Whole-exome Sequencing Reveals the Etiology of the Rare Primary Hepatic Mucoepidermoid Carcinoma

Ping Hou
Nanchang University Second Affiliated Hospital

Xiaoyan Su
Nanchang University Second Affiliated Hospital

Wei Cao
Nanchang University First Affiliated Hospital

Liping Xu
Nanchang university third affiliated hospital

Rongguyi Zhang
Nanchang University Second Affiliated Hospital

Zhihao Huang
Nanchang University Second Affiliated Hospital

Jiakun Wang
Nanchang University Second Affiliated Hospital

Lixiang Li
Nanchang University Second Affiliated Hospital

Linquan Wu
Nanchang University Second Affiliated Hospital

Wenjun Liao (liaowenjun120@163.com)
Nanchang University Second Affiliated Hospital

Research

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Abstract

Background Primary hepatic mucoepidermoid carcinoma (HMEC) is extremely rare and the molecular etiology is still unknown. Recently, The CRTC1-MAML2 fusion gene was detected in a primary HMEC which is often associated with MEC of salivary gland in the literature.

Methods a 64-year-old male was diagnosed with HMEC based on malignant squamous cells and mucus-secreting cells in immunohistochemical examination. Whole-exome sequencing (WES) and sanger sequencing were used to reveal the molecular characteristics of HMEC, and analysis with public datas among hepatocellular carcinoma, cholangiocarcinoma and salivary MEC. Meanwhile, The susceptibility genes were identified in pedigree investigation.

Result Significant somatic mutations in GNAS, KMT2C, ELF3 genes were identified in primary HMEC by WES and sanger sequencing. Meanwhile, through public data analysis, somatic GNAS gene altered in 2.1% hepatobiliary tumors, and typically GNAS occur at exon 8, in which Arg201 is converted to either a cysteine (R201C) or a histidine (R201H) related with cholangiocarcinoma associated with parasite infection. Furthermore, heterozygous germline mutations of FANCA, FANCI, FANCJ/BRIP1 and FAN1 genes were also identified. Pedigree investigation verified that mutation of susceptibility genes of Fanconi’s anemia were present in the pedigree.

Conclusions It was the first time to demonstrate the molecular etiology of the rare HMEC associated with germline Fanconi’s anemia mutations and somatic GNAS R201H mutation.

Background

Mucoepidermoid carcinoma (MEC) is a relatively common malignant neoplasm of salivary gland and mainly characterized by the pathological manifestation of mucous cells, intermediate cells and epidermoid malignant cells in intimately mixed nest. However, MEC rarely occurs at other sites of the body like the esophagus, thyroid gland, lacrimal gland, lung, thymus, breast, anal canal, uterine cervix, hepatobiliary and pancreatic system. Over 50–60% of salivary mucoepidermoid carcinoma (SMEC) are associated with a fusion gene, CRTC1/CRTC3-MAML2, which results from the chromosomal translocation t(11;19)(q21;p13). The CRTC1-MAML2 fusion gene was also observed in primary cervical MEC and bronchopulmonary MEC, which are different from adenosquamous carcinoma. However, pancreatic MEC did not show the CRTC1/3-MAML2 fusion and were a morphologic variant of pancreatic adenosquamous carcinoma. While it was controversy whether the CRTC1-MAML2 fusion gene was detected in primary hepatic MEC, otherwise MEC was also associated with hematopoietic stem-cell transplant in Fanconi anemia patients. So far, only 23 cases with primary hepatobiliary MEC (including hepatic, biliary and gall bladder MEC) that were pathomorphologically similar to SMEC have been reported in literatures (Table 1), but the molecular etiology of HMEC is still unknown. Here we revealed the genetic abnormality of a primary hepatic MEC by whole-exome sequencing (WES).
This study was approved by the Ethics Committee of the Second Affiliated Hospital of Nanchang University, and informed consent was obtained from the patient and his family. Parts of tumor tissues and corresponding non-tumor tissues were immediately conserved in a liquid nitrogen tank at -80°C. Some.

Methods And Case Material

A 64-year-old male was admitted to the Second Affiliated Hospital of Nanchang University on September 16, 2019 with a complaint of repeated abdominal distension over a period of one year. He had a previous history of icteric hepatitis and unknown biliary tract surgery 30 years prior and coronary artery stenosis disease with an implanted stent one year ago. No special abnormality was found in the physical examination. Most preoperative laboratory tests were in the normal range, including serum glutamic pyruvic transaminase, glutamic oxaloacetic transaminase, total bilirubin, electrolyte, glucose and platelet count, and serum alpha fetoprotein level. Hepatitis B surface antigen and hepatitis C antibody were both negative. Abnormal blood test results included C-reactive protein value, < 37 U/mL. Ultrasonography, magnetic resonance imaging and abdominal enhanced computed tomography (CT) revealed a mass lesion measuring approximate 10 cm in diameter at the left hepatic lobe and intrahepatic bile ducts with multiple stones (Supplementary Fig. 1A.B.C). The patient underwent resection with left hemihepatectomy and choledocholithotomy and T-tube drainage. Then, the patient was diagnosed with HMEC based on malignant squamous cells and mucus-secreting cells in pathological examination. There was gradually increased jaundice in the postoperative term, and the patient died of hepatic function failure at postoperative three months.

This study was approved by the Ethics Committee of the Second Affiliated Hospital of Nanchang University, and informed consent was obtained from the patient and his family. Parts of tumor tissues and corresponding non-tumor tissues were immediately conserved in a liquid nitrogen tank at -80°C. Some.
tissue samples were fixed in 10% formaldehyde and paraffin-embedded for pathological diagnosis and immunohistochemistry. Exome sequencing and Sanger sequencing were performed in fresh tumor tissue and corresponding non-tumor tissue. The WES datas from this study were deposited in NCBI Sequence Read Archive under accession SRA:SRP 266690.

**Histopathologica examination**

Hematoxylin eosin staining were detected for tumor samples and correspondent non-tumor samples. Strictly follow the standard procedure of immunohistochemistry. primary antibody diluted at 1:50. Antibody concluded MUC1 (Servicebio, mouse monoclonal, diluted 1:200), MUC5AC(Servicebio, mouse monoclonal, diluted 1:100), CK7 (Servicebio, mouse monoclonal, diluted 1:200), CK19 (Servicebio, rabbit polyclonal, diluted 1:500), and P63 (Servicebio, rabbit polyclonal, diluted 1:500). CEA(Servicebio, mouse monoclonal, diluted 1:200). Both of the two senior pathology specialists from Affiliated Hospital of NanChang University (XY Su and LP Xu) who examined the results suggested a diagnosis of HMEC.

**WES**

Exome capture was performed using 0.6 µg genomic DNA per sample. Sequencing libraries were generated using the Agilent SureSelect Human All Exon V6 kit (Agilent Technologies, CA, USA). The DNA libraries were sequenced on an Illumina HiSeq platform and 150 bp paired-end reads were generated. The original fluorescence image files were transformed to raw data by base calling and clean data was in turn recorded in FASTQ format. Valid sequencing data was mapped to the reference human genome (UCSC hg19) by Burrows-Wheeler Aligner software and results were stored in BAM format. Samtools mpileup and bcftools were used to do variant calling and identify single-nucleotide variant (SNV) and InDel. Functional annotation ANNOVAR was performed to do annotation for Variant Call Format and detect variants by 1000 Genome, dbSNP and other databases. Given the significance of exonic variants, gene transcript annotation databases such as Consensus CDS, RefSeq, Ensembl and UCSC were also included to determine amino acid alterations. Based on the paired samples of tumor tissue and corresponding non-tumor tissues, somatic SNVs were detected by MuTect, somatic InDels by Strelka and somatic CNVs by Control-FREEC. Software.SIFT, Polyphen/Polyphen2 and MutationAssessor were used to predict the impact of SNV on function; and truncated mutations (nonsense and frameshift) also resulted in high functional impact alterations.

**Public datas analysis**

The search and analytic strategy was performed using the public Datas (http://www.ncbi.nlm.nih.gov/). SNVs were examined in hepatocellular carcinoma (HCC) and cholangiocarcinoma (CHL). We also downloaded SMEC SNVs datas from published studies to examine the relation between alternations in the current case with downloaded data. We used an online tool to make Venn diagrams.

**Sanger sequencing and Pedigree survey**

Purified DNA was isolated from the proband’s fresh tumor tissue and corresponding non-tumor tissue and from peripheral blood lymphocytes of four family members using a TIANamp Genomic DNA Kit (Beijing Biotech, China). PCR was performed using the PTC-200PC instrument (BIO-RAD, USA). Primer sequences are shown in Supplementary table 2. The PCR products were examined using an ABI 3730XL DNA Analyzer (Applied Biosystems, USA).

**Results**

**Microscopic Features and Immunohistochemistry**

The mass was an irregular specimen measuring approximately 10 × 7 × 6 cm, revealing no fibrous capsule and tough solid texture, and resection from the middle edge showed grayish-white nodules without necrosis. Black pigment stones were also found in the small intrahepatic bile duct (Supplementary Fig. 1D). The resection edges were free of tumor. There was an absence of cirrhosis in the corresponding non-tumor sample, but massive inflammatory cell infiltration was observed in tumorous and portal areas. Hematoxylin and eosin staining revealed that the tumor cells were composed of epidermoid malignant cells, mucous cells and intermediate cells. Alcian blue staining revealed the mucin in mucin-producing cells. The distribution of epidermoid cell was not nest-like but was surrounded with intercellular bridges, with only a few glandular structures and occasional keratinization. Mucin-producing cells showed mainly abundant clear cytoplasm with eccentric pyknotic nuclei and predominantly showed intracellular mucus and rare extracellular mucus. The heterogeneity of epidermoid cells was obvious, with infiltrated stroma and invasion, but nerve, lymphovascular, blood invasion and necrosis were not apparent (Fig. 1A, B, C, D). Immunohistochemically, MUC5AC was only positive in the cytoplasm of mucous cells (Fig. 2A). Positive MUC1 was localized to the squamous and mucous tumor cell membrane (Fig. 2B). p63 was diffusely positive in the nuclei of malignant squamoid cells and mucinous cells (Fig. 2C), but cytoplasmic CK7 and CK19 were focal positive (Fig. 2D, E). CEA was negative on all tumor cells (Fig. 2F).

**Somatic variants in the proband**

The sequencing of tumor tissue and corresponding non-tumor tissue in proband provided average coverage depth on target of 175X and 112X, and >10X coverage of the targeted gene regions were 94% and 98%, respectively. The total somatic variants included 135 SNVs (53 missenses, 31 synonymous, 3 nonsense and others) and 4 Indels (2 frameshift insertions and 2 frameshift deletions). A total of 252 copy number variants (CNVs) (126 gain, 126 loss) were identified in 2591 genes (629 gene gain, 1691 gene loss). GNAS mutation (NM_000516:exon8:c.G602A:p.R201H) was detected in both tumor tissue and corresponding non-tumor tissue by WES, but the wild-type gene was determined in the corresponding non-tumor tissue by Sanger sequencing. In addition, protein-truncating genetic variants included frameshift indel mutation in ELF3 (c:909dupC:p.F303fs) and nonsense mutations in DOCK3(c:C2483A:p.S828X) and KMT2C (c.C1519T:p.Q507X) were detected in tumor samples by Sanger sequencing (Fig. 3D). The Circos map visually summarized the somatic genomic variations of SNVs, INDEL and CNVs of the HMEC (Supplementary Fig. 2).

**Bioinformatic**
A total of 1004 significant mutated genes (source: Onco KB) in SNVs from 8 studies contained 1487 HCC patients (supplementary Fig. 3A) and 654 significant mutated genes (source: Onco KB) in SNVs from 7 studies contained 445 CHL patients were downloaded (supplementary Fig. 3B) (www.cbioportal.org). We also obtained 312 significant mutated genes in SNVs from 18 SMEC patients from Kang T et al. study. There were six SNVs (STAT1, TGFB1R1, NTC1, KMT2C, ELF3, GNAS) in the HMEC that overlapped with primary liver tumors (HCC and CHL), only CHD3 gene in HMEC that overlapped with SMEC (Fig. 3A). Especially, somatic GNAS gene (included missense mutation, amplification, truncating mutation) altered in 2.1% (9/445) patients with primary hepatobiliary tumors included intrahepatic cholangiocarcinoma, extrahepatic cholangiocarcinoma and perihilar cholangiocarcinoma (Fig. 3B). In total of 3 patients underwent GNAS (p.R201H/C) missense mutation (putative driver) in exon 8. Importantly, Sample ID (W012, T026) were Opisthorchis viverrini-associated with cholangiocarcinoma (SRP007970) (Fig. 3C).

**Germline variants in the proband and pedigree analysis**

The 56 germline variants in 20 Fanconi’s anemia pathway genes included 19 missense variants and 25 synonymous variants and 12 UTR3 and UTR5 in the proband's non-tumor tissue (Table 2, Fig. 4A).

### Table 2

The sanger sequencing of germline variants in proband and family members

| Gene     | Locus | HGVS nomenclature | Alt | Rff | Proband | Sibling | Sibling | Son | Daughter |
|----------|-------|-------------------|-----|-----|----------|---------|---------|-----|----------|
| FANCJ    | 15q26.1| exon4:c.C257T:p.A86V | CT  | C   | heterozygous | Non     | Non     | Non | Non      |
|          |       | exon22:c.G2225C:p.C742S | CG  | C   | heterozygous | Non     | Non     | Non | Non      |
| FAN1     | 15q13.2| exon2:c.G698A:p.G233E | GA  | G   | heterozygous | heterozygous | heterozygous | Non | Non      |
| FANCA    | 16q24.3| exon16:c.G1501A:p.G501S | CT  | C   | heterozygous | homozygous | homozygous | homozygous | homozygous |
|          |       | exon9:c.A796G:p.T266A | TC  | T   | heterozygous | homozygous | homozygous | homozygous | homozygous |
|          |       | exon26:c.G2426A:p.G809D | TT  | C   | homozygous | homozygous | homozygous | homozygous | homozygous |
| FANCD1   | 13q13.1| exon14:c.T7397C:p.V2466A | CC  | T   | homozygous | homozygous | homozygous | homozygous | homozygous |
|          |       | exon2:c.C269A:p.T90N   | TT  | G   | homozygous | homozygous | homozygous | homozygous | homozygous |
| FANCF    | 17q10 | exon8:c.A2449G:p.T817A | CC  | T   | homozygous | homozygous | homozygous | homozygous | homozygous |

There was no family history of Fanconi’s anemia and solid tumor. The proband’s parents died of cardiovascular and cerebrovascular diseases 10 years ago. Only the old sibling showed hypertension and type 2 diabetes. The proband’s offspring did not show any disease. The same mutational variants in FANCD1/BRCA2, FANCF/RFWD3, and FAP100/C17orf70 were verified in the pedigree. Heterozygous variants in FANCA (c.G1501A:p.G501S), FANCA (c.A796G:p.T266A) and FANCI/BRIP1 (c.T2755C:p.S919P) were detected in the proband, but homozygous mutations were shown in the other family members. Heterozygous FAN1 variant were detected in the siblings and proband. Only heterozygous FAN1 variants (c.C257T:p.A86V and c.G2225C:p.C742S) were unique in the proband (Table 2, Fig. 4A).

**Discussion**

The correlation between two rare diseases is really difficulty for clinicians. The emergence of next-generation sequencing technologies has provided a simple and powerful approach for discovering de novo disease-associated genes, and these methods have furthered our understanding of the rare HMEC, which lacked the traditional high risk factors such as hepatitis B, hepatitis C, long-term alcohol intake history and cirrhosis, like most primary HMEC cases reported in the literatures (Table 1). In 2006, Zhu et al. reported an adult male patient, also absent for high risk factors who was first diagnosed with unusual HCC metastatic to right proximal ulna and metachronous esophageal squamous cell carcinoma resulting from the history of Fancon's anemia. Linares et al also report on a 31-year-old female patients with synchronous squamous cell carcinoma of esophagus and HCC associated with Fanconi's anemia. However, the present case only showed mild anemia, with an absence of the typical Fanconi’s anemia phenotype, only present of family Fanconi’s anemia gene carriers. In fact, increasing evidence has suggested that monoallelic carriers for Fanconi’s anemia genes were characterized by a adult-set predisposition to most solid cancers, especially squamous cell carcinomas from the epithelia of genitourinary tracts and the aerodigestive system. A previous study by Alter et al. reported that 20–30% of germline Fanconi’s anemia presents with unusual solid malignancies as the first clinic manifestation and shows absence of the phenotype of congenital Fanconi’s anemia. Interestingly, three patients were reported with HMEC, also absence of Fancon’s anemia, combined with synchronous hepatocellular carcinoma, synchronous squamous cell carcinoma in the cranial skin and metachronous gastric malignant carcinoma.

In fact, the phenotypes of germline Fanconi’s anemia mutation-related solid tumors were probably determined in mutation loads or a dose-dependent manner. So far, a total of 22 proteins (FANCA—FANCF/RFWD3 genes coding) and other FA network genes have been identified to function in Fanconi’s
anemia pathway. These proteins participate in pathways including genome maintenance processes and DNA repair in response to DNA damage, as well as interstrand cross-linking repair, homologous recombination and non-homologous terminal junctions. Compared with the 22 Fanconi's anemia genes, FAN1 has a mild role in interstrand crosslink repair and even patients with homozygous mutation in FAN1 were absent of Fanconi's anemia features. However, Lachaud et al. showed that FAN1 defects can cause cancers in knockin mice, and also FAN1 variant in high-risk pancreatic cancers abolishes recruitment by Ub-FANCD2, resulted in genetic instability without affecting interstrand crosslink repair. Otherwise germline FAN1 mutation, occurred frequently in high-risk pancreatic cancers and hereditary susceptibility to familial colorectal cancers. In the present case, we testified FAN1 mutations were present in the pedigree. Therefore, we propose that germline Fanconi's anemia mutations are probably predisposing factors to the occurrence of HMEC.

More importantly, Mutations in the GNAS gene typically occur at exon 8, in which Arg201 is converted to either a cysteine (R201C) or a histidine (R201H), lead to activation of the Gs subunit. This Gs constitutively activate the intracellular cyclin adenosine monophosphate (AMP) signal pathway. Otherwise, GNAS R201H/C missense mutation have been showed a cross-communication between JAK/STAT and cyclic-AMP pathways in rare subtypes of liver inflammatory tumorogenesis. Farges et al. also described an adult male patient with inflammatory hepatic adenoma associated with Fanconi's anemia and somatic GNAS mutation who developed malignant transformation of HCC. Consistent with inflammatory phenotype in the present HMEC, previous studies reported three HMEC patients with inflammation phenotype and inflammatory cell infiltration in the corresponding tumor-free liver tissue. Otherwise, somatic GNAS R201H/C mutation occurs frequently in secreting-mucous tumors like pancreatic intraductal papillary mucinous neoplasms (IPMN) as a bona fide precursor to carcinogenesis. GNAS R201 mutation was detected in pancreatic IPMN tissue, secretin-stimulated pancreatic juice and peripheral blood. The mucus production and carcinogenesis of the intestinal subtype of intraductal papillary neoplasm of the biliary ducts (IPNB), a counterpart of pancreatic IPMN, have also been connected with gain-of-function mutations of GNAS R201. Otherwise, The protein-truncating genetic variants in the present case included a frameshift insertion in ELF3 and nonsense mutation in KMT2C, which were also reported in biliary tumors as tumor suppressors, with consistently positive immunohistochemical results for CK7 and CK19 in the present case, which may suggest the possibility that HMEC originates from the biliary system. Moreover, GNAS and KMT2C were almost only mutated in O. viverrini-CHL compared non-O. viverrini-CHL, which consistent with Jarley K et al study that three HMEC patients were induced by biliary parasite infection and may explained that most HMECs(20/23) were reported in Asia areas with a high incidence of parasites.

In conclusion, the molecular characteristics of the current HMEC were more similar to primary liver biliary tumors rather than to SMEC. We present here for the first time the etiology of HMEC associated with germline Fanconi’s anemia mutations and somatic GNAS R201 mutation.

Abbreviations

MEC: Mucoepidermoid carcinoma, HMEC: hepatic mucoepidermoid carcinoma, SMEC: salivary gland mucoepidermoid carcinoma, HCC: hepatocellular carcinoma, CHL: cholangiocarcinoma, IPMN: intraductal papillary mucinous neoplasms, IPNB: intraductal papillary neoplasm of the biliary ducts, SNV: single-nucleotide variant, CNV: copy number variant. WES: whole-exome sequencing. Non: not have. NP: not provided, RL: right liver, LL: left liver, BD: bile duct, GB: gallbladder, S: surgery, R: Radiotherapy, C: Chemotherapy, Con: Conservative treatment, TACE: transcatheater arterial chemoembolization. PTCD: percutaneous transhepatic cholangial drainage. OS: overall survival.

Declarations

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Author contributions

Ping Hou and Xiaoyan Su who are the first co-author, contributed equally to the study. This study was designed by Wenjun Liao and Linquan Wu. Manuscript was written by Ping Hou and Xiaoyan Su. Pedigree analysis and clinical phenotype analysis was performed by Jiakun Wang and Zhihao Huang, with further review by Lixiang Li, Xiaoyan Su and Liping Xu. Whole exome sequencing and bioinformatic analysis by Wei Cao, Jiakun Wang, Rongguiyi Zhang, Zhihao Huang.

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Ethics approval and consent to participate

This study was approved by the Ethics Committee of the Second Affiliated Hospital of Nanchang University, and informed consent was obtained from the patient and his family.

Disclosed of potential conflicts of interest

The authors declare no conflict of interests. The authors declare no competing financial interest.

Consent for publication
The authors declare no competing financial interests.

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