

Advanced gastric glandular-endocrine cell carcinoma with 1-year survival after gastrectomy

HIROYUKI MATSUBAYASHI¹, SHINICHI TAKAGAKI¹, TAKAO OTSUBO¹, TAKAO IIRI¹, YUKA KOBAYASHI¹, TAKASHI YOKOTA¹, KIMITOSHI SHICHIJO¹, TETSUYA TADA², KEIICHI SATOH³, and MITSUYA IWAFUCHI⁴

¹Department of Gastroenterology, Tachikawa General Hospital, Nagaoka, 3-2-11, Kandachou, Nagaoka, Niigata 940-8621, Japan

²Department of Surgery, Tachikawa General Hospital, Nagaoka, Niigata, Japan

³Department of Pathology, Tachikawa General Hospital, Nagaoka, Niigata, Japan

⁴Department of Pathology, Medical Junior College of Niigata University, Niigata, Japan

Abstract

Primary gastric endocrine cell carcinoma (ECC) is extremely rare. In general, when it is advanced, gastric ECC causes extensive ulceration (type 2) and invades or metastasizes to other organs, frequently to the liver and sometimes to the lungs or bones, and carries a poor prognosis. We herein report a 67-year-old man with advanced gastric ECC of extensivepolypoid shape (type 1) but without distant metastasis, who underwent total gastrectomy and treatment with oral tegafururacil (UFT), and showed no sign of recurrence 1 year later.

Key words Endocrine cell carcinoma · Stomach · Polypoid type · Type 1

Introduction

Endocrine cell carcinoma (ECC) is rare in the gastrointestinal tract, accounting for about 1% of cancers in the esophagus, 0.2% in the colon, and 0.1%–0.4% in the stomach [1–3]. Gastric ECC shows various macroscopic types in the early stage [4], but in patients with advanced disease, the expansive-ulcerative type is seen in more than half of the patients and the polypoid type is a minority [5,6]. The biological behavior of gastric ECC is aggressive, as shown by frequent metastasis to liver and lymph nodes and the poor effectiveness of chemotherapy [4,5–7] (Table 1). Here, we report a patient with advanced gastric ECC of polypoid shape and large size, without any recurrence 1 year after gastrectomy.

Case report

A 67-year-old man, who had congenital hearing loss, diabetes mellitus, hypertension, and benign prostatic

hypertrophy, was referred to our hospital because of a polypoid gastric lesion first detected at another hospital after a complaint of epigastric discomfort. On admission, there was no enlargement of any superficial lymph node. Peripheral blood showed moderate anemia (RBC, 343×10^4 /mm³; hemoglobin [Hb], 8.9 g/dl; hematocrit [Ht], 27.3%), and serum showed hypoalbuminemia (albumin, 3.4g/dl), minimally high C-reactive protein (CRP; 1.4 mg/dl; normal range, <0.6 mg/dl), normal levels of tumor markers (alpha-fetoprotein [AFP], 1.8 ng/ml, carcinoembryonic antigen [CEA], 1.2 ng/ml, and carbohydrate antigen [CA]19-9, <10 U/ ml) and various levels of gastrointestinal hormones gastrin, 756 pg/ml; (normal range, 42-200 pg/ml); somatostatin, 23.0 pg/ml (normal range, 1-12 pg/ml); serotonin, 12.2µg/dl (normal range, 10-30µg/dl); glucagon, 43 pg/ml (normal range, 70–160 pg/ml); vasoactive intestinal peptide (VIP), <5 pg/ml (normal range, $\leq 100 \text{ pg/}$ ml); insulin, 6μ U/ml (normal range, $3-18\mu$ U/ml).

Gastric fluoroscopy (Fig. 1) showed a large polypoid lesion consisting of multiple irregularly shaped nodules extending from the cardia to the body. Gastric endoscopy (Fig. 2) showed a Type 1 (polypoid) lesion with an abrupt tumor margin and rough surface covered with whitish-yellowish exudate. The lesion was proven to be ECC by histological examination of a biopsy sample. Abdominal computed tomography (CT) (Fig. 3) showed an enhanced polypoid lesion and an enlarged lymph node adjacent to the posterior wall of the gastric body, but no liver metastasis.

Total gastrectomy (Fig. 4) and splenectomy was performed, without any complication. The surgical stage was IIIA (T3N1H0P0M0CY0) [48]. The histological diagnosis was gastric glandular-endocrine cell carcinoma, medullary-type, INF β , ss, ly1, v2, n1 (#3), pm(-), dm(-), 10 × 8.5 cm, type 1, Post, UM [48]. A histological adenocarcinoma component was recognized in the continuity of the ECC component (Fig. 5a–c). Immunohistological staining of the ECC was strongly and

Offprint requests to: H. Matsubayashi

Received: August 11, 2000 / Accepted: November 22, 2000

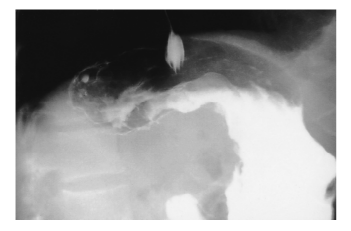


Fig. 1. Gastric fluoroscopy showed a large polypoid lesion, consisting of multiple irregularly shaped nodules, in the gastric cardia and body

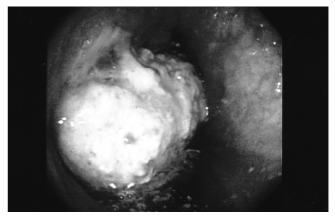


Fig. 2. Gastric endoscopy showed a polypoid lesion with an abrupt tumor margin and rough surface covered by a whitish-yellowish exudate



Fig. 3. Computed tomography demonstrated an enhanced polypoid lesion and an enlarged lymph node adjacent to the posterior wall of the gastric body, without liver metastasis



Fig. 4. Macroscopic view of resected stomach with a 10-cm rough polypoid lesion in the posterior wall of the gastric body and cardia

diffusely positive for Grimelius, chromogranin, neuronspecific enolase (NSE) (Fig. 5d), but negative for serotonin, gastrin, somatostatin, glucagon, insulin, pancreatic polypeptide, peptide YY, adrenocorticotropic hormone (ACTH), Fontana-Masson, MUC2, MUC5AC, and CD10. All of these immunostainings, except for Grimelius (focally positive) and CD10, were negative in the adenocarcinoma component. No hybrid cells, i.e., those with both adeno- and endocrine-cell characteristics, were recognized in the double-stained sections in which MUC2, MUC5AC, and CD10 were used for detecting adeno-characteristics and chromogranin and NSE were used for detecting endocrine characteristics. Immunostaining of p53 and Ki-67 [49] was strongly and diffusely positive in both the ECC and adenocarcinoma components. The yellowish-whitish

exudate that covered the tumor surface consisted of necrotic cancer cells, inflammatory cells, and fibrin.

The patient did well in his postoperative course and was soon discharged, taking oral tegafur-uracil (UFT), 300 mg/day. One year after the operation, he was examined by abdominal CT scan, which showed no recurrence or liver metastasis.

Discussion

Early in the twentieth century, endocrine cell tumors were categorized as "carcinoid" because of their histological resemblance to carcinoma [50]. Later, carcinoids were histologically divided into four types (A to D) by Soga et al. [51]. In Japan, the category of gastric endo-

 Table 1. Clinicopathological features of gastric endocrine cell carcinoma in the Japanese and English-language literature (1961–2001)

Case no.	Author	Year reported	Age/Sex	Location ^a	Size (cm)	Gross type ^b	Depth of invasion
Iananese	literature						
1	Hayashi [8]	1980	69/M	U (Post)	6×6	Type 3	se
2 3	Kazato [9] Ushiyama [10]	1983 1984	77/M 58/M	M (Ant) U	$\begin{array}{c} 1 \times 1 \\ 8 \times 7 \times 2.5 \end{array}$	III + IIc Type 1	sm se
4 5	Emi [11] Terasawa [12]	1984 1985	60/M 69/M	M (Less) EGJ	$\begin{array}{c} 3.1\times 2\\ 3.5\times 3\end{array}$	Type 3 Type 2	se mp
6 7	Kaketani [13]	1987 ″	58/M 69/F	L (Gre) ML (Less)	2.5×2.5 $15 \times 10 \times 11.5$	IIa + IIc Type 2	sm se
8 9 10 11 12 13 14	Ishihara [14] Iwafuchi [15] Takeshita [16] " Matsumoto [17] Maeta [18]	1988 1989 1990 " " 1990 1990	75/M 73/M 51/M 57/M 76/M 55/M 67/F	RS RS (Ant) M (Less) M (Ant) M (Less) ML (Less) L (Less)	$10 \times 7 \times 2.5$ 9.5 × 4.5 $6.5 \times 6.0 \times 1.4$ 6×5	Type 3 IIb IIC Type 2 Type 3 Type 2 Type 2	si se sm si si ss
15 16 17 18	Kuhara [19] Uesugi [20] Gotoh [21] Katsuyama [22]	1991 1991 1992 1992	69/M 64/M 48/M 73/M	ML (Less) L (Less) M (Gre) L (Gre)	13×12 2.6 × 2.5 6 × 5	Type 1 + 2 Type 2 Type 3	ss ss si si
19 20 21 22	Kamio [23] " Nakamoto [24] Masuyama [25]	1992 " 1992 1994	56/M 80/M 79/M 69/M	M (Less) RS (Post) L (Post) U (Less)	$7 \times 6.5 7 \times 6.5$	Type 5° Type 1 Type 2 Type 3	SS SS
23	Nagafuchi [26]	1994	57/M	L (Gre)	4.5 imes 3.5 imes 1.5	Type 2	si
24 25 26 27 28 29	Imamura [27] Maekawa [28] Kodama [29] Fudaba [30] Inoue [31] Sakai [32]	1995 1995 1995 1996 1996 1997	73/M 77/M 68/M 55/M 64/M 31/M	M (Post) U (Gre) L (Gre) L (Less) L (Ant) ML	4×2.5 2 $12 \times 8, 5 \times 3$	Type 1 Type 4 Type 2 Type 3 Type 2 Type 5 ^c	mp ss mp
30 31 32 33 34 35 36 37	Hiramoto [33] " " " " Iwasaki [34] " Yamamoto [35]	1998 " " " 1998 " 1999	58/M 75/M 63/M 62/M 64/F 58/M 69/M 56/F	L (Gre) M (Ant) M (Post) RS (Ant) M (Gre) M (Gre) LM (Less)	4 imes 4	(multiple) Type 3 Type 1 Type 3 Type 1 Type 3 Type 5 Type 2	
English-l 38 39 40 41 42	anguage literature Christodoulopoulos [36] " Vasudeo [37] Brodman [38] Parks [39]] 1961 " 1965 1968 1970	36/F 75/F 45/F 66/F 56/M	U (Post) L (Less) L U ML	$\begin{array}{c} 3.5 \times 2.5 \times 2 \\ 2.3 \times 1.8 \\ 6.5 \times 5 \\ 10 \\ 16 \times 8 \end{array}$	Type 2	mp si

Metastasis			Chemotherapy				
Liver	N	d Other	Operation ^e	and other treatment	Prognosis ^o		
_	+		STD + DP	_	9d, death of		
			012 01		other disease		
_	+	_	SG	_	NM		
+	+	Skin, bone, etc.	_	Radiation + ADM, MMC, MTX	2m, death		
_	+		SG		6m, death		
+	+		PG	TU ^f (orally), cyclophosphamide	4m, death		
—	+	Abdominal wall	SG	- MMC 5 EU ADM	2 y, no recurrence		
_	+	Abdominal wall, etc.	SG	MMC, 5-FU, ADM	5m, death		
+	+	Lung	—		2m, death		
+	+	Lt. adrenal $- \rightarrow$ Brain	– + (method NM ^o)	TU ^f , Ubenimex	4m, death		
+		\rightarrow Drain	+ (method NW [*])	_	1 y, death 4 m, death		
+			TG	_	2y, death		
_	+	_	SG	MFC	3y, no recurrence		
+	+	$- \rightarrow$ Peritoneum	_	CAI ^g : MMC, ADM→ MTX, 5-FU, MMC, leucovorin	3m, death		
_	+	$- \rightarrow Lung$	SG	_	2m, death		
—	+	-	SG	-	8m, no recurrence		
$- \rightarrow +$	_	Mediastinum, etc.	SG	+ (menu: NM)	5m, death		
$- \rightarrow +$	+		SG	MMC, 5-FU, OK-432, 5 KE	2m, death		
			TG	—	NM		
			– SG	- 5 FUL (arally)	NM 17m dooth		
$\rightarrow +$ +	+	_	3G TG	5-FU (orally) Res ^h : 5-FU, OK-432 2 KE \rightarrow 5-FU, TU ^f (orally), OK-432	17m, death 7m, alive		
_	+	_	SG	CDDP, etoposide \rightarrow etoposide (orally)	8m, no reccurence		
$- \rightarrow +$	+		SG	-	2y, death		
-	+	$- \rightarrow$ Peritoneum	TG	_	10m, death		
-	+		SG	FA-CDDP	5m, alive		
+	+	Peritoneum	Unresectable	MMC, 5-FU	9m, death		
_	+	Bone	_	Carboplatin, etoposide HDCT ⁱ , PBSCT ^j	7m, death 8m, alive		
$- \rightarrow +$	+	Peritoneum	SG	CDDP, 5-FU	6m, death		
$- \rightarrow +$ $- \rightarrow +$	+	rentoneum	$STG \rightarrow PH$	TU ^f (orally)	8m, alive		
_	+		STG	CDDP, TU ^f	(period NM), alive		
$- \rightarrow +$	+		TG	TU ^f	4m, death		
$- \rightarrow +$			RRS	Lentinan, $TU^{f} + HAE^{k}$	7 m, alive		
+	-	-	-	CAP-PVP ¹	3m, death		
+	-	_	-	CAP-PVP, TU ^f	10m, death		
+			TG	Res ^h : CDDP, Tegafur suppo	4m, death		
+			Simple excision	_	10m, alive		
			SG		9d, death		
_	+		SG	$\mathbf{N}\mathbf{M}^{\mathrm{p}}$	NM		
_	+	Omentum	EG + Sp	_	3 y 6m, alive		
—	+	Omentum, pancreas, spleen	SG + Sp		2m, death		

Case no.	Author	Year reported	Age/Sex	Location ^a	Size (cm)	Gross type ^b	Depth of invasion
43	Matsusaka [1]	1976	54/M	L		Type 5 [°]	mp
44	"	"	65/F	M (Less)	4×2.2	Type 2	ss
45	Chejfec [40]	1977	66/M	UÚ	$9 \times 8 \times 1$. 1	
46	"	"	79/M	(Gre)	$15 \times 9 \times 7$	Type 2	
47	Eimoto [41]	1980	66/M	Ù (Ánt)	$9 \times 7.5 \times 4$	Type 2	si
48	Abrams [42]	1980	33/F	UML	0.2-4.5	I (multiple)	sm
49	Shibuya [43]	1985	54/M	L (Gre)			si
50	Fukuda [44]	1988	74/M		0.8	IIc	sm
51	Hussein [45]	1990	42/M	L (Less)	3	Type 2	si
52	O'Byrne [46]	1997	54/M	U			
53	Sato [47]	1997	74/M	U (Less)	$5 \times 4 \times 1.5$	Type 2	SS
54	Current patient	2001	67/M	M (Post)	10×8.5	Type 1	SS

SMT, Submucosal tumor; ADM, adriamycin; MMC, mitomycin C; MTX, methotrexate; 5-FU, 5-fluorouracil; CDDP, cisplatin; VCR, vincristine ^aU, Upper-third stomach; M, middle stomach; L, lower third stomach; EGJ, esophgo-gastric junction; RS, remnant stomach; Ant, anterior wall; Post, posterior wall; Gre, greater curvature; Less, lesser curvature

^bJapanese classification of gastric carcinoma

°SMT-like gross type

d Lymph node

^eSG, Subtotal gastrectomy; TG, total gastrectomy; PS, pancreaticosplenectomy; STG, splenototal gastrectomy; PH, resection of left lateral lobe of the liver; RRS, resection of the remnant stomach; EG, esophagogastrectomy; Sp, splenectomy

^fTegafur-uracil

g Injection from celiac artery

^hUsing reservoir catheter

ⁱHigh-dose chemotherapy with CDDP + etoposide + CPA + VCR

Peripheral blood stem cell transplantation

^kEmbolization of hepatic artery

 $^{1}CPA + ADM + VCR + CDDP + etoposide$

 m Cyclophosphamide + doxorubicin + $\dot{V}CR$

ⁿCyclophosphamide + doxorubicin + etoposide

°Vincristine + chlorambucil + dexamethasone

^pNM, Not metioned; d, day(s); m, month(s); y, year(s)

crine cell carcinoma (ECC) was reported by Matsusaka et al. [1]. Endocrine cell tumors of the stomach have been classified as "classical-type carcinoid" and "endocrine cell carcinoma" (ECC) [2], with different cell origin [4,52], biological behavior, and prognosis [3,4]. Iwafuchi et al. [4] reported that most early gastric ECCs (76%) were located in the deep mucosa or submucosa and were adjacent to coexisting instramucosal differentiated adenocarcinomas. In our patient, an adenocarcinoma component was recognized in continuity with the ECC, which suggests ordinary histological carcinogenesis of an ECC or glandular-ECC.

In the development of gastric endocrine cell tumor, possible factors are chronic hypergastrinemia due to type A gastritis [52], pernicious anemia [53], and longterm treatment with proton pump inhibitors (PPI) [54]. In our patient, although there was no history of pernicious anemia or PPI administration and no gastrin production in the tumor, the serum gastrin level was revealed to be high. Although the origin of this high level was unknown, this may have played an important role in the progression of the tumor, because most gastrointestinal hormones, except for somatostatin, promote the proliferation of both the gastrointestinal epithelium and of cancer [3].

To date, 37 cases have been reported in the Japanese literature and 16 in the English-language literature, in terms of clinicopathology, therapies, and/or prognosis of gastric ECC (Table 1). According to previous studies with clinicopathological analysis [5–7,44], the macroscopic appearance of gastric ECC varies in its early stage. However, when it is advanced, type 2 (tumorulcerative) is dominant and type 1 (fungating or polypoid), as in our patient, is less frequent [5,6]. Matsui et al. [6] hypothesized that a crater-like ulceration develops, probably due to rapid proliferation, and that only a few tumors maintain a polypoid appearance in the advanced stage.

Metastasis					
Liver	\mathbf{N}^{d}	Other	Operation ^e	Chemotherapy and other treatment	Prognosis ^o
+			SG	_	1 y, death
				_	5 y, alive
_	+		Unresectable	_	1 m, death
+	_		_	_	2m, death
+	+	Peritoneum	Unresectable	5-FU, MMC	5m, death
_	+		TG	_	82 m, alive
+	+	Retroperitonium, omentum, pelvis	Unresectable	_	1m, death
$\rightarrow +$	+	_	TG	5-FU, OK-432	6m, death
+	+	Omentum	SG	$\begin{array}{c} \text{CAV}^{\text{m}} \rightarrow \text{CDDP}, \\ \text{VP-16} \rightarrow \\ \text{3-deazaguanine} \end{array}$	9m, death
_			_	$CDE^n \rightarrow carboplatin,$ MTX + CDE $\rightarrow VCD^o$	22m, death
_	+	_	TG	Tegafur \rightarrow MTX, CDDP, epirubicin, 5-FU	21 m, death
_	+	_	STG	TU ^f (orally)	1 y, no recurren

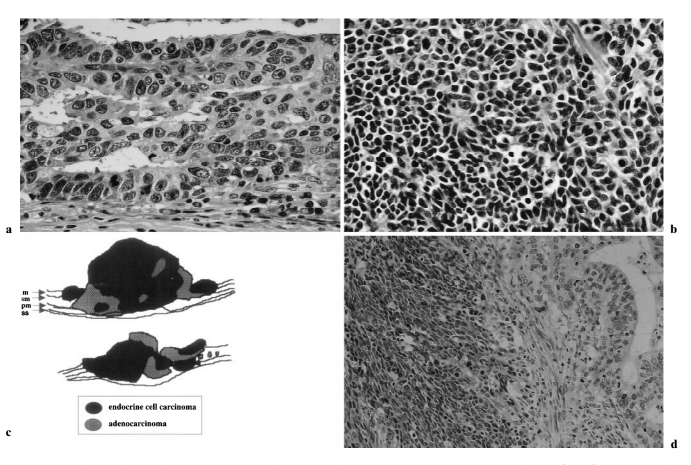


Fig. 5a–d. Histological findings of the tumor. This tumor consisted of components of **a** adenocarcinoma ($\times 100$) and **b** endocrine cell carcinoma ($\times 100$). **c** Schematic view of the main sections of the tumor. **d** Neuron-specific enolase (NSE) staining demonstrated strong positivity in the endocrine cell carcinoma component but not in the adenocarcinoma component ($\times 50$)

Concerning treatment, because of the tumor's aggressive biological characteristics, intensive chemotherapy, with or without operation, is recommended for gastric ECC at any stage [44]. Previously, several chemotherapeutic menus have been employed for gastric ECCs, but their effectiveness has been shown in only a few patients [32,46,55]. Sakai et al. [32] showed complete remission in a 31-year-old man with gastric ECC with multiple bone metastasis (case 29 in Table 1), with seven cycles of chemotherapy consisting of cisplatin (CDDP), etoposide, cyclophosphamide, epirubicin, and vincristine, followed by high-dose chemotherapy and peripheral blood cell transplantation. O'Byrne et al. [46] reported that a 54-year-old man with gastric ECC with severe epigastralgia and weight loss (case 52 in Table 1) was treated with six cycles of CDE (cyclophosphamide, doxorubicin, etoposide) chemotherapy, resulting in 11 months of complete remission. In patients with neuroendocrine carcinoma or undifferentiated small cell carcinoma, including lung small cell carcinoma and extrapulmonary small cell carcinoma, PE (cisplatin and etoposide) and CDE chemotherapies have been thought to be effective [56,57]. The tumor in our patient was resected curatively without distant metastsis, and this may explain the 1 year of survival in spite of the large tumor size and advanced stage. However, we thought it necessary to follow this patient carefully, because gastric ECCs show a high Ki-67 index, as in this patient [58], and some of them relapse after long intervals, such as 747 days [55].

Acknowledgments The authors are indebted to Dr. Geoffrey Barraclough of Kobe International Medical Association for his English-language review of this manuscript.

References

- 1. Matsusaka T, Watanabe H, Enjoji M. Oat-cell carcinoma of the stomach. Fukuoka Acta Medica 1976;67:65–73.
- Watanabe H, Jass JR, Sobin LH, editors. Histological typing of oesophageal and gastric tumours. WHO, 2nd edn. Berlin Heidelberg New York Tokyo: Springer-Verlag; 1990. p. 1– 108.
- 3. Tahara E. Endocrine tumors of the gastrointestinal tract: classification, function, and biological behavior. Dig Dis Pathol 1988;1:121–47.
- Iwafuchi M, Nishikura K, Watanabe H. Early endocrine cell carcinoma of the stomach and rectum (in Japanese with English abstract). Endosc Dig 1995;7:275–84.
- Iwafuchi M, Watanabe H, Ishihara N, Noda Y, Ajioka Y. Pathology of gastrointestinal carcinoid (in Janese). Clin Gasteroenterol 1990;5:1669–81.
- Matsui K, Kitagawa M, Miwa A, Kuroda Y, Tsuji M. Small cell carcinoma of the stomach: a clinicopathologic study of 17 cases. Am J Gastroenterol 1991;86:1167–75.
- Staren ED, Lott S, Saavedra VM, Jansson DS, Dezial DJ, Saclarides TJ, et al. Neuroendocrine carcinomas of the stomach: a

clinicopathologic evaluation. valuation. Surgery 1992;112:1039-47.

- Hayashi I, Horie A, Kuroda Y, Koto Y, Furusawa M, Nakahara K, et al. A case of gastric oat cell carcinoma (in Japanese). Jpn J Cancer Clin 1980;26:185–91.
- Kazato K, Kobayashi W, Kin T, Iida M, Uchida M. Gastric carcinoid with coexisting tubular adenocarcinoma in the same tumor, report of a case (in Japanese with English abstract). Stomach and Intestine 1983;18:245–53.
- Ushiyama H, Wataya T, Suzuki K, Nemoto N, Okano T, Uchida T, et al. An autopsy case of small cell carcinoma originating either in the cardia or esophagus (in Japanese with English abstract). Pathology and Clinical Medicine 1984;2:123–8.
- 11. Emi Y, Takahashi S, Yokose Y, Kinugasa T, Nakae D, Konishi Y. Malignant gastric endocrinoma coexisting with moderately differentiated adenocarcinoma in advanced gastric cancer (in Japanese with English abstract). J Nara Med Ass 1984;35:222–8.
- Terasawa K, Kawai K, Yokoe M, Sue K, Ihara K, Kitadai M, et al. A case of undifferentiated small cell carcinoma at the esophagocardiac junction (in Japanese with English abstract). Journal of Okayama Saiseikai General Hospital 1985;17:51–6.
- Kaketani K, Mitarai Y, Zeze K, Kuwahara A, Saito T, Kobayashi M. Two cases of oat cell carcinoma of the stomach (in Japanese with English abstract). Nihon Rinshogeka Gakkaishi 1987;48: 1687–92.
- 14. Ishihara T, Gondo T, Takahashi M, Uchino F, Kouchiyama T, Okazaki Y, et al. An autopsy case of synchronous small cell carcinoma in residual stomach and hepatocellular carcinoma containing small cell carcinoma foci (in Japanese with English abstract). Pathology and Clinical Medicine 1988;6:709–15.
- 15. Iwafuchi Y, Honda K, Ito T, Hasegawa A, Kunisada K, Kamimura A, Noda Y, et al. Endocrine cell carcinoma in remnant stomach (in Japanese with English abstract). ENDOSCOPIC FORUM for digestive disease 1989;5:203–7.
- Takeshita K, Kasahara M, Kuroda M, Mizoguchi Y, Horibe Y, Tashiro K, et al. Clinicopathological analysis of three cases of special gastric tumor (endocrine cell carcinoma) (in Japanese). Fujitagakuen Igakkaishi 1990;14:77–81.
- Matsumoto K, Sano M, Tobari M, Murakami T, Sugiyama Y, Yamagata N, et al. Small-cell carcinoma of the stomach: a case study (in Japanese with English abstract). Saishinigaku 1990;45: 2463–9.
- Maeta Y, Motoyama H, Uchikoshi Y, Tanaka Y, Boku S, Shibuya T, et al. A case of endocrine cell carcinoma of the stomach (in Japanese with English abstract). ENDOSCOPIC FORUM for digestive disease 1990;6:186–90.
- Kuhara T, Tsuchihashi K, Fujise Y. Small cell carcinoma of the stomach, report of a case (in Japanese with English abstract). Stomach and Intestine 1991;26:1059–65.
- 20. Uesugi H, Kiyohashi A, Sano H, Sakai T, Takagi S, Oikawa Y, et al. A case of the small cell carcinoma of the stomach (in Japanese with English abstract). Prog Dig Endosc 1991;39:323–6.
- Gotoh H, Nemoto N, Inamura H, Miyake K, Sakurai I, Suzuki T. A case of gastric small cell undifferentiated carcinoma with components of adenocarcinoma, squamous cell carcinoma and cartilaginous tissue (in Japanese). Pathol Clin Med 1992;10:1169–73.
- Katsuyama S. A case of small cell carcinoma of the stomach. A case of small cell carcinoma of the stomach (in Japanese with English abstract). Nihon Rinshogeka Gakkai Zasshi 1992;53:348–63.
- 23. Kamio T, Suko S, Kimura S, Kawazu R, Tanoue T, Hirota K, et al. Two cases of a gastric endocrine cell carcinoma (in Japanese with English abstract). Jpn J Cancer Clin 1992;38:1511–8.
- 24. Nakamoto M, Kawaguchi K, Nake S, Nishio Y, Urakawa T, Ioroi T, et al. A case of gastric endocrine cell carcinoma combined with adenocarcinoma in the sigmoid colon (in Japanese with English abstract). Nihonsholaligakkaishi 1992;25:2171–5.
- 25. Masuyama K, Oonishi Y, Sawataishi M, Suzuki S, Yamazaki K, Ishizawa S, et al. A case of small cell carcinoma of the stomach

with multiple liver metastases (in Japanese with English abstract). Jpn J Cancer Chemother 1994;21:2338–40.

- 26. Nagafuchi K, Nishihara K, Yamamoto H, Watanabe M, Hirose N, Miki T, et al. A case of the endocrine carcinoma of the stomach occurring concurrently with adenocarcinoma (in Japanese with English abstract). Nihon Shokakigeka Gakkaishi (Jpn J Gastroenterol Surg) 1994;27:1805–9.
- Imamura K, Fukuda M, Mori I, Kitamura T, Nakano M, Nakata T, et al. A case of the endocrine carcinoma of the stomach (in Japanese with English abstract). Jpn J Cancer Clin 1995;41:57– 60.
- Maekawa H, Nishimura K, Kobayashi S, Sakakibara N, Wada R. Gastric endocrine cell carcinoma coexisted with gastric tubular adenocarinoma — a case report — (in Japanese with English abstract). Nihon Rinshogeka Ikaishi 1995;56:1862–5.
- Kodama S, Nakatsuka H, Kushiro J, Fujitaka T, Taniyama K. A case report of gastrin producing endocrine cell carcinoma of the stomach. Shokakigeka (Gastroenterol Surg) 1995;18:1739– 44.
- Fudaba Y, Ohshiro H, Itamoto T, Nagano M, Ohshiro T, Kadoya T, et al. A case of gastric endocrine cell carcinoma (in Japanese with English abstract). Hiroshimakenritsubyoinishi 1996;28:111–5.
- Inoue F, Nishida O, Mizumoto T, Furukawa H, Saiga T. Small cell carcinoma of the stomach, report of a case (in Japanese with English abstract). Stomach and Intestine 1996;31:797–801.
- 32. Sakai K, Nomura H, Nogami T, Saeki T, Etoh Y, Imamura H, et al. A case of complete remission of gastric endocrine cell carcinoma with multiple bone metastasis by combination chemotherapy and high-dose chemotherapy with autologous peripheral blood cell transplantation (in Japanese with English abstract). Jpn J Cancer Chemother 1997;24:2277–80.
- Hiramoto Y, Onda M, Tokunaga A, Okino T, Lee Y, Ikeda K, et al. Endocrine cell carcinoma of the stomach; a report of five cases (in Japanese). J Jpn Soc Gastroenterol Carcinogen 1998;10:399– 401.
- 34. Iwasaki R, Nagahara A, Ohta K, Iijima K, Ohno Y, Ohkura R, et al. Chemotherapeutic effects on gastric endocrine cell carcinoma with multiple liver metastasis: report of two cases (in Japanese with English abstract). Gastroenterol Endosc 1998;40:1889–96.
- 35. Yamamoto S, Kawamura A, Ozaki S, Hiroi M, Araki K. A case of the endocrine carcinoma of the stomach finding markedly reduced liver metastasis by arterial of cisplatin (in Japanese). Gekachiryo (Surg Ther) 1999;81:124–7.
- Christodoulopoulos JB, Klotz AP. Carcinoid syndrome with primary carcinoid tumor of the stomach. Gastroenterology 1961; 40:429–40.
- Vasudeo PB, Mody AE, Vora MK, Dalal KA, Mascarenhas AFA. Malignant carcinoid tumor of the stomach. J Postgrad Med 1966; 12:57–60.
- Brodman HR, Pai BN. Malignant carcimoid of the stomach and distal esophagus. Am J Dig Dis 1968;13:677–81.
- Parks TG. Malignant carcinoid and adenocarcinoma of the stomach. Brit J Surg 1970;57:377–9.
- Chejfec G, Gould VE. Malignant gastric neuroendocrinomas. Ultrastructural and biochemical characterization of their secretory activity. Hum Pathol 1977;8:433–40.

- Eimoto T, Hayakawa H. Oat cell carcinoma of the stomach. Path Res Pract 1980;168:229–36.
- Abrams JS. Multiple malignant carcinoids of the stomach. Arch Surg 1980;115:1219–21.
- Shibuya H, Azumi N, Abe F. Gastric small cell undifferentiated carcinoma with adeno and squamous cell carcinoma components. Acta Pathol Jpn 1985;35:473–80.
- Fukuda T, Ohnishi Y, Nishimaki T, Ohtani H, Tachikawa S. Early gastric cancer of the small cell type. Am J Gastroenterol 1988;83: 1176–9.
- Hussein AM, Otrakji CL, Hussein BT. Small cell carcinoma of the stomach. Case report and review of the literature. Dig Dis Sci 1990;35:513–8.
- 46. O'Byrne KJ, Cherukuri AK, Khan MI, Farrell RJ, Daly PA, Sweeney EC, et al. Extrapulmonary small cell gastric carcinoma. A case report and review of the literature. Acta Oncol 1997;36: 78–80.
- Sato T, Sakuma H, Isobe T, Naka F, Ueda H, Matsubara F, et al. Concurrent small-cell carcinoma and adenocarcinoma of the stomach. Dig Surg 1997;14:61–4.
- Japanese Gastric Cancer Association. Japanese classification of gastric carcinoma. 2nd English ed. Gastric Cancer 1998;1:10– 24.
- 49. Matsubayashi H, Watanabe H, Nishikura K, Ajioka Y, Maejima T, Kijima H, et al. Advantages of immunostaining for DNA analysis using PCR amplification to detect p53 abnormality in long-time formalin-fixed tissues of human colorectal carcinomas. J Gastroenterol 1998;33:662–9.
- 50. Oberndorfer S. Karzinoid tumoren des dunndarms. Frankfurt Z Path 1907;1:426–32.
- Soga J, Tazawa K. Pathologic analysis of carcinoids. Histologic re-evaluation of 62 cases. Cancer 1971;28:990–8.
- 52. Itsuno M, Watanabe H, Iwafuchi M, Ito S, Yanaihara N, Sato K, et al. Multiple carcinoids and endocrine cell micronests in type A gastritis. Their morphology, histogenesis, and natural history. Cancer 1989;63:881–90.
- McGuigan JE, T McGuigan JE, Traudeau Wl. Serum gastrin concentration in pernicious anemia. N Engl J Med 1970;282:358– 61.
- Goldfain D, LeBodic MF, Lavergne A, Galian A, Modigliani R. Gastric carcinoid tumours in patients with Zollinger-Ellison syndrome on long-term omeprazole. Lancet 1989;I:776–7.
- 55. Kimura H, Konishi K, Kaji M, Maeda K, Yabushita K, Tsuji M, et al. Highly aggressive behavior and poor prognosis of small cell carcinoma in the stomach: flow cytometric and immunohistochemical analysis. Oncol Rep 1999;6:767–72.
- Gaast AVD, Verwey J, Prins E, Splinter TAW. Chemotherapy as treatment of choice in extrapulmonary undifferentiated small cell carcinomas. Cancer 1990;65:422–4.
- Moertel CG, Kvols LK, O'Connell MJ, Rubin J. Treatment of neuroendocrine carcinomas with combined etoposide and cisplatin. Cancer 1991;68:227–32.
- 58. Tanaka S, Mori H, Nakamura H, Tomita A, Umeno T, Ikeda S, et al. The evaluation of the biological aggressiveness of endocrine cell carcinoma of the stomach with proliferating cell nuclear antigen and Ki-67 labeling index (in Japanese with English abstract). Jpn J Gastroenterol Surg 1996;29:795–9.