Pancreatic Carcinoma with Prominent Mucin Production: A Clinicopathologic and Immunohistochemical Study of 9 Cases

Makoto Saito1, Hiroshi Nishihara2, Mishie Tanino1, Taichi Kimura1, Yoshihiro Matsuno3, Kazuo Nagashima1,4, Michio Shimizu5, Shinya Tanaka1*

1Department of Cancer Pathology, Hokkaido University Graduate School of Medicine, Sapporo, Japan; 2Department of Translational Pathology, Hokkaido University Graduate School of Medicine, Sapporo, Japan; 3Department of Surgical Pathology, Hokkaido University Hospital, Sapporo, Japan; 4Sapporo Higashi Tokushukai Hospital, Sapporo, Japan; 5Department of Pathology, Saitama Medical University, Saitama, Japan.

Email: *tanaka@med.hokudai.ac.jp

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ABSTRACT

Pancreatic carcinoma occasionally associated with prominent mucin production and this type of tumor designated as PCM (pancreatic carcinoma with prominent mucin production) was diagnosed depends on subjective estimation of the amount of mucous area, and there has been no report on a quantitative evaluation of the amount of mucinous area in the tumor. To examine the feature of PCM, we analyzed 9 cases of PCM among 243 cases of pancreas carcinoma and evaluated the amount of mucin by imaging analysis. Morphologically, 5 cases were classified as intraductal papillary mucinous neoplasms (IPMN)-derived PCM and 4 cases were as ductal adenocarcinoma (DA)-derived PCM. Mucous composition was found to be more than 50% in all IPMN-derived PCM cases, and that was 40% - 50% in DA-derived PCM cases with one exception. IPMN-derived PCM cases showed expansive growth with pancreatic duct dilatation filled with mucin, while DA-derived PCM cases possessed mucin infiltration into interstitial tissue. Immunohistochemically, three of 4 DA-derived PCM cases were MUC1(-)/MUC2(+), and the results of expressions for p16 and Dpc4 suggesting that DA-derived PCM was similar to IPMN-derived PCM rather than ordinary DA. Survival rate of DA-derived PCM cases was lower than that of IPMN-derived PCM cases. We advocate that DA-derived PCM may constitute a borderline group between IPMN and ordinary DA.

Keywords: Pancreatic Carcinoma with Prominent Mucin Production (PCM); Mucinous Non-Cystic Carcinoma (MNCC); Intraductal Papillary Mucinous Neoplasm (IPMN); Ductal Adenocarcinoma (DA); Immunohistochemistry

1. Introduction

Pancreatic carcinoma includes a certain population of those with prominent mucin production (PCM), including mucinous non-cystic carcinoma (MNCC), and few study have been reported because such cases were rare (1% - 3%) among invasive pancreatic carcinoma [1-3]. Recently, MNCC of the pancreas has been regarded as an advanced form of intraductal papillary mucinous neoplasm (IPMN) [4-6]. MNCC of the pancreas is defined in which mucin occupies more than 50% of tumor area by World Health Organization (WHO) [7], the Armed Forces Institute of Pathology (AFIP) [4], and the Japanese classification of pancreatic carcinoma [8]. However, the mucous area is subjectively estimated and no specific methods have been reported to quantify the mucous area in PCM. In addition, few molecular biological evaluations of MNCC have been performed [9].

Pancreatic intraepithelial neoplasia (PanIN) is precursor lesion of ordinary ductal adenocarcinoma (DA), which is different from IPMN [10]. It has been shown that MUC1 expression suggests the “aggressive” pathway in which PanIN progressed to DA, while MUC2 is a marker for “indolent” pathway for IPMN-derived MNCC formation. The mucinous phenotype serves as a marker for the typing of pancreatic cancer [5,6]. In molecular models of pancreatic multistep carcinogenesis, the inactivation of the p16 gene, the overexpression of Cytin D1, and the inactivation of the p53 and Dpc4 genes have been suggested as early, intermediate, and late events, respectively [11-15].
To analyze the clinicopathologic and immunohistochemical features of PCM, we analyzed 9 cases among 243 invasive pancreatic cancer cases. By the morphological analysis, we classified these 9 cases into two groups of PCM which derived from IPMN or DA, referred as IPMN-derived PCM and DA-derived PCM, respectively. The amount of mucin was measured by using imaging software. Immunostainings of MUC1/MUC2, p16, Cyclin D1, p53, and Dpc4 were performed. The clinicopathologic differences between IPMN- and DA-derived PCM were discussed.

2. Materials and Methods

2.1. Case Analysis

We reviewed 243 cases of pancreatic carcinoma which were surgically resected and diagnosed at the Laboratory of Cancer Research, Department of Pathology, Hokkaido University Graduate School of Medicine and the Department of Surgical Pathology, Hokkaido University Hospital from 1992 to 2003. For PCM lesions, 9 cases were selected in which conspicuous mucin was macroscopically seen diffusely over the tumor on cut surface. Cases with focal mucin or microscopically inside the cancer cells forming glandular lumens were excluded. Clinical stage was described based on pathological findings according to the Japanese classification [8].

PCM cases exhibiting intraductal papillary structure in dilated main and/or branch pancreatic duct with large amount of mucin, were regarded as being derived from IPMN [4,7,8]. All remaining PCM cases were regarded as being derived from DA. IPMN cases were further classified into intestinal type and pancreatobiliary type consisting of epithelial cells [16-18]. The protocol of this study conformed to the ethical guidelines of the Declaration of Helsinki and informed consent was obtained from each patients.

2.2. Estimation of the Mucous Area (Figure 1)

In all 9 PCM cases, the maximum cut surface of the tumor (1 - 9 slides) was almost in agreement with the site with the greatest mucin content. In lower magnification image of the H&E staining, tumor lesions were discriminated using Adobe® Photoshop®, and mucinous and non-mucinous parts were independently displayed with artificial colors as to avoid overlapping of either part (Figures 1(a) and (b)). The ratios in pixels of the mucinous part to the tumor lesion were obtained.

2.3. Immunohistochemical Analysis

The formalin-fixed, paraffin-embedded blocks obtained from the 9 PCM cases were performed for immunostaining with regular method described elsewhere (or site ref). Primary antibodies used are following: MUC1 (Clone Ma695, Novocastra Laboratories, Newcastle, UK; dilution 1:100), MUC2 (Clone Ccp 58, Novocastra Laboratories, Newcastle, UK; dilution 1:100), p16 (Clone ZJ11; NeoMarkers, Fremont, CA; dilution 1:50), Cyclin D1 (Clone SP4, Lab Vision, Fremont, CA; dilution 1:100), p53 (Clone DO-7, DAKO Corporation, Kyoto, Japan; dilution 1:500), and Dpc4 (Clone B8, Santa Cruz Biotechnology, Santa Cruz, CA; dilution 1:100).

Concerning MUC1, MUC2, p16, and Dpc4, the normal pancreatic tissue in each section served as an internal positive control. MUC1 and MUC2 were judged to be positive when 5% or more of the tumor cells were stained [16]. Cyclin D1 and p53 were judged to be overexpression when the nuclei of the tumor cells were homogeneously stained in >5% and >10%, respectively [19, 20]. Complete absence of nuclear expression of p16, or complete absence of cytoplasmic and nuclear expression of Dpc4, was judged to be negative [21,22]. Fifty ordinary DA cases were also examined as controls. Fisher’s exact probability test was performed, and p < 0.05 was judged to be significant.

3. Results

3.1. Clinicopathological Findings (Table 1)

In 9 PCM cases, 5 were derived from IPMN, and the
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Table 1. Clinicopathological findings of PCM cases.

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Location</th>
<th>Size</th>
<th>LN</th>
<th>pStage</th>
<th>Outcome</th>
<th>Mucin</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>71</td>
<td>M</td>
<td>B-T</td>
<td>5.6</td>
<td>(+)</td>
<td>IVa</td>
<td>Dead (17 M)</td>
<td>58%</td>
</tr>
<tr>
<td>2</td>
<td>75</td>
<td>M</td>
<td>H</td>
<td>8.4</td>
<td>(-)</td>
<td>III</td>
<td>Alive (67 M)</td>
<td>64%</td>
</tr>
<tr>
<td>3</td>
<td>54</td>
<td>M</td>
<td>H</td>
<td>11</td>
<td>(-)</td>
<td>IVa</td>
<td>Alive (48 M)</td>
<td>61%</td>
</tr>
<tr>
<td>4</td>
<td>56</td>
<td>M</td>
<td>H</td>
<td>7.2</td>
<td>(+)</td>
<td>IVa</td>
<td>Dead (6 M)</td>
<td>60%</td>
</tr>
<tr>
<td>5</td>
<td>70</td>
<td>M</td>
<td>H</td>
<td>3.3</td>
<td>(-)</td>
<td>III</td>
<td>Alive (69 M)</td>
<td>54%</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Case</th>
<th>Age</th>
<th>Sex</th>
<th>Location</th>
<th>Size</th>
<th>LN</th>
<th>pStage</th>
<th>Outcome</th>
<th>Mucin</th>
</tr>
</thead>
<tbody>
<tr>
<td>6</td>
<td>66</td>
<td>M</td>
<td>T</td>
<td>2.5</td>
<td>(+)</td>
<td>III</td>
<td>Dead (29 M)</td>
<td>56%</td>
</tr>
<tr>
<td>7</td>
<td>73</td>
<td>M</td>
<td>H</td>
<td>5.1</td>
<td>(+)</td>
<td>III</td>
<td>Dead (42 M)</td>
<td>46%</td>
</tr>
<tr>
<td>8</td>
<td>64</td>
<td>M</td>
<td>T</td>
<td>5.0</td>
<td>(+)</td>
<td>IVa</td>
<td>Dead (53 M)</td>
<td>41%</td>
</tr>
<tr>
<td>9</td>
<td>63</td>
<td>F</td>
<td>H</td>
<td>5.1</td>
<td>(+)</td>
<td>IVa</td>
<td>Dead (7 M)</td>
<td>40%</td>
</tr>
</tbody>
</table>

remaining 4 from DA. In DA-derived PCM cases, 2 cases were located in the tail, and the mean maximum diameter was smaller than IPMN-derived PCM (4.4 cm vs 7.1 cm). All of DA-derived PCM cases had lymph node metastasis, and died after a mean postoperative period of 32.8 months. The 4 year survival rate of PCM derived from IPMN and DA were 60% and 0%, respectively.

3.2. Mucin rate and Histopathologic Findings (Figure 2)

The mucin rate was more than 40% in all PCM cases, and it was higher than 50% (mean: 59.4%) in all IPMN-derived PCM. Five IPMN-derived PCM cases showed expansive growth with dilatation of pancreatic ducts filled with mucin (Figure 2(a)). Four cases were classified as intestinal type of IPMN (Figure 2(b)) and one case was recognized as pancreatobiliary type. In 4 DA-derived PCM cases (Cases 6-9), the mucin rate was less than 50% (mean: 45.8%), excluding 1 case (56%). The lesion of the exceptional one male case was clearly accompanied by cysts filling with mucin, but ovarian-type stroma as seen in mucinous cystic neoplasm was not present (Case 6). None of DA-derived PCM lesions showed gross dilatation of the pancreatic duct (Figure 2(c)). Histologically, all of DA-derived PCM cases were observed marked mucin infiltration into interstitial tissue and adenocarcinoma cells reside in a mucin pool (Figure 2(d)).

3.3. Immunohistochemistry (Table 2 and Figure 3)

For MUC1 and MUC2, 4 of 5 (80%) IPMN (intestinal type)-derived PCM cases showed MUC1(−)/MUC2(+) except 1 IPMN (pancreatobiliary type)-derived PCM with MUC1(−)/MUC2(−). Among the 4 DA-derived PCM cases, 3 (75%) were MUC1(−)/MUC2(+), while 1 was MUC1(+)/MUC2(−). In the 50 ordinary DA cases, 37 cases (74%) showed MUC1(+) and all were MUC2(−).

Figure 2. Representative features of PCM. (a) Macroscopic surface of IPMN-derived PCM (Case 2). Tumor filled with mucin expansively grew and directly invaded into the duodenal mucosa and the common bile duct; (b) Microscopic features of IPMN-derived PCM. Tumor cells were similar to colonic villous adenomas, and were classified as IPMN, intestinal type; (c) Macroscopic surface of DA-derived PCM (Case 7). Mucinous tumor was present in the head of the pancreas. Main pancreatic duct was unclear; (d) Microscopic features of DA-derived PCM. Conspicuous mucin was noted, and adenocarcinoma cells were floating in the mucin pool.
Table 2. Immunostaining of PCM cases comparison with ordinary ductal adenocarcinoma.

<table>
<thead>
<tr>
<th></th>
<th>IPMN-PCM</th>
<th>DA-PCM</th>
<th>Ordinary DA</th>
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<tbody>
<tr>
<td>MUC1</td>
<td>(0/5) 0%</td>
<td>(1/4) 25%</td>
<td>(37/50) 100%</td>
</tr>
<tr>
<td>MUC2</td>
<td>(4/5) 80%</td>
<td>(3/4) 75%</td>
<td>(0/50) 0%</td>
</tr>
<tr>
<td>p16</td>
<td>(5/5) 100%</td>
<td>(3/4) 75%</td>
<td>(13/50) 26%</td>
</tr>
<tr>
<td>CyclinD1</td>
<td>(1/5) 20%</td>
<td>(1/4) 25%</td>
<td>(22/50) 44%</td>
</tr>
<tr>
<td>p53</td>
<td>(2/5) 40%</td>
<td>(1/4) 25%</td>
<td>(23/50) 46%</td>
</tr>
<tr>
<td>Dpc4</td>
<td>(4/5) 80%</td>
<td>(3/4) 75%</td>
<td>(20/50) 40%</td>
</tr>
</tbody>
</table>

Not only IPMN- but also DA-derived PCM showed significant positivity for MUC2 which is different from ordinary DA.

For p16 (Figure 3(a)), Cyclin D1 (Figure 3(b)), p53 (Figure 3(c)), and Dpc4 (Figure 3(d)), positivity for p16 and Dpc4 tended to be higher in PCM derived either from IPMN or DA than those seen in ordinary DA. Overexpression of Cyclin D1 and p53 did not differ between PCM, derived either from IPMN or DA, and ordinary DA.

4. Discussion

The mucin contents of pancreas carcinoma have been evaluated by diverse criteria. Suda et al. suggested the importance of macroscopical observation as abundant mucin or mucino-nodules, and of presence of floating cancer cells in mucin lakes for diagnosis of MNCC [1]. Adsay et al. described that percentage mucin is 80% or higher to for MNCC distinguished from IPMN [2]. Generally, MNCC has been defined as that mucinous part occupying more than 50% of the tumor [3,4,7,8]. However, the evaluation of the mucous area has been made subjectively. For the evaluation of PCM, in this study, imaging software was employed, and we objectively determined the percentage of the mucinous component. In our study, 5 out of 9 cases (56%) of PCM arose from IPMN, in consistent with the previous report describing that 73% or 53% of mucin rich pancreas carcinoma was derived from IPMN by Suda et al. [1], or Adsay et al. [2], respectively. Reflecting on the histological features of IPMN-derived PCM which tends to grow expansively, mucin composition was higher than DA-derived PCM. With 50% as a borderline, PCM could be classified according to its derivation based on the mucin rate into DA-derived (40% - 50%) and IPMN-derived (≥50%) PCM (Figure 4).

Marked mucin infiltration into interstitial tissue was observed all of DA-derived PCM cases. While all of 50 ordinary DA cases were MUC2(−), 3 of 4 DA-derived PCM cases were MUC2(+) in DA-derived PCM cases, the expression patterns of p16 and Dpc4 were similar to those of IPMN-derived PCM rather than ordinary DA. Thus, despite the mucin rate was less than 50%, DA-derived PCM seemed to be included in MNCC. However, the cases in DA-derived PCM exhibited a worse 4-year survival rate compared with IPMN-derived PCM. Therefore, we propose a hypothesis that DA-derived PCM constitutes a borderline group between IPMN and ordinary DA. Further studies are required to clarify the biological relevance of DA-derived PCM.

In conclusion, we quantitate the mucinous area of PCM. PCM should have more than 40% of mucous area and especially IPMN-derived PCM contain the mucous area more than 50%. DA-derived PCM was similar to IPMN-derived PCM immunophenotypically rather than ordinary DA. This study provides the prototype of further analysis of PCM by evaluating large numbers of cases in the future.

5. Acknowledgements

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