A rare large duplication of \textit{MLH1} identified in Lynch syndrome

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Abstract

Background: The most frequently identified strong cancer predisposition mutations for colorectal cancer (CRC) are those in the mismatch repair (MMR) genes in Lynch syndrome. Laboratory diagnostics include testing tumors for immunohistochemical staining (IHC) of the Lynch syndrome-associated DNA MMR proteins and/or for microsatellite instability (MSI) followed by sequencing or other techniques, such as denaturing high performance liquid chromatography (DHPLC), to identify the mutation.

Methods: In an ongoing project focusing on finding Mendelian cancer syndromes we applied whole-exome/whole-genome sequencing (WES/WGS) to 19 CRC families.

Results: Three families were identified with a pathogenic/likely pathogenic germline variant in a MMR gene that had previously tested negative in DHPLC gene variant screening. All families had a history of CRC in several family members across multiple generations. Tumor analysis showed loss of the MMR protein IHC staining corresponding to the mutated genes, as well as MSI. In family A, a structural variant, a duplication of exons 4 to 13, was identified in \textit{MLH1}. The duplication was predicted to lead to a frameshift at amino acid 520 and a premature stop codon at amino acid 539. In family B, a 1 base pair deletion was found in \textit{MLH1}, resulting in a frameshift and a stop codon at amino acid 491. In family C, we identified a splice site variant in \textit{MSH2}, which was predicted to lead loss of a splice donor site.

Conclusions: We identified altogether three pathogenic/likely pathogenic variants in the MMR genes in three of the 19 sequenced families. The \textit{MLH1} variants, a duplication of exons 4 to 13 and a frameshift variant, were novel, based on the InSiGHT and ClinVar databases; the \textit{MSH2} splice site variant was reported by a single submitter in ClinVar. As a variant class, duplications have rarely been reported in the MMR gene literature, particularly those covering several exons.

Keywords: Genetic predisposition, Lynch syndrome, Mismatch repair genes, Whole-genome sequencing
Background
Familial cancer, here defined as two or more first-degree relatives diagnosed with the same cancer, accounts for some 15% of colorectal cancer (CRC) [1]. The most frequently identified strong cancer predisposition mutations for CRC are those in mismatch repair (MMR) genes in Lynch syndrome, which account for approximately 1% of CRCs in the population (depending on the population) [2]. A number of other high-risk genes are known but variants in these are very rare [3]. In addition, ever-increasing numbers (> 100) of low-risk gene variants have been described for CRC [4]; yet combined, the high and low-risk variants explain only a small proportion of the known familial risk and even less of the heritability estimated in twin studies [5, 6].

Clinical diagnostics of Lynch syndrome usually first considers family history based on the Amsterdam and Bethesda criteria [7]. These are not perfect as half of germline-confirmed Lynch syndrome patients fail to meet the Amsterdam II criteria and, although the Bethesda guidelines are sensitive, their specificity is low [7]. Diagnostic laboratory tests include testing tumors for immunohistochemical (IHC) staining of the Lynch syndrome-associated DNA MMR proteins and/or for microsatellite instability (MSI) [7]. While these tests alone have a sensitivity ranging from 55 to 90% of predicting Lynch syndrome, combining the two will reach a sensitivity over 90% [7]. The identification of mutations is done by sequencing, or by other techniques, such as denaturing high performance liquid chromatography (DHPLC) or multiplex ligation dependent probe amplification (MLPA) for structural variants [8]. More recently, next generation sequencing panels have become the golden standard in identification of pathogenic germline variants in hereditary cancer syndromes. In a recent study, a universal 83-gene next generation sequencing panel identified nearly double as many pathogenic germline variants related to hereditary cancer syndromes as the guideline-directed targeted testing in unselected cancer patients, leading to a treatment change for nearly 30% of these patients [9]. This highlights the usefulness of next generation sequencing in the clinical praxis and compensates the limitations of the clinical and guideline-based risk assessment.

We have been involved in a whole-exome/whole-genome sequencing (WES/WGS) project aimed at identifying Mendelian type cancer syndromes in families referred to the Hereditary Cancer Center, Szczecin. In three families fulfilling the Amsterdam II criteria of Lynch syndrome with negative results in DHPLC mutation screening of the Lynch syndrome-related MMR genes we identified a mutation in these genes using whole genome sequencing. Here, we report these variants, particularly a large duplication in the MLHI gene, as these types of large structural variants, particularly insertions are rarely described in Lynch syndrome [10–14].

Patients and methods
In several regions of Poland, population screening was performed mainly in years 2000–2014, in which questionnaires on cancer family history were collected systematically. Individuals with a positive CRC family history were invited to genetic outpatient clinics all over Poland and their more detailed family histories were taken through detailed face-to-face interviews. Nineteen families with strong CRC aggregation compatible with an autosomal dominant pattern of inheritance were recruited to the study. Each family had at least three pathologically confirmed CRC cases; 17 families had at least one case diagnosed below the age of 55 years. All 19 families had undergone DHPLC analysis for MMR variants with negative test results [15]. The ethical approval for this study design was obtained from the Bioethics Committee of the Pomeranian Medical Academy in Szczecin No: BN-001/174/05. Sample collection was performed following the guidelines proposed by this Committee. A written informed consent was signed by each participant in accordance with the Helsinki declaration.

WES on CRC patients and healthy family members of 5 families and WGS on 14 families was performed in the Illumina X10 platform using DNA extracted from the blood samples. WGS was carried out as paired-end sequencing with a read length of 150 bp. Sequences were mapped to the reference human genome (build hg19, assembly hs37d5) using BWA mem (version 0.7.8) and duplicates were marked using Picard (version 1.125). Single nucleotide variants and small indels were called by using Platypus (version 0.8.1) and annotated using ANNOVAR [16], dbSNP [17], 1000 Genomes phase III [18], dbNSFP v.2.9 [19], and ExAC [20], respectively. Variant filtering was carried out by considering a minimum of 5 reads coverage and a minimum QUAL score of 20. To check for family relatedness, a pairwise comparison of variants among the cohort was performed.

GATK gCNV module (version 4.1.7.0) was used to call germline copy number variants (gCNVs) from the WES/WGS samples individually against a background of 200 WGS samples sequenced from the sample platform. The gCNVs were called based on the best practice recommended by the GATK (https://gatk.broadinstitute.org/hc/en-us/articles/360035531152%2D%2DHow-to-Call-common-and-rare-germline-copy-number-variants). The major deviation from the above best practice was that the gCNVs cohort models were created only for the Gencode v19 exonic regions of WGS data by considering them as the target regions. The sequences of the samples from the CRC families were
compared against this model. This decreased the turnaround time for the analysis of gCNVs from the WGS data.

The resulting gCNV segments with QS score above 30 were selected and annotated with the subset of gnomAD structural variant (SV) data (version 2.1, variants with ‘PASS’ filter tags and ‘DUP’ or ‘DEL’ SV types) using vcfanno [21]. The segments with at least 80% overlap with a common gnomAD SV (popmax MAF > 0.1%) of same SV subtype were considered as common and removed. In addition, to consider a gCNV as rare, at least 50% of the targets (exons here) in the gCNV segments should have the denoised ploidies among the bottom (in the case of deletion) or top (in the case of duplication) 5% of denoised cohort ploidies from the background cohort samples. Subsequently, the candidate rare gCNVs were selected if they followed the disease inheritance pattern in the family. For the candidate gCNVs the genomic breakpoints were manually reviewed using the Integrative Genomic Viewer (IGV) [22] to determine the genomic coordinates of the gCNVs.

Sequencing data were visually inspected using IGV to exclude false positive variants. For variants causing a frameshift, we used the Translate tool (https://web.expasy.org/translate/) to translate the nucleotide sequence to a protein sequence. The effect of splice site variants on splicing was analyzed using NetGene2 (http://www.cbs.dtu.dk/services/NetGene2/). Combined Annotation-Dependent Depletion (CADD) score was used to evaluate the deleteriousness of the variants; the scores > 20 and > 30 are indicative of the top 1% and top 0.1% of deleterious variants, respectively [23]. The InSiGHT database available at the Leiden Open Variation Database (LOVD) v.3.0 [24, 25], ClinVar (https://www.ncbi.nlm.nih.gov/clinvar/) [26], gnomAD database (https://gnomad.broadinstitute.org/) and the recent publication on Chinese MMR variants were used as a reference [11].

IHC and MSI analyses were performed as reported previously [8, 27] in CRC samples from individuals with a MMR gene variant detected through WES or WGS.

**Results**

In three of the 19 families sequenced, pathogenic/likely pathogenic MMR gene variants were identified. The pedigree of family A is shown in Fig. 1. Several patients diagnosed with CRC were present in three generations. We sequenced the affected father (diagnosed at age 70 years) and his son (diagnosed at age 32 years). Additionally, two unaffected individuals were sequenced. The pedigrees of families B and C are found in Additional file 1: Fig. 1.

The detected variants are listed in Table 1. In family A, a structural variant, a duplication of chr3:37045366–37,071,869 covering exons 4 to 13 of MLH1 was identified. It was predicted to lead to a frameshift at amino acid 520 and a premature stop codon at amino acid 539. The duplication was identified in both patients and in an unaffected female relative who was 9 years older than her affected brother. In family B, a one base pair deletion was found in MLH1 which resulted in a frameshift and a stop codon at amino acid 491. In family C, the three affected individuals carried a splice site variant in MSH2, with a CADD score of 23.4 (Table 1). According to NetGene2, the MSH2 variant c.792 + 1G > C lead to a loss of a splice donor site.

The IHC and MSI results of the tumor samples from the Lynch syndrome patients are shown in Table 2. Tumor samples from patients from families A and B did not express MLH1 and PMS2 proteins while in family C the tumor sample was negative for MSH2 and MSH6 proteins. The results are in line with the mutation
analysis as MLH1 and PMS2 as well as MSH2 and MSH6 form heterodimers. Further in line, the MSI analysis showed MSI-high for families A and C; the analysis for family B failed. Capillary electrophoresis MSI diagrams for samples from families A and C are shown with pattern shifts for the monomorphic markers (Additional file 1: Fig. 2). The identical migration of the pentanucleotide markers confirms the sample identity.

The consequences of the MLH1 variants on the gene and protein structure are shown in Fig. 2. In family A, the large duplication of exons 4 to 13 covered a small section of the ATP binding domain (HATPase C domain) and the entire mismatch repair domain (MutL, i.e., MSH2-MLH1 heterodimer binding domain) as well as a small part of the MLH1 C-terminal domain (Fig. 2a). The duplication was predicted to lead to a frameshift at amino acid 520 and a premature stop codon at amino acid 539. Figure 2b shows the MLH1 frameshift variant at amino acid 425 in family B leading to a premature stop codon at amino acid 491. Both variants were predicted to lead to the deletion of the MLH1 C-terminal domain, which is needed for the MLH1-PMS2 heterodimerization.

**Discussion**

The present sequencing effort in families with a CRC family history suggestive of autosomal dominant inheritance identified two families with a pathogenic variant in the MLH1 gene and one family with a likely pathogenic variant in the MSH2 gene. In Poland, over 100 MMR gene point mutations have been identified, most of which are either frameshift or nonsense mutations leading to a truncated protein [28]. In over 60% of all Polish Lynch syndrome families a recurrent mutation is present. Two of the most frequent alterations are a substitution of A to T at the splice donor site of intron 5 of MSH2 and a missense change (A681T) of MLH1 [8]. In Polish patients, large deletions have been described particularly in the MSH2 gene [8].

The present three variants have so far not been reported in InSiGHT [24, 25]; only the MSH2 variant has been reported once in ClinVar [26]. However, the InSiGHT database lists similar MLH1 variants causing a frameshift and leading to a protein truncation at approximately the same position as our variants, both have a classification “pathogenic” [24, 25] (Table 3). The MSH2 variant is reported in ClinVar by a single submitter (accession number VCV000951452.1) and predicted to be “likely pathogenic” [26]. InSiGHT reports another nucleotide change at the same position, a c.792 + 1G > A variant, with a classification “pathogenic” (Table 3).

The present MLH1 variants were predicted to cause protein truncation and to be pathogenic, while the MSH2 splice site variant was predicted to be likely pathogenic. They add to the large collection of (likely) pathogenic variants in the MMR genes. Although all of these were unique, the duplication is of special interest as large duplications have rarely been reported for MMR genes. In the European literature somewhat over 10 exon level duplications have been reported, most of them in MSH2 and fewer in MLH1 [10, 12]. Similarly, in the recent Chinese literature survey on 34,000 individuals including both cancer cases and individuals without cancer, 540 MMR variants were found, but only 3 single exon duplications were reported for MLH1 and one for MSH2 [11]. In one of these papers the breakpoints implicated Alu mediated recombination as a mechanism and the duplication was predicted to create a premature

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**Table 1** Mismatch repair gene variants in three colorectal cancer families

| Family | Gene | CHROM_POS_REF_ALT | HGVS nomenclature | ANNOVAR annotation | Protein change |
|--------|------|-------------------|-------------------|-------------------|---------------|
| Family A | MLH1 | 3_37076355_AG_A | LRG_216t1:c.1274del | frameshift | p.(Arg425Serfs*66) |
| Family B | MLH1 | 3_37076355_AG_A | LRG_216t1:c.1274del | frameshift | p.(Arg425Serfs*66) |
| Family C | MSH2 | 2_47639700_G_C | LRG_218t1:c.792 + 1G > C | splicing | |

a Human genome build hg19, assembly hs37d5
b according to den Dunnen JT: HGVS Recommendations for the Description of Sequence Variants: 2016 Update, Hum Mutat 37:564

c ClinVar c.792 + 1G > C, likely pathogenic, review status: criteria provided, single submitter (accession number VCV000951452.1)

d ClinVar c.792 + 1G > C, likely pathogenic, review status: criteria provided, single submitter (accession number VCV000951452.1)

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**Table 2** Immunohistochemistry (IHC) and microsatellite instability (MSI) analysis on tumor samples of the colorectal cancer families

| Family | Variant | IHC MLH1 | IHC PMS2 | IHC MSH2 | IHC MSH6 | MSI |
|--------|---------|----------|----------|----------|----------|-----|
| Family A | MLH1 duplication exons 4–13 | negative | negative | positive | positive | high |
| Family B | MLH1 c.1274delG;p.Arg425Serfs*66 | negative | negative | positive | positive | failed |
| Family C | MSH2 c.792 + 1G > C | positive | positive | negative | negative | high |
stop codon and the formation of a truncated protein [12]. In the InSiGHT database, only 9 exon-level duplications in MLH1 are reported compared to 77 deletions; in MSH2 the numbers are 7 duplications and 84 deletions [24, 25] (Additional file 1: Table 1). While all the deletions had clinical classification “pathogenic”, only 2 duplications in MLH1 and 3 in MSH2 were classified as “pathogenic”. While large deletions most likely lead to non-functional proteins, the effect of large duplications may depend on whether the duplication is in-frame or not. The duplication in MLH1 we present here is predicted to cause a frameshift and a truncated protein.

The present duplication of exons 4 through 13 covered a small section of the ATP binding domain (HATPase C domain), the entire mismatch repair domain (MutL, i.e., MSH2-MLH1 heterodimer binding domain) and part of the MLH1 C-terminal domain [29–31]. The out-of-frame change at amino acid 520 was predicted to cause a stop codon further down-stream at amino acid 539. Thus, the resulting truncated protein is probably degraded by nonsense mediated decay as supported by the IHC results of lack of MLH1 protein in the tumor. The C-terminal end of MLH1 contains important binding sites for heterodimeric MMR proteins that contribute to the various key functions such as endonuclease activity [30, 31].

The fact that these three mutations were missed in the previous screening early 2000s may be due to the methodology used at that time, DHPLC. The DHPLC primers were designed to cover all exons and approximately 30–60 bp upstream and downstream of each exon. As the breakpoints of the large duplication in MLH1 were located 526 bp downstream of exon 4 and 1446 bp upstream of exon 13, it was missed. Also the splice site variant in MSH2 may have been missed, because its distance to the upstream primer for detecting exon 4 was only 2 bp. Only the frameshift variant in MLH1 may have been possible to detect. This calls for the recommendation that historically negative cases, assessed by inferior methods, should be re-considered for testing using up-to-date methodologies.

Conclusions
We identified three novel MMR gene variants that were predicted to lead to truncated proteins. The variants segregated with the disease and are expected to predispose to Lynch syndrome phenotypes, including CRC.

**Table 3** Examples of InSiGHT DNA and protein changes for variants causing similar DNA and/or protein changes as the variants identified in the Polish families

| Gene | CHROM_POS_REF_ALT | InSiGHT DNA change [protein change] | InSiGHT class |
|------|-------------------|------------------------------------|---------------|
| MLH1 | 3_37070422-37070423_G_GT | c.1557_1558insT [p.(Val520Cysfs*8)] | pathogenic |
| MLH1 | 3_37067349_TA_T | c.1261del [p.(Ser421Valfs*70)] | pathogenic |
| MSH2 | 2_47639700_G_A | c.792 + 1G > A | pathogenic |

* Human genome build hg19, assembly hs37d5
b nearby position with similar consequence as caused by the large duplication in Family A
c nearby position with similar consequence as caused by the frameshift variant in Family B
d same position as the splice site variant in Family C, different nucleotide change
Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s13053-021-00167-0.

Additional file 1: Fig. S1. (Family B) Pedigree of the colorectal cancer family with MLH1 frameshift variant. (Family C) Pedigree of the colorectal cancer family with MSH2 splice site variant. Fig. S2. Microsatellite instability (MSI) analysis of the tumor samples of two family members from Family A and one tumor sample from Family C. For each family, individuals with the tumor samples analyzed are indicated by an arrow and the MSI plots are shown for the corresponding germline and tumor samples. Table S1. Number of large deletions and duplications in the mismatch repair genes reported in the InSiGHT database and their clinical classification according to Mismatch Repair Gene Variant Classification Criteria by the InSiGHT Variant Interpretation Committee.

Abbreviations

CRC: colorectal cancer; MMR: mismatch repair; IHC: immunohistochemical; MSI: microsatellite instability; DHPLC: denaturing high performance liquid chromatography; MLPA: multiplex ligation dependent probe amplification; WES/WGS: whole-exome/whole-genome sequencing; gCNV: germline copy number variant; SV: structural variant; IGV: Integrative Genomic Viewer; CADD: Combined Annotation-Dependent Depletion

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Authors' contributions

Conceptualization KH, RS, JL, AF; data curation NP, MS; formal analysis AK, NP; funding acquisition KH, RS, JL; methodology AK, NP, MS, DD, KG; project administration KH, AF; resources NP, MS, DD, KG, MK, JL; supervision KH, AF; visualization AK, AF; writing – original draft KH, AF; writing – review & editing all authors. The author(s) read and approved the final manuscript.

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Availability of data and materials

The datasets generated and/or analysed during the current study are available from the corresponding author on reasonable request.

Ethics approval and consent to participate

The ethical approval for this study design was obtained from the Bioethics Committee of the Pomeranian Medical Academy in Szczecin No: BN-001/174/05. Sample collection was performed following the guidelines proposed by this Committee. A written informed consent was signed by each participant in accordance with the Helsinki declaration.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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