ENETS Consensus Guidelines for High-Grade Gastroenteropancreatic Neuroendocrine Tumors and Neuroendocrine Carcinomas

R. Garcia-Carbonero\textsuperscript{a} H. Sorbye\textsuperscript{b} E. Baudin\textsuperscript{c} E. Raymond\textsuperscript{d} B. Wiedenmann\textsuperscript{e} B. Niederle\textsuperscript{f} E. Sedlackova\textsuperscript{g} C. Toumpanakis\textsuperscript{h} M. Anlauf\textsuperscript{i} J.B. Cwikla\textsuperscript{j} M. Caplin\textsuperscript{k} D. O'Toole\textsuperscript{l} A. Perren\textsuperscript{m}

all other Vienna Consensus Conference participants

\textsuperscript{a}Medical Oncology Department, Hospital Universitario Doce de Octubre, Madrid, Spain; \textsuperscript{b}Department of Oncology, Haukeland University Hospital, Bergen, Norway; \textsuperscript{c}Institut Gustave Roussy, Villejuif, and \textsuperscript{d}Oncologie Médicale, Hôpitaux Universitaires Paris Nord Val de Seine, Paris, France; \textsuperscript{e}Department of Hepatology and Gastroenterology, Campus Virchow Klinikum, Charité Universitätsmedizin Berlin, Berlin, Germany; \textsuperscript{f}Department of Surgery, Medical University of Vienna, Vienna, Austria; \textsuperscript{g}Department of Oncology, First Faculty of Medicine and General Teaching Hospital, Prague, Czech Republic; \textsuperscript{h}Neuroendocrine Tumour Unit, Royal Free Hospital, London, UK; \textsuperscript{i}Institut für Pathologie und Zytologie, St. Vincenz Krankenhaus, Limburg, Germany; \textsuperscript{j}Department of Radiology, Faculty of Medical Sciences, University of Warmia and Mazury, Olsztyn, Poland; \textsuperscript{k}Neuroendocrine Tumour Unit, Royal Free Hospital, London, UK; \textsuperscript{l}NET Centre, St. Vincent’s University and Department of Clinical Medicine, St. James Hospital and Trinity College, Dublin, Ireland; \textsuperscript{m}Institute of Pathology, University of Bern, Bern, Switzerland

Introduction

Neuroendocrine carcinomas (NEC) are rare in the gastrointestinal (GI) tract, whereas they are frequent in the form of small cell carcinoma (SCLC) in the lung. Therefore, most of the suggested guidelines arise from analogy to SCLC. As for other extrapulmonary primary tumor locations, published data on NEC of the GI tract are scarce. This guideline encompasses all WHO grade 3 (G3) gastroenteropancreatic (GEP) neoplasms; however, in the future, G3 neuroendocrine neoplasms (NEN) will probably be separated according to differentiation, as explained below, and potentially according to organ of origin, such as for well-differentiated neuroendocrine tumors (NET) G1/G2.

According to the WHO classification of 2010, NEC are defined as poorly differentiated NEN with Ki-67 >20% and hence G3. Increasing evidence suggests that G3 NEN are not a homogenous entity and can be further subclassified into biologically relevant subgroups. A separation based on the proliferative index (Ki-67 >55%) was shown to have clinical implications regarding response to chemotherapy and prognosis: NEC with Ki-67 >55% responded better to platinum-based chemotherapy and, nevertheless, had a 4 months’ shorter median survival than G3 NEN in the lower proliferative range (20–55%) [1]. More recent publications show that morphological differentiation and Ki-67 are able to separate prognostic groups among G3 cases, and therefore a separation of well-differentiated G3 NET from poorly differentiated G3 NEC is emerging [2–4]. The exact criteria need to be defined both on the morphological and on the molecular level. The spectrum of mutations of well-differentiated...
pancreatic NET is different from that of pancreatic NEC [5], suggesting different ways of tumorigenesis. However, to date there are no solid data that adequately address the implications of these observations in terms of treatment effect of the different available regimens.

### Epidemiology

The GEP tract is the most common site of extrapulmonary NEC, accounting for 35–55% of all NEC originating from the lung. Only about 5% of all GI NEN have Ki-67 >20% [6, 7]. This frequency might differ by organ, with about 7% in the pancreas [7] and up to 40% in the colon [8]. GEP NEC are therefore very rare neoplasms representing <1% of all GI malignancies. Up to 85% have metastases at the time of diagnosis (65% distant) [1, 6]. Metastases are most frequently found in the liver (70%) followed by the lung (15%), bone (15%) and brain (4%) [1]. No gender difference has been identified. The mean age at diagnosis is 60 years [1].

### Clinicopathological Features

As the great majority of these tumors are not associated with a hormonal syndrome (<5%), and more than two thirds of all patients present with advanced disease, clinical presentation is dominated by tumor-derived site-specific symptoms and the constitutional syndrome characteristics of advanced cancer (anorexia, weight loss and fatigue). Depending upon tumor location, a wide variety of symptoms may occur. The neuroendocrine nature of these tumors is generally not suspected from the clinical presentation, although as in SCLC paraneoplastic syndromes may occur in some patients (i.e., Cushing or Inappropriate ADH Secretion Syndromes). A detailed anamnesis and physical examination are fundamental to appropriately guide diagnostic procedures.

According to the WHO 2010 classification, NEC are poorly differentiated highly aggressive neoplasms, sometimes with organoid features, marked nuclear atypia and multifocal necrosis [9]. A diffuse expression of neuroendocrine markers (diffuse for synaptophysin, focal for chromogranin A, and the latter may be absent) separates the entity pathologically from poorly differentiated carcinoma.

The grading introduced by ENETS in 2006 [10] of NEC is by definition G3, either based on a proliferation index >20% or >20 mitoses in 10/HPF. This proposition has been adopted by the WHO classification and was shown repeatedly to be clinically applicable in predicting a very aggressive subset of NEN [7].

### Prognosis and Survival

Survival is poor in NEC, ranging from 38 months for patients with localized disease to 5 months in the metastatic setting according to the SEER population registry data, which involved 2,546 patients diagnosed with GI NEC from 1973 to 2012 in the USA [11]. Median survival in the metastatic setting may be as short as 1 month for patients receiving only best supportive care, up to 12–19 months for those treated with best available therapy [1, 12]. Only 5% of all patients are long-time survivors [7]. Progression-free survival after cisplatin-based chemotherapy and overall survival differs according to the location of the primary tumor, with poorer reported outcomes in esophageal, colonic and rectal NEC compared to gastric and pancreatic ones in some large European series [1]. In contrast, survival of pancreatic NEC was poorer in Japanese patients [13]. A poor performance status, high proliferation rate, elevated baseline lactate dehydrogenase (LDH) and thrombocytosis are other factors that have also been associated with a worse prognosis.

### Diagnostic Procedures

#### Biochemical Tests

Plasma chromogranin A may be elevated in up to two thirds of patients with advanced NEC [1], although the levels are generally lower than those observed in well-differentiated tumors [14, 15]. In contrast, the levels of other tumor markers such as neuron-specific enolase (NSE) are higher in poorly differentiated tumors than in NET, and they are significantly associated with survival. However, the role of circulating tumor markers to predict and monitor outcome has not been properly assessed in extrapulmonary NEC. Screening for other hormonal markers is not justified unless clinically indicated.

#### Endoscopic and Imaging Procedures

Endoscopic examination of the primary tumor site is recommended, which is also useful to obtain a biopsy for histological diagnosis. If this is not feasible, endoscopic ultrasound-guided or percutaneous procedures can be useful. Once the histological diagnosis of a G3 NEC has been confirmed, complete staging using whole-body CT
scan or MRI should be performed to assess the extent of
disease and to design the most appropriate therapeutic
strategy. A lung primary should be reasonably excluded
(negative imaging studies of the lung). FDG-PET may be
useful if radical surgery is being pursued or if clarification
of equivocal findings on conventional imaging may
change the therapeutic approach. Radiolabeled soma-
tostatin analogue scans are not routinely recommended,
as poorly differentiated tumors generally do not express
somatostatin receptors. However, data from large series
indicate positive SRI findings in a substantial proportion
of patients with certain primary tumors (up to 45% of
pancreatic NEC), particularly those with proliferative in-
dexes in the low range of G3, and may differ by histologi-
cal subtype (45% of small cell vs. 32% of large cell NEC)
[1]. In the absence of neurological symptoms, brain CT
or MRI are not recommended, as the incidence of brain
metastasis in extrapulmonary NEC is rather low (<5%)
[1]. Bone scans are neither indicated if there is no clinical
or biochemical suspicion of bone metastasis. In the pre-
se resemblance of elevated LDH, peripheral blood leukoerythro-
blastosis or thrombocytopenia, a bone marrow biopsy
may be considered.

Minimal Consensus Statement

Clinical signs and symptoms should guide the appropriate
diagnostic procedures (summarized in fig. 1). Chromogranin A
and NSE testing is not mandatory, although they may be useful
if elevated at diagnosis. A proper assessment of their utility in
extrapulmonary NEC is, however, pending. Other hormone tests
are not routinely recommended.

A minimal diagnostic workup should include site-specific
endoscopic assessment with tumor biopsy, and whole-body CT
scan (and/or MRI) for tumor staging. In patients with metastatic
disease, an ultrasound-guided percutaneous biopsy may be per-
fomed if feasible. Somatostatin receptor scintigraphy is not rou-
tinely indicated but may be considered in tumors with prolifera-
tive indexes in the low range of G3 (Ki-67 <55%). Bone scans or
brain imaging (CT or MRI) should not be performed in the ab-
scence of site-specific symptoms. FDG-PET may be considered in
patients in whom radical surgery is being pursued or if clarifica-
tion of equivocal findings on conventional imaging may change
the therapeutic approach. FDG-PET may be useful in resectable
cases for whole body assessment.

Histopathology and Genetics of Poorly
Differentiated NEC

Histopathologically, NEC show a neuroendocrine
phenotype by immunohistochemistry, in large cell NEC
a positivity for synaptophysin is mandatory, chromo-
granin A staining is variable and may be weak or absent.
Rarely, both markers may be negative in small cell NEC
(<5% [1]). Other neuroendocrine markers such as NSE or
CD56 are less specific and must be used with caution. Ki-
67 is by definition >20% [10] and in half of the cases, it is
>55% [1]. Punctate or geographic necrosis is frequent.
Reporting of the immunohistochemical results above as
well as the proliferative index by mitosis is essential. So-
atostatin receptor 2A (Sstr 2a) immunohistochemistry
is optional [16, 17]. Over 90% of G3 NEC do not produce
hormones [17].

In the setting of a carcinoma of unknown primary, the
expression of transcription factors such as Ttf1, Cdx-1 or
Isl1 cannot be used to help localize the site of the primary
tumor [18].

Care must be taken to differentiate NEC from poorly
differentiated adenocarcinoma, especially in certain or-
gans such as the pancreas, where a differential diagnosis
with acinic cell carcinoma may be particularly challeng-
ing [19]. NEC are separated into large cell and small cell
types; however, no clear clinicopathological differences
between the two types have been shown for the pancreas
[19].

Pancreatic NEC show a genetic profile different from
NET with frequent mutations in p53 and RB [5] and a
much higher mutation rate (in review), similar to pulmo-
ary small cell carcinoma. Furthermore, up to 40% of all
NEC present a minor component of adenocarcinoma
(colon [20], stomach [21]) or squamous cell carcinoma
(esophagus, anus). If the non-endocrine component ex-
cedes 30%, the neoplasm is classified as mixed adeno-
endocrine carcinoma. Differentiation together with
proliferation and mutation spectrum will be important in
discriminating G3 NET from G3 NEC in the future [2–4].

Minimal Consensus Statement

A routine pathological report should include morphology
(large cell vs. small cell and differentiation), staining for chromo-
granin A and synaptophysin and Ki-67 estimate or/mitotic
count.

Treatment

Evidence to support treatment recommendations for
GEP G3 NEC is scarce and derives from limited retro-
spective series and very few small non-controlled clinical
trials. Most investigators, therefore, treat this entity
in analogy to the much more common SCLC due to

Garcia-Carbonero et al.
their histological and clinical resemblance. Bearing these caveats in mind, guidance is hereby provided (fig. 2). Nevertheless, generating prospective and preferably controlled data is greatly needed and encouraged in this setting.

**Surgery**

Curative surgery is usually attempted in localized disease, although retrospective series indicate that it is rarely curative as a sole therapeutic modality [22]. Given the high relapse rate observed following radical surgery, most clinicians would advocate platinum-based adjuvant therapy in this setting. Data reported by Casas et al. of a large series of esophageal small cell carcinomas support this approach [23]. In this study, survival was 20 months for patients who received systemic chemotherapy in addition to local treatment versus only 5 months for those who were treated with local therapy only, and the type of treatment was found to be an independent prognostic factor in multivariate analysis. Some authors propose neoadjuvant chemotherapy followed by definitive surgery, although data to support this approach are scarce [24]. In patients with important comorbidities or where the tumor’s anatomical site makes surgical resection not advisable due to high morbidity (i.e., esophagus), a definitive course of radiotherapy and chemotherapy is a reasonable treatment strategy.

In the context of advanced metastatic disease, debulking or cytoreductive surgery and surgical resection of metastasis are not recommended. Other ablative strategies of liver metastasis (i.e., radiofrequency ablation, TACE) are also discouraged.

**Medical Therapy**

Chemotherapy is an essential part of the multimodal approach for localized NEC and the mainstay of care in advanced disease. Survival of patients with metastatic NEC treated with chemotherapy varies widely (from 7 to 19 months) but shows a substantial improvement over that reported for patients that receive only best supportive care (1 month). No randomized studies, however, have properly addressed the magnitude of this effect so far. Swift referral for consideration of palliative chemotherapy is recommended as performance status deterioration may occur rapidly and preclude further therapy. Based on their established role in metastatic SCLC, cisplatin and etoposide (EP) have been one of the most widely used regimens in GEP NEC (table 1) [1, 13, 25–29], with response rates in the largest most recent series of ~30% and median survival of around one year. In one of the largest series published to date, Sørbye et al. [1] observed that Ki-67 was significantly associated with response to chemotherapy. Indeed, patients with Ki-67 values >55% had a greater response rate (42 vs. 15%) al-

---

**Fig. 1.** Diagnostic algorithm for NEC and G3 NET.
though poorer survival (10 vs. 14 months) than patients with Ki-67 values <55%. Other negative prognostic factors in this study were a poor performance status, primary colorectal tumors and elevated platelets or LDH levels, which were all associated with decreased survival.

Alternative regimens substituting carboplatin for cisplatin, or irinotecan for EP, have been validated in SCLC and seem at least equivalent in terms of efficacy in limited series of GEP NEC (table 1) [13, 30–35], with different toxicity profiles. In the context of advanced stage SCLC, a randomized study conducted in Japan demonstrated that the combination of irinotecan and cisplatin (IP) was associated with improved overall survival as compared to the standard cisplatin and EP combination [36]. Two subsequent randomized Western trials, however, failed to confirm this superiority. Both regimens produced comparable efficacy, with less hematological and greater GI toxicity with the IP combination (particularly diarrhea and vomiting) [37, 38]. Consistent with these findings, large retrospective data of systemic chemotherapy for advanced GEP NEC from 23 Japanese institutions documented that the IP regimen was associated with greater response rates (50 vs. 28%) and survival (13 vs. 7 months) than the EP regimen, and this difference was more remarkable in hepatobiliopancreatic NEC. Prognostic factors in this study included primary tumor site (those with hepatobiliopancreatic primaries having the worst prognosis) and elevated baseline LDH levels, whereas treatment schedule was not an independent predictive factor for survival. Three-drug regimens such as cisplatin, EP and paclitaxel do not seem to substantially improve efficacy and are significantly more toxic [39].

Evidence for salvage therapy in patients progressing on first-line platinum-based regimens is very limited (table 1) [1, 13, 40–45]. Overall, response rates are lower (18% in the NORDIC NEC study), although small series have documented response rates of 23–40% with oxaliplatin-based (XELOX, FOLFOX) or irinotecan-based (FOLFIRI, IP) regimens. Welin et al. reported a 33% response rate with temozolomide, alone or in combination with capecitabine and bevacizumab, in a cohort of 25 patients with poorly differentiated NEC (17 of GEP origin) [44]. A Ki-67 index <60% was predictive for response to treatment and survival. In contrast, no responses were observed in another series of 28 NEC cases treated with temozolomide monotherapy [45]. Re-treatment with platinum/EP may also be considered in patients that achieve good durable responses upfront and progress after a treatment break of at least 3 months, provided no cumulative toxicity (i.e., neurotoxicity, ototoxicity) precludes further treatment with platinum.
agents. Other agents tested include amrubicin, S-1 or taxanes (table 1).

**Other Treatment Options (Radiotherapy, PRRT)**

In contrast to recommendations for patients with limited-stage SCLC, prophylactic cranial irradiation is not indicated in patients with successfully treated localized GEP NEC, as the incidence of brain metastasis in patients with extrapulmonary NEC is rather low. Palliative radiotherapy may be considered for localized bone metastasis to control pain or to prevent skeletal complications.

Although a subgroup of NEC expresses somatostatin receptors, there are no data to support the use of somatostatin analogs in this context. Some case reports have communicated long-lasting responses to peptide receptor radionucleotide therapy (PRRT) in NEC with high expression of somatostatin receptors, but this therapeutic strategy is generally not successful in the majority of G3 tumors [46].

---

**Table 1. Series of patients with advanced G3 NEC of the GI tract treated with chemotherapy**

<table>
<thead>
<tr>
<th>First author</th>
<th>Patients, n</th>
<th>Primary site</th>
<th>Chemotherapy regimen</th>
<th>RR, %</th>
<th>Survival, months</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>First-line therapy</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mitry [26]</td>
<td>41</td>
<td>GEP (20), lung (10), H&amp;N (4), UKP (7)</td>
<td>Cisplatin/EP</td>
<td>42</td>
<td>15</td>
</tr>
<tr>
<td>Deutschbein [28]</td>
<td>18</td>
<td>G3 NEC (primary NR)</td>
<td>Cisplatin/EP +/– paclitaxel</td>
<td>17</td>
<td>NR</td>
</tr>
<tr>
<td>Iwasa [27]</td>
<td>21</td>
<td>Hepatobiliar and pancreas</td>
<td>Cisplatin/EP</td>
<td>14</td>
<td>6</td>
</tr>
<tr>
<td>Patta [29]</td>
<td>8</td>
<td>Colorectal</td>
<td>Cisplatin/EP</td>
<td>63</td>
<td>10</td>
</tr>
<tr>
<td>Sørbye [1]</td>
<td>252</td>
<td>GEP (69%), UKP (31%)</td>
<td>Cisplatin/EP</td>
<td>31</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Carboplatin/EP</td>
<td>30</td>
<td>11</td>
</tr>
<tr>
<td>Okita [31]</td>
<td>12</td>
<td>Gastric</td>
<td>IP</td>
<td>75</td>
<td>23</td>
</tr>
<tr>
<td>Nakano [32]</td>
<td>35</td>
<td>GEP (9), H&amp;N (18), GU/GYN (5), UKP (12)</td>
<td>IP</td>
<td>64</td>
<td>NR</td>
</tr>
<tr>
<td>Okuma [33]</td>
<td>12</td>
<td>Esophagus</td>
<td>IP</td>
<td>50</td>
<td>13</td>
</tr>
<tr>
<td>Lu [34]</td>
<td>16</td>
<td>GEP</td>
<td>IP</td>
<td>57</td>
<td>11</td>
</tr>
<tr>
<td>Kulke [35]</td>
<td>4</td>
<td>GEP/UKP</td>
<td>IP</td>
<td>25</td>
<td>NR</td>
</tr>
<tr>
<td>Yamasuch [13]</td>
<td>258</td>
<td>GEP NEC/MANEC</td>
<td>IP</td>
<td>50</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Cisplatin/EP</td>
<td>28</td>
<td>7</td>
</tr>
<tr>
<td><strong>Second- or third-line therapy</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Hentic [40]</td>
<td>19</td>
<td>GEP</td>
<td>FOLFIRI</td>
<td>31</td>
<td>18</td>
</tr>
<tr>
<td>Welin [44]</td>
<td>25</td>
<td>GEP (17), UKP (5), lung (3)</td>
<td>Temozolomide +/- capcitabine +/- bevacizumab</td>
<td>33</td>
<td>22</td>
</tr>
<tr>
<td>Olsen [45]</td>
<td>28</td>
<td>GEP (18), UKP (6), lung (1), GU (3)</td>
<td>Temozolomide</td>
<td>0</td>
<td>4</td>
</tr>
<tr>
<td>Bajetta [41]</td>
<td>13</td>
<td>GEP (58%)</td>
<td>XELOX</td>
<td>23</td>
<td>NR</td>
</tr>
<tr>
<td>Ferrarotto [42]</td>
<td>9</td>
<td>GEP (75%)</td>
<td>XELOX</td>
<td>29</td>
<td>NR</td>
</tr>
<tr>
<td>Hadoux [43]</td>
<td>20</td>
<td>G3 NEC (primary NR)</td>
<td>FOLFOX</td>
<td>29</td>
<td>10</td>
</tr>
<tr>
<td>Yamasuch [13]</td>
<td>25</td>
<td>GEP NEC/MANEC</td>
<td>Amrubin</td>
<td>4</td>
<td>8</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Platinum/EP</td>
<td>17</td>
<td>5</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Irinotecan</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>S-1</td>
<td>27</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>IP</td>
<td>40</td>
<td>9</td>
</tr>
<tr>
<td>Sørbye [1]</td>
<td>100</td>
<td>GEP, UKP</td>
<td>Various (Taxane-22; Tmz-35)</td>
<td>18</td>
<td>19</td>
</tr>
</tbody>
</table>

**GU** = Genitourinary; **GYN** = gynecological; **H&N** = head and neck; **MANEC** = mixed adenoneuroendocrine carcinoma; **NR** = not reported; **RR** = response rate; **UKP** = unknown primary.
For patients with localized disease, a combination of platinum-based chemotherapy with local treatment consisting of surgery, radiotherapy or both probably offers the greatest likelihood of long-term survival. Debulking or surgical resection of metastasis is not recommended. Systemic chemotherapy is indicated in advanced inoperable disease, provided the patient has adequate organ function and performance status, otherwise the patient should be rapidly referred for consideration of palliative chemotherapy. The combination of cisplatin and EP, or alternative regimens substituting carboplatin for cisplatin or irinotecan for EP, are recommended as first-line therapy. Since response rates of these regimens are lower in patients with a Ki-67 value in the lower range of G3 (20–55%), other treatment options may be explored in these patients (especially for NEC of GI origin). While second-line regimens have not been evaluated rigorously, options include temozolomide-, irinotecan- or oxaliplatin-based schedules as main alternatives. There are no data to support the use of somatostatin analogs or PRRT in patients with GEP NEC expressing somatostatin receptors. Prophylactic cranial irradiation is not indicated in patients with limited-stage disease in complete remission.

Follow-Up

Follow-up recommendations are based on expert opinion as there is no solid evidence to support the type and frequency of performance of specific procedures. Patients with localized G3 NEC who have undergone complete resection are recommended to be followed every 3–6 months during the first 2–3 years following surgery, and then every 6–12 months up to 5 years. Conventional imaging (CT scan or MRI) should be performed during these follow-up visits, but testing for general tumor markers (i.e., chromogranin A or NSE) is only indicated if elevated at diagnosis. Somatostatin receptor imaging procedures are generally not warranted in this setting, particularly if negative at diagnosis. FDG-PET may be indicated if equivocal findings are encountered on conventional imaging and/or if salvage surgery is being considered.

Follow-up of patients with advanced disease should be customized depending upon tumor kinetics (the Ki-67 proliferative index and the actual growth rate documented by serial CT scans), treatment strategy, side effects of therapy and general health condition. Clinical assessment visits should be scheduled frequently, as these patients generally present fast tumor kinetics, are highly symptomatic and/or receive toxic agents. Clinical judgement is advised to establish the appropriate assessment interval. Conventional imaging procedures are recommended to be performed every 2–3 months while on active therapy.

In patients with localized R0/R1 resected G3 NEC, conventional imaging (CT and/or MRI) and assessment of circulating tumor markers (if elevated at baseline) are recommended to be performed every 3 months during the first 2–3 years after surgical resection, and every 6–12 months up to 5 years following surgery. In patients with advanced disease G3 NEC, frequent clinical assessment visits should be performed, and conventional imaging is recommended every 2–3 months while on active therapy.

Please also refer to consensus guideline updates for other GEP NET [47–52, this issue].

Appendix

All Other Vienna Consensus Conference Participants

Bartsch, D.K. (Department of Surgery, Philippus University, Marburg, Germany); Capdevila, J. (Institute of Oncology, Vall d’Hebron University Hospital, Barcelona, Spain); Costa, F. (Centro de Oncologia, Hospital Sirio Libanês, São Paulo, Brazil); De Herder, W.W. (Department of Internal Medicine, Division of Endocrinology, Erasmus Medical Center, Rotterdam, The Netherlands); Delle Fave, G. (Department of Digestive and Liver Disease, Ospedale Sant’Andrea, Rome, Italy); Eriksson, B. (Department of Endocrine Oncology, University Hospital, Uppsala, Sweden); Falconi, M. (Department of Surgery, San Raffaele Hospital, Università Vita e Salute, Milan, Italy); Ferolla, P. (NET Center, Umbria Regional Cancer Network, Università degli Studi di Perugia, Perugia, Italy); Ferone, D. (Department of Endocrine and Metabolic Sciences, University of Genoa, Genoa, Italy); Gross, D. (Department of Endocrinology and Metabolism, Hadassah University Hospital, Mevasseret Tzion, Israel); Ito, T. (Pancreatic Diseases Branch, Kyushu University Hospital, Fukuoka, Japan); Jensen, R.T. (Digestive Diseases Branch, NIH, Bethesda, Md., USA); Kaltasas, G. (Department of Pathophysiology, Division of Endocrinology, National University of Athens, Athens, Greece); Kellestimur, F. (Department of Endocrinology, Erciyes University Medical School, Kayseri, Turkey); Kianmanesh, R. (Department of Surgery, CHU Robert Debré, Reims, France); Klöppel, G. (Institute of Pathology, Technische Universität München, Munich, Germany); Knigge, U. (Neuroendocrine Tumor Center of Excellence, Rigshospitalet, Copenhagen University Hospital, Copenhagen, Denmark); Koskudla, B. (Department of Endocrinology, Medical University of Silesia, Katowice, Poland); Krenning, E. (Department of Internal Medicine, Division of Nuclear Medicine, Erasmus Medical Center, Rotterdam, The Netherlands); Kwekkeboom, D. (Department of Internal Medicine, Division of Nuclear Medicine, Erasmus Medical Center, Rotterdam, The Netherlands); Öberg, K. (Department of Medical Sciences, Endocrine Oncology Unit, University Hospital, Uppsala, Sweden); O’Connor, J. (Department of Clinical Oncology, Institute Alexander Fleming, Buenos Aires, Argentina); Pape, U.-F. (Department of Hepatology and Gastroenterology, Campus Virchow Klinikum, Charité Universitätsmedizin Berlin, Berlin, Germany); Pascher, A. (Department of Visceral and Transplant Surgery, Campus VirchowKlinikum, Charité Universitätsmedizin Berlin, Berlin, Germany); Pavel, M. (Department of Hepatology and Gastroenterology, Campus Virchow Klinikum,

Garcia-Carbonero et al.
Charité Universitätsmedizin Berlin, Berlin, Germany); Ramage, J.K. (Gastroenterology Department, Hampshire Hospitals NHS Trust, Hampshire, UK); Reed, N. (Beaton Oncology Centre, Gartnavel General Hospital, Glasgow, UK); Rindi, G. (Institute of Anatomic Pathology, Policlinico A. Gemelli, Università Cattolica del Sacro Cuore, Rome, Italy); Ruszniewski, P. (Department of Gastroenterology, Beaujon Hospital, Clichy, France); Sundin, A. (Department of Radiology, Section for Molecular Imaging, University Hospital, Uppsala, Sweden); Taal, B. (Netherlands Cancer Centre, Lijn den, The Netherlands); Weber, W. (Department of Radiology, Memorial Sloan Kettering Cancer Center, New York, N.Y., USA); Zheng-Pei, Z. (Department of Endocrinology, Peking Union Medical College Hospital, Beijing, China).

References


19 Agaimy A, et al: ISL1 expression is not restricted to pancreatic well-differentiated neuroendocrine neoplasms, but is also commonly found in well and poorly differentiated neuroendocrine neoplasms of extrapancreatic origin. Mod Pathol 2013;26:995–1003.


High-Grade GEP Neuroendocrine Tumors and NEC

Neuroendocrinology 2016;103:186–194

DOI: 10.1159/000443172

193