

Significance of rare variants in genes involved in the pathogenesis of Lynch syndrome

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Abstract. The molecular characterization of patients with Lynch syndrome (LS) involves germline testing to detect a deleterious mutation in one of the genes of the mismatch repair (MMR) pathway. To date, however, a large proportion of patients with a clinical suspicion of LS who undergo genetic testing do not show a germline pathogenetic variant in these genes. Germline DNA from 73 patients with a clinical suspicion of LS was examined with next-generation sequencing methods, using a multigene custom panel designed and standardized by our research group, that targets a set of 15 genes. Deleterious variants were identified in 5.6% of index cases, while unclassified variants were identified in 80.3% of probands. To evaluate the pathogenicity of these uncertain variants, the American College of Medical Genetics and Genomics criteria was used, also considering wherever possible the microsatellite instability (MSI) status detected on tumor tissues as pathogenic criterion. In this manner, 8 of these uncertain significance variants were classified as likely pathogenic variants. Notably, some of these likely pathogenetic variants were also identified in the *MLH3* gene that is a gene not routinely analyzed for cases with a clinical suspicion of LS. The present study highlighted the importance of verifying the pathogenicity of the numerous variants of unknown significance identified in patients for whom heredity is already clinically confirmed suggesting the importance of considering the MSI-H status on the tumor of patients carrying an uncertain variant to evaluate its pathogenicity. Moreover, the present study also suggested analyzing other MMR genes, such as *MLH3*, in panels used for the molecular screening of LS.

Introduction

Lynch syndrome (LS) is an autosomal hereditary form of colorectal cancer (CRC) with high penetrance and an incidence of 3-5% of all CRC cases (1-3). This hereditary form of CRC is clinically distinguishable from familial adenomatous polyposis (FAP), another hereditary form of CRC that develops from numerous polyps, as FAP is associated with APC mutations (4). Individuals with LS are characterized by a high lifetime risk of development of CRC, as well as endometrial, gastric and ovarian cancers, and other extracolonic tumors (such as small intestine, brain, skin, hepatobiliary, and urinary tract) (5,6). This hereditary syndrome is due to germline mutations in DNA mismatch repair (MMR) genes, mainly *MLH1*, *MSH2*, *MSH6* and *PMS2* (7-11).

The molecular characterization of these patients involves germline testing to detect a deleterious mutation in *MLH1*, *MSH2*, *MSH6* or *PMS2* (11), by direct sequencing and multiplex ligation-dependent probe amplification (MLPA), commonly used for the detection of large rearrangements (12). Numerous pathogenic mutations have been detected in *MLH1* and *MSH2* (13,14), while germline mutations in *MSH6* and *PMS2* are responsible for the disease only in a minority of cases (15-18). DNA MMR deficiency determines the presence of microsatellite instability (MSI) or loss of MMR protein expression at the somatic level (19). To identify patients with LS suitable for genetic testing, MSI testing and/or the lack of MMR protein expression (dMMR) according to immunohistochemical staining of colorectal or endometrial tumoral tissues is recommended for universal tumor screening, and these should be conducted as the first step (20,21). However, the Amsterdam criteria (AC) still remain a valid tool to allow clinicians to raise a clinical suspicion of LS in young patients with CRC and a significant family history of cancer (22).

To date, however, a large proportion of patients that exhibit the MSI/dMMR phenotype and undergo genetic testing do not show germline pathogenic variants in any of the MMR genes (*MLH1*, *MSH2*, *MSH6*, or *PMS2*). Unfortunately, the molecular mechanisms underlying MSI/dMMR CRC remain poorly understood (23). With the introduction of a next-generation sequencing (NGS) approach (24,25) that allows the concurrent investigation of multiple genes (25), the evaluation of inherited CRC is changing. For example, previous high-throughput sequencing studies suggested that tumor-suppressed genes

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such as TP53 and cyclin-dependent kinase inhibitor 2A (CDKN2A) could be responsible for genomic instability in numerous sporadic cancers (26,27). In the last few decades, NGS applications have led to the discovery of mutations in several predisposing CRC genes and to LS-related cancers. Germline mutations in other DNA-repair genes, such as ATM, PALB2, MRE11, and CHEK2, expressed throughout the cell cycle in response to double-stranded DNA breaks, have been associated with susceptibility to CRC (28,29). Furthermore, variants in CDH1, a gene normally associated with gastric cancer, could be a risk factor for CRC (30); a reduction of its expression is associated with tumoral dedifferentiation, lymphatic vessel invasion, and metastatic processes in CRC (31). In several previous studies, common MSH3 polymorphisms and rare variants were found to be significantly associated with CRC and prostate cancer as low-penetrance risk alleles (32-35). Moreover, biallelic MSH3 germline mutations appear to cause an additional rare recessively-inherited subtype of colorectal adenomatous polyposis (36). Evidence for an involvement of MLH3 includes a recent publication by Olkinuora *et al* (37) showing that a biallelic MLH3-truncating variant causes classical or attenuated adenomatous polyposis and possibly extracolonic tumors, while rare heterozygous variants of the MLH3 gene are associated with the LS phenotype (38). Finally, individuals with constitutional MMR deficiency (CMMRD) often have a high risk of developing a broad spectrum of malignancies and frequently display features reminiscent of neurofibromatosis type 1 (NF1) (39).

However, despite the NGS applications involving the multiple and simultaneous investigation of the various genes *hitherto* associated with MSI/dMMR CRC, the number of pathogenic variants identified following these analyses remains almost unchanged. Instead the variants of uncertain significance (VUS) that are detected in these patients who exhibit the MSI/dMMR phenotype are increasingly numerous.

Therefore, the aim of the present study was to elucidate the molecular basis of predisposition to the development of hereditary LS-related cancers in a cohort of 73 patients with a clinical suspicion of LS, using an NGS multigene panel designed and standardized by our research group and evaluating the pathogenicity of the numerous VUS identified, by applying criteria well known in the literature in order to obtain conclusive interpretation.

Materials and methods

Patient selection. A total of 73 patients with suspected LS meeting AC and/or Bethesda guidelines (BG) were recruited from Federico II University Hospital, National Cancer Institute IRCCS G. Pascale Foundation, and Luigi Vanvitelli University Hospital, all in Napoli, in southern Italy between January 2006 and December 2019, after study of their clinical characteristics and MSI and/or MMR protein expression in tumors.

Personal and family histories were obtained from each proband and written informed consent was provided by all patients. The present study was approved (protocol no. 120/10) by the local ethics committee 'Comitato Etico per le Attività Biomediche 'Carlo Romano' ('Carlo Romano' Ethics Committee for Biomedical Activities) at the University of Naples Federico II (Napoli, Italy).

Genomic DNA extraction. Genomic DNA was isolated from 3 ml of peripheral blood lymphocytes using the Nucleon BACC2 kit (Cytiva), according to the manufacturer's protocol. DNA quantity was assessed using a NanoDrop OneC spectrophotometer (reading at 260 nm and ratio 260/280 and 260/230 nm) and an Invitrogen Qubit 4 fluorometer (both from Thermo Fisher Scientific, Inc.). DNA quality was evaluated by 1% agarose gel electrophoresis and visualized with ethidium bromide.

Mutational analysis of coding regions of MMR genes. Genomic rearrangements in the MMR genes were analyzed by MLPA using SALSA-MSH6 P072 and SALSA-PMS2 P008 C1 kits (MRC-Holland BV), according to the manufacturer's protocol.

Targeted NGS

Library construction. Patient DNA samples were examined using an AmpliSeq Custom Panel (Illumina, Inc.), targeting 15 genes (Table I) involved in the MMR pathway or associated with CRC and other well-characterized cancer syndromes. This panel was developed based on the literature (26-39) to include genes associated with an increased risk of developing colon cancer. The kit (cat. no. 20020495; Illumina, Inc.) includes 470 amplicon regions that cover 87,353 bp, all the exonic and flanking intronic regions of these genes.

DNA was diluted to a final concentration of 4 ng/ μ l using Low Tris-EDTA buffer (included in the kit) and re-quantitated with the fluorometric quantification method (Invitrogen Qubit 4). The standard input was 20 ng of DNA per sample. Briefly, the workflow involved multiplex PCR to amplify target regions of each DNA sample according to the procedure for two primer pools from an AmpliSeq Illumina Custom DNA Panel. FuPa reagent (included in the kit) was used to partially digest amplicons and each library was mixed and ligated with a unique index-specific combination. Subsequently, a second amplification step ensured sufficient quantity for the final sequencing analysis. The quality and quantity of the libraries obtained were assessed using a TapeStation 4200 System (Agilent Technologies, Inc.) with an Agilent DNA 1000 kit. The sequencing was performed on a MiSeqSystem (Illumina, Inc.) with a Nano V.2 flow cell (300 cycles) reagent kit according to the manufacturer's protocol. The raw data generated by this analysis are available on site Mendeley (Elsevier; <https://data.mendeley.com/datasets/mxp536twnw/draft?a=a642b2f6-f237-4e78-802c-c5cab21ef866>).

Bioinformatics analysis. The sequencing data was analyzed using a BaseSpace Sequence Hub (Illumina, Inc.). Primary data analysis involved the detection and analysis of raw data (signal analysis), targeting sequencing reads (base calling) and scoring base quality. FASTQ files (generated by MiSeq Reporter Software 1.3.17; Integrative Genomics Viewer) were the outputs from this primary analysis. A demultiplexing process was subsequently required to produce separated sequencing read files, according to the single index used for each sample. In a secondary data analysis, FASTQ files for each sample were aligned against an entire reference genome specified in the manifest file with a DNA Amplicon Analysis App on BaseSpace Sequence Hub (version 2.1.1).

Table I. Lynch syndrome full-exome panel.

Gene name	RefSeq	Band Chr	Genomic size	RNA size	Exons
<i>MLH1</i>	NM_000249.3	3p22.2	57497	2662	19
<i>MSH2</i>	NM_000251.2	2p21	80162	3226	16
<i>MSH6</i>	NM_000179.2	2p16.3	23872	4435	10
<i>PMS2</i>	NM_000535.5	7p22.1	35868	2851	15
<i>MLH3</i>	NM_001040108.1	14q24.3	37769	7911	13
<i>MSH3</i>	NM_002439.4	5q14.1	222168	4472	24
<i>EPCAM</i>	NM_002354.2	2p21	17881	1731	9
<i>CDH1</i>	NM_004360.3	16q22.1	98250	4815	16
<i>TP53</i>	NM_000546.5	17p13.1	19149	2591	11
<i>ATM</i>	NM_000051.3	1p34.1	11229	1945	16
<i>CHEK2</i>	NM_001005735.1	22q12.1	54092	1991	16
<i>PALB2</i>	NM_024675	16p12.2	38196	4069	13
<i>MRE11A</i>	NM_005591	11q21	76572	5141	20
<i>NF1</i>	NM_001042492	17q11.2	282751	12444	58
<i>CDKN2A</i>	NM_000077	9p21.3	7382	1267	3

In silico analysis of unclassified variants. The following variant calling step had the main objective of identifying variants using a post-processed BAM file (<https://basespace.illumina.com/analyses>; BAM metrics version 0.0.22). A default value of 10 was used to define the Variant Caller Depth Filter level. Lower filter values may cause further false positive variants to pass the filter.

Output VCard File (VCF) was finally used for downstream analysis on a Variant Interpreter App (<https://variantinterpreter.informatics.illumina.com>; version 2.16.0.235), integrated with a BaseSpace Sequence Hub, that provided variant classification and reporting.

The BaseSpace Variant Interpreter is a cloud-based platform that uses the following annotation sources: Single Nucleotide Polymorphism Database (dbSNP) (<https://www.ncbi.nlm.nih.gov/snp/?cmd=search>), Catalogue of Somatic Mutations in Cancer (COSMIC) (<https://cancer.sanger.ac.uk/cosmic>), ClinVar (<https://www.ncbi.nlm.nih.gov/clinvar/>), 1000 Genomes (<https://www.internationalgenome.org>), Exome Variant Server (EVS) (<https://evs.gs.washington.edu/EVS/>), (ExAC) (<https://gnomad.broadinstitute.org>), Polymorphism Phenotyping v.2 (PolyPhen-2), and Sorting Intolerant from Tolerant (SIFT). Such software classify germline variants as pathogenic, likely pathogenic, VUS, likely benign and benign. These categories follow The American College of Medical Genetics and Genomics (ACMG) guidelines (40). A total of 7 different complementary *in silico* programs were subsequently used for functional impact predictions of the identified variants.

Human Splicing Finder (HSF) (www.umd.be/HSF/) for silent and intron variants is a tool designed to predict the effects of mutations on splicing signals or to identify splicing motifs in human sequences. It contains all available matrices for auxiliary sequence prediction and presents a novel position weight matrix to assess the strength of 5' and 3' splice sites and branch points (41).

SIFT (<http://blocks.fhcrc.org/sift/SIFT.html>), PolyPhen (<http://genetics.bwh.harvard.edu/pph/>) (42), and PredictProtein server (<http://www.predictprotein.org>) (43) are prediction tools based on a combination of phylogenetic, structural and sequence annotation information characterizing a substitution with its position in the protein.

Mutation Taster (<http://www.mutationtaster.org/>) (44) and Align-Grantham Variation Grantham deviation (A-GVGD) (<http://agvgd.hci.utah.edu/>) (45) were employed in the study of missense variants. Briefly, Mutation Taster analyses comprise evolutionary conservation, splice-site changes, loss of protein features, and changes that may affect the amount of mRNA; moreover, the A-GVGD method can be used to identify sets of missense substitutions that are either enriched for deleterious variants or enriched for neutral variants.

Finally, Protein Variation Effect Analyzer (PROVEAN) is useful software for predicting whether nonsynonymous or indel variants are functionally important (<http://provean.jcvi.org/index.php>); its performance is comparable to that of SIFT or PolyPhen-2.

All variants identified were annotated according to the nomenclature recommendations from the Human Genome Variation Society (www.hgvs.org/mutnomen).

Variant analysis by Sanger sequencing. The coding regions corresponding to 22 variants (pathogenic variants and deleterious variants by *in silico* analysis) were amplified using customized primer sets (Table II). The PCR products were separated on a 1-2% agarose gel to check for unspecific amplicons. Subsequently, the PCR products were sequenced in both the forward and reverse directions using an ABI 3100 Genetic Analyzer (Applied Biosystems; Thermo Fisher Scientific, Inc.).

Results

A total of 73 patients with a clinical suspicion of LS were analyzed in the present study. The patients were selected

Table II. Primers, annealing PCR and amplicon sizes of PCR products corresponding to the pathogenic variants and variants resulted deleterious by *in silico* analysis.

Gene variants	Primer sequences (5'→3')	TM (°C)	Amplicon (bp)	Exon
MLH1 c.350C>T	F: GTGACCCAGCAGTGAGTTTT R: AGCCTCACTTTTACCCTCTCT	58	245	4
MSH6 c.3311_3312del	F: AGCCTCACTTTTACCCTCTCT R: TGGCTGACTTTTATGTAAGTGTG	58	398	5
MSH6 c.892C>T	F: TGGTGGCTCTGATGTGGAAT R: TTGCTTGTTTGGTGGCTGAG	59	206	4
ATM c.3802delG	F: TGCTACTGAACAAGGTCCCA R: CTCTCTTTGCTGTGCCATCC	59	448	26
MLH1 c.376T>A	F: GTGACCCAGCAGTGAGTTTT R: ACGTACTCAAGATCTCTGCCA	58	245	4
EPCAM c.332A>G	F: TGATGAAGGCAGAAATGAATGG R: ACAAGTAGTATAGGCAGCCCC	58	250	3
MSH3 c.1778G>A	F: CCTGGGCATTAGAGTGGGAA R: TGTCTCAAGCTGAAGAACAC	58	264	13
MLH3 c.470T>C	F: TATGGTTTCCGAGGAGAGGC R: CCAGTCTAGGGTCCATGCAT	59	232	2
MLH3 c.3440A>T	F: TCAGTTTGTGCAGAAAGAGGT R: GAGTTAGGTGGTACGATGTGT	58	250	3
ATM c.7475T>G	F: AGGAAGGTGTGTGAATTGCA R: CCTGACATCAAGGGGCTTATG	58	465	50
ATM c.1178G>T	F: GGCAACAACAGCGAAACTCT R: TGTCATGGCAATCACATATCCC	58	396	9
ATM c.8734A>G	F: ATTAGCTGTCAAACCTCCTAACT R: TGCCCAGCCCATGTAATTTT	58	221	60
CHEK2 c.911T>C	F: TGTCTTCTGTCCAAGTGCCT R: GGTCCCTCGATTTCTGCCTA	59	245	9
PMS2 c.1004A>G	F: AAAGTGAATTTGGCTGGGCG R: TGGCTGCTGACTGACATTTAGCTTG	59	498	10
PMS2 c.2249G>A	F: TCTCAGGAAGTTTGTGACACT R: CACCCAGCCGCTATAGTTCT	59	295	13
PMS2 c.1253C>T	F: GACCCTCTTCTCCGTCCAC R: GAGAGTCCACATGTTCCCTGC	58	491	11
MLH1 c.589-9_589-6delGTTT	F: GTTTGCTGGTGGAGATAAGGT R: ACGCCACAGAATCTAGGAGA	58	392	8
MRE11 c.1783+7A>G	F: ACTTTCTCCTTCTTCTCCCTCT R: TGTCAGAACTGCCTTAAAGACTG	59	410	15
CDH1 c.585A>C	F: TTCTCTGGGAGGGATTGGC R: CCCGGTGTCAACAAGCTTC	59	293	5
CDH1 c.344C>T	F: GAAGATTGCACCGGTGCGAC R: CAACAGCGAACTTCTCAGAAAA	58	250	3
NF1 c.4445T>C	F: CTGGGTGTATCTGGTGTGAAAA R: GGATCTATAACAATCTGCAAGCC	58	486	34
CHEK2 c.688G>T	F: CTTGAAGTGGACCCAGGAGT R: TGGGAAGTTATGAAGACGTGTT	58	242	6

F, forward; R, reverse.

as follows: 42 patients based on the AC (5), and 31 patients according to MMR deficiency detected in tumoral tissue with respect to the Bethesda guidelines (BG) (20). Individuals

undergoing analysis were all affected by CRC, and two had MMR-deficient endometrial cancer. These patients were also affected by ovarian, bladder, breast, prostate, melanoma and

Table III. Clinical and molecular characteristics of the 73-patient cohort.

Characteristics	Amsterdam criteria	Bethesda guidelines	Total
Sex			
Female	21	14	35
Male	21	17	38
Age at diagnosis, years (mean \pm SD)	33.47 \pm 11.96	47.59 \pm 16.93	44.89 \pm 14.26
Tumor type			
CRC	40	31	71
Breast	3	3	6
Endometrium	3	4	9
Other tumors	-	2	3
MSI status			
MSI	9	31	40
Unknown	31	-	31

MSI, microsatellite instability.

renal cancers. The mean age at diagnosis, MSI-high (MSI-H) status analysis and sex of patients are outlined in Table III.

All patients were previously shown to be negative for pathogenic variants in the MLH1 and MSH2 genes detected by single-gene analyses [denaturing high-pressure liquid chromatography (dHPLC) followed by Sanger sequencing] and large rearrangements (by MLPA).

Germline DNA samples from the patients were tested with NGS methods using a multigene custom panel developed by our research group that targets a set of 15 genes (Table I), MLH1, MSH2, MSH6, PMS2, MSH3, MLH3, CHEK2, MRE11, EPCAM, ATM, TP53, CDKN2A, PALB2, CDH1 and NF1. These genes are involved in the MMR pathway or associated with CRC and/or other well-characterized cancer syndromes (26-39).

Paired-end NGS generated an average of 705,000 reads, 93.93% of which mapped against the human genome (version GRCh38). Variants with 99.42% of exons covered were only selected, labelled as 'PASS' by the filter applied, with an estimated average amplicon depth of coverage of 1,350 reads. In this manner, an overall number of 724 variants were identified, of which only four (0.55%) were pathogenic variants already known, reported in the international databases Insight-Group and ClinVar (<http://insight-database.org>; <https://www.ncbi.nlm.nih.gov/clinvar/>) as pathogenic variants. Most of the remaining variants (87.8%) were already known in the literature as being benign or polymorphic variants; 86 variants (11.88%) had been classified as variants that are likely benign or of uncertain pathogenic significance.

Clinically significant variants: Pathogenic variants. At least one clinically significant variant was identified in four patients (5.6%) of the cohort: One nonsense variant in ATM, two nonsense variants in MSH6, and one missense variant in MLH1 (Table IV). These had not been previously found by traditional methods (dHPLC and Sanger sequencing). These variants were validated by Sanger sequencing (Fig. 1).

Unclassified and likely benign variants: Pathogenicity assessment. For the present study, 86 variants were considered as VUS. These were the variants that showed a minor allele frequency (MAF) that was very low (<0.1%) or not reported, some known in the international database (as ClinVar and Insight) and others novel. At least one of these VUS was identified in 57 patients (80.3%) in the cohort. These variants were distributed among the Lynch full-exome panel genes as follows: 4 in MLH1, 2 in MSH2, 4 in MSH6, 12 in PMS2, 12 in MSH3, 4 in MLH3, 3 in CHEK2, 5 in MRE11, 3 in EPCAM, 18 in ATM, 3 in TP53, 5 in PALB2, 5 in CDH1 and 6 in NF1. For each VUS identified in the present study, multiple bioinformatics analyses were performed using several software programs described in Table V and in the Materials and methods section. For silent and intronic variants, an *ad hoc in silico* analysis was used (as described in Table V and in the Materials and methods). Most of these variants presented discordant results from the computational algorithms applied. All lines of computational evidence supported a deleterious effect for 13 of these 86 variants. In addition, five others showed a deleterious effect in all but one or two of the computational algorithms used (Table V). These variants were validated by Sanger sequencing (Fig. 1).

Some of these rare variants are not present in healthy controls (as they are not reported in The Genome Aggregation Database, Exome Aggregation Consortium, or 1,000 Genome Projects database) and they are present in genes for which an association with a predisposition to developing colorectal tumors (or LS-related cancers) is well known. Therefore, the criteria reported in ACMG and the Association for Molecular Pathology guidelines were applied for the interpretation of sequence variants, (Table VI) (40). The MSI/dMMR status on tumor tissue was evaluated as a strong evidence of pathogenicity comparable with the results of well-established *in vivo* functional studies supportive of a damaging effect of variant on the gene or gene product only for the rare variants identified in the MMR genes. In this manner, eight of these 18 variants could be considered to be 'likely pathogenic' and for some of

Table IV. Clinically significant variants identified.

ID of patient	Gene	Variant	Variant classification	InSiGHT	ClinVar	PHENOTYPE (age onset)
07.19	MLH1	c.350C>T p. (Thr117Met)	Pathogenetic	Pathogenetic	Pathogenetic	Index case k-co (34), MSI-H; father k-co (37) and his father's brother succumbed to k-co (65)
07.13	MSH6	c.3311_3312del p. (Phe1104TrpfsTer3)	Pathogenetic	Pathogenetic	Pathogenetic	Index case k-co (49), MSI-H; father k-st (50), died k-co (58), his father's brother k-co (70), his cousin k-st (61) and colon polyps
14.07	MSH6	c.892C>T p. (Arg298Ter)	Pathogenetic	Pathogenetic	Pathogenetic	Index case k-end (32), MSI-H; father leukemia (72), and her father's mother succumbed to leukemia (45)
13.68	ATM	c.3802delG p. (Val1268Ter)	Pathogenetic	-	Pathogenetic	Index case k-co (36) and several colon polyps; Sister k-co (75), brother k-pro (60) and father died k-pan (70)

k-, cancer; -co, colon; -st, stomach; end, endometrium; -pan, pancreas; -pro, prostate.

these variants the analysis of segregation in family environment was also performed (Table VI). Of these, two variants, (in the MLH3 and ATM genes) are novel and not previously reported in the literature. For the remaining 78 variants, the criteria for being benign and pathogenic were contradictory; therefore, these variants remain classified as having an uncertain significance.

In Fig. 2, the percentage of these 'likely pathogenic variants' of the total uncertain variants identified for each gene analyzed, are presented. Notably, 50% of the gene variants identified in MLH1 and MLH3 are classified as likely pathogenic.

Discussion

The study of hereditary forms of CRC has increased the importance of genetic testing. However, the limited capacity of old genetic screening methods, due to their low sensitivity and small number of genes studied, has left numerous gaps in identifying variants conveying a predisposition to cancer. Indeed, more than half of CRC cases with a clinical suspicion of LS that are referred for genetic testing remain without a clear molecular diagnosis. In the present study, an NGS multigene custom panel was designed and standardized; it included, beyond the four MMR genes classically associated with LS, other MMR genes such as MLH3 and MSH3, and other genes that predispose to hereditary CRC or LS-related cancer according to the literature, (26-39) such as CHEK2, MRE11, EPCAM, ATM, TP53, PALB2, CDH1, NF1 and CDKN2A.

This NGS multigene panel was used to analyze 73 patients with a clinical suspicion of LS selected according to the BG and AC. The patients had already been shown to be negative for pathogenic variants in the MLH1 and MSH2 genes. This is probably the reason why the number of pathogenic variants clearly identified in the present study was so low. Only four pathogenic variants reported in the literature were identified in the cohort, of which one related to a gene not included among the MMR genes. Indeed, one patient (ID 13.68) was found to carry a variant, c.3802delG p. (Val1268Ter), in the ATM gene. This patient developed a right colon adenocarcinoma at an early age of onset (36 years) and was selected by AC due to a strong family history of early onset of colon tumors. Unfortunately for this patient, MSI could not be performed on the tumor tissue; however, it was certainly a case that could be confused with a clinical suspicion of LS. Variants in the ATM gene are associated in the recessive form with ataxia-telangiectasia and in the dominant form with breast cancer susceptibility and more generally with hereditary cancers (46). In the present study, this pathogenic variant, c.3802delG p. (Val1268Ter), appeared to be associated with CRC.

The remaining pathogenic variants were identified in MMR genes (MLH1 and MSH6) in patients who met the AC and showed a typical LS phenotype with MSI-H status in cancer tissues (Table III). No other variant that was already classified in the literature as pathogenic was identified in the cohort, although the patients were selected according to very specific criteria. However, numerous genetic variants beyond these four clearly pathogenic variants were identified in the present

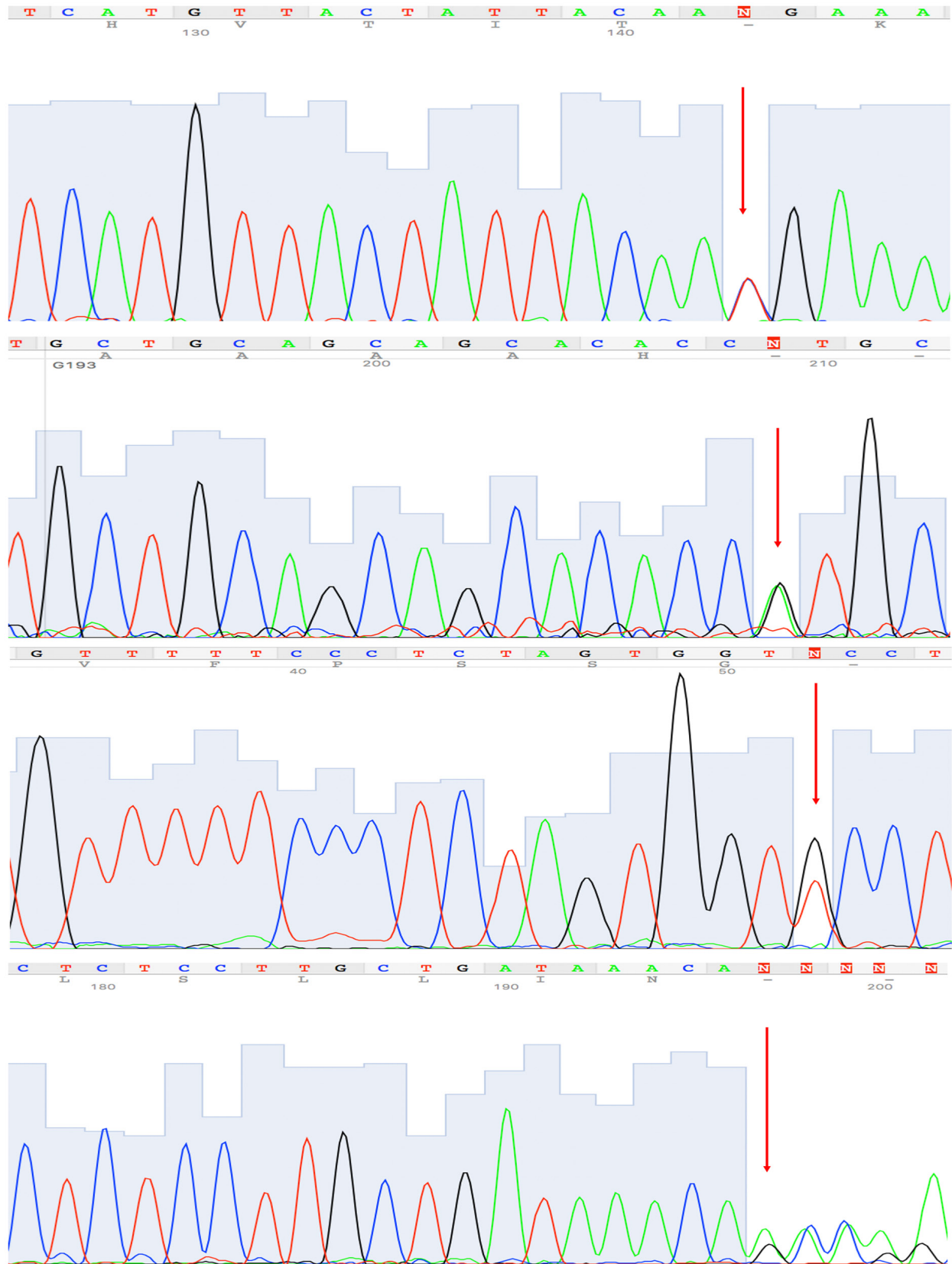


Figure 1. Certain representative electropherograms by Sanger-sequencing of variants identified in the present study. (1) c.350C>T, p. (Thr117Met), MLH1 gene. (2) c.2149G>A, p. (Val717Met), PMS2 gene. (3) c.688G>T, p. (Ala230Ser), CHEK2 gene. (4) c.589-9_589-6delGTTT, MLH1 gene.

study. The majority of these were benign or polymorphic variants, but numerous variants of unclear pathogenic significance were also found. Unfortunately, these variants were difficult

to clinically interpret, which poses a significant barrier to the broad utility of genetic testing and carrier screening. In LS, nearly 90% of the identified genetic variants that are not

Table V. Variants identified in the panel genes and predicted *in silico* to have a deleterious effect.

Variants	ID Cancer (age onset) selected criteria	MAF%	Mutation Taster	Poly Phen 2 [0-1]	Align GVGD	Predict protein	Provean (c. off 2.5)	Sift (c.off 0.05)	HSF	Source
c.376T>A p. (Tyr126Asn) <i>MLH1</i>	19.39 endometrial (26) BG	0.004%	Disease-causing	0.911 (POSS DAM)	CLASS C 65	95	DEL (-7.69)	DAM (0)		Insight-group ClinVar
c.332A>G p. (Asn111Ser) <i>EPCAM</i>	18.33 colon (40) AC	-	Disease-causing	1.00 (DAM)	CLASS	71	DEL	DAM	-	ClinVar
c.1778G>A p. (Arg593Gln) <i>MSH3</i>	14.42 breast (38) colon (38) AC	-	Disease-causing	1.00 (DAM)	C 45 CLASS C35	59	(-4.13) DEL (-3.90)	(0.004) DAM (0.001)	-	None
c.470T>C p. (Val157Ala) <i>MLH3</i>	18.17 breast (37) colon (38), melanoma AC	-	Disease-causing	1.00 (DAM)	CLASS C55	49	DEL (-3.66)	DAM (0)	-	None
c.3440A>T p. (Asn1147 Ile) <i>MLH3</i>	17.14 colon (52) AC	0.03%	Disease-causing	1.00 (DAM)	CLASS C65		DEL (-6.77)	DAM (0)	-	ClinVar
c.7475T>G p. (Leu2492 Arg) <i>ATM</i>	18.17 (as above) 06.10 colon (53) AC	0.02%	Disease-causing	1.00 (DAM)	CLASS C65	-	DEL (-4.53)	DAM (0.003)	-	None
c.1178G>T p. (Trp393Leu) <i>ATM</i>	19.24 colon (29) AC	0.002%	Disease-causing	1.00 (DAM)	CLASS C55		DEL (-4.86)	DAM (0)		None
c.8734A>G p. (Arg2912 Gly) <i>ATM</i>	18.08 rectum 16 BG	0.02%	Disease-causing	1.00 (DAM)	CLASS C65		DEL (-5.83)	DAM (0)		ClinVar
c.911T>C p. (Met304Thr) <i>CHEK2</i>	11.33 sigma (51) AC	0.0004%	Disease-causing	0.999 (DAM)	CLASS C 65	86	DEL (-3.76)	DAM (0.014)	-	ClinVar
c.1004A>G p. (Asn335Ser) <i>PMS2</i>	18.21 breast (49) colon (50) BG	0.1%	Disease-causing	1.00 (DAM)	CLASS C45	48	DEL (-4.47)	DAM (0.006)	-	Insight-group ClinVar
c.2149G>A p. (Val171Met) <i>PMS2</i>	18.36 colon (38) AC	0.08%	Disease-causing	0.982 (DAM)	CLASS C15	14	NEU (-1.79)	DAM (0.010)	-	ClinVar

Table V. Continued.

Variants	ID Cancer (age onset) selected criteria	MAF%	Mutation Taster	Poly Phen 2 [0-1]	Align GVGD	Predict protein	Provean (c. off 2.5)	Sift (c.off 0.05)	HSF	Source
c.1253C>T p. (Ser418Phe) <i>PMS2</i>	10.17 colon (49) AC	0.0008%	Disease-causing	0.977 (DAM)	CLASS C65	-6	DEL (-3.04)	DAM (0.004)		Insight-group ClinVar
c.589-9_589- 6delGTTT <i>MLH1</i>	18.08 (as above)	0.01 %								
c.1783+7A> G <i>MRE11</i>	09.10 colon (34) BG	0.004%	-	-		-	-	-		None
c.585A>C p. (Gln195His) <i>CDHI</i>	11.25 colon prostate (69) BG	-	Disease-causing	0.873 (POSS DAM)	CLASS C 15	-65	DEL (-2.59)	TOL (0.059)	-	ClinVar
c.344C>T p. (Thr115Met) <i>CDHI</i>	19.59 colon, kidney (36) AC	0.0024%	Disease-causing	0.992 (DAM)	CLASS C 15	-81	NEUT (-1.93)	TOL (0.080)	-	ClinVar
c.4445T>C p. (Ile1482Thr) <i>NF1</i>	06.08 colon (28) AC	-	Disease-causing	0.393 (BEN)	CLASS C65	-	DEL (-4.15)	DAM (0.002)	-	ClinVar
c.688G>T p. (Ala230Ser) <i>CHEK2</i>	18.30 sigma 24 BG	0.0008%	Disease-causing	0.076 (BEN)	CLASS C 65	64	DEL (-2.76)	DAM (0.006)	-	ClinVar

Poss Dam, possibly damaging; Dam, damaging; Del, deleterious; Neut, neutral; Tol, tolerated; Ben, benign.

Table VI. Variants defined as 'likely pathogenic'.

Variants	Gene	ID Cancer (age onset)		Somatic dMMR	The Genome Aggregation Database (GnomAD)	Exome Aggregation Consortium (ExAC)	1000 Genomes Project	Additional information	ACMG Criteria (a)	New classification variant	INSIGHT/CLINVAR
		Familial history (Segregation analysis)	(age onset)								
c.376T>A p.(Tyr126Asn)	<i>MLH1</i>	19.39 endometrial (26) Not affected- (N.D.) BG		<i>MSI-H NO</i> <i>MLH1 NO</i> <i>PMS2</i>	0.00004	0.00007	0.00020	This variant is located in the N-terminal ATPase domain	PS3, PM1, PP2, PP3, PP4	Likely pathogenic	Reported as VUS, as pathogenic and six times as benign/VUS
c.470T>C p.(Val157Ala)	<i>MLH3</i>	18.17 breast and colon (37, 38) other affected-breast and colon (N.D.) AC		<i>MSI-H</i>	No frequency	No frequency	No frequency	The valine residue is highly conserved.	PS3, PM2, PP2, PP3, PP4	Likely pathogenic	Not reported/VUS
c.3440A>T p.(Asn1147Ile)	<i>MLH3</i>	17.14 colon (52) other affected- endometrial and colon (N.D.) AC		<i>MSI-H</i>	0.00040	0.00044	0.00060	No relevant information	PS3, PP2, PP3, PP4	Likely pathogenic	Not reported/ conflicting interpretation
c.1178G>T p.(Trp393Leu)	<i>ATM</i>	19.24 colon (29) other affected- endometrial and gastric (N.D.) AC		<i>ND</i>	No frequency	No frequency	No frequency	No information	PM2, PM5, PP2, PP3, PP4	Likely pathogenic	Not reported
c.911T>C p.(Met304Thr)	<i>CHEK2</i>	11.33 sigma (51) other affected- colon (also in affected brother) AC		<i>MSI-H</i>	No frequency	No frequency	No frequency	The methionine residue is highly conserved. This variant has been reported to affect CHEK2 protein function (PMID: 30851065).	PM2, PP1, PP2, PP3, PP4	Likely pathogenic	Not reported/VUS

Table VI. Continued.

Variants	Gene	ID Cancer (age onset) Familial history (Segregation analysis)	Somatic dMMR	The Genome Aggregation Database (GnomAD)	Exome Aggregation Consortium (ExAC)	1000 Genomes Project	Additional information	ACMG Criteria (a)	New classification variant	INSIGHT/ CLINVAR
c.1253C>T p.(Ser418Phe)	PMS2	10.17 colon (49) other affected-colon (also in affected brother) AC	MSI-H NO MLH1 NO PMS2	No frequency	No frequency	No frequency	The seine residue is moderately conserved.	PS3, PM2, PP2, PP3, PP4	Likely pathogenic	Not reported/VUS
c.589-9_589- 6delGTTT	MLH1	18.08 Colon (18) other affected-colon and pancreas (also in affected father and two uncle) BG	MSI-H	No frequency	No frequency	No frequency	This variant could affect mRNA splicing (HSF software)	PS3, PM2, PP3, PP4	Likely pathogenic	Reported seven times as VUS/ conflicting interpretation
c.2149G>A p.Val717Met	PMS2	19.46 Colon (38) not affected-(N.D.) BG	MSI-H NO MLH1 NO PMS2	0.00076	0.00091	0.00020	The variant is located within the MutL C-terminal, and dimerisation functional domain.	PS3, PM2, PP3, PP4	Likely pathogenic	Not reported/ conflicting interpretation

PS, strong evidence of pathogenicity (class 1-4); PM, moderate evidence of pathogenicity (class 1-6); PP, Supporting evidence of pathogenicity (class 1-5); adapted from Ref (45). dMMR, deficient mismatch repair; N.D., not determined; BG, Bethesda guidelines; AC, Amsterdam criteria; VUS, variants of uncertain significance.

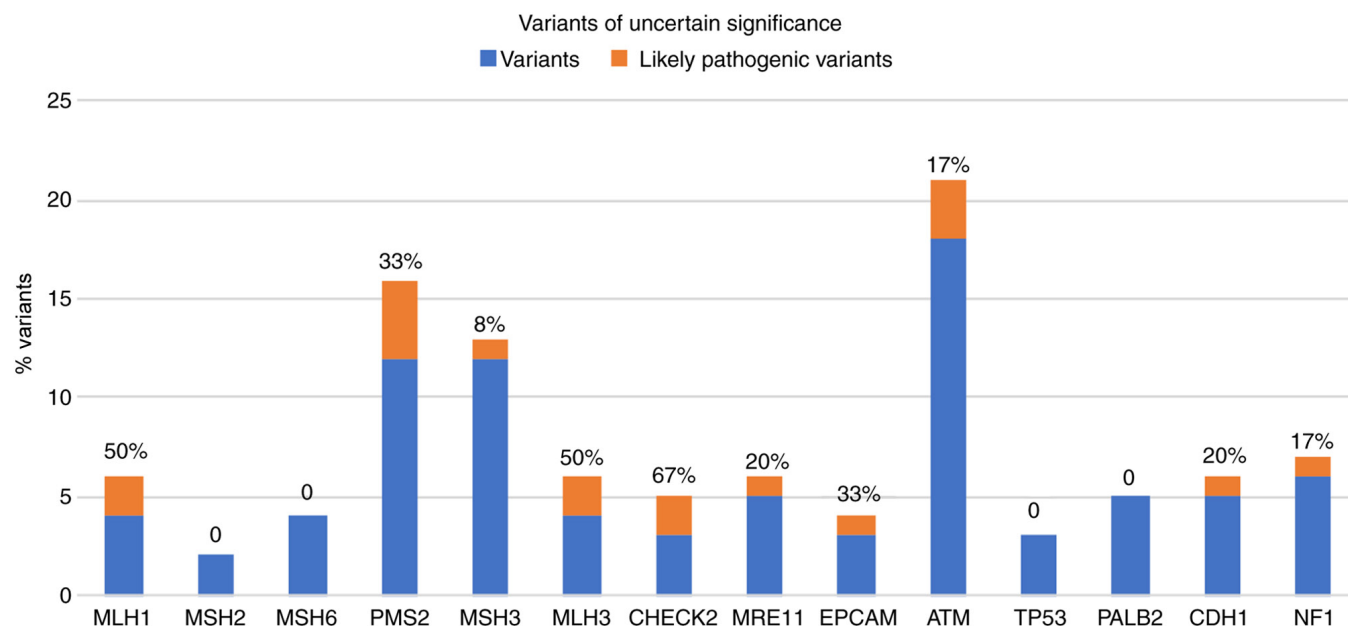


Figure 2. Graphical representation of the number of rare variants (minor allele frequency <0.1%) identified in the genes involved in the Lynch-Full-exome custom panel. Percent value (%) refers to the variants found in each gene analyzed that were predicted to be likely pathogenic by The American College of Medical Genetics and Genomics criteria.

included as nonsense or indel variants are deemed ‘variants of uncertain significance’ (47,48). To clarify the pathogenetic significance of such variants, it would have been useful to perform functional assays on proteins; recently, a massive parallel screen in human cells has been proposed to identify loss of function missense VUS in the MSH2 gene (49).

In the present study, 86 variants were identified that were already classified in international databases (Insight-group and/or ClinVar) as likely benign or of uncertain pathogenic significance, and four novel variants that showed a MAF that was very low or not reported. According to suggestions reported in the ACMG guidelines for the classification of variants (40), an interpretation of these variants was performed using multiple bioinformatics analyses, investigating seven different types of software for the pathogenic prediction of each variant. When the results of the *in silico* analyses were all in agreement as to the pathogenicity of the variants, it was examined whether these variants also respected the other ACMG criteria (population, computational, functional and segregation data) for establishment of the pathogenicity. Furthermore, the MSI/dMMR status (where this had been determined), found on the tumor tissues of the patients carrying these variants, was considered as a fundamental part for the interpretation of the pathogenicity. To the best of our knowledge, the MSI status in the majority of hereditary CRCs is associated to pathogenic variants in MMR genes. Thus, for the rare uncertain significance variants identified in the MMR genes that by bioinformatics analyses were resulted as pathogenic variants, it was hypothesized that the likely lack of function of corresponding protein at the somatic level, could be confirmed by the MSI/dMMR status showed on tumoral tissue of patients carrying these variants (Table VI). Therefore, our classification has arisen by the combination of the molecular and clinical data of each patient, in particular data from segregation and MSI analyses, applying the criteria provided from

the guidelines for the classification of variants established by the ACMG (40). In this manner, it was possible to classify eight of these 86 variants as ‘likely pathogenic’ variants (Table VI). Thus, applying the ACMG criteria (40) our variant interpretation differs from classifications reported in public databases, such as ClinVar which reports these variants in majority as uncertain significance. Surely, further studies are needed to establish the real pathogenic role of these variants; however, at present it can be hypothesized that these variants could be the cause of disease in eight patients of our cohort. Thus, in light of the results obtained in the present study, the importance of establishing for the variants identified in MMR genes, a correlation with a deficient MMR system at the tumor level, is suggested, thus strengthening the evaluation of pathogenicity (Table VI). The interpretation of the VUS represents a crucial step in clinical decision-making, improving risk assessment, and promoting appropriate medical management, including variant-specific cascade testing for relatives. Therefore, an accurate assessment of the predictions of the clinical significance of the VUS is needed. The rule-based classification of the ACMG as it was performed in the present study, can represent a valid alternative to functional studies of VUS, which remain the most reliable tool to support the pathogenicity or benignity of the variant studied.

Furthermore, it is interesting to note that two of the eight likely pathogenic variants were found in the MLH3 gene (Table VI), the c.470T>C p. (Val157Ala) and c.3440A> T p. (Asn1147Ile) variants. The first is a novel variant that was identified in a woman (ID 18.17) with adenocarcinoma *in situ* of the colon with an MSI-H phenotype and with breast cancer. This patient was also negative for pathogenetic variants in BRCA1 and 2, MutYH and APC. The second variant was identified in a woman (ID 17.14) who developed peritoneal adenocarcinoma with MSI-H at age 52. Both women were selected for the present study since they met the AC. Both these patients did not exhibit

MLH1 hypermethylation, and were also carriers of other two sequence variants in the MLH3 gene, the c.2476A>G and c.2531C>T, already described as benign in the ClinVar database. Finally, in these patients no other significant variants were identified in the genes included in the panel. The MLH3 gene is not routinely analyzed in patients with a clinical suspicion of LS. However, a previous study on MLH3-knockout mice highlighted the early onset of tumors in the abdominal sphere (50). Moreover, previous studies revealed a possible involvement of the MLH3 gene in LS (17,51). Loukola *et al* (51) and Wu *et al* (17) reported data on missense mutations and intronic substitutions in families meeting the AC, but without germline mutations in the MLH1 and MSH2 genes. Nonetheless, this gene is currently considered to be of low risk for a predisposition to the development of tumors on the LS spectrum. Unfortunately, functional assays were unable to be performed for either variant to clarify the pathogenetic effects, and no family segregation studies could be carried out due to a lack of interest from these patients. However, it is important to point out that both patients showed an MSI-H phenotype at the tumor level.

The results herein revealed that the use of the custom panel allowed the identification of variants in genes not routinely analyzed for cases with a clinical suspicion of LS, mainly variants in the MLH3 gene, but also rare variants identified in genes such as CHEK2, ATM, MSH3 and NF1. Although these results do not offer any evidence for a disease-causing role, they indicated the importance of deepening the study of all rare variants, to define their pathogenicity and to clarify the involvement of non-canonical genes in the pathogenesis of LS. The assessment of rare uncertain variants in genetic counseling could improve the risk estimate in those families that remain without a clear molecular diagnosis to provide precision medicine for this pathogenic condition (52).

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Availability of data and materials

The data generated and/or analyzed during the current study are available on site Mendeley (Elsevier; <https://data.mendeley.com/datasets/z9gkb9vd9j/draft?a=5466cd6c-f760-4e6d-ba6d-3033b599b6bd>).

Authors' contributions

FD conceptualized the study. FD, RL, ML and AN were involved in the literature search, in the data interpretation and critical reviewing of the manuscript. FD, AN and ML were involved in the preparation of the draft of the manuscript. FD, MDR and PI were involved in critically revising and editing the manuscript for important intellectual content. FD, RL, ML and AN confirm the authenticity of all the raw data. All authors have read and approved the final manuscript.

Ethics approval and consent to participate

The present study was approved (approval no. 120/10) by the local ethics committee, 'Carlo Romano' Ethics Committee for Biomedical Activities of the University of Naples, Federico II (Napoli, Italy). Written informed consent was obtained by all patients who participated in the present study.

Patient consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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