Extrapulmonary neuroendocrine small and large cell carcinomas: a review of controversial diagnostic and therapeutic issues.

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(Article begins on next page)
Extrapulmonary neuroendocrine small and large cell carcinomas: a review of controversial diagnostic and therapeutic issues

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Summary Extrapulmonary neuroendocrine carcinoma (EPNEC) is a heterogeneous and rare group of high-grade neoplasms occurring in different organs. They usually share a poor prognosis, but diagnostic and therapeutic options still include several controversial issues, due to the rarity of this condition and to differences in architecture and cell size, being some cases pure small cell carcinomas, other pure large cell neuroendocrine carcinomas and some others combined/mixed neuroendocrine carcinomas with a conventional non-neuroendocrine carcinoma. In addition, the therapeutic strategy varies in different organs (surgery and/or chemotherapy and/or radiation therapy and/or targeted treatments), and clinicians and pathologists are asked to interact to reach an accurate classification of every single case, as well as the most appropriate selection of the treatment options, even considering different time points of each EPNEC natural history. This overview highlights controversial pathological and clinical issues and summarizes possible solutions to most of such EPNEC-related problems.

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1. Introduction

In a recent commentary on diagnostic and classification problems of neuroendocrine (NE) tumors\cite{1}, several frequently asked questions (FAQ) were discussed, and some tentative answers offered to specific topics. Among them, the issue of high grade, poorly differentiated NE carcinomas (NEC) was briefly addressed (FAQ #4) with regard to their occurrence in several extrapulmonary organs, as either small cell carcinomas (SCC) or large cell NECs (LCNEC), apparently very similar to their well-known respective pulmonary counterparts\cite{2,3}. In the literature, the occurrence of extrapulmonary poorly differentiated NECs (EPNEC) has been reported in single case reports (see review\cite{4}), and only recently were relatively large series collected and analyzed from both clinical and pathological perspectives\cite{5}. However, the pathological aspects as well as the therapeutic strategies of EPNEC still seem rather heterogeneous in the literature, and the present review collects the available diagnostic and therapeutic information on this tumor group, highlighting crucial issues and possible solutions.
2. Relevant issues and possible solutions

2.1. Issue 0: organ distribution and incidence

EPNECs are defined as malignant epithelial tumors having a totally or predominant NE differentiated cell population and high-grade features. They are rare or extremely rare outside the lung and can develop in virtually any location. Relatively large series were reported in the gastroenteropancreatic (GEP) area, including the esophagus [6], stomach [7,8], colon and rectum [9], and pancreas [10]. In addition, EPNECs of the small or large type were investigated in the prostate [11,12], bladder [13] and uterine cervix [14,15]. Overall, EPNECs account for no more than 3% of all malignancies in individual anatomical sites.

NECs have also been described in the skin for more than 40 years, under the generic term of Merkel cell carcinoma. Although the morphology of classical small cell forms is not different from that of small cell lung cancer, Merkel cell carcinoma has heterogeneous morphological and molecular features, as well as variable and peculiar clinical behavior and therapeutic implications. For these reasons it has been excluded from the present review.

2.2. Issue 1

2.2.1. Problem: How to label these tumors?

2.2.1.1. Solution: Use the general term EPNEC; quote synonyms. A relatively long list of different terms are currently used to label these tumors. They refer either to cell size (extrapulmonary small or LCNECs), to tumor grade (high-grade NEC, G3) or to loss of differentiation (poorly differentiated NE/endocrine carcinomas). In addition, the occurrence of combined small and large cell NEC has been reported not only in the lung but also in some other locations (stomach, gallbladder, bladder, etc), although this distinction is not considered of clinical or prognostic relevance in these organs (including the lung) [5]. The terminology used for such tumor type(s) is itself heterogeneous, being only partially derived by their largely more common pulmonary counterpart. In fact, for LCNEC in extrapulmonary organs, the original terminology proposed by Travis and coworkers for the corresponding lung tumors has often been adopted [3], whereas when series of pure NECs or mixed/combined carcinomas are investigated in the stomach or colon rectum or bladder, the nomenclature is more confused and either SCC or poorly differentiated NEC terms are used, often interchangeably. In our view, this is not appropriate, since pure SCC generally do not differ from their classical pulmonary counterpart (oat cell type), but all other variants including intermediate cell type (a definition included in previous classifications), LCNEC and combined NE/non-NECs actually display more complex morphological features, which cannot be directly addressed to one histological type.

Referring to the available World Health Organization (WHO) classifications, the 2010 WHO classification of tumors of the digestive system restricted the term “NEC” to poorly differentiated, clinically aggressive neoplasms, which are graded as G3 by definition [16]. Similar to the lung, these include small and large cell variants (or a combination of the two), being relatively common in the stomach and very rare in the pancreas. By contrast, in other WHO classification schemes (namely of the urogenital tract and breast/female genital organs) [17,18] nomenclature per se is heterogeneous and include general terms such as NEC not otherwise specified or consider SCC type, only.

For practical purposes, in analogy to what is proposed in the GEP system, the nomenclature of EPNEC is advisable, with a subspecification of either small or large cell types. Such uncommitted terminology would allow to compare the pathological features and the real prevalence of EPNEC, as well as to address the issue of their most appropriate therapeutic strategy.

2.3. Issue 2

2.3.1. Problem: Differential diagnosis of small and large cell NEC

2.3.1.1. Solution: Identify architectural patterns, cytological features, and appropriate immunophenotype. EPNEC of the classical small cell type has uniform morphological features, irrespective of the organ in which they develop, overall similar to those of the pulmonary counterpart (Fig.). Conversely, other EPNEC forms, including pure LCNEC and combined small and large cell carcinomas, have peculiar features with differences in various locations (Table 1).

From a pathological point of view, the distinction between small and large cell tumors is of relevance for the purpose of a correct classification of the tumor itself and, above all, an accurate differential diagnosis from other...
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conditions made of either small or large cells in individual organs, which may be problematic especially in cytology [19] or tiny biopsy specimens. In practical terms, it was claimed that the distinction between SCC and LCNEC is an apparently useless exercise, not only due to the similar clinical behavior reported above but also because many cases (indeed probably the majority of them) actually contain mixed cell populations, all sharing more or less evident NE features but displaying various cell sizes. In any case, a full mixed cell populations, all sharing more or less evident NE features but displaying various cell sizes. In any case, a full range of options is possible, including pure SCC (former “oat cell”) cancers on the one side and LCNEC on the other. In between, various combinations of small and large cell carcinomas may occur, covering the whole spectrum of cell sizes and having variable tumor architecture and cytological features. As a matter of fact, similar to what is observed in the lung, small cell EPNEC are usually neoplasms with a diffuse (solid or noncohesive) growth and finely granular nuclei lacking evident nucleoli. By contrast, large cell EPNEC present a more structured architecture (solid, trabecular, or organoid) and frequently large nuclei with hyperchromatic nucleoli.

The occurrence of EPNEC may also create differential diagnostic problems with poorly differentiated forms of their respective nonendocrine carcinomas or other malignancies such as hematological and mesenchymal neoplasms. As an example, small cell EPNEC in the bladder may be easily misdiagnosed as urothelial carcinoma [20], also because often a combination of these two forms exists (see also below).

Concerning the differential diagnoses mentioned above, there are no markers useful to discriminate small cell from large cell cases. Both display an epithelial immunoprofile, with wide spectrum cytokeratin expression, which may occasionally have a paranuclear dot-like distribution, especially in the case of SCC. Conversely, high-molecular-weight cytokeratins (types 1, 5, 10, 14 of the Moll’s catalog) are typically absent in the vast majority of NE differentiated carcinomas irrespective of the location [21,22]. Among NE markers, chromogranin A (more often with a focal paranuclear dot-like pattern unmasked by heat-induced antigen retrieval procedures) and synaptophysin are the most reliable molecules supporting the morphological evaluation, while neuron-specific enolase (NSE) and CD56 are sensitive but much less specific. Interestingly, up to 80% of EPNEC express the thyroid- and lung-specific marker TTF-1, thus limiting its usefulness in the definition of the primary location in the cases of NEC of unknown primary origin [23,24].

An additional relevant challenge for pathologists is the differential diagnosis between high-grade NECs and well-differentiated, low-grade NE neoplasms (“carcinoid”). This is not an obvious exercise in the daily pathology practice, especially when facing small endoscopic biopsies or needle aspirates from metastatic sites (eg, liver). In fact, cases of well-differentiated NE tumors (NETs) made of small cells do exist as either the result of artifactual cellular size changes (frequent in small biopsies or cytological specimens) or, more rarely, as a special small cell variant of such tumors. In surgical specimens, the diagnosis is generally easy, in the presence of an organoid or trabecular growth of low-grade NETs but may not be straightforward in preoperative specimens. Although not specifically addressed for tumors in extrapulmonary locations, a study of 7 pulmonary carcinoid tumors wrongly interpreted as SCCs in the biopsy, but confirmed as low-grade NETs in the surgical specimen, showed that Ki-67 index is the most reliable marker for a differential diagnosis of high-grade carcinomas from carcinoid tumors [25].

In addition, cases exist of well-differentiated NETs having a borderline number of mitoses and/or extensive necrosis, which may induce the suspicion of an EPNEC. The usual approach is to perform an accurate mitotic count usefully supported by a Ki-67 immunostaining. However, with special reference to the GEP system, a grey zone (not well clarified by the WHO classification) exists in terms of proliferation between NEC (all Grade 3) and rare NETs showing a well-differentiated morphology but a Ki-67 labeling index higher than 20% (thus by definition G3, too). In this latter case, an additional useful method might be to consider the pattern and intensity of reactivity for chromogranin A, which is generally focal and/or located paranuclear in a dot-like fashion in high-

### Table 1
Comparison of general characteristic of pulmonary and extrapulmonary (divided into GEP and urogenital tract locations) NECs

<table>
<thead>
<tr>
<th>Location</th>
<th>Relative frequency as compared to NETs (carcinoids)</th>
<th>Most prevalent type (SCC vs LCNEC)</th>
<th>Frequency of mixed NE and non-NE features</th>
<th>Etiologic/ risk factors</th>
<th>Precursor lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lung GEP</td>
<td>More frequent</td>
<td>SCC</td>
<td>Very rare</td>
<td>Cigarette smoking</td>
<td>Unknown (possibly divergent differentiation from adenocarcinoma)</td>
</tr>
<tr>
<td></td>
<td>Less frequent</td>
<td>LCNEC</td>
<td>Frequent</td>
<td>Unknown</td>
<td>Unknown (possibly divergent differentiation from carcinoma subtypes)</td>
</tr>
<tr>
<td>Urogenital tract</td>
<td>More frequent</td>
<td>SCC</td>
<td>More than 50% of cases</td>
<td>Unknown</td>
<td>Unknown (possibly divergent differentiation from carcinoma subtypes)</td>
</tr>
</tbody>
</table>
grade EPNECs, while stronger and uniformly cytoplasmic in well-differentiated NETs.

2.4. Issue 3
2.4.1. Problem: What is the real role of Ki-67 index in small versus large cell NEC?

2.4.1.1. Solution: Mandatory use of Ki-67 index for grading GEP NEN, only. As a rule, Ki-67 index is to be assessed in all GEP NE neoplasms [16] to define tumor grading, with an accurate manual count rather than with an “eyeballing” approach [26]. However, Ki-67 evaluation is not valuable to distinguish small from large cell NECs, and there are no specific recommendations for Ki-67 investigation in sites other than the GEP system.

A literature review of the reported Ki-67 values in EPNEC is summarized in Table 2, based on published papers clearly reporting Ki-67 values for small and/or large cell components in EPNECs. Despite the difficulty of comparing the various figures obtained by different authors (also due to different classification criteria for small and large cells, as well as different counting methods), it seems that the mean proliferative index of small cell patterned NECs is slightly higher than that of morphologically proven LCNECs originated from pancreas, larynx, bile ducts, gastrointestinal tract, parotid, and urogenital tract (70.9% versus 62%).

2.5. Issue 4
2.5.1. Problem: How to classify EPNEC combined with adeno-, squamous, or urothelial carcinoma?

2.5.1.1. Solution: Heterogeneous criteria exist in different organs; favor morphological parameters and appropriate immunophenotypes. One of the most challenging (and also difficult to identify and treat) situation is the combination of NEC with a more or less well-represented non-NE component. Such component may be totally separated from the EPNEC (collision tumor) but more often is intermingled with the NEC population and indeed may not be readily apparent. This is especially true in cases of EPNEC combined with poorly differentiated non-NE components (for example basaloid squamous carcinoma of the uterine cervix or undifferentiated urothelial carcinoma).

For the pathological diagnosis of such cases, a first clue is to identify the NE cell population as such, and then the relative proportions of the exocrine and endocrine components are to be determined [49].

In the GEP area, this step is mandatory for the purpose of identifying so-called MANECs (mixed adeno-NEC), being at least 30% of each component required for rendering such a diagnosis in any given mixed tumor. In the stomach, mixed tumors follow a behavior intermediate between pure LCNEC (the most aggressive subgroup) and conventional gastric adenocarcinoma [7]. In the colon and rectum, the extent of

<table>
<thead>
<tr>
<th>Ref</th>
<th>No. of cases</th>
<th>Location</th>
<th>Ki-67 index (%) in different components</th>
</tr>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td>Small cell</td>
</tr>
<tr>
<td>Nagao et al 2000 [27]</td>
<td>2</td>
<td>Parotid</td>
<td>-</td>
</tr>
<tr>
<td>Papotti et al 2000 [28]</td>
<td>2</td>
<td>Gallbladder</td>
<td>-</td>
</tr>
<tr>
<td>Crafa et al 2003 [29]</td>
<td>1</td>
<td>Rectum</td>
<td>-</td>
</tr>
<tr>
<td>Soriano et al 2004 [30]</td>
<td>10</td>
<td>Bladder</td>
<td>33 (15-70)</td>
</tr>
<tr>
<td>Sugawara et al 2004 [31]</td>
<td>1</td>
<td>Ampulla of Vater</td>
<td>54</td>
</tr>
<tr>
<td>Fernandez-Figueras et al 2005 [32]</td>
<td>23</td>
<td>Bladder/lung</td>
<td>64.7</td>
</tr>
<tr>
<td>Stachs et al 2005 [33]</td>
<td>1</td>
<td>Endometrium</td>
<td>50</td>
</tr>
<tr>
<td>Lee et al 2009 [34]</td>
<td>1</td>
<td>Bladder</td>
<td>-</td>
</tr>
<tr>
<td>Miyamoto et al 2006 [35]</td>
<td>1</td>
<td>Rectum</td>
<td>-</td>
</tr>
<tr>
<td>Malhotra et al 2008 [36]</td>
<td>1</td>
<td>Liver</td>
<td>90</td>
</tr>
<tr>
<td>Kozyrakis et al 2009 [37]</td>
<td>1</td>
<td>Bladder</td>
<td>70</td>
</tr>
<tr>
<td>Yamaguchi et al 2009 [38]</td>
<td>1</td>
<td>Breast</td>
<td>85</td>
</tr>
<tr>
<td>Lewis et al 2010 [39]</td>
<td>10</td>
<td>Larynx</td>
<td>-</td>
</tr>
<tr>
<td>Righi et al 2010 [40]</td>
<td>11</td>
<td>Breast</td>
<td>58 (40-75)</td>
</tr>
<tr>
<td>Stojic et al 2010 [41]</td>
<td>1</td>
<td>Ampulla of Vater</td>
<td>-</td>
</tr>
<tr>
<td>Terada 2010 [42]</td>
<td>1</td>
<td>Endometrium</td>
<td>-</td>
</tr>
<tr>
<td>Terada 2011 [43]</td>
<td>1</td>
<td>Esophagus</td>
<td>100</td>
</tr>
<tr>
<td>Albisinni et al 2012 [44]</td>
<td>1</td>
<td>Prostate</td>
<td>100</td>
</tr>
<tr>
<td>Benkel et al 2012 [45]</td>
<td>1</td>
<td>Gallbladder</td>
<td>70</td>
</tr>
<tr>
<td>Jianu et al 2012 [46]</td>
<td>1</td>
<td>Stomach</td>
<td>-</td>
</tr>
<tr>
<td>Samad et al 2012 [47]</td>
<td>1</td>
<td>Bile ducts</td>
<td>-</td>
</tr>
<tr>
<td>Yachida et al 2012 [10]</td>
<td>19</td>
<td>Pancreas</td>
<td>67 (n = 9) (55.1-85.8)</td>
</tr>
<tr>
<td>Yamamoto et al 2012 [48]</td>
<td>1</td>
<td>Pancreas</td>
<td>80</td>
</tr>
<tr>
<td>Mean values</td>
<td>93 cases</td>
<td></td>
<td>70.9</td>
</tr>
</tbody>
</table>
the NE component (within the 30% rule) is not a predictor of behavior, and even a minor high-grade NE cell population was found able to metastasize [50]. In such location, the prognosis was comparable between MANEC (mixed adeno-neuroendocrine carcinoma) and NEC, with a better prognosis in cases whose NE component was made of large cells [9]. Very few cases of concurrent adenocarcinoma and NEC in the hepatobiliary organ were reported [51].

In other locations, there are either different diagnostic rules or none. NE breast carcinoma is one of the several variants of this neoplasm and is defined as a tumor having NE morphological features necessarily associated with the immunohistochemical expression of NE markers in at least 50% of the tumor [18]. Such tumor type may display variable morphological patterns, including high-grade NEC of the large or small cell type. These latter are generally pure forms of primary breast EPNECs and therefore the NE cell population largely exceeds the required 50% and reaches 90% to 100%. The other NE breast carcinomas are rather classical ductal or lobular carcinomas combined with a prominent (>50%) NE cell component having mucinous, trabecular, alveolar, or solid patterns, which morphologically have little to do with the currently discussed small or large cell NE cancers [40].

In the thymus, the definition of combined exocrine-NEC, as part of thymic carcinoma (type C), is rather generic, being restricted to the recognition of “two distinct areas, each corresponding to one of the histological types, including NEC” [52].

In the urogenital tract, combined carcinomas are as common as pure forms of EPNEC. In the prostate, EPNECs occur as either pure SCC (57%) or combined with an adenocarcinoma having a high (≥8) Gleason score, according to the largest published series of 95 cases [12]. In the bladder, no criteria are mentioned for the definition of mixed NEC: SCCs combined with urothelial, squamous, or adenocarcinoma occur more frequently (70%) than pure forms [53], and the trend is similar for the even rarer LCNEC [54].

Similarly, in the ovary and in the uterine cervix there are no specific rules. In the latter, the NE component in an otherwise squamous cell carcinoma has been described to range from focal (17%) to half a tumor, but NE differentiation was associated with an adverse prognosis irrespective of the extension [55].

2.6. Issue 5

2.6.1. Problem: Does molecular signature keep separate small and large cell EPNEC?

2.6.1.1. Solution: There are insufficient published data on this issue. Published genetic data are mostly dealing with mixed/combined exocrine-NECs, while separate analyses of large and small cell components within the same NET or comparison of genetic profiles in SCC versus LCNEC are lacking or are restricted to the pulmonary location, where a genetic similarity between SCC and LCNEC components within individual tumors is seen [56]. On the contrary, different genetic abnormalities in chromosomes 3p and 5q were found between pulmonary small and LCNECs, with only some overlapping features [57]. Moreover, data on a large series of lung SCC and LCNEC demonstrated that gene expression profiles are heterogeneous in the 2 forms and cluster analysis was unable to separate the two entities, but rather identified clinically distinctive subgroups [58].

In extrapulmonary locations, only a single recent study [10] analyzed 19 pancreatic NECs (9 small and 10 large cells), as well as 11 well differentiated NETs, for alterations of KRAS, CDKN2A/p16, P53, SMAD4/DPC4, DAXX, ATRX, PTEN, Bcl2, and Rb1 genes. Small and large cell NECs had genetically similar profiles but distinct from those observed in well-differentiated NETs. Other studies are restricted to single case reports, such as esophageal [43] and endometrial [42] EPNECs, that were investigated for KIT and PDGFR genes and found no mutations. Finally, some cytogenetic differences were reported between primary pulmonary and extrapulmonary SCC [59]. Overall, the current knowledge does not support or disprove a molecular separation of small and large cell pulmonary nor extrapulmonary NECs.

2.7. Issue 6

2.7.1. Problem: Any clinical meaning of distinguishing small and large cell NEC (in terms of behavior or therapeutic strategy)?

2.7.1.1. Solution: Yes, for differential diagnosis purposes. Probably yes for tuning chemotherapy protocols; in the future, personalized treatments are expected. Having defined the criteria for taking small cell apart from large cell carcinoma forms, the next question arises if this really matters. The reason for this question relies on the comparable overall survival for surgically resected pulmonary SCC and LCNEC, with figures of approximately 35% and 40%, respectively [60]. In this scenario, however, relatively recent reports seem to indicate a unique response rate of EPNEC to chemotherapy protocols conventionally used for pulmonary SCC [61].

Running through therapy-oriented studies on EPNEC available in the literature, it can be concluded that the treatment of EPNEC of the small cell type does not differ from the pulmonary counterpart [62–64] with response rates and global prognosis in general overlapping with those reported for lung SCC, even though some authors reported poorer results [65]. Since LCNECs of the lung, at least in their advanced stage, are also treated similarly to SCC [66], the whole spectrum of EPNEC generally undergoes the same chemotherapy approach. Indeed, some reports indicate that despite that the chemotherapy protocols are quite similar, response to therapy is different and the stability of disease after completion of chemotherapy may follow a different course [61,63,67].

The first report on EPNEC therapy goes back to 1991 when Moertel and coworkers [68] firstly proposed platinum-
etoposide regimens in NEC and identified a 67% response rate with a median survival of 19 months. These apparently excellent results were subsequently confirmed in 41 EPNEC patients who displayed a 42% response rate and a 15-month median survival [69]. Nowadays, according to the North American Neuroendocrine Tumor guidelines [64], first-line systemic chemotherapy with a platinum agent (cisplatin or carboplatin) and etoposide is recommended for most EPNEC patients with metastatic-stage disease; however, response durations are often short. Sequential or concurrent chemotherapy is recommended for patients with locoregional disease [70].

Unfortunately, more recent studies reported less favorable clinical results. In 21 unresectable or recurrent hepatobiliary and pancreatic EPNECs, combination chemotherapy with cisplatin and etoposide provided a 14% response rate and a median overall survival of 5.8 months [71]. Comparing the efficacy of a platinum-containing regimen in 41 advanced EPNECs with that of SCC of the lung, the response rates were much worse in the former (31% versus 78%) [61]. This remarkable discrepancy might be due to the different nature of the 2 tumors (classical pulmonary SCCs are highly chemo- and radiosensitive undifferentiated neoplasm, while EPNEC include more heterogeneous cancer subtypes), or to the anatomical location of the tumors, as partially confirmed by the worse response of hepatobiliary and pancreatic NECs (12% versus 57% for the other locations).

In a review of over 500 reported NETs with an unknown primary, despite the heterogeneity in terms of histology, grade, anatomic site, and tumor biology, it was found that most cases were managed with platinum-based regimens. The 294 patients with follow-up information had a median survival of 15.5 months, comparable to that reported for other locations (25 months). Unfortunately, in this study no morphological review of the cases was performed, thus leaving the question open as to the prevalent histological type of tumors associated with a better response.

With regard to molecular targets of chemotherapeutic agents, it has been reported that pulmonary SCC are not responsive to pemetrexed drugs [73]. This was supported by the observation that high-grade NEC of the small and large cell type and of both pulmonary and GEP origin express high levels of thymidylate synthase, the most important target molecule of antifolates [74], thus supporting the usefulness of assessing the intratumoral expression levels of known targets, to better define the therapeutic strategy and/or predict response to different agents. Recent data are emerging with regard to the activation of intracellular signaling pathways as possible targets of specific treatments. A high phospho-mammalian target of rapamycin expression was reported in 9 poorly differentiated (large cell type) GEP NEC [75], suggesting that mammalian target of rapamycin could be explored as a possible therapeutic target in this subtype, as already well known for low-grade NETs.

2.8. Issue 7

2.8.1. Problem: Any predictive factor of response to therapy in EPNEC?

2.8.1.1. Solution: Assess proliferation index in all cases; in the future, personalized treatments are expected. In the large series of GEP NEC analyzed in the NORDIC study [63] a negative prognostic role of performance status, colorectal primary, elevated platelet, and lactose dehydrogenase (LDH) levels was observed. Proliferation index differed in terms of impact on survival and response to chemotherapy. In fact, it was found that Ki-67 index at a cut-off of >55% was a positive predictor of response to platinum-based regimens (15% versus 42% comparing <55% versus >55% Ki-67, respectively), but patients with Ki-67 <55% had a significantly longer survival compared with patients with higher Ki-67 levels (14 versus 10 months). Unfortunately, in this study no morphological review of the cases was performed, thus leaving the question open as to the prevalent histological type of tumors associated with a better response.

With regard to molecular targets of chemotherapeutic agents, it has been reported that pulmonary SCC are not responsive to pemetrexed drugs [73]. This was supported by the observation that high-grade NEC of the small and large cell type and of both pulmonary and GEP origin express high levels of thymidylate synthase, the most important target molecule of antifolates [74], thus supporting the usefulness of assessing the intratumoral expression levels of known targets, to better define the therapeutic strategy and/or predict response to different agents. Recent data are emerging with regard to the activation of intracellular signaling pathways as possible targets of specific treatments. A high phospho-mammalian target of rapamycin expression was reported in 9 poorly differentiated (large cell type) GEP NEC [75], suggesting that mammalian target of rapamycin could be explored as a possible therapeutic target in this subtype, as already well known for low-grade NETs.

3. Conclusions

i. based on both clinical and pathological findings and from a practical point of view, a common terminology is advisable for pure extrapolumonary high-grade NECs, using the acronym EPNEC and a subsequent distinction into small and large cell subtypes;

ii. similar to the lung, small and large cell EPNECs differ histologically by cell size, architecture, and nuclear features, although morphologically intermediate cases exist and the general genetic background, as well as the clinical behavior, are similar in the 2 forms;

iii. recognition of mixed NE/non-NE forms in extrapolumonary locations is based on morphology and appropriate immunophenotype; due to its relatively high frequency, it should be excluded by extensive sampling, and the relative proportions on NE and non-NE components should be reported in any case to better understand the impact on clinical behavior and response to treatment(s);

iv. immunohistochemistry is useful to distinguish EPNEC from poorly differentiated non-NECs (using NE markers) and from well-differentiated NETs (using Ki-67, especially in small tissue fragments/cytological samples), but not to differentiate small and large cell EPNEC forms;

v. the pathogenesis of EPNEC is unclear and possibly different from that of pulmonary NECs;

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vi. platinum-based chemotherapy represents the most commonly used therapeutic approach in EPNEC (both small and large cell types), based on similarity between EPNEC and pulmonary SCC. However, some increasing evidence seem to indicate that response to therapy and prognosis are different between pulmonary and extrapulmonary NEC and possibly between EPNEC of the small and large cell types.

References


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