targeted NGS including all known genes (88) involved in POI to date (supplementary material and methods and Table S1)^{7,8} was performed. WES was performed by IntegraGen SA (Evry, France) in case of consanguineous or familial cases (See supplementary material and methods).^{7,9} We only considered variants classified as pathogenic or likely pathogenic according to ACMG

guidelines.⁶ Genes of unknown significance not previously related to Mendelian phenotype and identified in this study have been prioritized using *in silico* prediction softwares, ovarian expression, segregation studies, animal models, and interactions with other known genes causing POI.

- b) Copy Number Variation (CNV) were studied using two methods:

- For patients studied with WES, we used the Bioconductor DNACopy package (DNACopy 1.32.0) and compared control exome data to reference samples pool (DOI: 10.18129/B9.bioc.DNACopy). It implements the circular binary segmentation (CBS) algorithm to segment DNA copy number data. All changes were annotated with the catalog of the Database of Genomic Variants (DGV) to provide a comprehensive summary of structural variation in the Human genome.
- For patients studied with targeted NGS, we used an in-house coverage-based pipeline to detect CNV.¹⁰ This method is based on the Read Depth (RD) or the Read Count (RC), which reflects the abundance of the genome segment. RD is defined as the number of reads covering a given position in the sequence and RC is defined as the number of reads in a given region. Using these concepts and hypothesizing that the RD or RC is proportional to the initial abundance of the genetic material used for sequencing, we can detect potential CNV.

Chromosomal analysis

Hypersensitivity to the chromosome-breaking effect of crosslinking agents is a reliable marker of homologous recombination (HR) efficiency.¹¹ For POI genes involved in DNA repair and not expressed exclusively in germ cells, chromosomal breakage studies were performed in cultures of peripheral lymphocytes obtained from the patient, the mother when available, and a healthy woman as a control. The experiments were performed at the Gustave Roussy Institute (Villejuif, France), following a standard in-house protocol.^{7–9,12} Fresh peripheral blood lymphocytes were cultured under standard conditions for karyotyping. DNA damage was induced by treating the cells with mitomycin C (MMC; Sigma) for 48 h to examine cellular hypersensitivity to DNA crosslinking agents. Three treatment conditions were used: without MMC to analyze spontaneous DNA damage and with 150 or 300 nM MMC as recommended.¹¹ Chromosome breaks were scored by an experienced cytogeneticist using at least 20 metaphases.

Role of the funding source

The study funders had no role in the study design of this study, data collection, data analysis, data interpretation, or report writing.

Results

Cohort characteristics

The characteristics of the patients recruited in this study are summarized in Figure 1, Figure S1, and Table S2. The cohort of POI comprises 375 patients (344 index patients and 31 affected mothers or sisters): 26.7% were familial and 73.3% were sporadic, 24.8% of patients had PA, 65.3% SA, and 9.9% SP. Targeted NGS was performed in 345 patients and WES in 30 patients. Twelve patients (12/375; 3.2%) had syndromic POI with additional extra-ovarian features (Figure S1, Table S3).

Molecular findings

In our whole cohort, we identified 216 variants in 215 patients (out of 375) (Figure 2, Tables S4-S7). In fourteen patients, we found strong evidence of pathogenicity for nine genes not previously related to Mendelian phenotype or POI (Table 1). Using the ACMG criteria for genes known to be implicated in POI, pathogenic (PV) or likely pathogenic (LPV) were identified in 110 patients. The diagnostic performance of our NGS study with the ACMG criteria⁶ including only PV/LPV was 29.3% (110/375) for the whole cohort and 26.3% (61/232) for European patients ($n = 232$, 61.9% of the cohort) (Table S4). For isolated POI it was 28.4% (103/363 patients), and 58.3% for syndromic POI (7/12) (Table S3).

The diagnostic yield of targeted NGS is 28.7% (99/345) in the whole cohort, and 25.8% (57/221) in the European population. The diagnostic yield of WES is

36.7% (11/30) in the whole cohort and 36.4% (4/11) in the European population.

Most genes ($n = 24$) were identified in single patients. Only nine genes were found in two different patients/families and seventeen “recurrent” genes were found in at least three patients/families (Figure S2) confirming the high genetic heterogeneity in POI.

In 25 other patients (25/375; 6.7%), only heterozygous PV/LPV were found in known autosomal recessive POI genes (Figure 2). This may either indicate a new mode of inheritance for these variants (as for *BMP15* or *FOXL2* known to have both autosomal dominant and recessive inheritance¹), or most likely correspond to a carrier status (Table S7).

The genes displaying PV/LPV are involved in different pathways (Figure 2). Remarkably, 37.4% of genes are involved in meiosis/DNA repair or mitosis making this family the major family involved in POI, 35.4% are involved in follicular growth, 19% in metabolism and mitochondrial functions, Ovarian development (6.1%), NF- κ B pathway (1.4%), Autophagy (0.7%). (Figure 2).

We detected a CNV in a patient with POI and primary amenorrhea (ID 6). It was a 3.2kb-heterozygous deletion encompassing exons 10-17 of *STAG3* (Figure S3). The patient carried a heterozygous pathogenic variant of *STAG3*: NM_012447.4:c.3046delT:p.Ser1016-ProfTsTer16. Parental samples were not available to perform segregation studies. We thus considered this patient as a presumed compound heterozygous.

For all other patients, especially those identified as carriers in our study, carrying a pathogenic or likely

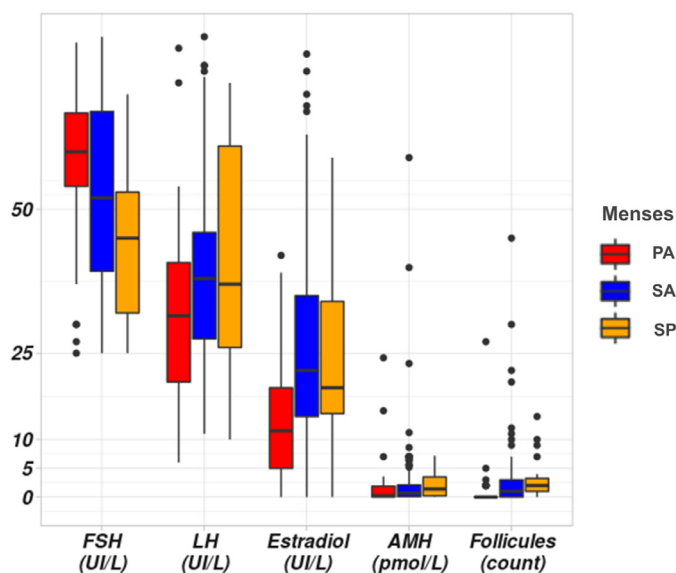


Figure 1. Hormonal and ultrasonographical datas in our cohort of patients with POI. Hormonal assays (FSH, LH, Estradiol and AMH) in patients according to menses. PA: Primary Amenorrhea, SA: Secondary Amenorrhea, SP: Spaniomenorrhea. The Y scale unit is variable according to each hormonal assay represented in the X-axis. Normal range: AMH (7-20.7 pmol/l); Respectively follicular, ovulatory, luteal phases and menopause: FSH (IU/L): (2.9-12), (6.3-24), (1.5-7), (17-95); LH: (IU/L) (1.5-8), (9.6-80), (0.2-6.5), (8-33); E2 (ng/l): (19.5-144.2),(63.9-356.7),(55.8-214.2),(< 32.2).

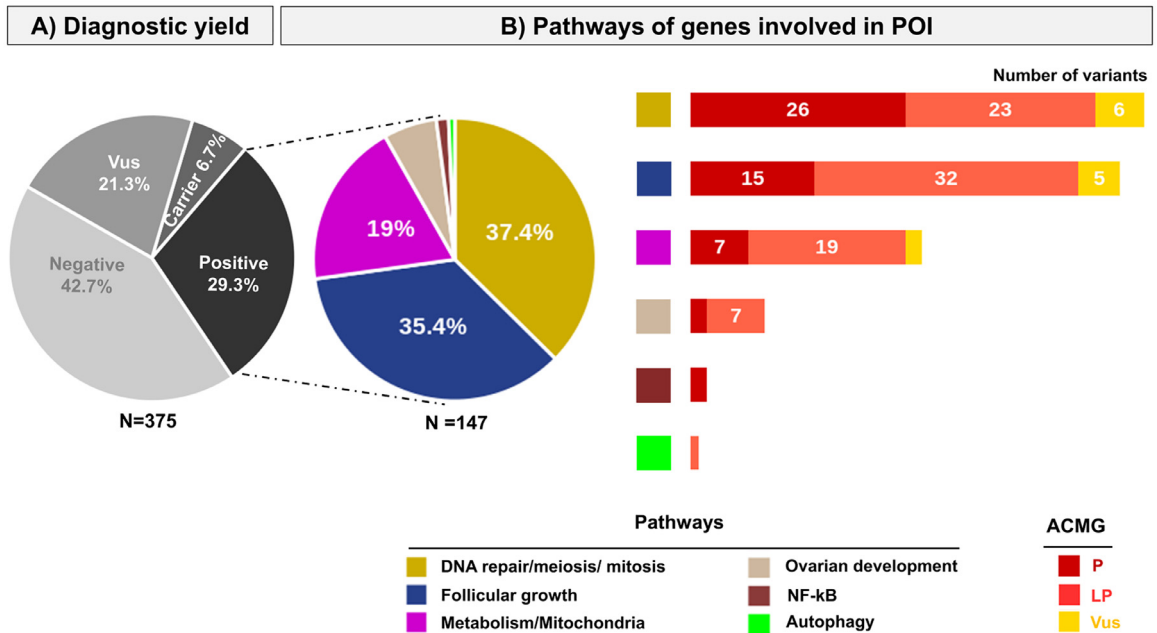


Figure 2. Genetic studies of the cohort of patients with POI. Patients were studied by a custom-made targeted NGS comprising 88 genes or by whole exome sequencing (see methods). **A) Diagnostic yield using ACMG criteria:** Variants are classified according to the American College of Medical Genetics (ACMG) guidelines. N= 375: the whole cohort comprises 375 patients with POI. Vus: Variant of unknown significance. Carrier: patients harbouring a heterozygous pathogenic variant in a known autosomal recessive POI gene. Positive: the diagnostic yield corresponds to patients carrying pathogenic (P) or likely pathogenic (LP) variants and is 29.3%. **B) Pathways of genes involved in POI.** The different pathways are indicated with different colors. The pie chart represents the proportion of patients with P or LP variants in a specific pathway: DNA repair meiosis and mitosis (37.4%), Follicular growth (35.4%), Mitochondria and Metabolism (19%), Ovarian development (6.1%), NF-kB (1.4%), Autophagy (0.7%). The histograms show the number and type of variants detected in each pathway.

pathogenic heterozygous variant in known autosomal recessive POI genes (n=19), no CNV was detected.

I. Identification of genes and pathways not previously related to Mendelian phenotype or POI. Twelve candidate genes were identified in our cohort. Nine candidate genes are not previously related to a Mendelian phenotype or POI. For *XPNPEP2*, there is to date only a balanced translocation (X; 12) presumed to interrupt this gene that has been described in a POI patient and her affected mother.¹³ The variants are mentioned in [Table 1](#) and the family pedigrees are shown in [Figure 3](#). We previously reported two genes, *FANCM* and *BRCA2*, involved in the RAD51 pathway causing non-syndromic POI.^{9,12}

a) NF-kB signalling and regulation of inflammatory responses. *NLRP11*: In a Tunisian patient with POI and SP at the age of 20 years, we identified two compound heterozygous variants of *NLRP11* encoding the Nod-like-receptor protein (NLRP)11: a nonsense substitution in exon 6 (NM_145007.4: c.1867A>T), introducing a stop codon (p.Arg623Ter) and a missense substitution in

exon 8 (NM_145007.4: c.2206G>A: p.Glu736Lys). The NLRP protein family regulates the immune response to infection through NF-kB and type I interferon signalling pathways.¹⁴ NLRP11 is highly expressed in the testes and ovary as are many other NLRP proteins.¹⁴ Remarkably, NLRP5 has been involved in POI in human and mice.¹⁵ Moreover, a very recent study based on mouse models highlights the important role of inactivated NF-kB in POI.¹⁶

b) RNA binding protein and posttranscriptional/translational regulations.

ELAVL2: In a Polish family, two sisters developed POI with SA at 20 and 24 years. WES revealed compound heterozygous variants in the RNA binding domain of *ELAVL2* (NM_001351472.2): c.448C>T, Arg150Cys and c.313C>T, Leu105Phe, the latter transmitted by the mother. *ELAVL2* encodes a RNA binding protein, mainly expressed in ovaries, testes and neurons. It is involved in post-transcriptional and post-translational regulations.¹⁷ Mice deficient in *Elavl2* are sterile and ovaries are devoid of follicle.¹⁷

c) Meiosis/ DNA repair genes/ mitosis.

C17orf53 (HROB): In two sisters from a consanguineous Turkish family, both presenting PA and streak gonads,

ID	Ethnicity	Age	Menses	Gene	variant	Status	Pathway	Expression	Animal or in vivo models
258	Turkish	15	SA	C18orf53 (HROB)	NM_024032.5: c.502delG:p.Glu168ArgfsTer35	Hom	DNA repair & Meiosis	Ubiquitous	C18orf53-/- mice are infertile ¹⁸
311	Turkish	16	PA						
310	North-African	24	SA	HELQ	NM_133636.5: c.3095delA: p.Tyr1032SerfsTer4	Hom	DNA repair & Meiosis	Ubiquitous	Helq-/- mice are infertile ¹⁹
373	North-African	20	SA						
317	North-African	16	PA	SWI5	NM_001318092.1: c.261-1G>C:p.?	Hom	DNA repair & Meiosis	Ubiquitous	Impaired meiosis in in yeast ²⁰
302	Asian	29	SA	CENPE	NM_001813.2: c.2023C>T:Q675Ter	Het	Cell Cycle	Ubiquitous	Embryonic lethality of Cenpe-/- mice ²³
303	Asian	14	SA						
304	Asian	34	SA						
311	North-African	16	PA	SPATA33	NM_153025.2: c.34dupT:p.Cys12LeufsTer2	Hom	Autophagy	Gonadal	Spata33 knockout suppresses mitophagy in germ cells ²⁴
270	North-African	20	SP	NLRP11	NM_145007.4: c.1867A>T: p.Arg623Ter/ c.2206G>A:p.E736K	Comp Het	Immunity-Inflammation	Ubiquitous	Reduced fertility in Nrlp11-/-mice ¹⁶
306	North-African	15	SA	CCDC150	NM_001080539.2: c.291_292delTG :p.Cys97Ter	Hom	Unkown	Gonadal	No available animal model
314	Turkish	17	SA	CCDC185	NM_152610.3: c.1174C>T :p.Gln392Ter	Hom	Unkown	Gonadal	No available animal model
318	European	20	SA	ELAVL2	NM_001351472.2:c.448C>T / :p. Arg150Cys / c.313C>T :Leu105Phe	Comp Het	Post-transcriptional regulation	Gonadal & neurons	Female Elavl2-/- mice are infertile devoid of follicles ¹⁷
319		24	SA						

Table 1: Variants in genes not previously related to Mendelian phenotype or POI.

PA: Primary amenorrhea; SA: secondary amenorrhea; Hom: homozygous; Het : heterozygous; Comp Het : compound heterozygous.

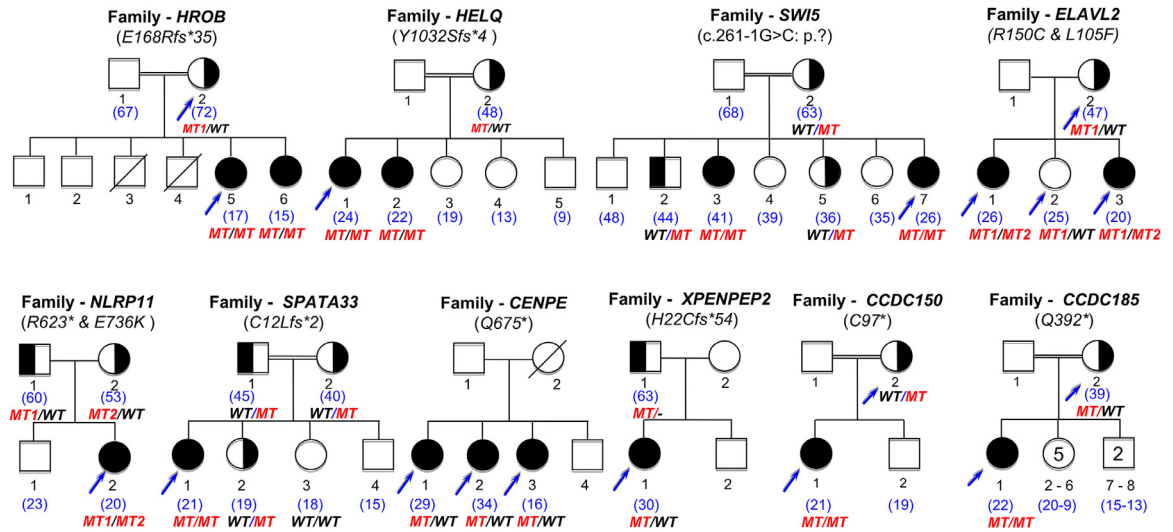


Figure 3. Pedigrees of the families with genes not previously involved in Mendelian phenotype or POI. On top of each family, the name of the gene is indicated, with the variant found. Double lines indicate consanguineous unions. The age of the patient at diagnosis is indicated in blue between brackets. Blue arrows indicate the patients and relatives studied with NGS (see text and Methods chapter). All family members available were studied by Sanger Sequencing. The segregation of the variants is shown under each individual sequenced. MT: mutated. MT1: first mutation. MT2: second mutation. WT: wild type. Family CCDC185: the number inside each symbol indicates the number of siblings in the family, either sisters (5) or brothers (2).

we identified a homozygous truncated variant of *HROB* (NM_024032.5: c.502delG; p.Glu168ArgfsTer35), found also homozygous in her affected sister and heterozygous in both parents. *HROB* (C17orf53) encodes a newly identified protein involved in homologous recombination (HR).¹⁸ It corresponds to a factor containing an oligonucleotide/oligosaccharide (OB) fold that recruits MCM8 and MCM9—two proteins that, when defective, cause POI—to the site of DNA damage. Mice deficient in *Hrob* are infertile with small uteri and ovaries completely devoid of follicle.¹⁸ Chromosome breakage studies revealed high chromosomal instability with spontaneous breaks, enhanced in the presence of mitomycin (MMC), and numerous radial figures, similar to what is observed in patients with Fanconi Anemia (FA) strongly supporting a defect in HR (Figure 4, Table S8).

HELQ: A patient born to consanguineous Moroccan parents presented SA at the age of 20 years. WES identified a homozygous truncated variant in exon 17 of *HELQ* (NM_133636.5: c.3095delA; p.Tyr1032SerfsTer4), heterozygous in her mother, who still had regular menses at the age of 50 years and homozygous in her affected sister. *HELQ* encodes a DNA helicase that plays a pivotal role in DNA replication, recombination and inter-strand crosslink repair.¹⁹ *HELQ* deficient mice exhibit subfertility and tumour predisposition.¹⁹ The orthologues of this protein in *Drosophila melanogaster* and *Caenorhabditis elegans* are required for meiotic double-strand break repair.¹⁹ Concordantly, chromosomal studies in the patient revealed spontaneous chromosomal breaks markedly enhanced with MMC (Figure 4, Table S8).

SWI5: The probanda (25 years) and her sister (41 years), born to consanguineous Moroccan parents, had POI with PA and streak ovaries devoid of follicles. The mother had menopause at the age of 52 years. WES in the probanda identified a homozygous splice variant in the canonical GT acceptor splice site of intron 3 of *SWI5* (NM_001318092.1: c.261-1G>C), also homozygous in the affected sister and heterozygous in unaffected siblings and parents. This variant is absent in all available databases. *In silico* predictions with all available softwares (Human Splice Finder, dbSNV, Alamut and SpliceAI) of the splice variant predicts a high impact on splicing. *SWI5* encodes a protein part of an auxiliary complex, Swi5-Sfr1 that regulates HR through the RAD51 pathway, synergistically with the HOP2-MND1 complex.²⁰ Interestingly, *PSMC3IP* (encoding HOP2) and *MND1* cause POI in humans.^{1,21} Furthermore, an orthologue of SWI5-SFR1 in yeast (*Schizosaccharomyces pombe*)—*Sae3-Mei5*—is required for meiotic HR.²⁰ Concordantly, the patient’s cells exhibit high MMC-induced chromosomal breaks (Figure 4, Table S8). All available studies on Swi5-Sfr1 have only been performed *in vitro* or in yeast, and so far, it is not known whether SWI5 plays a role in mammalian HR and meiosis.^{20,22} The family we report here provides a unique opportunity to observe the *in vivo* effect of a high impact bi-allelic molecular defect of *SWI5*, confirming its functional importance in humans. The role of SWI5 in meiosis is consistent with the phenotype of both sisters in our family, with early POI and ovarian atrophy. SWI5 could also act in a complex with SFR1 in humans, since the interaction between the two molecules is predicted to be

A)

	Gene	Genotype	Aberrant metaphases			Breaks/ metaphase			Radial figures		
			Untreated	MMC (150 nM)	MMC (300 nM)	Untreated	MMC (150 nM)	MMC (300 nM)	Untreated	MMC (150 nM)	MMC (300 nM)
Control	WT	WT	0/50 (0%)	1/50 (2%)	1/50 (2%)	0	1/50 (0,02)	7/50 (0,14)	0/50 (0%)	0/50 (0%)	0/50 (0%)
Patient 1	HELQ	Hom	3/50 (6%)	23/50 (46%)	41/50 (82%)	4/50 (0,08)	64/50 (1,28)	199/50 (3,98)	1/50 (2%)	8/50 (16%)	26/50 (52%)
Patient 2	HROB	Hom	5/50 (10%)	43/50 (86%)	19/19 (100%)	6/50 (0,12)	190/50 (3,8)	309/19 (16,26)	0/50 (0%)	numerous	extremely numerous
Patient 3	SWI5	Hom	0/50 (0%)	19/50 (38%)	45/50 (90%)	0	40/50 (0,8)	450/50 (9)	0/50 (0%)	4/50 (8%)	49/50 (98%)
Patient 4	Fanconi anemia	Hom	4/50 (8%)	48/50 (96%)	30/30 (100%)	4/50 (0,08)	437/50 (8,74)	>30	0/50 (0%)	numerous	extremely numerous

WT: wild type, Hom: Homozygous

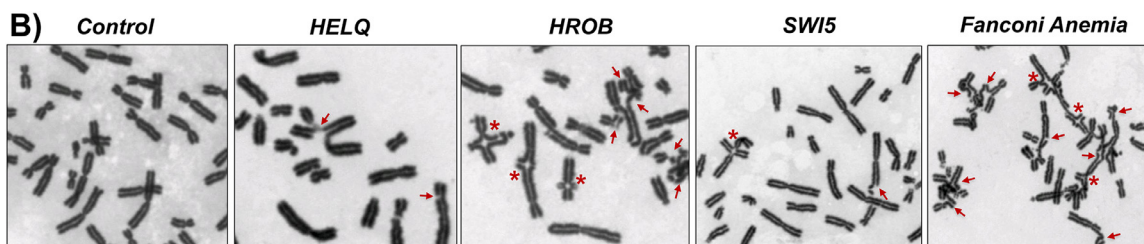


Figure 4. Chromosomal instability in patients with molecular defects of DNA repair genes. A) Chromosomal breaks analysis of the patient's lymphocytes, a control (a WT fertile woman), and a patient with Fanconi anemia, in the presence of increasing concentration of mitomycin (MMC). **B)** Selected figures of metaphases from the patients with molecular defects of SWI5, HELQ, HROB compared to Fanconi anemia and control cells in the presence of 300 nM of MMC. Chromosomal breaks are shown with red arrows and radial figures with asterisks.

In the absence of MMC, while no spontaneous breaks are observed in cells of the patient with the SWI5 homozygous splice variant, respectively 6% and 10% of cells of the patients with homozygous truncated variants of HELQ and HROB presented increased breaks, similarly to cells of the patient with Fanconi anemia (8%). In the presence of 150nM MMC, 86% of cells with the HROB pathogenic variant presented breaks with 3.8 breaks per metaphase very similarly to cells of the patient with Fanconi anemia (96%) and radial figures were observed in numerous cells of both types. Increased breaks were also observed in 46% of cells with the HELQ pathogenic variant (1.28 breaks per metaphase) and 38% of cells with the homozygous SWI5 variant (0.8 breaks per metaphase). Radial figures were observed in 16% and 8% of cells respectively. At 300 nM MMC, HROB-cells mimic Fanconi anemia cells, all presenting breaks (16.26 breaks per metaphase) with numerous radial figures, while 90% of SWI5-cells presented breaks with a mean of 9 breaks per metaphase and 82% of HELQ-cells with a mean of 3.98 breaks per metaphase. HROB-cells mimic Fanconi anemia cells with spontaneous chromosomal instability and marked hypersensitivity to MMC while HELQ-cells and SWI5-cells have a milder phenotype, but clearly with increased breaks when compared to control cells, and with radial figures, a pathognomonic feature of Fanconi anemia.

abolished in our patients. Our results thus reinforce the crucial role of the Swi5-Sfr1 complex in meiosis and provide a strong argument for extending its role to the human species.

CENPE: This study includes a family with mixed origin (Asian and European) in which three sisters presented POI with SA at 16, 29 and 34 years. WES identified a heterozygous, stop-gain variant of *CENPE* (NM_001813.2: c.2023C>T; Q675Ter). *CENPE* encodes the centromeric protein E, a kinesin-like motor protein essential for correct chromosome segregation in both meiosis and mitosis and in genome stability. It is highly expressed in germ cells. In mice this protein has an essential role in meiosis and in maturing oocytes.²³

d) Mitochondrial autophagy (Mitophagy).

SPATA33: In a patient with POI and PA born to consanguineous Algerian parents, WES studies identified a

homozygous insertion of a single nucleotide in the exon 1 of *SPATA33* (NM_153025.2: c.34dup; p.Cys12LeufsTer2), heterozygous in her parents and one unaffected sister. The other unaffected sister inherited both wild-type alleles. *SPATA33* is a protein exclusively expressed in mitochondria germ cells.²⁴ Recent *in vitro* studies performed on mouse testes revealed that this protein is localized in mitochondria and has been described as a novel autophagy mediator for mitophagy. This suggests a potential role of mitophagy in POI (mitochondrial autophagy).²⁴

e) Gene involved in peptide processing.

XPNPEP2: In a French patient presented POI and SA at the age of 30 years, we identified a paternal heterozygous frameshift deletion in exon 2 of *XPNPEP2* (NM_003399: c.63_94del; p.His22CysfsTer54). *XPNPEP2* encodes APP2, an aminopeptidase, ubiquitously expressed that removes N-terminal amino-acids

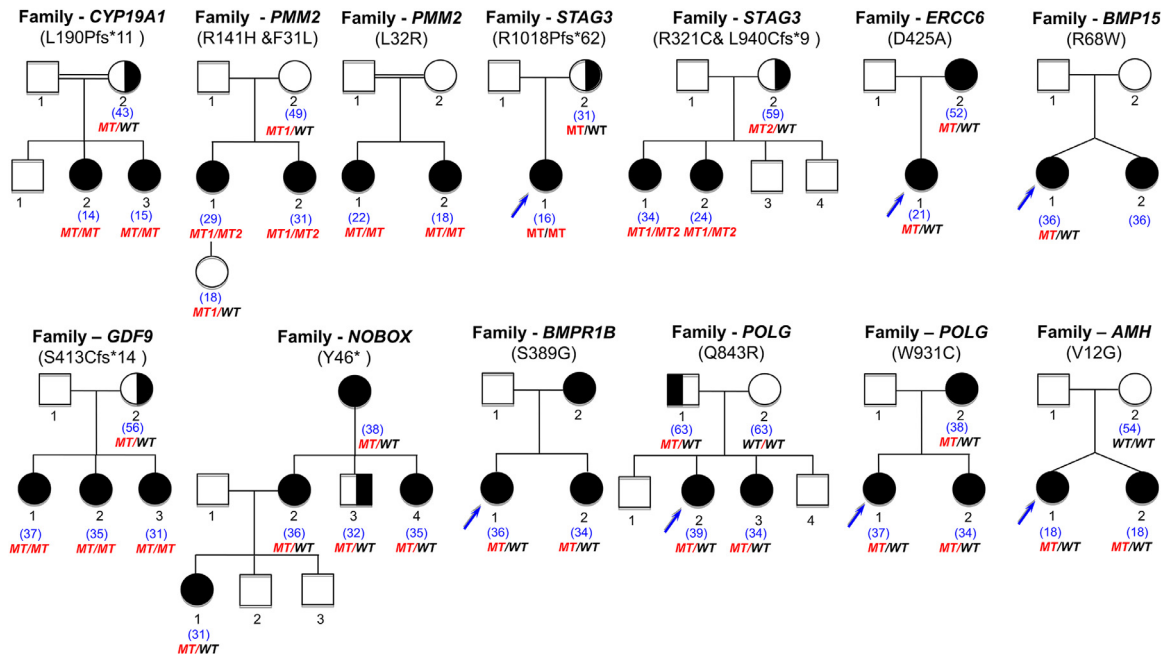


Figure 5. Pedigrees of the family confirming the causal role of genes in POI. Double lines indicate consanguineous union. Blue arrows indicate the patients and relatives studied with NGS, The other family members available are studied by Sanger sequencing. The segregation of the variant is shown under each sequenced individual. MT: Mutated, MT1: first mutation, MT2, second mutation, WT: wild type. The age of patients at diagnosis is indicated in blue between brackets.

from peptides with a penultimate prolyl residue (as Arg-Pro-Pro)²⁵ notably collagen that regulate follicle growth.²⁵ In the rat ovary, it was suggested that XPNPEP2 regulates collagens formation.²⁵ To date, there is only a single report of a balanced translocation (X;12) that interrupts *XPNPEP2* in a patient and her mother, who both presented with SA and POI.¹³ This point PV of this gene that we identified here validates its causal role in POI.

f) Genes with unknown function. Interestingly two genes belonging to the coiled-coil family with unknown function were identified in two unrelated patients.

CCDC150: In a consanguineous Moroccan patient with primo-secondary amenorrhea at the age of 15 years, WES identified a homozygous two-base-pairs deletion in exon 2 of *CCDC150* (containing 28 exons): NM_001080539.2: c.291_292delTG: p.Cys97Ter. This gene is highly expressed in the testes, according to the GTex database. Of note, only adult ovaries are included in GTex. Other genes exclusively expressed in testes according to GTex were in fact also expressed in foetal ovaries such as *STAG3*.¹ Very little is known about this protein. However, a strong interaction is predicted between *CCDC150* and *RAD50*, a major protein involved in homologous recombination, using String software (<https://string-db.org/>).

CCDC185 (C1orf65): A Turkish patient born to consanguineous parents had POI and early SA at the age of 17 years. WES identified a homozygous stop codon in

the unique exon of *CCDC185* (NM_152610.3: c.1174C>T: p.Gln392Ter), heterozygous in the mother. This gene is exclusively expressed in the testes according to the GTex database.

II. Involvement of genes responsible for syndromic POI in patients with apparently isolated forms.

Unexpectedly, in 32 (8.5%) patients initially presenting as isolated POI, we identified PV/LPV in genes known to cause syndromic POI (Table S7, S9). One patient with POI and early SA had the G188Q homozygous PV of *GALT*, the most frequent mutation causing galactosemia.²⁶ It is a severe, life-threatening metabolic disease often diagnosed at birth and fatal without specific regimen. Enzymatic assays confirmed the diagnosis. Reverse phenotyping revealed only a cataract during infancy treated by surgery. Our patient survived to adulthood without any specific medical care unlike all previously reported adult patients with an early childhood diagnosis who had a continuous adapted regimen. Eight patients inherited genetic defects in genes causing pulmonary arterial hypertension (PAH): *CAV1* (n=1), *BMPR2* (n=2) or *BMPR1B* (n=5).²⁷ *BMPR2* and *BMPR1B* encode a transmembrane receptor, member of the bone morphogenetic protein (BMP) receptor family and *CAV1* encodes caveolin that has a crucial role in signal transduction and vesicular trafficking.²⁷⁻²⁹ All these proteins are expressed both in lungs and ovaries especially in granulosa cells and display functional roles in both tissues.^{1,27,28} Deficient *bmp15*^{-/-} mice are subfertile with decreased ovulation and fertilization rates while *gdf9*^{-/-}

mice are sterile.²⁸ *Cav1*^{-/-} null mice show reduced fertility.²⁹ The absence of PAH in particular in patients with variants of *BMPR2* may be explained by several mechanisms: i) the type of the variant; indeed, 50 to 75 % of the *BMPR2* variants detected in patients presenting PAH were truncated variants while in POI only missense variants are found^{30,31}; ii) the localization of the variant: the majority of the *BMPR2* variants detected in patients with PAH are located in the kinase domain of the protein (encoded by exon 6-10)³⁰ while three out of the four *BMPR2* variants detected in POI patients are localized in the cytoplasmic tail of *BMPR2* (the L656S and R597Q variants described here and the previously reported S987F variant).³² It was reported that patients carrying a mutation affecting the cytoplasmic tail of *BMPR2* were characterized by an older age at diagnosis compared with other *BMPR2* mutation carriers³³ and iii) the low penetrance of the mutation observed for PAH genes in particular for *BMPR2*.^{27,31} The penetrance of *BMPR2* mutation is estimated to be between 20 and 43%.^{27,31} A 20-year-old patient had a LPV in *BMPR1A*. This gene predisposes to hamartomatous polyposis, absent in our patient. Seven patients had genetic defects in *POLG* encoding the mitochondrial DNA polymerase, without associated neurological or ocular symptoms. We report four supplementary variants of *TP63* (one splice variant and three missense variants), a gene causing both isolated and syndromic POI with dominant inheritance.³⁴ *TP63* variants described in POI are usually located in the C-terminal region.³⁴ However, missense and truncated variants in the different domains of *TP63* causing POI have been described.³⁵ Our data confirm that pathogenic or likely pathogenic variants in both the 5' end (c.192+2T>G) of *TP63* or in the N-terminal (p.Ile225Thr, p.Lys214-Glu) or the C-terminal (p.Thr566Met) regions of *TP63* cause POI. One patient had a heterozygous LPV of *TWINK*, a mitochondrial helicase implicated in progressive external ophthalmoplegia with both dominant and recessive mode of inheritance.³⁶ In all these cases, a comprehensive evaluation of patients and their families was recommended with appropriate follow-up in a multidisciplinary team.

III. Confirmation of the causal role of genes in POI. In 26 patients, we identified PV/LPV in thirteen POI genes previously described in single patients/families (Figure 5, Table 2). This confirmed their role in the pathogenesis of POI worldwide. We showed recently that a patient harbouring a homozygous *SPIDR* truncating variant exhibited a strong MMC-induced chromosomal fragility. This supported a role of *SPIDR* in the RAD51 pathway and HR.⁸

IV. Syndromic POI

- Known syndromic Genes identified

In our cohort, 12 patients (12/375 =3.2%) had syndromic POI (Table S3). We identified the genes involved in seven patients (7/12; 58.3%). In all these cases, genetic diagnosis led to a complete screening for other signs potentially associated with each syndrome, and to familial segregation studies.

- Possible syndromic POI not previously described

Interestingly, our series included four patients with new syndromic phenotypes indicating previously unreported syndromes (Table S3). They include: 1) keratoconus, 2) horseshoe kidney and epilepsy, 3) Marfanoid habitus, scoliosis and hyperlaxity and 4) single kidney, Hashimoto's thyroiditis, ptosis, and scoliosis. Genes causing these syndromic POI phenotypes remain to be discovered.

V. Patients with variants in two different genes. In a small proportion of patients (8/375; 2.1%), we identified P/LPV in two different genes (Table S10). In these patients, however, one of the mutated genes alone was sufficient to cause POI. Therefore, we did not find evidence of di/multigenic inheritance of POI in our cohort. Moreover, each exome is known to contain at least hundreds of rare variants, and it is not surprising to find heterozygous pathogenic or likely pathogenic variants in more than one POI gene. A recent study based on 6,447 exomes of healthy, genetically unrelated Europeans of two distinct ancestries estimates that each individual is a carrier of at least two pathogenic variants in currently known autosomal recessive genes.³⁷

Discussion

Pathways and candidate genes involved in POI not previously described in humans

Our study shows the powerful approach to identify genetic etiology and pathways not previously implicated in POI. This allows better understanding of the pathophysiology of POI and opens the way to the development of novel therapeutic targets.

The immune pathway has a major role in the ovary and may explain 4–31% of POI cases.¹ However, no specific gene has been found in isolated POI. AIRE causes the APECED syndrome (Autoimmune polyendocrinopathy candidiasis ectodermal dystrophy).¹ We involve in this study *NLRP11* a gene of the immune NF-κB pathway in isolated POI not previously implicated. Environmental factor especially phthalates, bisphenol A, pesticides have a negative impact on ovarian function, yielding increased follicular depletion.¹ Interestingly, the NF-κB pathway is involved in the production of TNF-α and IL-6 in response to bisphenol A and may also be a therapeutic target.³⁸

We also describe novel mechanisms in POI revealed by the genetic defect of *ELAVL2*, involving post-

ID	Ethnicity	Menses (age)	Gene	Variant	Status	ACMG	Pathway
321	North-African	PA	BRCA2	NM_000059.4:c.8350C>T :p.Arg2784Trp	Hom	P	DNA repair & Meiosis
167	European	SA (18)	FANCM	NM_020937.4:c.5101C>T :p.Gln1701Ter / c.1528G>A : p.Gly510Ser	Pres Comp Het	P/LP	DNA repair & Meiosis
			ATM	NM_000051.4:c.6596_6597delCT : p.Ser2199Ter	Het	P	DNA repair & Meiosis
192	Finnish	PA	FANCM	NM_020937.4:c.5101C>T :p.Gln1701Ter	Hom	P	DNA repair & Meiosis
305	Finnish	PA	FANCM	NM_020937.4:c.5101C>T :p.Gln1701Ter	Hom	P	DNA repair & Meiosis
306	Finnish	PA	FANCM	NM_020937.4:c.5101C>T :p.Gln1701Ter	Hom	P	DNA repair & Meiosis
326	European	SP (20)	FANCM	NM_020937.4:c.5101C>T :p.Gln1701Ter / c.575A>T : p.Gln192Leu	Comp Het	P/LP	DNA repair & Meiosis
323	European	SP (17)	MSH4	NM_002440.4:c.2198C>A : p.Ser733Ter	Hom	P	DNA repair & Meiosis
59	Asian	PA	SPIDR	NM_001080394.4: c.814C>T: p.Arg272Ter	Hom	LP	DNA repair & Meiosis
55	North-African	SA (39)	BNC1	NM_001717.4: c.2319C>A:p.Asn773Lys	Het	LP	DNA repair & Meiosis
88	European	SP (18)	ERCC6	NM_000124.4: c.1996C>T : p.Arg666Cys	Het	LP	DNA repair & Meiosis
109	European	SP (27)	ERCC6	NM_000124.4: c.1996C>T : p.Arg666Cys	Het	LP	DNA repair & Meiosis
79	European	SA (19)	ERCC6	NM_000124.4: c.1274A>C :p.Asp425Ala	Het	LP	DNA repair & Meiosis
105	European	SA (30)	ERCC6	NM_000124.4: c.1274A>C :p.Asp425Ala	Het	LP	DNA repair & Meiosis
211	Middle East	SA (37)	ERCC6	NM_000124.4: c.1274A>C :p.Asp425Ala	Het	LP	DNA repair & Meiosis
140	European	PA	BMPR1A	NM_004329.3: c.1327C>T : p.Arg443Cys	Het	LP	Follicular growth
155	European	SA (22)	BMPR1B	NM_001256793.2: c.761G>A : p.Arg254His	Het	LP	Follicular growth
156	European	SA (22)	BMPR1B	NM_001256793.2: c.761G>A : p.Arg254His	Het	LP	Follicular growth
267	North-African	SA (33)	BMPR1B	NM_001256793.2: c.1165A>G : p.Ser389Gly	Het	LP	Follicular growth
268	North-African	SA (35)	BMPR1B	NM_001256793: c.1165A>G:p.Ser389Gly	Het	LP	Follicular growth
			PCCB	NM_000532: c.646A>G:p.Met216Val	Het	LP	Metabolism
38	European	SA (16)	BMPR1B	NM_001256793.2: c.836A>T:p.Gln279Leu	Het	LP	Follicular growth
163	European	SA (21)	BMPR2	NM_001204.7: c.1967T>C : p.Leu656Ser	Het	LP	Follicular growth
275	European	PA	BMPR2	NM_001204.7:c.1790G>A : p.Arg597Gln	Het	LP	Follicular growth
			AMH	NM_000479.5:c.35T>G :p.Val12Gly	Het	LP	Follicular growth
325	European	SP (24)	ESR1	NM_001437.2:c.335C>A :p.Ser112Ter	Het	LP	Follicular growth
112	Turkish	SA (14)	CAV1	NM_001172895.1: c.1A>G: p.Met1Val	Het	P	Follicular growth
272	European	PA	RCBTB1	NM_018191.4: c.1271T>G :p.Phe424Cys / c.962C>T : p.Pro321Leu	Pres Comp Het	LP/LP	Follicular growth
168	North-African	SA (31)	ATG7	NM_006395.3: c.1478A>G : p.Lys493Arg	Het	LP	Autophagy

Table 2: Variants confirming the causal role worldwide of genes reported in single patients/families.

PA: Primary amenorrhea; SA: secondary amenorrhea; Hom: homozygous; Het: heterozygous; Comp Het: compound heterozygous; Pres Comp Het: Presumed compound heterozygous (parents not available); ACMG: Classification of variant according to American college of Medical genetic; P: Pathogenic, LP, Likely pathogenic. Patient 105 is the mother of patient 79; Patient 267 and 268 are sisters.

transcriptional and translational regulations in the ovaries, known to play a major physiological role.³⁹ *Elavl2*-mice are infertile. Interestingly, these two candidate genes (*ELAVL2* and *NLRP11*) have been recently associated with the age of natural menopause (ANM)⁴⁰ reinforcing the genetic link between POI and the physiological variance of the ANM (see further).

We found strong evidence of a causal role in POI for four DNA repair/meiosis/mitosis genes based on mice/yeast models, and/or chromosomal instability. A high chromosomal fragility was found in our patients with defects in three new DNA repair genes: *HROB*, *HELQ*, *SWI5*, the highest found for the homozygous truncation of *HROB*, similar to FA cells (Figure 3, Table S8). However, there was no sign of FA in our patients (anaemia,

cancer microcephaly, small size or café-au-lait spots). This chromosomal fragility may lead to an increased risk of malignancies and therefore a long-term follow-up by a multidisciplinary team is indicated.

In our study, we only performed MMC-induced chromosome breakage studies in patients with possible genetic alteration of double-strand DNA repair genes like *HELQ*, *SWI5* and *HROB* (*C17orf53*). Indeed, MMC-induced chromosomal breakage study only detects defects in homologous DNA repair and does not allow evaluating defects in other DNA repair pathways.⁴¹ Thus, we suggest a combined two-step approach in POI patients using sequentially an NGS study followed by a chromosomal breakage study in patients with variants in a gene known or supposed to be involved in

homologous recombination. This can improve variant classification.

Interestingly, we identified other candidate genes, *CCDC155* and *CCDC185*, exclusively expressed in gonads but with unknown function. Mice models are needed to unravel unknown pathophysiological mechanisms involved in POI. *SPATA33* has been recently suggested to be involved in autophagy. If confirmed in rodent models, it would support the implication of this crucial pathway in human follicular physiology and provides a future perspective on curing autophagy-related reproductive infertility diseases as for other linked diseases.⁴²

Performance of genetic diagnosis and impact on patients' management

Despite the large-scale use of NGS analysis in the era of genomic medicine, the study of genes involved in POI is often limited to the screening of the *FMR1* premutation (3.5% of cases).¹ We show here in our large cohort of 375 patients that NGS is an effective diagnostic tool for patients with POI. Using ACMG criteria, the rate of detection of pathogenic variants was 29.3% in the whole cohort and 26.3% in the European population (232 patients) (including only P/LP variants in known POI-genes. Variants in genes not previously related to POI or a Mendelian phenotype are considered of unknown significance-VUS as recommended⁶). This high yield supports a thorough genetic study of all unexplained POI patients in clinical practice. Very few studies were previously reported including either a small cohort (20–60 patients), a very limited number of POI genes (n=18),^{43–45} or reporting only heterozygous variants, the pathogenicity of which could not be formally demonstrated. Indeed no familial segregation studies were performed⁴⁶ and the rodent models deficient in these genes have a recessive mode of transmission.⁴⁶ Moreover, in these studies, authors also considered VUS or likely benign variants that cannot be used for a genetic diagnosis.^{43–47} Only variants classified as pathogenic or likely pathogenic can guide management of patients. Accordingly, we could not confirm the high involvement described up to 6.2% for some POI genes notably *NOBOX* or *BMP15*.^{43–45} Of note, the *R117W-NOBOX* variant is classified benign according to ACMG criteria and has a high Minor Allele Frequency-MAF up to 8.17% in the African population in GnomAD exome and genome databases. Efforts to improve adherence to these guidelines will be important to prevent erroneous misclassification of non-pathogenic variants in POI genetic testing and inappropriate diagnosis.

To date, both homozygous and heterozygous variants of *NOBOX* have been reported in patients with POI, some of which have been validated with *in vitro* functional studies.^{1,45,48–50} We report an additional truncating variant of *NOBOX* (Y46X) in two unrelated

patients (ID 143 and 331) (Figure 5). Unlike previous reports, we were able to perform a large familial transmission study in three generations in one family (family of the probanda with ID 331). The patient's mother, grandmother and maternal aunt, all with POI and secondary amenorrhea, harboured the Y46X variant of *NOBOX*. This strongly supports an autosomal dominant transmission of this truncated variant. Altogether these data argue that inheritance pattern of *NOBOX* is either autosomal dominant¹ or autosomal recessive^{49,50} with a genotype-phenotype correlation. Biallelic truncated variants of *NOBOX* cause POI and primary amenorrhea with a recessive mode of inheritance while heterozygous variants of *NOBOX* yield POI and secondary amenorrhea with an autosomal dominant transmission or haploinsufficiency.

The identification of the genetic cause has a major impact on genetic counselling and personalized patient care, with:

- i) *Psychological consequences for patients and their family*: as it reveals the causal mechanism for their infertility. Furthermore, offering a presymptomatic genetic testing in siblings may enable fertility preservation, if appropriate.
- ii) *Important information on the fertility prognosis by predicting a possible residual OR*. There is no direct method to evaluate this OR. AMH assays are negative when only small follicles are present in the ovaries.¹

We show that in 60.5% of our patients the genetic defect allows prediction of a persistent OR at an early stage: this includes patients with defects in genes involved in follicular growth (35.4%), metabolism and mitochondrial functions (19%), ovarian development (6.1%), autophagy (1 patient). Indeed, previous studies have reported positive AMH values and even pregnancies in patients with defects of these gene families, and the OR is also conserved in the corresponding rodent models.^{1,51} It should be noted that, concordantly, previous histological study revealed the presence of an OR in 57% of patients with POI.⁵²

For instance, we identified a mild form of galactosemia with a homozygous PV of *GALT* in a patient initially presenting with isolated POI. Such patients have 48% of chances of being spontaneously pregnant in two years²⁶ (90% in the general population). The patient stopped the oocyte donation program she had started. Other patients with complete *FSHR* or *BMP15* inactivation are known to have a preserved OR, although a rapid decrease can occur.^{1,52,53} Similarly, patients with POI and mild mutations of some meiosis/DNA repair genes: *FANCM*, *ERCC6*, *BNC1* may have spontaneous pregnancies, as found in our cohort.^{1,9} This may be due to a hypomorphic mutation or alternative compensatory pathways or to the fact that heterozygous mutations are

found with possible escape of oocytes to DNA damage. The genetic diagnosis of POI is thus a medical priority as soon as all other causes have been eliminated. Fertility preservation in a multidisciplinary team should be discussed without delay in these patients and their affected siblings when appropriate to avoid the rapid follicular atresia that may occur in such patients.^{1,53} The prediction of a residual OR could allow these patients to benefit from new in vitro activation- IVA techniques in the near future.⁵⁴ Variable results were obtained in small cohorts of unselected POI with a yield of 3.7 up to 21% of healthy newborns.⁵⁴ A genetic pre-screening of patients could markedly improve the success of IVA by excluding patients whose ovarian reserve is predicted to be absent or severely impaired.

Similarly, in male infertility⁵⁵ it has been proposed that a genetic diagnosis could predict positive testicular sperm extraction (TESE) or, on the contrary, may avoid unnecessary surgical interventions in case of major defects of DNA repair and meiosis genes.⁵⁵

By contrast, in 29.4% of our patients, often with PA, major defects in meiosis/DNA repair genes are found. A major alteration of the OR is predicted with a very poor fertility prognosis, as in previously reported patients.⁴ This is also confirmed by the corresponding animal models.⁵¹ A rapid referral of these patients to egg donation or adoption could avoid repeated IVF failures and unnecessary delay in their parental desire.

iii) *Prevention or treatment of possible comorbidities.* In our cohort, 45.9% of patients may have, or are at risk to develop, associated comorbidities, requiring a comprehensive assessment by a multidisciplinary team:

- α) Other organic defects are found in genetically revealed syndromic POI (8.5% of cases): in 32 patients initially referred as isolated POI, we identified pathogenic variants of genes causing syndromic POI. Similarly, Alhathal et al found that in 3.1% (9/275 patients presenting male infertility with azoospermia or severe oligozoospermia), infertility can be the sole or major phenotypic expression of genes that are known to cause multisystemic manifestations in humans.⁵⁶ In the same way, Kraucz et al found that 3/29 patients presenting infertility with Sertoli cell-only syndrome had biallelic variants of *FANCA* with only subtle haematological abnormalities corresponding in fact to occult FA.⁵⁷ The existence of potential defects of other organs, undiagnosed, requires complete evaluation of the patients and most often a long-term follow-up of patients and their families.
- β) Tumour/cancer in POI linked to meiosis/DNA repair genes (37.4% of cases): 37.4% of patients had alteration of meiosis/DNA repair genes or

mitosis. Although this gene family is the main contributor to POI, unexpectedly only six genes so far are known to cause POI and tumours/cancers in humans: *BRCA2*, *FANCM*, *MCM8/9*, *PSMC3IP*, *STAG3*.^{1,7,9,12} The majority of the corresponding deficient mice, however, developed tumours.⁵¹ Prospective studies in larger cohorts of patients will allow a more precise evaluation of the risk of tumours/cancers in these patients. Genetic counselling should be offered to patients and their families with long-term follow-up when adequate.

The identification of this latter subpopulation of patients with POI due to PV/LPV variants in cancer susceptibility genes is therefore a medical and scientific priority.

Overlap between genes implicated in azoospermia and POI

There is a remarkable overlap between genes involved in azoospermia and POI.⁵⁶ It is interesting to note that genes belonging to the meiosis/DNA repair family including genes of the synaptonemal complex is the leading family shared by POI and male infertility with azoospermia. This is the case for novel variants in genes identified in our study: *STAG3*,^{58–60} *SPIDR*,^{8,56,61} *PSMC3IP*,^{62,63} *HFM1*,^{56,64} *FANCM*.^{9,65} We can therefore expect the discovery of other genes causing both male and female infertility in the near future, belonging especially to the meiosis and DNA repair gene family.

Link between genes involved in POI and genes involved in the physiological reproductive lifespan

Epidemiological studies in various populations have evidenced a link between the age of natural menopause and lifespan or the odd ratio of all-cause mortality, including cancer.^{1,4,5} WES and genome-wide association studies (GWAS) have highlighted a genetic link between POI and the ANM, through the DNA repair/meiosis gene family, the latter also being involved in aging.^{4,5} Recently, a large GWAS study involving more than 200 000 women worldwide identified 290 loci involved in the ANM with an effect ranging from 3.5 weeks to 74 weeks (~1.2 years).⁴⁰ Very interestingly, three genes involved in POI in our study: *HELQ*, *ELAVL2* and *NLRP11* were also found to be associated with the ANM. This markedly supports the role of these candidate genes and pathways in the age of physiological and pathological menopause/POI, especially *ELAVL2* and *NLRP11*.

Conclusion

We describe here a highly successful genetic screening in a large cohort of patients with POI, allowing a genetic diagnosis in 29.3% of the patients. We showed that in 8.5% of cases, POI is the only phenotypic expression of

a complex genetic disorder affecting other organs. This led to an adapted genetic counselling and a personalized management of patients and families either for prevention/treatment of potential comorbidities (other organ defects, or tumours/cancers when DNA repair genes are involved) or for early fertility preservation when an OR can be predicted in the light of the future development of innovative treatments of IVA. Strong evidence involved 9 new genes in POI and, three previously undescribed pathways that may provide new therapeutic targets for the infertility of POI in the future. Identifying the causes of POI will greatly expand our knowledge of the mechanisms controlling fertility in women.

Limitations

This work involves the largest cohort of patients with POI studied by next generation sequencing to date. However, due to the relatively high prevalence of this condition (1 to 3.7% of women before the age of 40),^{1,2} a larger cohort could be studied in the future to better define the monogenic part of POI, ~30 % as shown in this study. This will: i) confirm the proportion of POI that can benefit from genetic diagnosis, ii) definitively implicate the nine genes identified here as candidate genes in POI. Indeed, these genes had never been associated with POI or a Mendelian disease, iii) to study the impact of a genetic diagnosis on the management of patients. Our study could encourage physicians to systematically prescribe NGS studies to all patients without an identified etiology, and not be limited solely to karyotyping and screening for the FMR1 premutation, the only genetic studies currently performed in most centres across the world. The Guidelines of International Societies in the field of Reproduction and Fertility may be updated in the light of these results. It should be noted that in our study, the segregation of the identified variants could only be achieved in 40% of cases because the parents were not available for the other patients. The absence of available trios has hampered the assessment of *de novo* mutations that might explain a subset of POI as has been observed in infertile men⁶⁶ or other conditions⁶⁷ and may improve the diagnostic performance of NGS studies in POI.

Contributors

Conceptualization: MM, CO. Formal analysis: AH, MM, NA. Funding acquisition: MM. Data Collection and Patient's Recruitments: AR, CO, SDC, IB, LBR, NF, CB, RR, KM, SL, MMa, ICD, AC, MP, BDH, SP, CD, CNB, HE, MK, CRL, MF, ELB, PL, EG, AZ, FA, FP, EBB, AB, LG, MBe, NG, SO, BGD, BI, LL, SH, AMGj, ABP, HF, MP, HL, SE, PBG, SW, SH, KA, SCJ. Resources: JP, CC. Clinical Investigation: AH, AMG, KA. Data Curation: AH, MBr, SC, MLRS. Methodology: AH, MM. Project administration: AH, MM, CO. Software: AH. Supervision: MM. Validation: MM. Visualization: AH. Writing-original draft: AH, MM.

Writing-review & editing: AH, MM and KA. Verification of the underlying data: AH, MM. All authors read and approved the final version of the manuscript.

Data sharing statement

Non-identifying data supporting the claims made in this paper are available upon request after its publication. Requests should be addressed to the corresponding authors of this paper: Prof Micheline Misrahi (micheline.misrahi@aphp.fr).

Declaration of interests

The authors declare no conflict of interest.

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Supplementary materials

Supplementary material associated with this article can be found in the online version at doi:[10.1016/j.ebiom.2022.104246](https://doi.org/10.1016/j.ebiom.2022.104246).

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