ORIGINAL ARTICLE

Endoscopic and histological features of Helicobacter pylori-negative differentiated gastric adenocarcinoma arising in the antrum

Maiko Takita,* Ken Ohata,* Rin Inamoto,* Marie Kurebayashi,* Syunya Takayanagi,* Yoshiaki Kimoto,* Yuichiro Suzuki,* Rindo Ishii,* Kohei Ono,* Ryoji Negishi,* Yohei Minato,* Eiji Sakai,*† Takashi Muramoto,* Nobuyuki Matsuhashi‡ and Shin Ichihara§

Departments of *Gastrointestinal Endoscopy, ‡Gastroenterology, NTT Medical Center Tokyo, Tokyo, †Department of Gastroenterology, Yokohama Sakae Kiosai Hospital, Yokohama and §Department of Surgical Pathology, Sapporo Kosei General Hospital, Sapporo, Japan

Key words
diagnosis, differentiated-type cancer, endoscopic resection, gastric cancer, Helicobacter pylori.

Accepted for publication 22 February 2021.

Correspondence
Ken Ohata, Department of Gastrointestinal Endoscopy, NTT Medical Center Tokyo, 5-9-22 Higashi-gotanda Shinagawa-ku, Tokyo 141-8625, Japan.
Email: ken.ohata1974@gmail.com

Declaration of conflict of interest: None

Funding support: NTT Medical Center Tokyo

Abstract

Background and Aim: With the increasing prevalence of persons without Helicobacter pylori (HP) infection, cases of HP-negative gastric cancer are increasing. Although rare, cases of differentiated adenocarcinoma of the antrum have been reported in HP-negative patients. We collected cases with such lesions and investigated their endoscopic and histological features.

Methods: Of 1965 consecutive patients with early gastric cancer who underwent endoscopic resection between January 2009 and December 2017, we extracted 9 cases of HP-negative differentiated adenocarcinoma located in the antrum (HPN-DAA). The clinical data, endoscopic findings, and histopathological findings were reviewed.

Results: Of the nine patients with HPN-DAA, seven were male, and the median age was 53.8 years. The tumor arose from the pyloric gland mucosa in all cases. According to the endoscopic findings, the lesions were flat-elevated or depressed, mimicking varioliform gastritis. Magnifying endoscopy with narrow-band imaging showed the absence of a clear demarcation line or an irregular microvessel/surface pattern. As for the histopathological findings, eight of the nine lesions were diagnosed as high-grade dysplasia/intraepithelial neoplasia, while the remaining case was diagnosed as tubular adenocarcinoma with submucosal infiltration. The findings of immunohistochemistry confirmed that three cases were of the intestinal mucin phenotype and six were of the mixed gastric and intestinal mucin phenotype.

Conclusion: HPN-DAA is a very rarely occurring cancer that had never been recognized earlier. They belong to the new category of HP-negative cancers, and there seems to be a certain number of such cases.

Introduction

Chronic gastritis caused by persistent Helicobacter pylori (HP) infection is known to be strongly associated with the development of gastric cancer, and HP infection has been certified as a “definite carcinogen” by the World Health Organization.1 With the high prevalence of HP infection, Japan is well known as a country with a high incidence of gastric cancer.2 Uemura et al. reported in their study that there were no cases of gastric cancer in the group without HP infection in their case series.3 However, in clinical practice, we sometimes encounter a small number of gastric cancers in patients without HP infection. As the number of persons without HP infection increases, cases with gastric cancer in this population is also expected to increase. These are recognized as cases of HP-negative gastric cancer. The reported incidence of HP-negative gastric cancer is in the range of 0.5–3.1%, which is quite a significant number.4–6

HP-negative gastric cancers can be classified into several phenotypes. The most common type is signet-ring cell carcinoma, which is a discolored lesion, frequently identified in the middle or lower third of the stomach.7 Meanwhile, a variety of differentiated-type gastric cancers has also been identified in patients without HP infection (e.g. gastric adenocarcinoma of the fundic gland type8; low-grade differentiated cancer of the gastric phenotype9; and foveolar-type dysplasia, endoscopically showing a raspberry-like appearance and recognized as a low-grade differentiated type of adenocarcinoma in Japan10). More recently, although rare, a differentiated type of adenocarcinoma mimicking varioliform gastritis has been reported as occurring in the gastric antrum of persons without HP infection. There are only a few
case reports, and the characteristics of these lesions remain unclear.11–13

In the present study, we collected a number of cases of HP-negative differentiated adenocarcinoma located in the antrum (HPN-DAA) and revealed the associations between their endoscopic and histological features.

Methods

Study design and patients. This retrospective cohort study was conducted to investigate the endoscopic and histopathological features of HPN-DAA. Among 1965 consecutive patients with early gastric cancer who underwent endoscopic resection at the NTT Medical Center Tokyo between January 2009 and December 2017, we extracted cases without HP infection. HP-negative status was confirmed by the following criteria: (i) no atrophy or gastritis as assessed by endoscopy, (ii) no histologic evidence of HP infection, (iii) negative results of two consecutive tests for HP infection, and (iv) negative history of HP eradication therapy. The endoscopic findings were reviewed using the Kimura-Takemoto Classification14 and Kyoto classification of gastritis.15 HP-negative patients met these inclusion criteria: no mucosal atrophy and histological confirmation of a regular arrangement of collecting venules on the lesser curvature of the lower gastric body. Resected specimens were histologically evaluated according to the updated Sydney system.16 A patient was defined as HP-negative when there was no mucosal atrophy, intestinal metaplasia, presence of HP, or infiltration by neutrophils and mononuclear cells. With regard to the test for HP infection status, we use the anti-HP serum IgG antibody kit (BML, INC., Tokyo, Japan) and 13C-labeled urea breath test (UBIT® tablets 100 mg, Otsuka Pharmaceutical Co. Ltd., Tokyo, Japan). HP-negative patients met the criteria of a negative result in both the former (<3.0 U/ml) and the latter (<2.5%). Finally, 39 cases (2.0%) were identified as cases of HP-negative cancers. Then, we divided the cases into several types according to their endoscopic findings: (i) signet-ring cell carcinoma; (ii) gastric adenocarcinoma of the fundic gland type; (iii) low-grade differentiated cancer of the gastric phenotype and foveolar-type dysplasia, endoscopically showing a raspberry-like appearance; and (iv) HPN-DAA. Finally, 19 patients with HPN-DAA were enrolled in this study. The clinical data, including age; gender; and endoscopic findings such as the tumor location, tumor size, tumor morphology, preoperative diagnosis, as well as histopathological findings, were reviewed. To elucidate HPN-DAA characteristics, we extracted patients with HP-positive differentiated adenocarcinomas located in the antrum (HPP-DAA) who underwent endoscopic resection during the same period and compared HPN-DAA and HPP-DAA clinicopathological findings. We obtained written informed consent for the endoscopic resection and investigation of the long-term outcomes at the time of the initial treatment. This study was conducted with the approval of the ethics committee of NTT Medical Center Tokyo (20-22).

Procedure of endoscopy. We use magnification video endoscope systems (GIF-H260Z or GIF-H290Z; Olympus Medical Systems Co. Ltd., Tokyo, Japan) and a standard optical video endoscope system (EVIS LUCERA ELITE system; Olympus Medical Systems). We obtained the images of the lesions, not only by white-light endoscopy (WLE) but also by magnifying endoscopy with narrow-band imaging (ME-NBI). The diagnostic strategy adopted for the findings of ME-NBI was in accordance with the Magnifying Endoscopy Simple Diagnostic Algorithm for early Gastric cancer (MESDA-G).17 To ensure precise results of the pathological examination, all the lesions were resected by endoscopic submucosal dissection (ESD). All the procedures were performed by endoscopists certified by the Japan Gastroenterological Endoscopy Society with 10 years of experience or longer in endoscopy.

Pathological examination. The tumor size was determined by measuring the resected specimen prior to tissue fixation in formalin. The specimens were processed and cut into sections at a thickness of 4 μm, which were stained with hematoxylin and eosin. Then, the differentiation degree, invasion depth, presence/absence of lymphovascular invasion, and the lateral and vertical resection margins were assessed. Cell differentiation (based on the staining status for MUC2, MUC5AC, MUC6, CDX-2, and CD10), proliferation (based on the Ki-67 index), and genetic status (based on the staining status for MLH1, MSH2, MSH6, and PMS2) were evaluated immunohistochemically using monoclonal antibodies to MUC2 (Novocastra), MUC5AC (Novocastra/Leica Biosystems, Newcastle, UK), MUC6 (Novocastra), CDX-2 (Dako/Agilent Technologies, Glostrup, Denmark), CD10 (Novocastra), Ki-67 (clone MIB-1; Dako), MLH1 (FALCO biosystems, Kyoto, Japan), MSH2 (FALCO), MSH6 (FALCO), and PMS2 (FALCO). The histopathological examinations were performed by an experienced pathologist specialized in pathology of the gastrointestinal tract (S.I.).

Statistical analyses. The statistical significance of differences in clinical parameters were evaluated using the χ2 test or unpaired Student’s t-test. A P value <0.05 was considered statistically significant.

Results

Clinical features and endoscopic findings. The clinic features and endoscopic findings of the patients are shown in Table 1. Among the nine patients, seven were male, and the median age was 53.8 years (range 32–64 years). None of the patients had a history or family history of gastric cancer. None of the patients showed atrophy or gastritis in the background mucosa. The lesions in all cases were located in the antrum. Morphologically, they were flat-elevated or depressed-type lesions, like varioliform gastritis (Fig. 1). ME-NBI showed the absence of a clear demarcation line and of the irregular microvessel/surface pattern (IMVP/IMSP) in almost all the cases. Hence, according to MESDA-G, all the lesions were diagnosed as noncancer. Biopsy specimens had been obtained in all the cases previously. Only three patients were categorized as group 5 (Carcinoma) in advance, while the remaining were classified as group 2 (Indefinite for neoplasia; material for which diagnosis of neoplastic or nonneoplastic lesion is difficult), group 3 (Adenoma), or group 4 (Neoplastic lesion that is suspected to be carcinoma) based on the “Group classification.”18 Furthermore, despite the lesions having been diagnosed several years
Table 1   Clinicopathological features of *Helicobacter pylori*-negative differentiated adenocarcinoma located in the antrum

| Case | Gender | Age | Background mucosa | Location | Morphology | Size (mm) | Preoperative diagnosis | Differentiation | Depth |
|------|--------|-----|--------------------|----------|------------|-----------|------------------------|----------------|-------|
| 1    | Male   | 51  | Pyloric            | Less     | IIa + IIc  | 10        | Group 4 tub1            | tub1           | M     |
| 2    | Male   | 57  | Pyloric            | Ant      | IIa        | 6         | Group 3 tub1            | tub1           | M     |
| 3    | Male   | 56  | Pyloric            | Post     | IIa        | 12        | Group 2 tub1            | tub1           | M     |
| 4    | Male   | 51  | Pyloric            | Gre      | IIa        | 14        | Group 3 tub1            | tub1           | M     |
| 5    | Male   | 32  | Pyloric            | Gre      | IIa        | 8         | Group 5 tub1            | tub1           | M     |
| 6    | Female | 48  | Pyloric            | Gre      | IIc        | 10        | Group 3 tub1            | tub1           | M     |
| 7    | Female | 61  | Pyloric            | Gre      | IIa + IIc  | 18        | Group 5 tub1            | tub1           | SM1   |
| 8    | Male   | 64  | Pyloric            | Post     | IIa        | 2         | Group 2 tub1            | tub1           | M     |
| 9    | Male   | 64  | Pyloric            | Gre      | IIa        | 3         | Group 5 tub1            | tub1           | M     |

Ant, anterior wall; Gre, greater curvature; Less, lesser curvature; M, intramucosal cancer; Post, posterior wall; SM1, invasion depth < 500 μm from muscularis mucosa.

Figure 1  White-light endoscopy of 10 *Helicobacter pylori* (HP)-negative differentiated adenocarcinoma located in the antrum cases. The lesions were recognized as a single erosion in the antrum of the stomach in the absence of HP infection. Unlike typical gastric cancers, the border line was unclear. A–I corresponds to cases 1–9 in Tables 1 and 2.
earlier in three of the cases, the patients had been followed up as the lesions were diagnosed as not being neoplastic. Hence, endoscopic resection was performed for the purpose of diagnosis in some of the cases. Representative WLE and ME-NBI images are shown in Figure 2.

Pathological findings. All the lesions were resected by ESD without any complications. The histopathological findings of the nine cases are shown in Tables 1 and 2. Even though the lesions in all cases showed similar endoscopic findings, their pathological findings were not identical. With regard to the background mucosa, the tumors arose from the pyloric gland mucosa in all cases. All the lesions were diagnosed as well-differentiated tubular adenocarcinoma based on the glandular structure and cellular atypia using the Japanese gastric cancer treatment guidelines.19 Except for one case, the remaining eight cases were of intramucosal cancer (“carcinoma in situ”). These were classified as noninvasive high-grade neoplasia (category 4) according to the Vienna classification20 and as high-grade dysplasia/intraepithelial neoplasia according to the WHO classification.21 As for the single case, namely, case 7, the lesion showed submucosal invasion (250 μm from the muscularis mucosa,
diagnosed as SM1 according to the Japanese gastric cancer treatment guidelines), which was diagnosed as invasive neoplasia (category 5) according to Vienna classification and tubular adenocarcinoma by the WHO classification. In addition, they often had nonneoplastic epithelium on the surface layer or within the lesion. Lymphovascular invasion was not confirmed in any of the cases, and all cases were resected with negative lateral and vertical margins. Interestingly, it was notable that all cases had fibromuscular obliteration, such as in the case of rectal mucosal prolapse syndrome, of the lamina propria around the lesions. Immunohistochemistry revealed that three of the cases were of the intestinal type, and six were of the mixed type (Table 2). Of the six cases of mixed-type HPN-DAA, two were gastric mucin phenotype dominant (MUC5AC(+), MUC6 (+)), although they also showed CDX2(+), while four cases were intestinal mucin phenotype dominant (MUC2(+), CDX2 (+)), and they were also MUC6(+). The Ki-67 labeling index of the tumor cells was over 90%, but the superficial nonneoplastic foveolar epithelial cells were negative for Ki-67, with a Ki-67 labeling index in the range of 49.4%–66.8%. With regard to the expression of mismatch repair (MMR) proteins such as MLH1, MSH2, MSH6, and PMS2, no down-regulation in the expression of any of these proteins was observed in any of the cases. Representative pathological images are shown in Figure 3.

**Comparison of HPN-DAA and HPP-DAA.**
Clinicopathological findings of HPN-DAA and HPP-DAA are summarized in Table 3. Patients with HPN-DAA were significantly younger than those with HPP-DAA. No gender difference was found. Average HPN-DAA tumor size was significantly smaller than that of HPP-DAA. With respect to morphology, HPN-DAA were more often observed to protrude compared with HPP-DAA. No difference between groups was found in the prevalence of submucosal invasion.

---

**Figure 3** Histopathological images of *Helicobacter pylori*-negative differentiated adenocarcinoma located in the antrum (case 6). (a) The lesion arose from the pyloric glands in the absence of intestinal metaplasia. Fibromuscular obliteration of the lamina propria is observed in the background mucosa. (b,c) The neoplastic gland showing irregular glandular arrangement with low-grade cellular atypia. There is some nonneoplastic epithelium on the surface layer of the lesion: (b) low power field and (c) high power field. (d–i) Immunohistochemistry: The neoplastic cells showing negative staining for MUC5AC (d) and positive staining for MUC6 (e), MUC2 (f), CDX-2 (g), and CD10 (h). (i) Ki-67 labeling index is 59.2%.
cinoma arising from the antrum, although there have been three reports of differentiated adenocarcinoma located in the antrum (HPN-DAA) and H. pylori-positive (HPP-DAA) patient characteristics. 

| HPN-DAA (n = 9) | HPP-DAA (n = 558) | P value |
|----------------|------------------|--------|
| Age, years     | 53.8 ± 10.0      | 71.3 ± 8.6 | <0.01 |
| Gender, male   | 7 (77.8%)        | 411 (73.7%) | 0.56 |
| H. pylori status | N/A             |        |        |
| Negative       | 9 (100%)         | 0       |        |
| Positive       | 0                | 242 (43.4%) |        |
| Eradicated     | 0                | 316 (56.6%) |        |
| Diameter, mm   | 9.2 ± 5.1        | 13.8 ± 9.7 | 0.01 |
| Macropscopic morphology | Protruded | 4 (44.4%) | 195 (34.9%) | 0.39 |
|                | Flat, depressed  | 5 (55.6%) | 363 (65.1%) |        |
| Tumor depth    | 1 (11.1%)        | 11 (2.0%) | 0.15 |
| <500 μm        | 8 (88.9%)        | 533 (95.5%) |        |
| ≥500 μm        | 1 (11.1%)        | 11 (2.0%) |        |
| Histological type | tub1            | 9 (100%) | 453 (81.2%) | 0.36 |
|                | tub2             | 0       | 85 (15.2%) |        |
|                | pap              | 0       | 20 (3.6%) |        |

Values are mean ± standard deviation or n (%). pap, papillary adenocarcinoma; tub1, well-differentiated adenocarcinoma; tub2, moderate-differentiated adenocarcinoma.

Discussion

With the decline in the number of people with HP infection owing to improved sanitary environments and spread of eradication therapy, the incidence of gastric cancers in persons without HP infection is expected to show a relative increase in Japan. There have been some previous reports of HP-negative cancers, such as signet-ring cell carcinoma; gastric adenocarcinoma of the fundic gland type; low-grade differentiated cancer with the gastric mucin phenotype; and foveolar-type dysplasia, endoscopically showing a raspberry-like appearance. These lesions are usually known to arise from the fundic gland area. Other types of gastric cancer are rarely experienced in HP-negative patients. In contrast, the lesions in our cases arose from the pyloric glands in the gastric antrum. Although there have been three reports of differentiated adenocarcinoma arising from the antrum, this is the first report to examine a certain number of such cases in detail.

As for our cases, or cases of HPN-DAA, the lesions were observed as superficially elevated or depressed lesions close to the pyloric ring in all cases. Unlike typical gastric cancers, the border line was unclear. In addition, even with ME-NBI, the demarcation line or IMVP/IMSP was not observed in most of the lesions. Hence, they were diagnosed as nonneoplastic lesions according to conventional diagnostics. Moreover, as it was difficult to make a definite diagnosis due to the low-grade atypia, only three cases (33.3%) were diagnosed as cancer in advance; another three cases with lesions that were not diagnosed as being neoplastic underwent follow-up examinations for several years (4–11 years). Fortunately, none of them showed rapid growth. Of the nine cases, eight were categorized as intramucosal cancer. However, if a single erosion, like varioliform gastritis, in the antrum of the stomach in the absence of HP infection is noted in the future, it is important to be aware that, although unlikely, the lesion could be a cancer. In that case, described above, as it is difficult to diagnose HPN-DAA endoscopically, it is thought that biopsy should be actively performed. Even if the result was indefinite for neoplasia, the possibility of cancer cannot be ruled out, so endoscopic resection for the purpose of diagnosis is considered to be acceptable.

As for the histopathological diagnosis, HPN-DAA showed weak atypia—not only structural atypia but cellular atypia as well. In addition, they often had a nonneoplastic epithelium inside the lesion as in gastric cancers detected after HP eradication. These histopathological features may make the endoscopic diagnosis of these lesions more difficult, and as a result, the lesions were often diagnosed as nonneoplastic lesions even by ME-NBI. Besides, interestingly, all lesions showed fibromuscular obliteration of the lamina propria in the background mucosa. Owing to this, the background mucosa around the lesion often protruded; therefore, it was considered that, morphologically, HPN-DAA mimicked varioliform gastritis. This fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.

As for the mucin phenotype, it has been reported that HPN-DAA are of the intestinal mucin phenotype. As most HP-negative cancers, such as gastric adenocarcinoma of the fundic gland type, low-grade differentiated cancer of the gastric phenotype, and foveolar-type dysplasia, are known to be of the gastric mucin phenotype, this fibromuscular obliteration was usually demonstrated in the distal antrum, and we thought that this phenomenon was due to gastric peristalsis (like in rectal mucosal prolapse syndrome). While fibromuscleosclerosis was not a specific finding of HPN-DAA, this feature is considered important for understanding the endoscopic findings of HPN-DAA.
fundic gland type. As for tumors with microsatellite instability, although they are often known to occur in female patients and arise from the antrum, no decrease in the expression of MMR was observed in the lesions in this study. However, it would still be difficult to rule out the possibility of involvement of some genetic abnormality in the carcinogenic pathway of HPN-DAA.

With regard to other risk factors for carcinogenesis, in addition to genetic mutations, EBV infection and autoimmune gastritis are also known risk factors. As for cancer associated with EBV infection, it has the characteristics of moderate to poorly differentiated cancer with lymphoid cell invasion, most commonly involving the upper to middle third of the stomach. We did not perform tests for the presence of EBV infection as the lesions in this study showed characteristics that were clearly different from the above-described features. Similarly, gastric cancers associated with autoimmune gastritis are assumed to show atrophic gastritis in the background mucosa, thus being obviously different from the lesions in this study.

In addition, considering that there was one case which showed submucosal invasion, HPN-DAA does appear to have the potential to transform into invasive cancer. With regard to HP-negative cancer, reports of intramucosal cancer have accounted for the majority so far; however, there have been a few reports of advanced cancers. Examination of a larger number of cases of HPN-DAA in the future is warranted for a more precise elucidation of their characteristics.

Compared to HPP-DAA, HPN-DAA tumor size was smaller. Similar results have been reported for HP-negative signet-ring cell carcinoma. Further investigation is needed to clarify whether this is due to differences in tumor proliferative capacity or whether HPN-DAA just happened to be discovered at early stages, supported by the lack of gastritis or intestinal metaplasia in the background mucosa. Considering that several cases of HPN-DAA did not change significantly over several years of follow-up, they may be slowly progressing lesions. In addition, patients with HPN-DAA were younger than those with HPP-DAA. Several similar reports have observed that patients with HP-negative cancer are younger than those with HP-positive cancer. If further studies confirm that patients with HP-negative cancer are younger than those with HP-positive cancer, we may expect the average age of gastric cancer patients to decrease as the HP-negative population increases.

The present study had several limitations. First, the sample size was very small as this study was a retrospective observational analysis performed at a single institution besides the fact that the lesion is rare. Further investigation in a larger population, such as through a multicenter analysis, is necessary. Second, the effects of bile acids were not investigated in this study. It is known that bile acids can damage DNA or cause carcinogenesis. A more detailed investigation of this point would be desirable. Third, although the endoscopic findings were similar, it is difficult to exclude the possibility that the disease may not have been the same in all cases as they had different mucin phenotypes. A more detailed examination in a larger population is warranted in the future to elucidate the characteristics, as well as pathogenesis, of these tumors in the future.

In conclusion, HPN-DAA is a very rare cancer that has never been recognized previously. The lesions have the characteristics of well-differentiated carcinoma and lack any specific endoscopic findings; these are some of the reasons they may have been overlooked so far. They belong to the new category of HP-negative cancers, and there seems to be a certain number of such cases. With the estimated increase in the population without HP infection, more attention will need to be paid to the possibility of such lesions when performing endoscopic examinations.

Acknowledgments

The authors thank all their colleagues at NTT Medical Center Tokyo who supported the study. They also thank IMIC (https://www.imic.or.jp/) for editing a draft of this manuscript.

References

1. IARC Working Group on the Evaluation of Carcinogenic Risks to Humans. Schistosomes, liver flukes and Helicobacter pylori. In: IARC Monogr Eval Carcinog Risks Hum 1994, Vol. 61. Lyon: Publication: IARC; 1994; 177–240.
2. Kobayashi T, Kikuchi S, Lin Y et al. Trends in the incidence of gastric cancer in Japan and their associations with Helicobacter pylori infection and gastric mucosal atrophy. Gastric Cancer. 2004; 7: 233–9.
3. Uemura N, Okamoto S, Yamamoto S et al. Helicobacter pylori infection and the development of gastric cancer. N. Engl. J. Med. 2001; 345: 784–9.
4. Kato S, Matsukura N, Tsukada K et al. Helicobacter pylori infection-negative gastric cancer in Japanese hospital patients: incidence and pathological characteristics. Cancer Sci. 2007; 98: 790–4.
5. Kakinoki K, Kushima R, Matsubara A et al. Reevaluation of histogenesis of gastric carcinomas: a comparative histopathological study between Helicobacter pylori-negative and H. pylori-positive cases. Dig. Dis. Sci. 2009; 54: 614–20.
6. Ono S, Kato M, Suzuki M et al. Frequency of Helicobacter pylori-negative gastric cancer and gastric mucosal atrophy in a Japanese endoscopic submucosal dissection series including histological, endoscopic and serological atrophy. Digestion. 2012; 86: 59–65.
7. Yamamoto Y, Fujisaki J, Omae M, Hirasawa T, Igarashi M. Helicobacter pylori-negative gastric cancer: characteristics and endoscopic findings. Dig. Endosc. 2015; 27: 551–61.
8. Ueyama H, Yao T, Nakashima Y et al. Gastric adenocarcinoma of fundic gland type (chief cell predominant type): proposal for a new entity of gastric adenocarcinoma. Am. J. Surg. Pathol. 2010; 34: 609–19.
9. Yamada A, Kaise M, Inoshita N et al. Characterization of Helicobacter pylori – Naïve Early Gastric Cancers. Digestion. 2018; 98: 127–34.
10. Shibagaki K, Fukuyama C, Mikami H et al. Gastric foveolar-type adenoma endoscopically showing a raspberry-like appearance in the Helicobacter pylori-uninfected stomach. Endoscopy International Open. 2019; 07: E784–91.
11. Ozaki Y, Suto H, Nosaka T et al. A case of Helicobacter pylori-negative intramucosal well differentiated gastric adenocarcinoma with intestinal phenotype. Clin J Gastroenterol. 2015; 8: 18–21.
12. Kotani S, Miyaoka Y, Fujiwara A et al. Intestinal-type gastric adenocarcinoma without Helicobacter pylori infection successfully treated with endoscopic submucosal dissection. Clin J Gastroenterol. 2016; 9: 228–32.
13. Yoshii S, Hayashi Y, Takehara T. Helicobacter pylori negative early gastric adenocarcinoma with complete intestinal mucus phenotype mimicking verrucous gastric. Dig. Endosc. 2017; 29: 235–6.
14. Kimura K, Takemoto T. An endoscopic recognition of the atrophic border and its significance in chronic gastritis. Endoscopy. 1969; 3: 87–97.
15 Kamada T, Haruma K, Inoue K, Shiotani A. Helicobacter pylori infection and endoscopic gastritis -Kyoto classification of gastritis. Nihon Shokakibyo Gakkai Zasshi. 2015; 112: 982–93 (in Japanese, Abstract in English).
16 Dixon MF, Genta RM, Yardley JH, Correa P. Classification and grading of gastritis: the updated Sydney System. International workshop on the histopathology of gastritis, Houston 1994. Am. J. Surg. Pathol. 1996; 20: 1161–81.
17 Muto M, Yao K, Kaise M et al. Magnifying endoscopy simple diagnostic algorithm for early gastric cancer (MESDA-G). Dig. Endosc. 2016; 28: 379–93.
18 Japanese Gastric Cancer Association. Japanese classification of gastric carcinoma: 3rd English edition. Gastric Cancer. 2011; 14:1 01–12.
19 Japanese Gastric Cancer Association. Japanese gastric cancer treatment guideline 2010: 3rd English edition. Gastric Cancer. 2011; 14: 113–23.
20 Schlemper RJ, Riddell RH, Kato Y et al. The Vienna classification of gastrointestinal epithelial neoplasia. Gut. 2000; 47: 251–5.
21 Lauwers G, Carneiro F, Graham D et al. WHO Classification of Tumors of the Digestive System, 4th edn. JARC: Lyon, 2010.
22 Shiota S, Murakami K, Suzuki R, Fujioka T, Yamaoka Y. Helicobacter pylori infection in Japan. Expert Rev. Gastroenterol. Hepatol. 2013; 7: 35–40.
23 Ito M, Tanaka S, Takata S et al. Morphological changes in human gastric tumours after eradication therapy of Helicobacter pylori in a short-term follow-up. Aliment. Pharmacol. Ther. 2005; 21: 559–66.
24 Owen DA. Lamina propria, Stomach. In: Histology for the Pathologists, 5th edn. Philadelphia: Wolters Kluwer, 2020; 607.
25 Cancer Genome Atlas Research Network. Comprehensive molecular characterization of gastric adenocarcinoma. Nature. 2014; 513: 202–9.
26 Funakoshi T, Miyamoto S, Kakiuchi N et al. Genetic analysis of a case of Helicobacter pylori-uninfected intramucosal gastric cancer in a family with hereditary diffuse gastric cancer. Gastric Cancer. 2019; 22: 892–8.
27 Nomura R, Saito T, Mitoni H et al. GNAS mutation as an alternative mechanism of activation of the Wnt /β-catenin signaling pathway in gastric adenocarcinoma of the fundic gland type. Hum. Pathol. 2014; 45: 2488–96.
28 Sugimoto R, Sugai T, Habano W et al. Clinicopathological and molecular alterations in early gastric cancers with the microsatellite instability-high phenotype. Int. J. Cancer. 2016; 138: 1689–97.
29 Fukayama M. Epstein-Barr virus and gastric carcinoma. Pathol. Int. 2010; 60: 337–50.
30 Okano A, Kato S, Ohana M. Helicobacter pylori-negative gastric cancer: Advanced-stage undifferentiated adenocarcinoma located in the pyloric gland area. Clin J Gastroenterol. 2017; 10: 13–17.
31 Horiiuch Y, Fujisaki J, Ishizuka N et al. Study on clinical factors involved in Helicobacter pylori-uninfected, undifferentiated-type early gastric cancer. Digestion. 2017; 96: 213–19.
32 Matsuo T, Ito M, Takata S, Tanaka S, Yoshihara M, Chayama K. Low prevalence of Helicobacter pylori-negative gastric cancer among Japanese. Helicobacter. 2011; 16: 415–19.
33 Bernstein H, Bernstein C, Payne CM, Dvorakova K, Garewal H. Bile acids as carcinogens in human gastrointestinal cancers. Mutat. Res. 2005; 589: 47–65.