Gastric neuroendocrine tumors: management and challenges

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Abstract: Gastric neuroendocrine tumors derive from enterochromaffin-like cells in the stomach mucosa. Based on histologic, serologic, and endoscopic findings, they may be further differentiated into types I, II, and III, with varying degrees of aggressiveness. In this article, diagnostic and classification strategies are reviewed, as are endoscopic, systemic, and surgical modalities for management. A multidisciplinary approach is advocated to provide the most effective patient care.

Keywords: neuroendocrine, tumor, carcinoid, gastrin, stomach

Background

First reported by Askanazy in 1923, gastric neuroendocrine tumors (NETs) comprise approximately 1.8% of all gastric tumors 1 and develop from enterochromaffin-like (ECL) cells in the gastric mucosa. Gastric NETs are also referred to as gastric carcinoids. The term karzinoide, or carcinoma-like, was coined by Oberndorfer in 1907 to describe this class of tumors, which behaves in a relatively benign fashion compared to adenocarcinomas. 2 Our understanding of NETs as a whole has evolved over time, and the World Health Organization (WHO) classification system now employs the term neuroendocrine tumor instead of carcinoid. As such, gastric NETs are part of a broad category of gastroenteropancreatic-neuroendocrine tumors (GEPNETs), which encompass well-differentiated NETs arising from the gastrointestinal tract. In an effort to standardize the system and assist clinicians to accurately predict clinical outcomes, GEPNETs are graded histologically based on mitotic count and/or Ki67 index. The 2010 WHO histologic classification describes well-differentiated NETs as having a Ki67 index <3% and <2 mitoses per 10 high-power fields (HPFs) (G1), moderately differentiated NETs with a Ki67 index 3–20% or 2–20 mitoses per 10 HPFs (G2), and poorly differentiated NETs with a Ki67 index >20% or >20 mitoses per 10 HPFs (G3). 3,4

The prevalence of gastric NETs is difficult to establish due to a lack of uniform data collection from cancer registries worldwide. A study published in 2015 analyzing national cancer registries in 10 European countries, the US, and Japan determined the prevalence of gastric NETs per 10,000 population to be 0.32 in Europe, 0.17 in the US, and 0.05 in Japan. 5 The authors of this study did note that their values may be underestimations due to a tendency of cancer registries to reflect the aggressive tumors that require treatment as opposed to benign tumors.

Clinically, gastric NETs are categorized into types I, II, and III (Table 1). The basis for these subtypes is rooted in gastric pathophysiology. 6 Gastrin is produced by...
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Risk of metastasis is slightly higher in type II gastric NETs. Small, asymptomatic, and well-differentiated tumors. The potential to metastasize varies from 2–5% to 10–30% to 50–100%.

Serum gastrin
- High
- High
- Normal

Associated disease
- Atrophic gastritis
- Gastrinoma
- None

Type II gastric NETs represent 5–10% of gastric NETs. Like type I disease, they are also typically multiple, small, asymptomatic, and well-differentiated tumors. The risk of metastasis is slightly higher in type II gastric NETs compared to type I lesions but is still low. Overall, tumor invasion beyond the submucosa into the muscularis propria or lymph node and liver involvement occurs in 5–12% of cases.

Type II gastric NETs are also associated with hypergastrinemia, but the feature that distinguishes them from type I tumors is their association with ZES and multiple endocrine neoplasia type I (MEN1) syndrome, which are hypersecretory states. Up to 30–50% of patients with MEN1 syndrome will develop gastric NETs, especially if ZES is present. Type II gastric NETs unsurprisingly have a high rate of loss of heterozygosity at the MEN1 gene locus of 75–100%. Interestingly, 17–73% of type I gastric NETs and 25–50% of type III gastric NETs carry the same mutation, suggesting that MEN1 mutations cannot be used to distinguish among the 3 types of gastric NETs.

Type II gastric NETs are typically nonfunctioning tumors, and symptoms upon presentation are usually secondary to peptic ulcer disease and ZES. While other genetic syndromes such as type 1 neurofibromatosis, von Hippel–Landau disease, and tuberous sclerosis complex are associated with NETs, gastric involvement is rare.

Approximately 10–15% of gastric NETs are categorized as type III tumors. These lesions typically exist as solitary larger tumors, often >2 cm in size. Histopathologically, they vary from well- to poorly differentiated tumors, and their overall prognosis is relatively poor. Upon initial presentation, the incidence of concurrent metastasis is >50%, typically with hepatic involvement. Unlike type I and II gastric NETs, type III lesions do not have any associated predisposing conditions. Fasting gastrin levels are normal without any G cell or ECL cell hyperplasia. While type I and II gastric NETs tend to be nonfunctional tumors, type III NETs with hepatic metastases may be associated with carcinoid syndrome, although this occurrence is rare.

**Diagnosis and evaluation**

Studies used to diagnose and differentiate gastric NET disease may be broadly divided into endoscopic, biochemical,
histopathologic, and imaging studies. As symptoms related to carcinoid syndrome are rare, endoscopy is the gold standard for diagnosing gastric NETs. During esophagogastroduodenoscopy (EGD), aspiration of gastric fluid may be performed to assess a gastric pH, though this value can be artificially elevated by PPI use. Endoscopic ultrasound (EUS) may be performed on larger lesions to evaluate the depth of gastric wall involvement and for lymphadenopathy. In type I disease, tumors are often found in the gastric fundus and described as subcentimeter and multifocal. Biopsies of the rest of the stomach may detect atrophic gastritis or Helicobacter pylori (HP). Type II disease may also appear as multifocal subcentimeter polypoid lesions, but there may be concurrent peptic ulcer disease in the setting of a hypersecretory state. The gastric pH is generally high (>4) in type I disease and low (<2) in type II tumors. Tumors in type III disease typically appear as solitary larger lesions that may be ulcerated with hemorrhage. Gastric pH is typically normal, and there is an association with HP infection but not atrophic gastritis or peptic ulcer disease.

Biochemical testing is often performed to differentiate between the subtypes of gastric NETs. Gastrin levels are elevated in type I and II gastric NETs, while they are normal in type III. However, concurrent or recent PPI use may elevate gastrin levels; therefore, PPIs should be withdrawn at least 1 week before obtaining accurate fasting gastrin levels. In cases of ZES, abrupt PPI withdrawal can lead to serious consequences, including gastrointestinal perforation, and a careful PPI wean may be recommended when this entity is suspected. During weaning, PPIs are replaced by histamine H2-receptor antagonists such as ranitidine 1–2 weeks before formal testing. The H2-receptor antagonist should be dosed as 450–750 mg every 6 hours and then stopped in the final 24–30 hours before testing. Antacids may be taken as needed until the midnight before testing. Patients should be given explicit instructions to seek medical attention during this period should they develop diarrhea, nausea, vomiting, or severe abdominal pain. Further differentiation between type I and II gastric NETs may be established by incorporation of the gastric pH as gastrin level alone cannot determine type I or type II disease. Finally, given the association of chronic atrophic gastritis with type I gastric NETs, a low serum vitamin b12 and positive parietal cell and/or intrinsic factor antibodies may be found. While genetic testing for MEN1 may be considered when there is suspicion of type II disease, it is not considered a diagnostic tool for gastric NETs. A small case series of type II gastric NET in the setting of a confirmed germline MEN1 mutation has recommended screening for parathyroid and pituitary tumors.

Conventional cross-sectional imaging modalities such as computed tomography (CT) and magnetic resonance imaging (MRI) are commonly used to assess for disease spread and for staging of advanced lesions. However, their value is limited in type I and II diseases as these are frequently characterized by small tumors. Similarly, functional imaging with somatostatin receptor scintigraphy, fluoro-D-glucose positron emission tomography, Ga-DOTATOC PET/CT, or MRI is of limited use in smaller tumors. That said, 68Ga-DOTATOC may be helpful in localizing occult gastrinomas in the setting of type II gastric NETs. In type III disease where tumors are larger with a tendency to metastasize, conventional CT or MRI studies are helpful for tumor detection and staging. Larger studies are needed to determine the precise role of functional imaging in gastric NETs, though early data on Ga-DOTATOC imaging are promising.20,21

Endoscopic management

Endoscopic management is predominantly utilized in type I and localized type II disease. In type III disease, endoscopy plays a smaller role given the higher likelihood of concurrent metastatic disease. The lack of consensus surrounding the management of gastric NETs is highlighted by discrepancies between the published guidelines of the National Cancer Comprehensive Network (NCCN) and European Neuroendocrine Tumor Society (ENETS).22,23 Beyond a conventional EGD, EUS may also be performed to evaluate tumor depth (Figure 1C), though the cutoff size to prompt this procedure is not yet defined. Generally, lesions smaller than 8–10 mm are not amenable to fine needle biopsy during EUS and lesions smaller than 5 mm may be challenging to visualize endoscopically. The potential benefits of EUS include determination of any muscularis propria invasion that may preclude a complete endoscopic resection and evaluation of lymphadenopathy. The NCCN recommends EUS in type III disease to evaluate for lymphadenopathy and tumor depth, while an EUS is recommended in type I and II disease as clinically indicated. The ENETS guidelines mention the role of EUS for staging tumors but are less specific with respect to subtype of gastric NETs, and emphasize the need for further investigation to determine the cutoff size for endosonographic examinations of tumors.

In type I gastric NETs, the cutoff size for a tumor to harbor metastatic potential is thought to be 10 mm.24 Assuming there is no muscularis propria or lymph node involvement, endoscopic resection of lesions >10 mm is favored in the ENETS guidelines as this is considered the least invasive approach.23 The 2017 NCCN guidelines are less specific and
recommend resection of only “prominent” tumors. There is no consensus regarding the treatment of subcentimeter lesions, though surveillance annually or every 2 years is favored by both guidelines. The ENETS guidelines recommend performing an EGD with polyp resection or biopsy every 1–2 years. While more aggressive approaches that entail resecting all visible lesions or selectively removing lesions that are at least 5 mm in size have been described, there have not been any studies comparing these strategies. Recognizing a lack of data from large studies and published series demonstrating tumor recurrence rates of 18–63.6%, a prudent approach would be to continue endoscopic surveillance even if resection is performed.

EUS has a larger role in type II than type I gastric NETs in order to rule out pancreatic lesions and determine if the gastric tumor is a primary or secondary lesion. Endoscopic resection of a primary type II gastric NET is a feasible primary treatment option according to both NCCN and ENETS guidelines. However, if there are duodenal or pancreatic lesions, a patient-individualized multidisciplinary approach at a neuroendocrine center of expertise may be necessary.

In cases of type III disease in which metastasis has been ruled out, an EUS may assist in evaluating for lymphadenopathy. If negative, endoscopic or surgical resection is recommended, although NCCN guidelines favor endoscopic resection exclusively for superficial subcentimeter lesions with “low-grade” histology.

If indicated, endoscopic resection is chiefly accomplished by 2 techniques: endoscopic mucosal resection (EMR) and endoscopic submucosal dissection (ESD). EMR typically involves injecting a saline solution into the submucosal space underneath the lesion to achieve proper lifting and delineation of the lesion’s borders, followed by resection with a cautered snare. ESD requires specialized endoscopic instruments to penetrate a lesion’s surrounding submucosal space followed by cautery to complete an en bloc resection. This technique is effective for lesions not thought amenable to EMR and is practiced chiefly in Asia and is less available in Western countries. The literature on ESD for gastric NETs is scant as most publications focus on use of ESD for resection of rectal lesions. Nevertheless, it is considered safe and associated with high rates of complete resection of gastric lesions if done at high-volume centers. EMR and ESD are both associated with a low, but tangible, risk of perforation or bleeding and thus should be performed by an experienced endoscopist. One study from Japan reported on the use of ESD for 42 patients with gastrointestinal NETs (of which most were rectal); there were 2 (5%) cases of postoperative bleeding and 2 (5%) perforations.

**Medical management**

In type I and II gastric NETs, somatostatin analogs (SSAs) have been shown to decrease levels of gastrin and have an antiproliferative effect on ECL cells. Limited studies, including a few small prospective studies, have demonstrated regression or complete disappearance of tumors and marked decrease in serum gastrin, lasting up to several years. SSAs (e.g. octreotide and lanreotide) can be considered in cases in which endoscopic resection is not feasible due to extensive multifocal disease, or submucosal/lymph node involvement, as well as recurrent disease after repeated endoscopic resection. However, small studies have shown that this antiproliferative effect is not durable, since a rebound in serum gastrin to pretreatment levels and tumor recurrence are possible after cessation of therapy. A repeat cycle of SSA after disease recurrence may again induce tumor regression. Given the lack of randomized clinical trials with SSAs for this indication, their high cost, and the comparatively benign course of type I/II gastric NETs with endoscopic resection, ENETS recommends their use be restricted to cases of metastatic type I gastric NETs with proven somatostatin receptor 2 (SSTR2) expression and a low Ki67 index, and NCCN guidelines recommend consideration of SSAs only for type II gastric NET cases in which the primary tumor has not been resected (i.e. to control gastrin secretion), in conjunction with endoscopic surveillance and resection of prominent tumors.

As in type I and II gastric NETs, systemic therapy in type III is warranted only in locoregional disease that is unresectable or metastatic. Management in these cases is identical to that of all unresectable or metastatic gastrointestinal NETs and is based on a multitude of factors, including the patient’s symptoms, tumor grade, tumor burden, and progression of disease during imaging-guided surveillance.

Systemic options for well-differentiated NETs typically include SSAs and everolimus; interferon alfa-2b is less commonly employed. Treatment is often multimodal, particularly in cases of liver metastases, for which liver-directed therapies such as hepatic arterial embolization are commonly employed. Peptide receptor radionuclide therapy is an emerging tool for somatostatin receptor-expressing tumors, but is not approved for use in the US. The role of systemic chemotherapy in the treatment of well-differentiated gastric
NETs is ill-defined. Unfortunately, there are no randomized clinical trials or prospective data comparing the efficacy of one therapy to another, or determining the most efficacious sequence of therapies for well-differentiated NETs. Platinum-based combination chemotherapy is typically reserved for poorly differentiated neuroendocrine carcinomas.22,49-51

Surgical management
Type I gastric NETs are indolent but frequently present with multifocal primary tumors and often recur.6,52 However, lymph node metastases are rare (~10%), and disease-specific mortality is exceedingly uncommon.53 Therefore, careful patient selection for surgery is of paramount importance. Surgical resection of type I gastric NETs may be considered when primary tumors are not amenable to endoscopic resection due to the depth of larger size lesions involving the muscularis propria. Surgery may also be warranted when there is a concern for regional lymph node involvement based on cross-sectional imaging or EUS. Finally, surgery clearly has a role when there is biopsy-proven or suspected coexisting gastric adenocarcinoma arising in the setting of chronic atrophic gastritis.53,54

Surgery for type I gastric NETs not amenable to endoscopic resection may entail resection of the primary tumors combined with antrectomy to remove the source of gastrin production.53 A regional lymphadenectomy should be done for staging. After antrectomy for type I gastric NETs, gastrin levels typically normalize, and regression of primary tumors occurs in most patients. Recurrent disease may result if a complete antrectomy is not performed and gastrin-producing cells are left behind.53 Disease may also recur if gastric NETs develop gastrin-independent growth. Therefore, endoscopic surveillance should be considered after antrectomy to look for recurrent NETs and to screen for gastric adenocarcinoma, which can arise in patients with chronic atrophic gastritis.54

Type II gastric NETs arise almost exclusively in patients with a gastrin-producing NET of the duodenum or pancreas (gastrinoma, ZES) and MEN1.6,55,56 In patients with ZES/MEN1, 13–37% develop type 2 gastric NETs.56 Most (80–90%) of type II gastric NETs are not invasive, and the primary gastrinoma should be resected, if feasible.22,56 If resection of the primary gastrinoma is not done, type II gastric NETs can often be managed endoscopically and/or with SSAs. In patients with long-standing hypergastrinemia in the setting of ZES/MEN1, advanced serotonin-producing type II gastric NETs have been observed, requiring total gastrectomy and hepatectomy to control extensive local tumor burden (gastric obstruction) and the carcinoid syndrome, respectively.56

Type III gastric NETs may be well differentiated (G1, G2) or poorly differentiated (G3).57 Well-differentiated type III gastric NETs are frequently invasive and metastasize to regional lymph nodes; therefore, patients are typically managed with an oncologic resection of the primary tumor and regional lymph nodes. In carefully selected patients with type III gastric NETs with low-risk features (<2 cm, confined to submucosal layer, no lymphovascular invasion), favorable results with endoscopic resection (EMR, ESD) have been reported in South Korea.58 The outcome of patients with poorly differentiated type III gastric NETs is extremely poor, resembling that of patients with small-cell lung cancer; therefore, the role of surgery is quite limited.59 A reasonable approach is to treat patients with poorly differentiated type III gastric NETs with upfront platinum-based systemic therapy, and perhaps consider surgical resection in only those who have locoregional disease.

Conclusion
Gastric NETs are clinically categorized into types I, II, and III. A combination of fasting serum gastrin levels, gastric pH, and endoscopic and imaging findings is necessary to differentiate among the types of gastric NETs as their prognoses all vary. While some gastric NETs are indolent and can be managed by endoscopic resection and surveillance, refractory disease may require treatment with an SSA. Liver-directed therapy and/or systemic therapy with everolimus or chemotherapy is typically reserved for advanced disease. Surgical resection is reserved for type I and type II NETs that are endoscopically unresectable, or carefully selected patients with well-differentiated type III disease. While the NCCN and ENETS provide guidelines, the diagnosis and management of gastric NETs remains challenging in some areas, and a multidisciplinary approach is preferred to ensure consideration of all therapeutic options.

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References


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