

Research Article

Correlation between Immunohistochemical and Histomorphological Features of Ampullary Carcinomas: A Study on 72 Cases from a Tertiary Health Care Center

Usha Mary Abraham¹ and Subramaniam Ramkumar¹

¹Department of Pathology, Amala Institute of Medical Sciences, Amala Nagar PO, Thrissur, 680555 Kerala, India ²Department of Pathology, Woodland Hospital, Dhankheti, East Khasi Hills, Shillong, Meghalaya 793003, India

Correspondence should be addressed to Subramaniam Ramkumar; drramkumar@woodlandpathology.com

Received 2 September 2019; Accepted 22 October 2019; Published 6 January 2020

Academic Editor: Oronzo Brunetti

Copyright © 2020 Usha Mary Abraham and Subramaniam Ramkumar. This is an open access article distributed under the Creative Commons Attribution License, which permits unrestricted use, distribution, and reproduction in any medium, provided the original work is properly cited.

Tumors involving the ampulla could be arising primarily in the ampulla or extending from the adjacent. When a neoplasm is centered primarily in the ampulla with or without periampullary mucosal involvement, it is considered a primary ampullary carcinoma. These tumors generally have a better prognosis than duodenal and pancreaticobiliary neoplasms secondarily involving the ampulla. Distinguishing between the two primary types has prognostic implications, as the pancreaticobiliary-type ampullary carcinomas are said to have a poorer prognosis than the intestinal-type. Morphological analysis alone may not suffice in this context. The role of immunohistochemistry has therefore been explored by various groups of workers. The purpose of the present study was to determine the role of morphology and cytokeratin profile in accurate typing of ampullary carcinomas as intestinal and pancreaticobiliary.

1. Introduction

Junctions of two different types of epithelia can give rise to tumors that may show features of either the involved epithelia or a mixture or an intermediate amount of both. One such junction is the ampulla of Vater [1,2]. Primary ampullary carcinomas generally have a better prognosis than duodenal and pancreaticobiliary neoplasms that secondarily involve the ampulla [3, 4]. On the basis of microscopic appearance, primary ampullary carcinomas are classified as intestinal, pancreaticobiliary, or mixed types [3, 4]. Distinguishing between the first two types is important: The pancreatobiliary-type carries a poorer prognosis than the intestinal-type [5, 6]. Morphological analysis alone may not be sufficient for diagnosis. The role of immunohistochemistry has therefore been explored by various groups of workers. The pattern of cytokeratin (CK) expression, CK7–/CK20+ in the intestinal-type and CK7+/CK20- in the pancreaticobiliary-type, correlated with morphological characteristics in some of these studies [7, 8]. However, results of other studies contradicted these observations [9, 10]. The purpose of this study was to determine the role of cell structure and CK profile in accurate typing of ampullary carcinomas as intestinal and pancreaticobiliary.

2. Materials and Method

Seventy-two specimens of ampullary carcinoma were studied in the Department of Pathology, in a tertiary health care center in India, over a period of 8 years. Of these, 25 were obtained through pancreaticoduodenectomy and 47 through endoscopic biopsy. After the relevant paraffin sections and blocks were screened, 30 cases were selected for this study, according to the following criteria:

- (1) Histopathological features were unequivocally those of a pure, invasive adenocarcinoma
- (2) The neoplasm involved primarily the ampulla of Vater, with or without periampullary extension, as

judged by gross examination of the resection specimens and by the endoscopic appearance

Specimens with an exclusively periampullary pattern of growth were not included in the study. Clinical data included the mode of presentation and endoscopic or operative findings, obtained from case files and pathology request forms. Gross, microscopic, and immunohistochemical findings were recorded in the pathological examination.

2.1. Gross Examination

2.1.1. Resection Specimens. For each specimen, we recorded the location and pattern of growth (ampullary or mixed ampullary/periampullary); size of the lesion; gross morphological type (i.e., ulcerative, polypoid, or stenosing); and involvement of the head of the pancreas, duodenal wall, common bile duct, and lymph nodes.

2.1.2. Endoscopic Biopsy Samples. Endoscopic observation yielded information regarding the pattern of growth, size, and gross morphological type. This information was collected through pathology request forms or from patient files.

2.2. Microscopic Examination. We examined the architecture of the neoplasm (e.g., tubular, papillary, or mixed) cell structure (including cell height), the presence of nuclear atypia, and the nature of the stroma. We attempted to classify each of the tumors as intestinal or pancreaticobiliary or of mixed-type.

2.3. Immunochemistry. Immunostaining for CK7 and CK20 was performed on sections from all 30 blocks by means of the following technique [11]. First, two specimens of colonic carcinoma and one of cholangiocarcinoma were chosen as controls for CK7 and CK20, respectively. From the chosen paraffin blocks, $4 \mu m$ thick sections were made and then stained with hematoxylin and eosin and visualized by microscopy. To detect CK7 and CK20 expression, we performed immunohistochemical studies with horseradish peroxidase (HRP), a supersensitive polymer, along with appropriate controls. We used 3,3'-diaminobenzidine tetrahydrochloride as the chromogen. By using a pressure cooker for 10 min and citrate buffer at a pH of 6.0, we retrieved the antigens to CK7 and CK20 (Bio-Genex, San Ramon, CA). The sections were then assessed microscopically. Scoring of nuclear immunoreactivity was performed in the well-stained area with no interference by nonspecific staining background. Immunoreactivity was graded according to the percentage of cells expressing the marker: 0 (no cells expressed the marker), 1 (<25% of neoplastic cells showed positivity), 2 (25%-50% of neoplastic cells showed positivity), and 3 (>50% of neoplastic cells showed positivity). The intensity of staining was also assessed, although in a subjective manner, and graded from 1+ to 3+ by two of the authors independently (UA and VN), and their results were compared.

3. Results

Of the specimens studied, 21 showed correlations between morphological type and CK7/CK20 staining pattern. Of these, two were of the intestinal-type of ampullary carcinoma (Figure 1(a)-1(e)), 14 were of the pancreaticobiliary-type (Figure 2(a)-2(e)), and five showed mixed intestinalpancreaticobiliary morphological type and CK profile (Tables 1 and 2; Figure 3(a)-3(d)). In the remaining nine cases, the CK profile did not match cell structure (Tables 3 and 2).

The assessment of CK7 and CK20 expression revealed that of the nine metastatic tumors, 100% were CK7+, but only 44% (four) were CK20+. Among the six tumors without metastasis in our series, five (83%) expressed CK20, and four (67%) expressed CK7. This is in agreement with the earlier observations that colorectal adenocarcinomas with lymph node metastasis showed more CK7 expression than tumors without metastasis [4, 6].

3.1. Lymph Node Metastasis and Tumor Types. A total of 15 specimens obtained from Whipple's resections were included in this study, of which nine showed metastatic deposits in the lymph nodes. Of the nine specimens, only seven had a morphological type that was correlated with CK profile, which enabled subtyping of the tumors. Of the seven carcinomas with lymph node metastasis, four were of pancreaticobiliarytype and three were of mixed-type. Pure intestinal carcinomas did not show lymph node involvement. Of the two cases with no morphological-immunohistochemical correlation, which had also metastasized to lymph nodes, one had a pancreaticobiliary CK profile and an intestinal morphological appearance, and the other had a mixed CK profile and a pancreaticobiliary morphological appearance. Regardless of morphological subtypes, we observed that of the nine tumors with lymph node metastasis, all expressed CK7, whereas only four (44%) expressed CK20. Of the six tumors without metastasis, five (83%) expressed CK20, and four (67%) expressed CK7.

In this study, we attempted to find correlations of individual histological features (such as architecture, predominant cell type, nuclear grade, and degree of desmoplasia) with CK7 and CK20 expressions. We found that tumors with an exclusively tubular architecture always expressed CK7, but not CK20.

With regard to cell height, CK20 expression was correlated strongly with tall columnar cells; all CK20+ tumors had a tall columnar cell component.

3.2. Cell Height and Cytokeratin Expression. Twelve of the neoplasms assessed consisted of cuboidal or low columnar cells or both types. Eleven of these had a CK7+/CK20- profile, and the other had a CK7-/CK20- profile. Seven of these had a CK7+/CK20- profile, eight had a CK20+/CK7 + profile, and three had a CK7-/CK20+ profile. Although CK7 expression showed no significant correlation with cell height, all 11 CK20+ tumors had a considerable proportion of tall columnar cells.



FIGURE 1: (a) Pancreatobiliary-type carcinoma with a tubular pattern (hematoxylin-eosin, 40x). (b) Pancreatobiliary-type carcinoma with marked nuclear atypia (hematoxylin-eosin, 400x). (c) Pancreatobiliary-type carcinoma with desmoplasia (hematoxylin-eosin, 100x). (d) Pancreatobiliary-type carcinoma with CK7 positivity (immunohistochemistry, 400x). (e) Pancreatobiliary-type carcinoma with CK20 negativity (immunohistochemistry, 100x).



FIGURE 2: (a) Intestinal-type carcinoma with a tubulopapillary pattern (hematoxylin-eosin, 100x). (b) Intestinal-type carcinoma with minimal nuclear atypia (hematoxylin-eosin, 400x). (c) Intestinal-type carcinoma with the absence of desmoplasia (hematoxylin-eosin, 400x). (d) Intestinal-type carcinoma with CK20 positivity (immunohistochemistry, 100x). (e) Intestinal-type carcinoma with CK7 negativity (immunohistochemistry, 100x).

The distribution of the different nuclear grades did not differ between CK7+ and CK20+ tumors. However, 100% of high nuclear grade (+++) neoplasms expressed CK7, and 20% expressed CK20. This suggests a correlation between CK7 expression and high grade; however, the majority of tumors in this study were either pancreatobiliary or of mixed-type and were CK7+. Observations about desmoplasia also followed a similar pattern. The results should be viewed with regard to the fact that majority of the tumors were CK7+.

3.3. Nuclear Grade and Cytokeratin Profile. All of the ten tumors with +++ nuclear grade showed CK7 expression. Of the 11 tumors with CK20 expression, nine had a low to medium nuclear grade (Tables 4 and 2). Among the 12 tumors with grade 2 to 3+ desmoplasia, CK7 expression

Morphological type	Cytokeratin profile	No. of cases
Intestinal	CK7-/CK20+	2
Pancreaticobiliary	CK7+/CK20-	14
Mixed	CK7+/CK20+	5
Total		21

 TABLE 1: Morphological and immunohistochemical correlations in the subtypes of ampullary carcinoma.

was present in 11 (92%), and CK20 was present in 5 (42%). In tumors with 0-1+ desmoplasia, 83% expressed CK7 (15/18) and 33%, CK20 (6/18; Tables 5 and 2).

Our findings are important because accurate histological typing of ampullary carcinomas would provide information that guides staging and planning of operative procedures. It would also provide a solid foundation for analyzing pathogenetic mechanisms of these tumors by more advanced methods, such as molecular studies.

4. Discussion

Ampullary carcinomas constitute 5% of gastrointestinal carcinomas and usually occur in people older than 60 years. The clinical symptoms and signs include jaundice, itching, loss of appetite, loss of weight, pale colored stools, and other signs of biliary obstruction [12]. Ampullary carcinomas, although usually small at the time of diagnosis, present with early obstructive symptoms thereby resulting in early detection.

Most of these tumors are present as a small mass projecting into the duodenal lumen or as periductal thickening or bulging of the papilla. Intra-ampullary tumors extending into the duodenal mucosa can present as an exophytic or ulcerative growth [13, 14]. The spread of these tumors may be invasive or noninvasive. Noninvasive spread includes intramucosal spread to the duodenum as well as the proximal areas of the common bile and pancreatic ducts. The invasive tumors spread via the musculature into the adjacent duodenal and/or pancreas. The lymph nodes involved are the peripancreatic group. Distant metastasis is usually to the liver, also to the peritoneum, lungs, and pleura. Vascular and perineural invasions are also known to occur in these tumors [15, 16].

Pathologic staging is difficult due to the complexity of the anatomy of the ampulla of Vater. Ampullary carcinomas have been staged according to the AJCC Cancer Staging Manual 6th edition [13]. Based on the primary tumor (T), there are seven stages. Pancreaticoduodenectomy is the treatment of choice and the mainstay of therapy [3, 4, 17].

They can arise in the ampullary duodenal aspect, lined by intestinal mucosa, or in the deeper aspect, lined by pancreatic or bile duct mucosa. Morphologically, the two types resemble their colonic and pancreatic counterparts [11]. The intestinal variant consists of elongated tubular units lined by cells ranging in height from tall columnar to cuboidal, with basal nuclei and apical mucin. The immunohistochemical marker profile of these tumors is similar to that of colonic carcinomas (CK20+, CK7-, and mucin 2 positivity (MUC2+)) [13]. The pancreatobiliary variant has simple or branching glands lined by epithelium ranging in height from cuboidal to low columnar. Its immune profile resembles that of pancreatic carcinomas (CK7+, CK20-, and MUC2-) [13]. The intestinal variant has a better prognosis than pancreatobiliary subtypes [18].

However, the ampulla has complex histomorphological features, and the transition from pancreatobiliary ductal epithelium to intestinal epithelium is very subtle. Morphological overlaps are common and pose challenges in tumor classification. Further staging (T1/T2 or T3) is difficult due to mucosal invasion by these tumors and the occurrence of rare pancreatic acinar in the walls of the ampulla. Hence, a multifaceted diagnostic correlational approach among tumor morphological type, immunohistochemistry (CK and mucin immunoprofiling), and imaging studies can help circumvent the pitfalls in diagnosing ampullary carcinomas. Several studies with this multifaceted approach have been conducted; their focus was on the histomorphological features in varying combinations.

Zhou et al. were the first to show agreement between the histological classification and immunohistochemical characterization based on CKs. They found a high specificity of CK7 for pancreatobiliary carcinomas and of CK20 for intestinal carcinomas [7]. Similar findings were confirmed by the studies of Le Pessot et al. [8] and Roh et al. [15]. Furthermore, the findings of Westgaard et al., who studied 114 resection specimens, indicated similar moderate agreement between morphological type and CK immunoprofiling. They found that histological typing significantly discriminates between the pancreatobiliary and intestinal types of pancreatic head adenocarcinomas, which is important for determining prognosis [18]. CK7 and CK20 expressions could also have some bearing on the prognosis of carcinomas, regardless of subtypes. At least in one series on colorectal carcinomas, CK20 was expressed more in low-grade tumors and CK7 was expressed more in high-grade tumors with lymph node metastasis [17].

This study was an attempt to find correlations between the morphological type and CK profile of ampullary adenocarcinomas. Both CK7 and CK20 are simple CKs with restricted tissue and neoplastic distribution. In this study, 70% of ampullary carcinomas showed ampullary adenocarcinomas are correlated to an extent with CK profile, and morphological-immunohistochemical subtyping is possible in 70% of cases. Lymph node metastasis occurs more frequently with CK7+ tumors than with CK20+ tumors. However, the above fact is hypothesized due to the limitation of the number of cases evaluated. This can be further evaluated by performing relevant studies. Among the microscopic features, an exclusively tubular architecture is associated with CK7 expression. Tall columnar cell component is unequivocally related to CK20 expression.

The correlations between some of the individual microscopic features, such as architecture and cell height with CK7 and CK20 expressions, are greater than those between intestinal and pancreatobiliary morphological types and combined CK7-CK20 profile.

TABLE 2

rrelation	resent	resent	×	resent	resent	resent	resent	resent	resent	resent	resent	resent
co. Co:	uliary P	iliary P	iliary	uliary P	al I	iliary P	uliary P	al	al/ iliary p	al/ iliary P	iliary P	iliary/ _P al
Inferen in IHC	Pancreato	Pancreatot	Pancreatot	Pancreatot	Intestir	Pancreatot	Pancreatot	Intestir	Intestin pancreatob	Intestin pancreatob	Pancreatot	Pancreatob intestin
Intensity	I	I.	I	I.	I	I	I.	ŧ	‡	‡	I	+
och emistry Extent	I	I	I		m	I	I.	5	6	ŝ	I	г
Intensity	‡	ŧ	ŧ	ŧ	I	ŧ	ŧ	I	‡	ŧ	ŧ	ŧ
Extent	ņ	e,	ñ	ŝ	I	ñ	ŝ	I	m	ŝ	ŝ	5
Impression	Pancreatobiliary	Pancreatobiliary	Intestinal	Pancreatobiliary	Intestinal	Pancreatobiliary	Pancreatobiliary	Intestinal	Intestinal/ pancreatobiliary	Intestinal/ pancreatobiliary	Pancreatobiliary	Pancreatobiliary/ intestinal
Stromal desmoplasia	1	I		+	+	ŧ	+		ŧ	ŧ	+	+
findings Nuclear atypia	I.	+	+ to +++	ı.	+	ŧ	+ to++	÷	ŧ	+ to ++	ŧ	ŧ
Microscopic Cell height	Low columnar	Tall columnar	Tall columnar	Cuboidal	Tall columnar	Low columnar	Low columnar to tall columnar	Tall columnar	Low columnar to tall columnar	Cuboidal to tall columnar	Low columnar	Tall columnar
Architecture	Tubule, trabeculae	Tubule, papillae, solid pattern	T ubules, papillae	Tubules	T ubules, papillae	T ubules, cords, and groups	Tubules, papillae, focal cribriform	Tubules, papillae	Tubules, papillae	Tubules, cribriform and solid pattern	Tubules	Tubules, papillae
spread Lymph node involvement	1/6		I			1/4		i	2/8			3/6
Extent of f Local extent of spread	Infiltration into pancreas seen	Infiltration into wall of duodenum and bile duct	Infiltration into wall of duodenum and bile duct		Infiltration into duodenal wall and focally in to pancreas	Infiltration into pancreas and duodenal wall		i	I	Infiltration in to duodenal wall		Infiltration in to duodenal wall and
Morphology	Nodular mass	Nodular mass	Nodular mass		Ulceroproliferative growth	Friable nodular mass		Nodular mass	Polypoid pedunculated mass	III circumscribed firm bulky mass		Ulcer with raised margins
findings Size	1×1 cm	0.8 × 0.5 cm	$3 \times 1 \mathrm{cm}$		4×3 cm	2 × 3 cm		1 B	5 × 3 cm	1.9 × 1.5 cm		1.5 × 1 cm
Gross Pattern of growth	Intra- ampullary	Intra- ampullary	Periampullary	Intra- ampullary	Periampullary	Intra- ampullary	Periampullary	Ampullary	Periampullary	Ampullary		Periampullary
Type of specimen	Whipple's specimen	Whipple's specimen	Whipple's specimen	Endoscopic biopsy	Whipple's specimen	Whipple's specimen	Endoscopic biopsy	Endoscopic biopsy	Whipple's specimen	Whipple's specimen	Endoscopic biopsy	Whipple's specimen
Radiological findings	CT shows a small enhancing lesion in D2. Endoscopy shows bulky ampulla with small ulcer	Endoscopy shows fleshy ampulla	Endoscopy shows nodular mass	Endoscopy shows prominent ampulla	CT shows periampullary mass with dilated CBD	Endoscopy showed a friable prominent ampulla	Endoscopy showed a periampullary growth	Endoscopy large ampullary growth	Endoscopy polypoid pedunculated mass	USG-mass in the head of the pancreas, endoscopy- prominent ampulla	Endoscopy- friable ulcerated ampulla	Endoscopy- duodenal ulcer with a mass
Clinical details	Obstructive jaundice for 6 months	Obstructive jaundice for 3 months	Jaundice - 3 weeks	Obstructive jaundice	Jaundice for 3 weeks	Jaundice 1 month	Jaundice 2 weeks. Known case of tuberculosis	Recurrent pancreatitis, abdominal pain 6 months	Jaundice 2 weeks	Obstructive jaundice, abdominal pain 1 month	Loss of weight, jaundice 1 month	Jaundice, abdominal pain 1 month
Age/sex	48/F	58/F	44/M	55/F	59/F	40/M	72/male	57/male	38/male	78/male	68/male	57/male
Biopsy number	2414/10	2041/10	2208/09	1134/10	1937/10	1883/07	8707/10	8764/10	S810/10	S1184/10	S1444/10	S188/09
Serial no.	-	7	m	4	ŝ	٥	14	∞	6	10	=	12

Gastroenterology Research and Practice

Correlation	Present	Present	×	Present	Present	Present	Present	×	Present	×	×
Inference in IHC	Pancreatobiliary/ intestinal	Pancreatobiliary	Intestinal	Pancreatobiliary	Pancreatobiliary/ intestinal	Pancreatobiliary	Pancreatobiliary	Pancreatobiliary/ intestinal	Pancreatobiliary	Intestinal	Pancreatobiliary
Intensity	+	I	I	I	+	I	I	+	I.	I	I.
och emistry Extent	-	I	I	I	-	I	I	-	I.	I	I.
Intensity	+	+	‡	I	I	I	I	I	I		+
Extent	m	-	m	-	г	7	°.	m	7	I	-
Impression	Pancreatobiliary/ intestinal	Pancreatobiliary	Intestinal	Pancreatobiliary	Intestinal	Pancreatobiliary	Pancreatobiliary	Pancreatobiliary	Pancreatobiliary	Pancreatobiliary	Intestinal
Stromal desmoplasia	ŧ	‡	‡	‡	T	+	+	+	+	+	I
findings Nuclear atypia	+	ŧ	+	‡	‡	+	‡ ‡	* *	ŧ	ŧ	+
Microscopic Cell height	Low columnar to tall columnar	Cuboidal to low columnar	Tall columnar	Low columnar	Tall columnar	Low columnar	Low columnar	Tall columnar	Low columnar	Cuboidal to low columnar	Tall columnar
Architecture	Tubules, sheets, and cords	Tubules	Tubules, papillae	Tubules, cribriform	Tubules, cribriform	Tubules	Tubules, cribriform	Papillary, cribriform	Tubules, papillae	Tubules	Tubules, papillae
pread Lymph node nvolvement	3/5	1/5	1/1	I				5/10			
Extent of s Local extent of spread ii	pancreatic parenchyma Infiltration in to duodenal wall	Infiltration in to wall of duodenum, CBD, pancreas	I	I				Involve adjacent duodenum pancreatic parenchyma and CBD			
Morphology	Ulcer in duođenal lumen	Edema of ampullary mucosa, stricture with occlusion of CBD lumen	Nodular projection at ampulla	A mpullary prominence			,	Ulceroproliferative growth	Ampullary prominence		
findings Size	4 × 2 cm	1.5 cm	1.5 cm	I				I			
Gross 1 Pattern of growth	Perianpullary	Ampullary	Ampullary	Intra- ampullary	Periampullary	Periampullary	Periampullary	Periampullary and ampullary	Intra- ampullary	Periampullary	Periampullary
Type of specimen	Whipple's specimen	Whipple's specimen	Whipple's specimen	Endoscopic biopsy	Endoscopic biopsy	Endoscopic biopsy	Endoscopic biopsy	Whipple's specimen	Endoscopic biopsy	Endoscopic biopsy	Endoscopic biopsy
Radiological findings	in the ampullary region Endoscopy- ulcer in ampullary region	CBD showed an area of hard stricture with occlusion of lumen	Endoscopy- ampullary prominence	CT-dilated CBD and pancreatic duct	CT-1HBR dilatation, mass lesion in periampullary region of CBD	USG-dilated CHD, CBD up to terminal end	ERCP- periampullary growth	Endoscopy- growth in D2, dilated CBD	Endoscopy ampullary growth	USG-mass in pancreas. Endoscopy ampullary prominence	CT-dilated main pancreatic duct and CBD up to
Clinical details	Jaundice, loss of weight 1 month	Abdominal pain and jaundice for 2 weeks	Jaundice, fever, vomiting for 3 months	Obstructive jaundice 4 months	Jaundice, vomiting 15 days	Abdominal pain, loss of weight, jaundice 2 months	Obstructive jaundice 2 months	Itching, loss of appetite 2 months	Obstructive jaundice 4 months	Itching, abdominal pain, and jaundice for 1 month	Obstructive jaundice
Age/sex	52/male	55/male	50/male	65/male	53/male	47/male	57/male	66/male	60/male	60/male	52/male
Biopsy number	\$1264/07	S2691/08	S3074/09	S4966/09	S2724/09	S853/09	S1281/08	S4369	S2193/16	1640/09	558/07
Serial no.	13	14	15	16	17	18	19	50	21	22	23

TABLE 2: Continued.

6

Gastroenterology Research and Practice

TABLE 2: Continued.

Serial	Biopsy	Age/sex	Clinical details	Radiological findings	Type of	Gross Pattern of growth	s findings Size	Morphology	Extent of Local extent of spread	spread Lymph node	Architecture	Microscopic Cell height	findings Nuclear atvnia	Stromal	Impression	I Extent	mmunohistc Intensity	ochemistry Extent	Intensity	Inference in	Correlation
01	TOTINI .		crimina .	eSumm	mannade	Brown			n oprau	involvement		mgm	arypia	mendomeon						IHC	
24	4267/09	70/male	Obstructive jaundice	ampulla. No mass lesion Endoscopy- friable growth ampullary region	Endoscopic biopsy	Ampullary					Tubules, papillae	Low columnar to tall columnar	ŧ	ŧ	Pancreatobiliary	ň	‡	I.	I	Pancreatobiliary	Present
25	4084	51/male	Recurrent pancreatitis 6 months	Endoscopy ulceration in the periampullary region	Whipple's specimen	Periampullary	1 × 2 cm	Ulcerated lesion with raised borders			Tubules, cribriform, papillae	Tall columnar to cuboidal	+	+	Intestinal	-	+	5	‡	Pancreatobiliary/ intestinal	Present
26	4675/10	56/male	Pancreatitis, cholecystitis 1 month	Dilated IHBR	Whipple's specimen	Ampullary	1.8×0.8 cm	Ampullary prominence	,	2/8	Tubules	Cuboidal to low columnar	ŧ	÷	Pancreatobiliary	n	ŧ	I	I	Pancreatobiliary	Present
27	3826/10	65/male	Obstructive jaundice 1 month	Moderately dilated intra- and extrahepatic biliary channels and pancreatic duct	Endoscopic biopsy	Perianpullary		,		,	Tubules, cribriform	Low columnar to cuboidal (mucinous)	ŧ	+ +	Pancreatobiliary	m	‡	i.	1	Pancreatobiliary	Present
28	3995/10	55/male	Obstructive jaundice	ERCP- periampullary growth	Endoscopic biopsy	Periampullary		Periampullary nodular growth	·		Tubules, papillae	Cuboidal to tall columnar	+	I	Pancreatobiliary/ intestinal	ŝ	ŧ	I	I	Pancreatobiliary	x
59	680/11	55/male	Obstructive jaundice	Endoscopy- periampullary growth	Whipple's specimen	Periampullary	4 × 2.5 cm	Ulceroproliferative growth			T ubules, papillae, cribriform	Tall columnar to cuboidal	‡	+ + +	Pancreatobiliary/ intestinal	3 in papillary pattem and negative in tubules	‡ ‡	2 in papillary to 3 in tubules	++ to +++	Pan creatobiliary/ intestinal	Present
30	2845/11	50/male	Obstructive jaundice	Endoscopy- periampullary ulceroproliferative growth	Endoscopic biopsy	Periampullary	1	Ulceroproliferative growth			Papillae, tubules	Tall columnar	‡	‡	Pancreatobiliary/ intestinal	I	I	ŝ	+	Intestinal	×

Gastroenterology Research and Practice



FIGURE 3: (a) Tall columnar cells in mixed-type carcinoma (hematoxylin-eosin, 400x). (b) Cuboidal cells in mixed-type carcinoma (hematoxylin-eosin, 400x). (c) CK7 positivity in mixed-type carcinoma (immunohistochemistry, 100x). (d) Ck20 positivity in mixed-type carcinoma (immunohistochemistry, 400x).

TABLE 3: Specimens	lacking corre	lation	between	morph	ological	type
and immunohistoch	emical findii	ngs.				

Morphological type	Cytokeratin profile	No. of cases
Intestinal	CK7+/CK20– (pancreaticobiliary)	3
Intestinal	CK7+/CK20+ (mixed)	2
Pancreaticobiliary	CK7+/CK20+ (mixed)	1
Pancreaticobiliary	CK7+/CK20- (intestinal)	1
Pancreaticobiliary	CK7-/CK20- (undefined)	1
Mixed	CK7+/CK20– (pancreaticobiliary)	1
Total		9

TABLE 4: Number of cases by nuclear grade and cytokeratin profile.

Cytokoratin profile		Nuclear grade	2
	+	++	+++
CK7+/CK20-	4	6	8
CK7-/CK20-	0	1	0
CK7+/CK20+	1	5	2
CK7-/CK20+	2	1	0
Total		20	10

TABLE 5: Number	of cases	by o	desmoplasia	grade	and	cytokeratin
profile.						

Cutakaratin profile		Des	smoplasia g	grade	
Cytokeratin prome	+	++	+++	•	Total
CK7+/CK20-	7	5	2	4	18
CK7-/CK20-	1	0	0	0	1
CK7+/CK20+	3	3	1	1	8
CK7-/CK20+	1	1	0	1	3
Total	12	9	3	6	30

Data Availability

The data used to support the findings of this study were from the laboratory archives of the Department of Pathology, PSG Institute of Medical Sciences and Research, Coimbatore, Tamil Nadu, India. The data is available from the corresponding author upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

References

 H. F. Frierson Jr., "The gross anatomy and histology of the gallbladder, extrahepatic bile ducts, vaterian system, and minor papilla," *The American Journal of Surgical Pathology*, vol. 13, no. 2, pp. 146–162, 1989.

- [2] J. Rosai, "Pancreas and ampullary region," in *Rosai and Acker-man's Surgical Pathology*, pp. 1092–1095, Elsevier, St. Louis, Missouri, 9th edition, 2004.
- [3] M. Cecilia, F. Preiser, A. E. Noffsinger, G. E. Stemmerman, P. E. Lantz, and P. G. Isaacson, "Epithelial tumors of the small intestine," in *GI Pathology An Atlas and Text*, pp. 482–485, Wolters Kluwer, Lippincott Williams & Wilkins, Philadelphia, Baltimore, New York, London, Beunos Aires, Hong Kong, Sydney, Tokyo, 3rd edition, 2008.
- [4] R. K. Yantiss and D. A. Antonioli, "Polyps of the small intestine," in *Surgical Pathology of the GI tract, Liver, Biliary tract, and Pancreas*, R. R. Goldblum and R. R. Odze, Eds., pp. 447– 480, Elsevier Inc, Philadelphia, PA, 2nd edition, 2009.
- [5] J. R. Howe, D. S. Klimstra, R. D. Moccia, K. C. Conlon, and M. F. Brennan, "Factors predictive of survival in ampullary carcinoma," *Annals of Surgery*, vol. 228, no. 1, pp. 87–94, 1998.
- [6] W. Kimura, N. Futakawa, S. Yamagata et al., "Different clinicopathologic findings in two histologic types of carcinoma of papilla of Vater," *Japanese Journal of Cancer Research*, vol. 85, no. 2, pp. 161–166, 1994.
- [7] H. Zhou, N. Schaefer, M. Wolff, and H. P. Fischer, "Carcinoma of the ampulla of Vater," *The American Journal of Surgical Pathology*, vol. 28, no. 7, pp. 875–882, 2004.
- [8] F. Le Pessot, M.-L. Ranty, M.-F. Hellot et al., "Cytokeratins 7 and 20 immunohistochemistry in ampullary carcinomas," *Annales de Pathologie*, vol. 24, no. 4, pp. 312–318, 2004.
- [9] N. S. Goldstein and D. Bassi, "Cytokeratins 7, 17, and 20 reactivity in pancreatic and ampulla of vater adenocarcinomas. Percentage of positivity and distribution is affected by the cut-point threshold," *American Journal of Clinical Pathology*, vol. 115, no. 5, pp. 695–702, 2001.
- [10] J. V. Duval, L. Savas, and B. F. Banner, "Expression of cytokeratins 7 and 20 in carcinomas of the extrahepatic biliary tract, pancreas, and gallbladder," *Archives of Pathology & Laboratory Medicine*, vol. 124, no. 8, pp. 1196–1200, 2000.
- [11] F. Gheza, E. Cervi, G. Pulcini et al., "Signet ring cell carcinoma of the ampulla of Vater: demonstration of a pancreatobiliary origin," *Pancreas*, vol. 40, no. 5, pp. 791–793, 2011.
- [12] P. Schirmacher and M. W. Büchler, "Ampullary adenocarcinoma – differentiation matters," *BMC Cancer*, vol. 8, no. 1, 2008.
- [13] H. P. Fischer and H. Zhou, "Pathogenesis of carcinoma of the papilla of Vater," *Journal of Hepato-Biliary-Pancreatic Surgery*, vol. 11, no. 5, pp. 301–309, 2004.
- [14] N. V. Adsay, "Gall bladder, extrahepatic biliary tree, and ampulla," in *Sternberg's Diagnostic Surgical Pathology*, S. E. Mills, D. Carter, J. K. Greenson, V. E. Reuter, and M. H. Stoler, Eds., pp. 1636–1643, Lippincott Williams & Wilkins, Philadelphia, 5th edition, 2010.
- [15] Y. H. Roh, Y. H. Kim, H. W. Lee et al., "The clinicopathologic and immunohistochemical characteristics of ampulla of Vater carcinoma: the intestinal type is associated with a better prognosis," *Hepato-Gastroenterology*, vol. 54, no. 78, pp. 1641–1644, 2007.
- [16] J. T. Painter, N. P. Clayton, and R. A. Herbert, "Useful immunohistochemical markers of tumor differentiation," *Toxicologic Pathology*, vol. 38, no. 1, pp. 131–141, 2010.

- [17] R. Bayrak, S. Yenidünya, and H. Haltas, "Cytokeratin 7 and cytokeratin 20 expression in colorectal adenocarcinomas," *Pathology, Research and Practice*, vol. 207, no. 3, pp. 156– 160, 2011.
- [18] A. Westgaard, S. Tafjord, I. N. Farstad et al., "Pancreatobiliary versus intestinal histologic type of differentiation is an independent prognostic factor in resected periampullary adenocarcinoma," *BMC Cancer*, vol. 8, no. 1, 2008.



The Scientific World Journal

Journal of Immunology Research



Research and Practice











BioMed Research International



PPAR Research

Journal of Ophthalmology



Computational and Mathematical Methods in Medicine



International



Behavioural Neurology



Evidence-Based Complementary and Alternative Medicine







Research and Treatment





Oxidative Medicine and Cellular Longevity



Submit your manuscripts at www.hindawi.com