Updates and management algorithm for neuroendocrine tumors of the uterine cervix

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ABSTRACT

Neuroendocrine carcinomas of the cervix account for less than 2% of all invasive cervical cancers and are classified as low-grade (carcinoid, atypical carcinoid tumor) or high-grade (known as small- and large-cell) neuroendocrine carcinomas. There are increasing data showing that cervical neuroendocrine carcinomas may be associated with the human papillomavirus (HPV), especially HPV18, and most will stain positive for p16. Immunohistochemistry markers such as synaptophysin and CD56 are the most sensitive markers. Although there are no commonly associated mutations, PIK3CA, KRAS, and TP53 are the most frequently found mutations in neuroendocrine tumors. Neuroendocrine cervical carcinomas are exceedingly aggressive tumors with a high tendency for nodal involvement and distant metastases. Age, lymph node metastases, smoking, pure small-cell histology, and tumor size are independent prognostic factors. Overall, the 5-year survival rate is 36% and the median overall survival ranges between 22 and 25 months. Treatment options are often extrapolated from small-cell lung cancer and limited retrospective studies. The preferred treatment is a multimodal approach of surgery, chemoradiation, and systemic chemotherapy. The most common chemotherapy regimen used as initial therapy is a combination of cisplatin and etoposide. In the setting of recurrent disease, a combination of topotecan, paclitaxel, and bevacinumab has demonstrated favorable outcomes. Multicenter tumor registries, such as the Neuroendocrine Cervical Tumor Registry (NeCTuR), are an opportunity to evaluate patterns of disease treatment and oncologic outcomes.

BACKGROUND

Neuroendocrine tumors are a rare entity of the female genital tract, with cervix being the most common primary site. First described by Alboros-Saavedra in 1972, these tumors account for 1.4% of all invasive cervical cancers and nearly 200 new cases are diagnosed in the United States each year.2 3 The 2014 World Health Organization (WHO) Classification2 categorizes cervical neuroendocrine tumors as low-grade (previously referred to as carcinoid tumor and atypical carcinoid tumor) or high-grade neuroendocrine carcinomas (previously referred to as small-cell carcinoma or large-cell neuroendocrine carcinoma) (Table 1). In the cervix, small-cell neuroendocrine tumor is the most common (80%) followed by large-cell neuroendocrine carcinoma (12%), and other histologic types such as undifferentiated neuroendocrine tumors (8%). Unlike squamous and adenocarcinoma subtypes, which spread primarily by local extension, high-grade neuroendocrine tumors have a high rate of lymphatic and hematogenous metastasis even when disease is clinically limited to the cervix. These tumors frequently have lymph vascular space involvement, strong association with HPV18, and an increased risk of nodal metastases at diagnosis, with 40% of clinical stage IB1 tumors diagnosed with positive pelvic nodes, in comparison to 10%–15% node positivity seen in clinical stage IB1 squamous cell carcinoma.3 4 Patients with small-cell carcinoma are more likely to be diagnosed at late stage.5 6 Extra-pelvic disease at the time of diagnosis is also common, especially in the lungs and liver, and is often associated with high risk of relapse.7 8 The median overall survival for patients with small-cell neuroendocrine tumors is less than 2 years.5 6 9–11

There is a total of 3,538 cases of neuroendocrine cervical cancers reported in 147 studies in the literature to date and only nine studies included more than 50 patients.3 The lack of prospective studies limits our ability to define concrete chemotherapy regimens, and the primary treatment is a multimodal approach including surgery, radiation, and chemotherapy.12 13

At MD Anderson Cancer Center we have a registry of neuroendocrine cervical carcinomas with patients treated not only at our institution but also elsewhere. All neuroendocrine cervical carcinomas are treated by the same small group of physicians that also serve as consultants for doctors from around the world. In this review we aim to summarize the latest and most relevant information regarding this disease and also provide our algorithms for diagnosis and treatment. In addition, we will discuss key points to help physicians around the world provide better treatment for this uncommon disease.

ETIOLOGY

The association between squamous cell carcinoma, adenocarcinoma, and adenosquamous carcinoma of the cervix with human papillomavirus (HPV) has been well established and documented.14 15 For cervical neuroendocrine carcinomas, the link with HPV has not
been as definitively established but emerging data suggest a likely association. Alejo et al. explored HPV DNA detection and genotype distribution and their relation to histologic and immunohistochemical features in 49 cervical neuroendocrine tumors. The authors found that 86% of neuroendocrine tumors had HPV DNA. Single infection (one HPV type) was reported in 98% of cases. HPV16 was found in 55% of neuroendocrine tumors, HPV18 in 41%, and 4% of tumors were positive for other HPV types. HPV18 was four times (41%) more frequent in neuroendocrine tumors compared with other HPV types. They also found that neuroendocrine tumors were more frequently associated with concomitant glandular (rather than squamous) lesions. Neuroendocrine carcinomas showed marked lymphatic permeation, a feature that is particularly characteristic of HPV18-related tumors.

Almost all squamous cell carcinomas (97%) and half of adenocarcinomas show p16-positive staining. In the study by Alejo et al., the authors found that p16 staining was over-expressed in 86% of cases. All carcinoid, atypical carcinoid, and large-cell neuroendocrine carcinomas were p16-positive, while 79% of small-cell carcinomas showed p16-positive staining. Overall, concordant results of p16 and HPV detection were observed in 89% of cases. In a study by Kuji et al., the authors evaluated 37 high-grade neuroendocrine cases and found that 72% had high-risk HPV infections (HPV16 in 14% and HPV18 in 86%). Castle et al. published a metaanalysis of 32 studies, including 403 small-cell and nine studies including 45 large-cell neuroendocrine tumors. The authors found that 85% of small-cell carcinomas were HPV-positive (HPV16 and/or HPV18: 78%). In a subanalysis of five studies, including 75 patients with small-cell carcinoma, 93% were positive for p16 by immunohistochemistry and 100% were HPV-positive. Large-cell neuroendocrine carcinomas were HPV-positive in 88% (HPV16 and/or HPV18: 86%).

The possible relationship between cervical neuroendocrine carcinomas and HPV infection is a very important finding as these cancers may be prevented by prophylactic HPV vaccines. It also opens the possibility of administering immunotherapy or therapeutic vaccines as a treatment for women with this disease.

**IMMUNOHISTOCHEMISTRY**

Immunohistochemistry stains are frequently used to help diagnose neuroendocrine tumors. The most frequently used neuroendocrine stains are chromogranin A, synaptophysin, CD56, and neuron-specific enolase. For large-cell neuroendocrine carcinomas, in addition to morphology by hematoxylin and eosin stain, chromogranin A is the most specific marker to help confirm the diagnosis. Marker positivity in small-cell neuroendocrine tumors ranges from 33% to 100%. In the latest systematic review published by Tempfer et al., the most frequently observed immunohistochemistry markers were synaptophysin (79%), neuron specific enolase (69%), chromogranin (66%), and CD56 (61%). Synaptophysin and CD56 are the most sensitive neuroendocrine markers, but CD56 lacks specificity. Chromogranin is the most specific neuroendocrine marker but lacks sensitivity, with approximately 50% positivity for small-cell neuroendocrine carcinoma.

In the study by Alejo et al., 65% were positive for at least one immunohistochemical marker (chromogranin, CD56, and/or synaptophysin) with CD56 being the most frequently positive (62%) followed by chromogranin (39%) and synaptophysin (26%). Similarly, other authors found CD56 to be the most sensitive marker of neuroendocrine differentiation (62%) followed by chromogranin A (39%) and synaptophysin (26%).

Chromogranin positivity may be very focal with punctuate cytoplasmic immunoreactivity, which is only visible on high-power magnification. Small-cell neuroendocrine carcinoma may be only focally positive (often punctuate cytoplasmatic staining) or even negative with broad-spectrum cyto-keratins. As previously mentioned, most cervical high-grade neuroendocrine carcinomas are diffusely positive for p16 due to the presence of high-risk HPV.

Insulinoma-associated protein 1 (INSM1) may be more specific for neuroendocrine tumors than chromogranin A or synaptophysin. In a recent publication, 37 high-grade neuroendocrine cases were stained with neuroendocrine markers and INSM1. Chromogranin A and synaptophysin were each expressed in 86% of cases. INSM1 was detected in 95% of cases. INSM1 seems to be a useful new neuroendocrine marker and may be associated with the development of high-grade neuroendocrine carcinomas.

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Investigators at MD Anderson Cancer Center studied the immunohistochemical expression and prognostic role in survival of HER-2/neu, epidermal growth factor receptor (EGFR), vascular endothelial growth factor (VEGF), cyclooxygenase-2 (COX-2), estrogen receptor, and progesterone receptor in small- and large-cell neuroendocrine cervical carcinomas in 24 patients. Twenty-three cases (96%) expressed VEGF, eight (33%) expressed EGFR, 10 (42%) expressed HER-2/neu, and seven (29%) expressed COX-2. No significant differences in the expression of these factors were found between small- and large-cell tumors. Only HER-2/neu expression was associated with survival, showing that patients with negative HER-2/neu expression tumors had significantly shorter survival than those whose tumors were positive, 14.2 months versus 33 months, respectively (p=0.03).

Immunohistochemical expressions may be helpful when trying to determine the site of origin of metastatic small-cell carcinoma. Some studies have shown that 33%–84% of small-cell cervical
carcinomas exhibit diffuse nuclear positivity for thyroid transcrip-
tion factor-1 (TTF-1).21 25–27 Liu et al28 compared different expres-
sion of neuroendocrine markers, TTF-1, p53, and Ki-67 in 23 cases of cervical small-cell and 56 cases of pulmonary small-cell
carcinomas using immunohistochemistry. The TTF-1 expression of small-cell carcinoma of the lung showed significantly higher diffuse
and strong positivity in tumor cell nuclei than the cervical counter-
part (p=0.003). Although both tumors had similar morphological
features, they have different immunohistochemical panel. Both had
similar positivity for CD56 and chromogranin A, but the expression
of the synaptophysin in cervical small-cell was significantly higher
than in small-cell lung cancer (p=0.007).

Immunohistochemistry is important at the time of diagnosis but
one should be aware that positive staining could be as low as 33%
using standard markers and that stains may be only focally positive.
New markers should be considered if the diagnosis is still unclear
as they shown higher positivity rates, as in the case of INMS-1 and
VEGF.

MOLECULAR AND IMMUNE PROFILE TESTING

Although small-cell cervical cancer is a rare tumor without a
unifying mutational event, identifying genetic alterations that are
amenable to targeted therapy provides an opportunity to individ-
ualize therapy especially in relapses where the spectrum of treat-
ments is limited. While high-risk HPV may be involved at an early
stage of oncogenesis in many tumors, additional driving events
have been postulated to facilitate the progression of small-cell
carcinomas. Identification of oncogenic drivers has allowed for
a better understanding of the natural history of neuroendocrine
tumors. In a recent systematic review, the most common mutations
were in p53 (26%), KRAS (12%), PIK3CA (18%), and c-myc (53%)
genes. Loss of heterozygosity was found in 30% of cases.13

Frumovitz et al29 identified 44 patients with pure or mixed small-
cell cervical cancer. All patients underwent mutational analysis
using next-generation sequencing of mutational hotspots in 50
cancer-related genes. Thirty-five mutations were identified in 24
patients (55%). Most patients (63%) had one mutation, 29% had
two mutations, and 8% had 3 mutations. In all 44 patients, the
most common mutations were in PIK3CA (n=8; 18%), KRAS (n=6,
14%), and TP53 (n=5, 11%). No other mutation was found in >7%
of specimens. Of the 24 patients who had a mutation, 21 (88%) had
at least one alteration for which there currently exists a class of
biological agents targeting that mutation. Although in patients with
small-cell cervical cancer no common mutation was found, almost
half had at least one actionable mutation for targeted therapy.
This is of great importance for patients with recurrent disease where
treatment options are limited. Mutational analysis may help to
select patients for phase I trials or even off-label therapies.

Our group at MD Anderson Cancer Center recently evaluated the
presence of PD-L1 receptors as well as mismatch repair proteins
(MHL1, MSH2, MSH6, and PMS2) in specimens of high-grade
neuroendocrine cervical cancer. In 23 samples tested for PD-L1
expression, 22 (96%) were negative for the receptor. In 25 spec-
ims evaluated for microsatellite instability, all (100%) showed
intact expression of mismatch repair proteins. These findings
suggest a low likelihood of response to PD-1/PD-L1 inhibitors in
the recurrent setting.30

DIAGNOSIS, STAGING, AND IMAGING

Cervical neuroendocrine carcinomas are staged using the same
International Federation of Gynecology and Obstetrics (FIGO)
staging system used for other cervical histologies. The 2018 FIGO
staging system allows for imaging and pathologic findings to modify
tumor stage.31 This is a crucial change for cervical neuroendocrine
carcinomas given the higher rate of nodal and distant metastasis at
the time of diagnoses. The Society of Gynecologic Oncology (SGO)
guideline recommends that given the high rate of distant meta-
static disease in neuroendocrine tumors, imaging evaluation should
include either a computed tomography (CT) or positron emission
tomography (PET)/CT scan.32

For disease confined to the cervix both clinically and radiologi-
cally, pelvic magnetic resonance imaging (MRI) is considered the
best imaging method for tumors greater than 10 mm to evaluate
tumor size and local extension of the disease.33 For women with
seemingly early-stage disease, we recommend obtaining a pelvic
MRI in order to determine if a patient is a candidate for surgical
treatment. Another option is transrectal ultrasonography, although
this is not commonly utilized at most centers. For other histo-
logic subtypes of cervical cancer (squamous, adenocarcinoma, or
adenosquamous), when performed by expert radiologists, the accu-
racies of transrectal ultrasound compared with MRI in detecting para-
metrial infiltration was 98.9% and 94.7% (p<0.021), respectively. It
was considered superior to MRI in detecting small tumors (<1 cm2)
(90.5% vs 81.1% (p=0.049), respectively). Transrectal ultrasound
was also found to be superior to MRI when evaluating residual
tumors after conization (93.7% vs 83.2%, respectively; p<0.006).34

Our approach at MD Anderson Cancer Center is to conduct a
thorough gynecologic examination, including a rectovaginal exam,
and a PET/CT scan for initial radiologic staging. For patients with
disease seemingly limited to the cervix, we recommend either an
MRI or ultrasound pre-operatively to evaluate local disease exten-
sion. For those patients dispositioned to primary radiation therapy, a
pelvic MRI is frequently performed for radiation planning.

We only perform brain imaging in the presence of lung or liver
metastases or neurologic symptoms. We strongly encourage physi-
cians to stage patients using the FIGO 2018 system as it might
classify patients in a more ‘real’accurate stage according to tumor
size, nodal spread, and distant metastases. This allows more
tailored treatment and provides patients with a more definitive
discussion on prognosis when taking into account nodal status and
distant organ involvement (Figure 1).

DISEASE PROGNOSIS

Several studies have reported age,5 lymph node metastases,11 35–37
race,8 smoking,38 pure small-cell histology, LUSI,11 and tumor
size35 36 38 as independent prognostic factors for neuroendocrine
cervical carcinomas. However, stage was the most commonly cited
poor prognostic factor in a majority of studies.7 5 35–37 39 Overall, the
prognosis of women with neuroendocrine cervical cancer remains
poor despite multimodal treatment plans, with a 5-year survival
rate of 36% and a median overall survival between 22 and 25 months.5 When considering early (I-II) versus advanced stage (III-IV), the 5-year overall survival rates are 31%–51% and 0%–7%, respectively7 38 40 (Table 2).

PRIMARY TREATMENT

Early-Stage Disease

In a recently published systematic review that included 3,538 neuroendocrine patients, the most common primary treatment was radical surgery combined with chemotherapy, either as neoadjuvant or adjuvant (40/48 studies).13 There was no standard chemotherapy regimen but platinum and etoposide was the most commonly used treatment (24/40 studies). Radiotherapy-based treatment schemes were also commonly utilized in the upfront setting for early-stage disease (15/48 studies).

Both the SGO32 and the Gynecologic Cancer Intergroup (GOG)61 recommend multimodal therapy for all stages of neuroendocrine tumors of the cervix, and the majority of patients receive some combination of surgery, radiation, and chemotherapy. For early-stage disease (tumors ≤4 cm) and negative nodes on imaging, radical hysterectomy and pelvic lymphadenectomy followed by chemotherapy with platinum and etoposide is the primary management recommended with consideration for additional radiotherapy.32

Surgery

There are no prospective studies comparing surgery with chemoradiation therapy for early-stage, resectable tumors. A study from Ishikawa et al11 of 93 patients with stage I-II high-grade neuroendocrine carcinoma of the cervix showed a median overall survival of 111 months and a disease-free survival of 47 months. There was significant variability as to how patients were treated, but the majority of patients (88%; 95%) underwent radical surgery, and five (5%) had definitive radiotherapy. Of the patients receiving surgery, 37 (40%) underwent radical surgery and pelvic lymphadenectomy followed by post-operative chemotherapy, 14 (15%) received only surgery, and 25 (27%) received adjuvant radiation or chemoradiation therapy after surgery with or without neoadjuvant chemotherapy before surgery. The hazard ratio (HR) for death for patients who did not undergo surgery as part of their primary treatment was 4.74 (95% CI 1.01 to 15.9). Patients who underwent radical surgery had a better overall survival than those who received definitive radiotherapy (p=0.043). The authors found that most clinicians favored radical surgery followed by adjuvant chemotherapy with an etoposide–platinum or irinotecan–platinum regimen as the optimal treatment of choice for stages I-II high-grade neuroendocrine cervical carcinoma. They also concluded that even in early-stage disease, both local recurrence and distant metastasis occurred frequently, and therefore a more effective treatment strategy is required.

Wang et al42 reviewed 146 patients with stage I-II disease. Of these, 116 (79%) underwent surgery as part of their primary treatment (primary surgery with or without adjuvant therapy, neoadjuvant chemotherapy plus radical hysterectomy, or peri-operative chemotherapy plus radical hysterectomy). The remaining 30 patients did not undergo surgery but rather had radiation therapy with or without chemotherapy. The authors found there was a trend of worse
failure-free survival (41% vs 61%, p=0.086) and cancer-specific survival (48% vs 62%, p=0.122) for those women who had radical surgery compared with those who did not undergo surgery.

From our study of patients with stage I-IIA clinically node-negative disease, those who received definitive chemoradiation had significantly better median event-free survival than those who underwent surgery (median not reached vs 18 months, p=0.04).38 Based on the conflicting results from multiple retrospective studies, the role of surgery for early-stage neuroendocrine tumors seems unclear, but certainly surgery alone without adjuvant chemotherapy and/or radiation is not appropriate for any patient with high-grade neuroendocrine carcinomas of the cervix.

Chemotherapy
Unlike the role of surgery for patients with neuroendocrine tumors, the role of chemotherapy is well established. There is no standard chemotherapy regimen, but due to its pathologic appearances and clinical behaviors similar to small-cell lung cancer, almost all patients with small-cell cervical cancer receive platinum and etoposide as part of their primary therapy.13 For early-stage disease, SGO and GCIG guidelines recommend that patients with complete surgical resection undergo adjuvant chemotherapy.32 41 Zivanovic et al7 reported on 11 patients with early-stage disease (stage IA2-IB2). Of the 11 patients, seven recurred with 86% of patients having distant recurrence. The 3-year distant recurrence-free survival rate was 83% for patients who received chemotherapy and 0% for patients who did not receive chemotherapy as part of their initial treatment (p=0.03). The estimated 3-year overall survival rate was 83% for patients who received chemotherapy and 20% for patients who did not receive chemotherapy as part of their initial treatment (p=0.36). Ishikawa et al11 found that there was an improved disease-free survival in 41 patients who received adjuvant chemotherapy with etoposide-platinum or irinotecan-platinum (HR 0.27, 95% CI 0.10 to 0.69). Also, adjuvant chemotherapy after surgery reduced extra-pelvic recurrences with an OR of 0.37 (95% CI 0.13 to 0.99, p=0.047). A trend toward improved overall survival was also observed when adjuvant chemotherapy was given, but was not statistically significant (HR 0.39, 95% CI 0.15 to 1.01). Lee et al43 reported on a trend in favor of post-operative chemotherapy, and concluded that primary radical surgery followed by adjuvant chemotherapy is the preferred treatment modality for patients with early-stage disease.

Regarding the number of cycles of adjuvant chemotherapy, Pei et al44 retrospectively evaluated 92 patients with stage I-II small-cell carcinomas and found that adjuvant chemotherapy with cisplatin and etoposide for at least five cycles was associated with improved 5-year recurrence-free survival compared with other treatments (68% vs 21%, p<0.001). On multivariate analysis, nodal disease (p<0.003), parametrial extension (p<0.03), and cycles of...
etoposide-cisplatin (cisplatin-etoposide <5 cycles; p<0.001 and no chemotherapy; p<0.004) were independent prognostic factors for disease recurrence.

Pelvic Radiation
A retrospective study including 68 patients with stage IB1-IIA disease reported by Lee et al. suggested that radical hysterectomy followed by adjuvant chemotherapy might be sufficient, as the patients who received chemoradiation did not seem to have a better outcome. In a Japanese multicenter study, the risk of pelvic recurrences after surgery were lower if patients received post-operative radiation (16%) versus patients who did not undergo radiation (25%) but the difference did not reach significance (OR 0.61, 95% CI 0.16 to 2.01). Data from MD Anderson Cancer Center showed that the most common sites for first recurrence are outside the pelvis: 38% in the lungs, 34% in the liver, and 25% presenting as carcinomatosis. Brain recurrence was seen in 25% of patients, and always in the setting of concurrent lung or liver metastases. If patients received radiation therapy, only 21% had in-field recurrences.

There seems to be agreement for early-stage disease that chemotherapy is a key component of treatment, but there are conflicting data on the role of combined surgery and chemoradiation. Protocol at MD Anderson Cancer Center for early-stage disease (tumors ≤4 cm) and negative nodes on pre-operative imaging is open radical hysterectomy and sentinel lymph node (SLN) biopsies. Post-operatively, our treatment recommendations include adjuvant chemoradiation with concurrent cisplatin and etoposide. Chemotherapy regimen consists of cisplatin 60 mg/m² on day 1 and etoposide 100 mg/m² on days 1–3 every 21 days. We decrease the dose of etoposide from 120 to 100 because of the potential toxicity with concurrent radiation. Patients receive two cycles as concurrent chemotherapy, and two to four additional cycles after radiation is completed. Ideally, we aim for patients to be able to receive a total of six cycles. In order to decrease the treatment morbidity associated with post-operative radiotherapy, we only perform SLN mapping and not a full lymphadenectomy when bilateral mapping is achieved (Figure 2).

Prophylactic Cranial Radiation
As treatment for small-cell cervical cancer is derived from the literature on small-cell lung cancer where prophylactic brain irradiation is commonly used for occult metastases, the use of this approach in cervical small-cell tumors has been explored. Weed et al. reported in 2003 that 25% (2/8) of patients with apparent early-stage small-cell cervical cancer developed brain metastases and therefore they proposed prophylactic cranial radiation. However, neither routine brain CT nor cranial radiation are recommended by the SGO guidelines on initial evaluation of asymptomatic patients for small-cell tumors. In their 14-year retrospective experience Hoskins et al. found no cranial metastases. Data from MD Anderson Cancer Center showed that brain metastases were mostly found (7/8 patients, 86%) when other organ metastases were found.

Figure 2  High-grade neuroendocrine cervical carcinoma primary treatment algorithm."FIGO 2018 stage **If possible six cycles of chemotherapy, with a minimum of four cycles total. PLND, pelvic lymph node dissection; SLN mapping, sentinel lymph node mapping.
As part of our primary treatment plan, we do not perform prophylactic whole-brain radiation for asymptomatic patients.

Fertility-Sparing Surgery
With a median age at diagnosis for cervical neuroendocrine tumors of 37 (range 24–77) years, fertility preservation may be a concern both for patients and physicians. The two primary issues in this patient population are ovarian function (both hormonal and reproductive function) and uterine preservation. As it pertains to the question of ovarian preservation and maintaining ovarian function in patients with locally advanced disease, one may consider ovarian transposition. However, even when doing so, after external beam pelvic radiation and/or brachytherapy, ovarian preservation is only maintained in 65% of patients. In addition, particularly in patients with neuroendocrine tumors, there are theoretical concerns regarding residual microscopic disease in the ovary.

There are no data to support the consideration of fertility preservation, such as simple conization or radical trachelectomy, in patients with early-stage disease. Although fertility-preserving surgeries have been reported in women with early-stage small-cell cervical cancer, uterine preservation fertility-sparing treatment is not recommended by the National Comprehensive Cancer Network (NCCN) guidelines for cervical neuroendocrine tumors. Given that most patients are likely to undergo post-operative radiation therapy it would be unusual to consider fertility-sparing surgery.

Our approach at MD Anderson Cancer Center is to delay starting treatment for 2 to 3 weeks in order to allow for ovarian stimulation and egg retrieval. However, ovarian transposition at the time of radical surgery when the plan is for post-operative radiation or prior to definitive chemoradiation for locally advanced disease seems reasonable if the ovaries appear grossly normal intra-operatively. For young women (<50 years old) who are ultimately made menopausal by surgical and/or radiation therapies, we recommend starting hormone replacement therapy. We strongly recommend against any uterine-preserving surgeries such as simple conization, simple trachelectomy, or radical trachelectomy.

Locally Advanced Disease
The SGO guideline recommendation for locally advanced disease or non-surgical candidates is a combination of chemotherapy (platinum and etoposide) and radiation. The most commonly used regimen of cisplatin and etoposide is usually given with cisplatin at 60 to 80 mg/m² on day 1 and etoposide at 80 to 120 mg/m² on days 1 to 3 every 21 to 28 days. Carboplatin may be substituted for cisplatin. Other regimens such as paclitaxel and cisplatin, paclitaxel and carboplatin, and the combination of vincristine, cisplatin, and bleomycin have also been reported.

Hoskins et al. first reported that cisplatin and etoposide could be safely administered concurrently with radiation, with additional cisplatin and etoposide given after completing radiation (chemoradiation followed by chemotherapy). A combined-modality approach of chemoradiation followed by chemotherapy showed that for stage IB–IVA, concurrent chemoradiation with ≥5 cycles of cisplatin-etoposide was associated with improved 5-year disease-free survival (63% vs 13%, p=0.025) and overall survival (75% vs 17%, p=0.016). Distant disease recurrence was the most common type of recurrence (28%) and clinical outcome correlated with initial disease extent. The 3-year disease-free survival rate was 55% and recurrences were uncommon after 3 years.

An abstract presented at56 by Bajaj et al. reported on 73 patients with small-cell cervical carcinoma stage IB2–IVA treated with chemoradiation at seven centers in the United States. The median follow-up time was 19 months with 66% of patients having had a recurrence. Median time to recurrence was 10 months. Recurrence was associated with current smoking (HR 3.32, p<0.01), total equi-effective dose (EQD2) <50 Gy vs 71–80 Gy (HR 3.3, p=0.07), and no brachytherapy (HR 1.5, p=0.25). Brain recurrences were seen in 15% of patients. A decreased hazard of brain recurrence was associated with brachytherapy (HR 0.5, p<0.01), total equi-effective dose (EQD2) >75 Gy (HR 0.11, p=0.04), and cisplatin and etoposide versus cisplatin alone (HR 0.35, p=0.23). Overall survival for all patients was 48%, favoring concurrent chemoradiation followed by adjuvant chemotherapy versus concurrent chemoradiation only (HR 0.49, p=0.10). The number of cycles (cisplatin-etoposide) was associated with improved overall survival (HR 0.45, p=0.01) and decreased recurrence (HR 0.67, p=0.07). Patients receiving cisplatin-etoposide had improved overall survival (59% vs 44%) and a lower recurrence rate (65% vs 74%) compared with those receiving cisplatin only. Robin et al. identified 100 patients in the National Cancer Data Base (NCDB) with locally advanced non-metastatic neuroendocrine cervical cancer that were treated with definitive chemoradiation between 2004 and 2012. In multivariate analysis, the addition of brachytherapy, compared with external beam radiotherapy alone, was associated with an improved median survival of 49 vs 22 months (HR 0.48, 95% CI 0.25 to 0.88). There was no difference in overall survival for patients treated with neoadjuvant chemotherapy versus patients who received chemotherapy started concurrently with radiation (HR 0.85, 95% CI 0.48 to 1.50).

For women with stage IB3–IVA (FIGO 2018) disease or non-surgical candidates, our recommendation at MD Anderson Cancer Center is chemoradiation with cisplatin and etoposide followed by additional chemotherapy with cisplatin and etoposide for a total of four to six cycles (two cycles with radiation and two to four cycles after radiation completed) with a goal of six total cycles. For women with stage IVA disease, palliative chemotherapy with cisplatin and etoposide is recommended (Figure 2).

RECURRENT DISEASE
First Recurrence
For patients with recurrent disease, there is limited consensus on the optimal treatment approach with no standard treatment protocols, and both the SGO and GCIG recommend individualized treatment. Often the standard choices include single-agent topotecan, irinotecan, paclitaxel, or docetaxel as these regimens are commonly used to treat recurrent small-cell lung cancer. Our experience at MD Anderson Cancer Center showed that these single-agent regimens had very low activity in patients with recurrent neuroendocrine cervical cancer.

Our regimen of choice for patients with recurrent or progressive disease who have already been treated with a platinum and etoposide combination is a triplet regimen including topotecan, paclitaxel, and bevacizumab given in the same manner as in Gynecologic Oncology Group (GOG) 240, a phase III study
in patients with recurrent cervical cancer (squamous cell carcinoma, adenocarcinoma, and adenosquamous carcinoma). The rationale for this schema is that first this regimen is tolerable in women who have previously undergone definitive chemoradiation. In addition, as single-agent paclitaxel or topotecan are active and frequently used in recurrent small-cell lung cancer, the combination would presumably be equally as active, and potentially more active than the single-agent regimens. Third, small-cell cervical cancers express the vascular epithelial growth factor receptor over 95% of the time supporting the addition of bevacizumab as an active agent. Finally, all three drugs have been approved by the US Food and Drug Administration for the treatment of recurrent cervical cancer, thus reimbursement is usually not a concern.

Our results with the three-drug combination were published in the largest series of chemotherapy treatments for women with recurrent neuroendocrine cervical carcinoma. Thirteen patients who received topotecan, paclitaxel, and bevacizumab were compared with 21 patients receiving other regimens, mostly platinum-based regimens with or without taxane. The triplet regimen was associated with a significant improvement in outcome. Median progression-free survival was 8 months versus 4 months for the triplet regimen and other regimens, respectively (HR 0.21, 95% CI 0.09 to 0.54). Median overall survival was 9.7 months for the triplet regimen and 9.4 months for patients receiving other regimens (HR 0.53, 95% CI 0.23 to 1.22). For the triplet regimen, 62% of patients received treatment for >6 months and 31% for >12 months versus 19% and 10% of those who received other regimens, respectively.

For first recurrence, patients at MD Anderson Cancer Center are typically treated with the three-drug regimen topotecan (0.75 mg/m² on days 1–3), paclitaxel (175 mg/m² on day 1), and bevacizumab (15 mg/kg on day 1 on a 21-day cycle) (the ‘Texas Cocktail’). As part of our strategy for treating patients with recurrent disease, we order molecular testing, PD-L1, and mismatch repair proteins (MLH1, MSH2, MSH6, and PMS2) testing at the time of first recurrence. We use our own institutional platform or other outside providers that offer similar mutational analyses (eg, Foundation Medicine or Caris). As most patients with recurrence will eventually have progressive disease, this testing allows us to triage patients to a targeted therapy either on-protocol or off-label (Figure 3).

Multiple Recurrences

Immune checkpoint inhibitors and targeted therapies may be beneficial when patients have suffered multiple recurrences; however, the literature is limited in this setting with only three case reports published to date. Paraghamian et al used nivolumab in a patient with recurrent, metastatic, programmed cell death ligand-1 (PD-L1)-negative small-cell neuroendocrine cervical carcinoma, who experienced a complete response. Sharabi et al reported a patient with metastatic, chemotherapy-refractory neuroendocrine carcinoma with bowel obstruction due to a large tumor burden. Liquid biopsy demonstrated a high number of tumor mutations. The patient was treated with radiotherapy combined with nivolumab and experienced a near-complete systemic resolution of disease for at least 10 months. Our experience with checkpoint inhibitors, however, has been disappointing with no responders (article in preparation). As mentioned above, high-grade neuroendocrine cervical tumors are almost always PD-L1-negative and microsatellite-stable so checkpoint inhibitors should be used with caution.
Lyons et al. used the mitogen-activated protein kinase 1 (MEK)-inhibitor trametinib in a woman with recurrent small-cell neuroendocrine cervical carcinoma whose tumor was found to have a KRAS mutation, and had a complete radiologic response after three cycles. At MD Anderson Cancer Center, for patients whose disease progresses after treatment with topotecan, paclitaxel, and bevacizumab, the recommendation is targeted therapy on a clinical trial. When patients progress after targeted or immunotherapy treatment, we recommend a phase I clinical trial or hospice care.

SURVEILLANCE

The recommended surveillance for squamous, adenocarcinoma, and adenosquamous carcinomas consists of a physical examination, including a pelvic exam, every 3 to 6 months for 2 years and then every 6 to 12 months for an additional 3 years. In addition, a Papanicolaou test should be performed annually. For cervical neuroendocrine tumors, the SGO guidelines recommend physical examination and symptoms review with periodic full-body imaging with either CT or PET/CT scan. The guideline does not give any recommendation on the frequency of such follow-up and no data exist to make definitive recommendations.

Given the aggressive nature of cervical neuroendocrine tumors, our surveillance plan at MD Anderson Cancer Center consists of a physical examination (with a pelvic exam) every 3 months for the first 2 years and then every 4 to 6 months for the next 3 years. In addition, we order routine radiologic imaging of the chest, abdomen, and pelvis, with either a CT scan or PET/CT scan at each visit.

TUMOR REGISTRY AT MD ANDERSON CANCER CENTER

Large studies in patients with uncommon or rare tumors are difficult with limited prospective data with which to guide decisions. For women with small- and large-cell neuroendocrine cervical carcinoma there currently exists a rich network on social media sites, making recruitment to protocols much more feasible. In 2013, we established a Neuroendocrine Cervical Tumor Registry (NeCTuR) at The University of Texas MD Anderson Cancer Center where data are prospectively entered. To date, we have enrolled 252 patients with 110 patients from outside MD Anderson. We continue to enroll and follow patients from around the world. Interested providers may inquire about participation by emailing nectur@mdanderson.org.

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REFERENCES


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