RESEARCH ARTICLE

Effect of Xeroderma Pigmentosum Complementation Group F Polymorphisms on Gastric Cancer Risk and Associations with H. pylori Infection

Ji-Shun Zhang*, Chuan Zhang, Xue-Yan Yan, Zhi-Fang Yuan, Zhuo-Yang Duan, Hui Gao

Abstract

We conducted a hospital case-control study by genotyping four potential functional single nucleotide polymorphisms (SNPs) to assess the association of Xeroderma pigmentosum complementation group F (XPF) with gastric cancer susceptibility, and role of XPF polymorphisms in combination with H. pylori infection in risk definition. A total of 331 patients with gastric cancer and 355 controls were collected. Four SNPs of XPF, rs180067, rs1799801, rs2276466 and rs744154, were genotyped by Taqman real-time PCR method with a 7900 HT sequence detector system. The gastric cancer patients were more likely to have smoking habit, a family history of cancer and H. pylori infection. We did not find any significant difference in the genotype distributions of XPF rs180067, rs1799801, rs2276466 and rs744154 between cases and controls. However, multivariate logistic analysis showed a non-significant decreased risk in patients carrying rs180067 G allele, rs1799801 T allele or rs2276466 T allele genotypes. A non-significant increased risk of gastric cancer was found in individuals carrying the rs744154 GG genotype. A non-significant increased risk of gastric cancer was found in individuals carrying the rs744154 GG genotype. Stratification by H. pylori infection and smoking was not significantly different in polymorphisms of XPF rs180067, rs1799801, rs2276466 and rs744154. The four XPF SNPs did not show significant interaction with H. pylori infection and smoking status (P for interaction was 0.35 and 0.18, respectively). Our study indicated that polymorphisms in rs180067, rs1799801, rs2276466 and rs744154 may affect the risk of gastric cancer but further large sample size studies are needed to validate any association.

Keywords: Xeroderma pigmentosum complementation group F - SNPs - gastric cancer - H. pylori

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Introduction

Worldwide, gastric cancer is the second leading cause of death from cancer, with an estimated one million new cases in 2008 (988,000 cases), accounting for 8% of all cancer-related death worldwide. More than 70% of all gastric cancer cases occurred in developing countries, and approximately half of all cases occur in China (IARC, 2008). Although several epidemiologic studies suggest that Helicobacter pylori (H. pylori) infection is one of the most important risk factors for gastric cancer. It is estimated almost 30% of the world’s population are infected with H. pylori, but only about 1% of them occur gastric cancer (Graham et al., 1991; Parsonnet et al., 1997). Therefore, the genetic and environmental factors may influence the susceptibility to gastric cancer (Ghoshal et al., 2007; Ghoshal et al., 2008).

DNA repair play a key role in keeping the function of the stability and integrity of human genome. It is well known that there are five main DNA repair pathways with more than 130 genes, of which the nucleotide excision repair (NER) pathway plays a crucial DNA repair mechanism by removing various DNA lesions caused by UV radiation and some chemical agents, such as bulky adducts, cross links, oxidative DNA and alkylating damage and thymidine dimers (De Silva et al., 2000; Friedberg, 2001; Wood et al., 2001). It is reported that xeroderma pigmentosum (XP) complementation groups in human, XPA to XPG, represent the rate-limiting proteins in the NER pathway (Cleaver, 2000).

The XPF gene is located on chromosome 16p13.12, contains 11 exons and spans approximately 28.2 kb, and plays a key role in the 5' incision made in the NER pathway, and function as removing DNA interstrand cross-links and DNA double-strand breaks (Mueser et al., 1996; Friedberg, 2000). It is reported that polymorphisms in several SNPs of XPF alter genetic susceptibility to cancer, such as colorectal cancer, breast cancer, lung cancer and prostate cancer as well (Crew et al., 2007; Chang et al., 2008; Agalliu et al., 2010; Gil et al., 2011). However, there is still no evidence on the association between XPF polymorphisms and gastric cancer risk in Chinese
population (He et al., 2012). Therefore, we conducted a case-control study by genotyping four potential functional SNPs to assess the association of XPF polymorphisms with gastric cancer susceptibility, and role of XPF polymorphisms in combination with H. pylori infection and smoking status in the risk of gastric cancer.

Materials and Methods

All the subjects were collected from the Affiliated Beijing Chaoyang Hospital of Capital Medical University between December 2008 and November 2011. A total of 402 patients with newly histopathologically confirmed primary gastric cancer, including gastric cardia adenocarcinoma and non-cardia adenocarcinoma, were included in our study. Of 402 patients, 331 patients were willing to participate into our study, with a participation rate of 82.3%. All the cases were selected from the First Affiliated Hospital of Zhengzhou University. We excluded from those who suffered from secondary or recurrent tumors. A total of 377 controls were selected from the same hospital during the same time period, and 355 controls wanted to participate into our study. Controls were outpatients from Surgical Department, Plastic Surgery Department and ENT Department as well. Controls that had a history of cancer and digestive tract disorders were excluded. All patients were asked to provide 5 ml blood samples for DNA extraction.

The H. pylori infection status was determined by the method of enzyme linked immunosorbent assay (ELISA) from 5 ml blood. H. pylori IgG antibodies (HpIgG Ab) were measured using commercially available kit (Genesis Diagnostics, Cambridgeshire, UK) according to the manufacturer’s instructions.

We selected potential functional SNPs of interested XPF from Database of single nucleotide polymorphisms (SNPs) of NCBI(http://www.ncbi.nlm.nih.gov/) and SNPinfo (http://snpinfo.niehs.nih.gov/) with the following criteria: (1) the minor allele frequency should not less than 5% of the Chinese population; (2) influencing the microRNA binding sites activity. Genomic DNA was extracted using the buffy coat fraction of each blood sample and 5 ml blood samples for DNA extraction.

We did not find significant differences in the ages or sex distributions between cases and controls (P > 0.05). Of 331 cases with gastric cancer and 355 controls, 95% confidence intervals (95% CI) was used to assess the association between the polymorphisms in selected SNPs and gastric cancer risk by multivariate logistic regression model. The interaction of these SNPs with H. pylori infection and smoking status was also estimated by spearman correlation analysis. Statistical significance was set at P<0.05 and all tests were two-sided.

Results

The demographic and clinic characteristics of the selected cases and controls were shown in Table 1. The mean ages of the 331 cases with gastric cancer and 355 controls were 55.7±8.3 and 56.3±8.5 years, respectively. We did not find significant differences in the ages or sex distributions between cases and controls (P > 0.05). There was no significant difference in the drinking status between the two groups. However, the gastric cancer patients were more likely to have smoking habit, a family history of cancer and H. pylori infection (P < 0.05). Of the cancer cases, 54.6% were intestinal, and 40.8% were diffuse, 54.3% of the cases were early gastric cancer.

The genotype distributions of XPF rs180067, rs1799801, rs2276466 and rs744154 in cases and controls were summarized in table 2. All the genotype frequencies of selected SNPs in controls were in line with the Hardy-Weinberg equilibrium. However, we did not find the significant difference in the genotype distributions of XPF rs180067, rs1799801, rs2276466 and rs744154 between cases and controls. Multivariate logistic analysis was conducted to evaluate the effect of XPF rs180067, rs1799801, rs2276466 and rs744154 on the risk of gastric cancer. Using the wide-type genotype as the reference genotype, we found a non-significant decreased risk in patients carrying rs180067 G allele, rs1799801 T allele or rs2276466 T allele genotype. A non-significant increased frequency distributions of the selected SNPs between cases and controls were evaluated by χ² test. χ² test was used to analyze the Hardy-Weinberg equilibrium in controls. Adjusted odds ratios (OR) and their 95% confidence intervals (95% CI) was used to assess the association between the polymorphisms in selected SNPs and gastric cancer risk by multivariate logistic regression model. The interaction of these SNPs with H. pylori infection and smoking was also estimated by spearman correlation analysis. Statistical significance was set at P<0.05 and all tests were two-sides.

Table 1. Distributions of Demographic and Clinic Characteristics

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Cases %</th>
<th>Controls %</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, yr (Mean±SD)</td>
<td>55.7±8.3</td>
<td>56.3±8.5</td>
<td>0.82</td>
</tr>
<tr>
<td>Male</td>
<td>214</td>
<td>210</td>
<td>0.14</td>
</tr>
<tr>
<td>Family history of cancer</td>
<td>18</td>
<td>2</td>
<td>0.6</td>
</tr>
<tr>
<td>Smoking status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ever</td>
<td>108</td>
<td>86</td>
<td>0.015</td>
</tr>
<tr>
<td>Never</td>
<td>223</td>
<td>269</td>
<td>0.75</td>
</tr>
<tr>
<td>Drinking status</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ever</td>
<td>123</td>
<td>113</td>
<td>0.215</td>
</tr>
<tr>
<td>Never</td>
<td>208</td>
<td>242</td>
<td>0.683</td>
</tr>
<tr>
<td>H. pylori infection (positive)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Intestinal</td>
<td>181</td>
<td>151</td>
<td>0.54</td>
</tr>
<tr>
<td>Diffuse</td>
<td>135</td>
<td>151</td>
<td>0.408</td>
</tr>
<tr>
<td>Mixed</td>
<td>15</td>
<td>151</td>
<td>0.46</td>
</tr>
<tr>
<td>Stage of gastric cancer</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Early gastric cancer</td>
<td>180</td>
<td>151</td>
<td>0.543</td>
</tr>
<tr>
<td>Advanced gastric cancer</td>
<td>151</td>
<td>151</td>
<td>0.457</td>
</tr>
</tbody>
</table>

The demographic and clinic characteristics were compared between cases and controls by χ² test or student’s test. Differences in
infection and smoking status modified the association between the selected SNPs and gastric cancer risk. Although the published genome-wide associated study has not showed ERCC3 variants was an susceptibility loci, it is difficult to evaluate the effects of the genotypes for the potentially functional SNPs in XPF in the genome-wide associated study due to not enough samples. In the present study, we found the polymorphisms in rs180067, rs1799801, rs2276466 and rs744154 was not associated with gastric cancer risk, and no interaction was found between *H. pylori* and smoking.

The XPF rs2276466 and rs3136038 was found to be associated with a reduced risk of squamous cell carcinoma of the head and neck in a previous study (Yu et al., 2012). This study indicated the GF genotype of rs2276466 was significantly associated with a decreased risk of head and neck cancer, and TT genotype of rs3136038 showed a borderline significant decreased cancer risk (Yu et al., 2012). A previous meta-analysis indicated that XPF-rs1799801 may be associated with a reduced cancer risk in Caucasian population, and XPF-rs1800067 was related to a decreased risk of pancreatic cancer. However, a few studies have investigated the association between XPF SNPs and risk of gastric cancers, and only two studies reported their association (He et al., 2012; Zhou et al., 2012). One study conducted in China analyzed the two XPF functional SNPs, rs2276466C>G and rs6498486A>C, with risk of gastric cancer, and reported that no functional XPF SNPs contribute to risk of gastric cancer (He et al., 2012). Another study also conducted in China showed no significant association of XPF-357A/C with the risk of gastric cancer in a population of a high-incidence region of China (Zhou et al., 2012). Our study also found no significant association between polymorphisms in rs180067, rs1799801, rs2276466 and rs744154 and gastric cancer risk, which are similar to the previous results. However, our findings need to be confirmed by large-scale studies.

There were some limitations in our study. First, the patients in our study were selected from one hospital, which may be lack of presentation, and have selection and information bias. Second, we only analyze four polymorphisms in *H. pylori* infection and smoking status. Finally, we did not validate their association.

To the best of our knowledge, our study is the first attempt to evaluate the potential association between the four selected SNPs of XPF and risk of gastric cancer in Asian population. We did not find the significant association between the four selected SNPs and risk of gastric cancer. Moreover, we did not find the *H. pylori* infection and smoking status modified the association between the selected SNPs and gastric cancer risk. Although the published genome-wide associated study has not showed ERCC3 variants was an susceptibility loci, it is difficult to evaluate the effects of the genotypes for the potentially functional SNPs in XPF in the genome-wide associated study due to not enough samples. In the present study, we found the polymorphisms in rs180067, rs1799801, rs2276466 and rs744154 was not associated with gastric cancer risk, and no interaction was found between *H. pylori* and smoking.

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### Discussion

To the best of our knowledge, our study is the first attempt to evaluate the potential association between the four selected SNPs of XPF and risk of gastric cancer in Asian population. We did not find the significant association between the four selected SNPs and risk of gastric cancer. Moreover, we did not find the *H. pylori*

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