Genetic and epigenetic profiling of a solitary Peutz–Jeghers colon polyp

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Abstract Colon polyps represent precursor lesions of colon cancers and their malignant potential varies according to histological subtype. A rare subtype of colon polyps is the Peutz–Jeghers (PJ) polyp. PJ polyps mostly occur in the context of Peutz–Jeghers syndrome, which is characterized by the development of multiple polyps in the intestinal tract and hyperpigmentation of oral mucosa and lips. Peutz–Jeghers is an autosomal dominant disorder caused by pathogenic variants of the serine threonine kinase STK11. PJ polyps very rarely occur outside of the syndrome and are then referred to as solitary PJ polyps. Contrary to the situation in Peutz–Jeghers, the genetic basis and the malignant potential of solitary PJ polyps are currently unknown. Here we describe a detailed and comprehensive genetic profile of a solitary PJ polyp. Pathological examination revealed a high tissue homogeneity with >80% epithelial cells. Whole-genome sequencing failed to identify any clonal mutations but demonstrated a significant number of subclonal mutations. No somatic or germline mutations were found at the STK11 locus, suggesting that solitary PJ polyps are genetically distinct from Peutz–Jeghers polyps. In addition, methylome analysis revealed global hypomethylation and CpG island hypermethylation, two features that have been described as hallmarks of the colorectal cancer epigenome. These results provide an example of a premalignant lesion that is defined by epigenetic, rather than genetic changes. Furthermore, our findings support the notion that solitary PJ polyps constitute neoplastic tissue with malignant potential that should be removed for cancer prevention.

INTRODUCTION

Colon polyps are outgrowths of the colon mucosa and can progress to colon cancer. Polyps with malignant potential are categorized as neoplastic polyps, whereas polyps that are considered to have no cancer risk are categorized as nonneoplastic. Each of these categories is further subdivided into multiple histological subtypes. The Peutz–Jeghers (PJ) polyp constitutes a rare subtype with a characteristic histological appearance showing tree-like branching of smooth muscle cells and lobular organization of colonic crypts (Tse et al. 2013). PJ polyps almost exclusively occur in the context of the Peutz–Jeghers syndrome, which is characterized by mucocutaneous hyperpigmentation and the development of multiple intestinal polyps in the first two to three decades of life. Peutz–Jeghers is an autosomal dominant disorder caused by pathogenic variants of the STK11 gene located on human Chromosome 19p13 (Yoo et al. 2002).
Even though PJ polyps are categorized as nonneoplastic, the disorder is associated with a 93% cancer risk (Mehenni et al. 2006), and the risk of colon cancer in Peutz–Jeghers patients has been estimated at 39% (van Lier et al. 2010). Also, neoplastic transformation of PJ polyps has been documented (Defago et al. 1996). Contrary to the polyps associated with Peutz–Jeghers syndrome, however, the molecular pathology and malignant potential of solitary PJ polyps is unknown. Only one study reported molecular analysis of a solitary PJ polyp finding no loss of heterozygosity at the 19p13.3 locus and no mutation of STK11 (Kitaoka et al. 2004), suggesting that solitary PJ polyps constitute a separate entity. To further characterize the molecular pathology and malignant potential we here conducted the first comprehensive DNA sequencing and DNA methylation analysis of a solitary PJ polyp.

RESULTS

Clinical Presentation and Family History
The patient complained of intermittent midabdominal pain that had started ~6 mo prior to admission. In addition, he reported a recent change of bowel habits with intermittent loose stools, no loss of weight, and no rectal bleeding. Past medical history included coronary heart disease and chronic obstructive pulmonary disease. The family history was negative for gastrointestinal tumors. Outpatient colonoscopy had shown a pedunculated polyp located in the distal section of the sigmoid colon and sigmoid diverticulosis. The patient was then referred to our clinic for polypectomy. The age of the patient at the time of polypectomy was 77 yr.

At admission clinical examination and laboratory studies were unremarkable and mucocutaneous hyperpigmentation was not detectable. Repeat colonoscopy in our hospital confirmed sigmoid diverticulosis and a pedunculated polyp of the distal sigmoid colon located at 20 cm from the anocutaneous line with a diameter of ~3 cm. No other polyps were detected by colonoscopy. The polyp was removed by an endoscopic snare and submitted for pathological assessment. Hematoxylin and eosin staining showed the characteristic lobular grouping of crypts surrounded by branching smooth muscle fibers (Fig. 1). The fraction of epithelial cells in the polyp was estimated to be >80%.

Genomic Analyses
Whole-genome sequencing was performed at 70× genome coverage, respectively, for the polyp sample and for a blood sample from the same patient (Table 1). Data analysis identified 2079 high-confidence somatic single-nucleotide variants (SNVs), most of which correspond to C > T substitutions in an ACG trinucleotide context. Allele frequencies show a sharp peak at 0.1, indicating that most of the mutations are nonclonal. This pattern is reflected in the subset of 17 missense variants (Table 2). Notably, the ERBB3 p.P306H mutation has been described in the COSMIC (Catalogue of Somatic Mutations in Cancer) database and was also identified in a non-small-cell lung adenocarcinoma in our in-house tumor cohorts. Located in the growth factor receptor/furin-like cysteine-rich domain, it is annotated as pathogenic by various functional impact prediction tools.

Analysis of mutational signatures (Alexandrov et al. 2013) identified AC1 as the dominant contributor. This signature corresponds to the spontaneous deamination of 5-methylcytosine, correlating with the age of the patient. We did not find any signatures related to DNA mismatch repair, which is in line with the stable genomic profile. Copy-number analysis based on read depth plots and structural variation detection tools failed to reveal any reliable focal or large-scale somatic aberrations. We observed a subclonal loss of the Y Chromosome in both the blood and, to a lesser extent, the polyp. This phenomenon can be explained by age-related loss of Y in a fraction of white blood cells (Pierre and Hoagland 1972; Forsberg et al. 2014).
From the 42 high-confidence somatic insertions and deletions (indels) that were identified genome-wide, none fall into a coding gene or splice site, probably because of allele frequencies below the default reliability threshold. Lowering the threshold identified 243 indels, three of which were located in coding regions (Table 2). The \textit{FLT3} frameshift insertion is located in the protein kinase domain, just 30 amino acids from the hotspot position at D835.

Notably, neither somatic nor germline mutations or copy-number alterations were found to be present at the \textit{STK11} locus, even though this gene is often found mutated in Peutz–Jeghers patients (Yoo et al. 2002). The only deviation from the reference sequence is the common intronic variant rs4807008, for which the patient is homozygous. We could also not identify any rare or private coding germline SNV or indel in known cancer-predisposing genes. These results are consistent with the findings from the only other published study investigating the \textit{STK11} mutation status of a solitary PJ polyp (Kitaoka et al. 2004).

DNA Methylation Analyses

Because our genomic analysis did not detect any clonal mutations, we also analyzed the epigenome of the polyp. To this end, we used array-based methylation profiling of 450,000 methylation sites in the human genome (Infinium HumanMethylation450 arrays). The same platform was also used by The Cancer Genome Atlas (TCGA) for the analysis of 373 colorectal tumors and 45 control colon mucosa samples. Principal component analysis of the combined data sets showed that the solitary PJ polyp clustered separately from the majority of the normal colonic mucosa samples (Fig. 2A), thus indicating an aberrant methylome in the adenoma sample.

To further analyze these methylation changes, methylation values of individual probes were compared between the polyp and the TCGA normal data set. This identified 306

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**Table 1.** Sequencing coverage of whole-genome sequencing

| Sample | Read pairs sequenced | Reads aligned | PCR duplicates | Average read coverage |
|--------|----------------------|---------------|----------------|-----------------------|
| Polyp  | 1,680,950,965        | 99.73%        | 8.70%          | 69.97×                |
| Blood  | 1,764,374,344        | 99.56%        | 8.50%          | 73.81×                |

PCR, polymerase chain reaction.
| Chr | Position | dbSNP ID | Ref. base | Alt. base | Gene   | Variant type | Predicted effect | Coding DNA exchange | Amino acid exchange | Genotype | Allele freq. | COSMIC ID  |
|-----|----------|----------|-----------|-----------|--------|--------------|------------------|---------------------|---------------------|-----------|--------------|------------|
| 1   | 38082222 | rs200218812 | C         | T         | RSPO1  | Substitution | Missense         | c.G220A             | p.V74I              | heter.   | 0.07         |            |
| 1   | 158622275 |          | C         | A         | SPTA1  | Substitution | Missense         | c.G3357T            | p.K1119N            | heter.   | 0.06         |            |
| 3   | 73432532 |          | G         | A         | PDZRN3 | Substitution | Missense         | C3185T              | p.S1062             | heter.   | 0.07         |            |
| 4   | 156860666 | rs199672156 | C         | T         | CTSO   | Substitution | Missense         | c.G409A             | p.V137M            | heter.   | 0.10         |            |
| 5   | 52097492 |          | CGGA      | C         | PELO   | Deletion     | No frameshift    | c.977_979del        | p.326_327del        | heter.   | 0.05         |            |
| 5   | 52097497 |          | C         | CT        | PELO   | Insertion    | Frameshift       | c.982_983insT       | p.R328fs            | heter.   | 0.05         |            |
| 5   | 148753978 |          | C         | T         | IL17B  | Substitution | Missense         | c.G497A             | p.R166H            | heter.   | 0.08         |            |
| 6   | 102124509 |          | T         | A         | GRIK2  | Substitution | Missense         | c.T553A             | p.L185M            | heter.   | 0.06         |            |
| 7   | 95906629 |          | C         | T         | SLC25A13 | Substitution | Missense         | c.G91A              | p.G315              | heter.   | 0.06         | COSM3883588 |
| 8   | 25745417 |          | C         | T         | EBF2   | Substitution | Missense         | c.G823A             | p.G275R             | heter.   | 0.07         |            |
| 8   | 67355035 |          | C         | T         | ADHFE1 | Substitution | Missense         | c.C100T             | p.P34S              | heter.   | 0.07         |            |
| 11  | 73179461 |          | A         | G         | FAM168A | Substitution | Missense         | c.T59C              | p.M20T              | heter.   | 0.09         |            |
| 12  | 56482369 |          | C         | A         | ERBB3  | Substitution | Missense         | c.C917A             | p.P306H             | heter.   | 0.09         | COSM403445 |
| 13  | 27257046 | rs37084396 | G         | A         | WASF3  | Substitution | Missense         | c.G1286A            | p.R429Q             | heter.   | 0.11         |            |
| 13  | 28589790 |          | T         | FL3       |    | Insertion    | Frameshift       | c.2597_2598insGACCACCAG | p.G866fs            | heter.   | 0.07         |            |
| 13  | 50141395 |          | C         | A         | RCBTB1 | Substitution | Missense         | c.G21T              | p.W7C               | heter.   | 0.10         |            |
| 14  | 24045075 |          | C         | A         | JPH4   | Substitution | Missense         | c.G970T             | p.D324Y             | heter.   | 0.10         |            |
| 19  | 50393133 |          | C         | T         | CAPB5  | Substitution | Missense         | c.G518A             | p.R173H             | heter.   | 0.09         | COSM299924 |
| 19  | 52147182 |          | G         | A         | SIGLEC14 | Substitution | Missense         | c.G1498A            | p.A500T             | heter.   | 0.09         |            |

*dbSNP, Database for Short Genetic Variations; COSMIC, Catalogue of Somatic Mutations in Cancer; het., heterozygous.*
hypermethylated probes and 11,276 hypomethylated probes (Fig. 2B). Interestingly, the hypermethylated probes were strongly associated with CpG islands, whereas the hypomethylated probes were often associated with intergenic regions (Fig. 2B). These features have been described as the defining hallmarks of the colon cancer methylome (Berman et al. 2012). We also identified “tumor prone CpG islands” (Berman et al. 2012) using the TCGA colorectal cancer data set and subsequently compared their methylation with the polyp data (see Methods). This identified several CpG islands that showed tumor-associated hypermethylation in the solitary PJ polyp (Fig. 2C). Of note, no methylation changes were detected at the STK11 locus (Fig. 2D), indicating that the locus is not affected by a cancer-specific epigenetic mutation.

DISCUSSION

Solitary PJ polyps are a rare type of hamartomatous polyps of unknown etiology. Our report provides the first genome sequence of a solitary PJ polyp and reveals a number of genetic variants with relatively low allele frequencies (~10%) and in genes that are not commonly
associated with colorectal carcinogenesis. Consistent with a previous report (Kitaoka et al. 2004), no genetic alterations were detected at the STK11 locus, which is commonly associated with Peutz–Jeghers. These findings indicate that the molecular profile of solitary PJ polyps differs from Peutz–Jeghers syndrome. Our report also provides the first methylome analysis of a solitary PJ polyp and reveals moderate epigenetic alterations that are similar to the methylation changes of colorectal tumors. Together, our findings establish the first comprehensive genetic profile of a solitary PJ polyp.

Colorectal polyps are precursor lesions of colorectal cancer and are generally considered to evolve clonally from the colorectal mucosa (Fearon et al. 1987). The absence of clonal mutations in the solitary PJ polyp therefore suggests that the initial signal for clonal expansion is not provided by genetic changes in epithelial cells but by other mechanisms. Indeed, our data can be interpreted to reflect a substantial intrapolyt heterogeneity for subclonal mutations, which is consistent with the “Big Bang” model of colorectal carcinogenesis (Sottoriva et al. 2015). The acquisition of a tumor-driving mutation in a specific subclone could then represent the initiating event for tumor growth.

Interestingly, our methylome analysis revealed additional molecular alterations of the polyp genome. A comparative methylation pattern analysis with data sets from normal colorectal mucosa and colorectal tumor samples revealed moderate but clearly detectable epigenetic changes in the polyp that resemble the methylation changes commonly associated with colorectal cancer (Berman et al. 2012). More specifically, we identified “tumor-prone CpG islands” that showed a similar degree of hypermethylation in the polyp and in colorectal cancer. These findings are consistent with the notion that epigenetic changes represent early events during colorectal carcinogenesis (Lao and Grady 2011).

It is well established that Peutz–Jeghers patients carry a significant cancer risk, and it has been shown that cancer can develop in PJ polyps (McGarrity et al. 2000). Accordingly, surveillance colonoscopy is recommended in affected individuals starting at age 8, and removal of polyps of >5 mm is recommended (Syngal et al. 2015). Contrary to the polyps from Peutz–Jeghers patients, solitary PJ polyps are considered to have no malignant potential (Oncel et al. 2003). However, because solitary PJ polyps constitute a rare entity it is not possible to reliably determine malignant potential by clinical studies. Rather, cancer risk has to be extrapolated from molecular data. Our finding of tumor-like CpG island hypermethylation and genomic hypomethylation in a solitary PJ polyp supports the notion that such polyps do carry a cancer risk and should be removed.

**METHODS**

Pathology and DNA Isolation

The polyp was removed by colonoscopy using an endoscopic snare. Subsequently, a representative part of the sample was removed and placed into RNAlater solution for stabilization of nucleic acids. The remaining portion of the polyp was formalin fixed and paraffin embedded for pathological analysis. DNA was isolated using the QIAGEN Blood and Tissue Kit. Following hematoxylin and eosin staining of paraffin embedded sections pathological assessment of polyp morphology was conducted according to current guidelines (Quirke et al. 2011).

Genomic Analyses

Each sample was sequenced on two lanes of an Illumina X-Ten platform (see Table 1 for sequencing coverages). Reads were mapped to the 1000 Genomes phase 2 assembly of the human reference genome GRCh37 (NCBI build 37.1, downloaded from ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/reference/phase2_reference_assembly_sequence) using
a hardware-accelerated version of BWA (Burrows–Wheeler alignment) (Li and Durbin 2009) version 0.7.8 mem with option -T 0. The sambamba package (Tarasov et al. 2015) version 0.4.6 and 0.5.9, respectively, was used for sorting and to mark polymerase chain reaction (PCR) duplicates during merging the per-lane BAM (binary alignment) files.

For detection of SNVs we used our in-house pipeline. The pipeline is based on SAMtools mpileup and BCFTools (Li et al. 2009) version 0.1.19 with parameter adjustments to allow calling of somatic variants with low allele frequency using heuristic filtering as described previously (Jones et al. 2013). We applied Platypus (Rimmer et al. 2014) version 0.7.4 to identify short indels with a similar reliability scoring as for SNVs. After not detecting any coding indels with the default high-confidence threshold of 8, the value was lowered to 7 to allow also medium-confidence events with variant frequency of <10% and allele bias at the same time. All mutations were annotated with ANNOVAR (Wang et al. 2010) version November 2014, using the GENCODE version 19 gene model. Additional information was retrieved by overlapping the genomic positions with dbSNP (Database for Short Genetic Variations) version 141, 1000 Genomes phase1 integrated calls 20101123, COSMIC version 71, and dbNSFP (Database for Nonsynonymous SNPs’ Functional Predictions) version 2.9 (Liu et al. 2011, 2013).

A supervised analysis of the genome-wide somatic SNVs for mutational signatures (Alexandrov et al. 2013) was conducted with the R package YAPSA. A detailed description of the algorithm is provided elsewhere (Dieter et al. 2016). We searched for structural variants with CREST (Wang et al. 2011) and applied in-house tools for copy-number identification.

**DNA Methylation Analysis**

Available Infinium 450K raw data idat files from colon adenocarcinoma, rectal adenocarcinoma, and corresponding normal samples were downloaded from The Cancer Genome Atlas repository and preprocessed using Minfi (Aryee et al. 2014). Probes located on the sex chromosomes, or probes that failed detection P-value testing in at least one sample, were removed from the analysis. Probes showing cross-reactivity to at least one distant site, and thus allowing a mismatch of one base in the probe sequence, were also eliminated. Finally, probes containing a SNP with an allele frequency of at least 0.01 and located a maximum of 5 bases away from the single-base extension (SBE) site were also removed. β values were extracted from the remaining probes.

Principal component analysis using the R package FactoMiner (Lê et al. 2008) was performed with the extracted and merged β values of the TCGA data set and the PJ polyp data. Differentially methylated probes between the TCGA normal data set and the PJ polyp were addressed by identifying the probes in the PJ polyp data with an absolute β-value difference of ≥0.1 from the mean β value of the TCGA normal data set and with an absolute β value greater than the mean β value plus three times the standard deviation of the TCGA normal data set.

Hypermethylated tumor-prone CpG islands were identified in the TCGA data set by averaging the β values of all probes per CpG island and comparing the result of the tumor and normal data using a two-sided t-test. CpG islands showing a β-value difference of ≥0.1 and a P-value of <0.01, were considered hypermethylated in colorectal tumors.

**ADDITIONAL INFORMATION**

**Data Deposition and Access**

Genome-sequencing data are available from the NCBI Sequence Read Archive (SRA; https://www.ncbi.nlm.nih.gov/sra) under the accession numbers SRR5028731 (polyp) and
SRR5028722 (blood). Methylation data are available from the EBI-EMBL ArrayExpress database (https://www.ebi.ac.uk/arrayexpress/) under the accession number E-MTAB-5255.

**Ethics Statement**
Written consent for tissue sampling and analysis was obtained from the patient and was conducted as approved by the ethics committee of the University of Heidelberg (reference number S-101/2010). Also, permission in writing was obtained from the patient to allow for publication of genetic data.

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**Author Contributions**
H.L. provided patient samples. H.L. and F.L. designed the research. F.B. and B.H. performed the research and analyzed the data. B.B. contributed to the data analysis. H.L., F.B., B.H., and F.L. wrote the manuscript. All authors made significant contributions to, reviewed, and approved the final version of the manuscript.

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