Imaging of Merkel Cell Carcinoma: What Imaging Experts Should Know

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Abbreviations: FDG = fluorodeoxyglucose, MCC = Merkel cell carcinoma, MCPyV = Merkel cell polyomavirus, NCCN = National Comprehensive Cancer Network, NET = neuroendocrine tumor, SLNB = sentinel lymph node biopsy, SSTR = somatostatin receptor

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Merkel cell carcinoma (MCC) is a rare and aggressive cutaneous neuroendocrine tumor with a higher mortality rate than melanoma. Approximately 40% of MCC patients have nodal or distant metastasis at initial presentation, and one-third of patients will develop distant metastatic disease over their clinical course. Although MCC is rare, its incidence has been steadily increasing. Furthermore, the immunogenicity of MCC and its diagnostic and therapeutic application have made MCC one of the most rapidly developing topics in dermatology and oncology. Owing to the aggressive and complex nature of MCC, a multidisciplinary approach is necessary for management of this tumor, including dermatologists, surgeons, radiation oncologists, medical oncologists, pathologists, radiologists, and nuclear medicine physicians. Imaging plays a crucial role in diagnosis, planning for surgery or radiation therapy, and assessment of treatment response and surveillance. However, MCC is still not well recognized among radiologists and nuclear medicine physicians, likely owing to its rarity. The purpose of this review is to raise awareness of MCC among imaging experts by describing the epidemiology, pathophysiology, and clinical features of MCC and current clinical management with a focus on the role of imaging. The authors highlight imaging findings characteristic of MCC, as well as the clinical significance of CT, MRI, sentinel lymph node mapping, fluorine 18 fluorodeoxyglucose PET/CT, and other nuclear medicine studies such as bone scintigraphy and somatostatin receptor scintigraphy.

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SA-CME LEARNING OBJECTIVES

After completing this journal-based SA-CME activity, participants will be able to:

■ Describe the epidemiology, pathophysiology, unique clinical manifestation, and current treatment strategy of MCC.
■ Discuss the important role of imaging in management of MCC.
■ List the unique imaging characteristics of MCC as an aggressive cutaneous NET with somatostatin expression.

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Introduction

Merkel cell carcinoma (MCC) is a rare and highly aggressive cutaneous malignant neuroendocrine tumor (NET). MCC was named on the basis of its ultrastructural and immunophenotypic resemblance to sensory Merkel cells in the base of the epidermis to the dermis, which serve as touch receptors (Fig 1) (1,2). However, the cell of origin for MCC is not yet fully understood. MCCs are most frequently found in the dermis but can manifest clinically in any layer of the skin, from intraepidermal to subcutaneous tissues (3).

Although MCC is a rare skin cancer, its incidence has been increasing strikingly. In 2013, the incidence of MCC in the United States was reported to be approximately 2500 cases per year. This is expected to rise to 2800 cases in 2020 and 3200 cases in 2025 owing to the aging baby boomer population (4). The increasing incidence is noticeable even compared with that of other neoplasms. During 2000–2013, the total number of MCC cases was reported to have increased by 95%, while all solid tumor cases increased by 15% and melanoma by 57% (4).
89% of MCC patients present with three or more of the AEIOU characteristics: asymptomatic or lack of tenderness, expanding rapidly, immune suppression, older than 50 years, and ultraviolet radiation–exposed site on a person with fair skin (13). However, the dermatologic manifestations of MCC are nonspecific and could be initially interpreted as a benign cutaneous lesion such as a cyst, folliculitis, or a lipoma. A review of 195 MCC patients reported that the majority of MCC lesions (56%) were presumed to be benign at biopsy (13). Therefore, MCC-specific management is usually initiated after a surprising pathology result has been received by the clinician.

Causative Factors
The Merkel cell polyomavirus (MCPyV) was discovered in 2008 (12). Although MCPyV infection itself is common and does not directly cause
problems; it is causally linked to 80% of MCC cases in the United States, whereas the remaining 20% are caused by extensive ultraviolet radiation mutations (12).

**MCPyV Serology Test**

Studies have shown the clinical utility of a serology test detecting antibodies to MCPyV oncoproteins (T antigens). About half of MCC patients produce antibodies to MCPyV oncoproteins at diagnosis (16). A baseline MCPyV oncoprotein antibody test is useful for all newly diagnosed MCC patients to assist in following the patient during his or her clinical course. The clinical practice guidelines of the National Comprehensive Cancer Network (NCCN) recommend considering quantification of MCPyV oncoprotein antibodies as part of the initial workup for patients with MCC (17).

If patients are seropositive at diagnosis, a rising titer may be an early indicator of recurrence (Fig 3). For seropositive patients, this test may be used for surveillance and may reduce imaging studies, decreasing the cost and radiation exposure of imaging and the toxic effects of contrast material (16). If patients are seronegative at initial diagnosis, their risk of recurrence is 42% higher than that of seropositive patients, and close surveillance with more frequent imaging is needed (16). The antibody test is not used beyond this baseline determination in patients who do not produce antibodies.

**Staging and Prognosis**

The American Joint Committee on Cancer (AJCC) TNM staging system, eighth edition, is the most widely used staging system (9). Each stage is divided into a clinical stage and a pathologic stage. Clinical detection of disease may be via inspection, palpation, or imaging. Pathologic detection of nodal or distant metastatic disease may be via sentinel lymph node biopsy (SLNB), lymphadenectomy, or fine-needle or standard biopsy of the suspected metastasis.

Stage I tumors are equal to or smaller than 2 cm in maximum dimension. Stage II tumors have
A maximum dimension of greater than 2 cm or bone, muscle, fascia, or cartilage invasion. Stage III tumors have regional lymph node metastasis. Stage IV tumors have distant metastasis.

Estimated 5-year overall survival is 51%, 35%, and 14% for local, nodal, and distant disease, respectively (9). In the eighth edition AJCC staging system, patients with stage pIII disease with no identifiable primary or an unknown primary are included in the stage pIIIA subgroup because they have a significantly better prognosis than patients with nodal disease and a known primary (stage pIIIB) (9). This may be because improved antitumor immunity eliminated the primary tumor and also targets residual disease (18).

**Role of Imaging**

MCC is a unique skin cancer that has many interesting imaging features. Radiology and nuclear medicine play an essential role in its management. Figure 4 is a flowchart for management of MCC from the NCCN clinical practice guidelines (17).

**Imaging of Primary or Regional MCC**

The role of imaging in MCC management is primarily to detect regional nodal or distant metastases, as many patients are referred for imaging for staging. The primary lesion may or may not have been resected at the time of imaging. In locally advanced disease or distant metastatic disease, imaging plays an important role in determining the location of the lesion and identifying local-regional invasion to surrounding organs to guide surgical or radiation therapy planning. This is especially essential in head and neck disease, owing to its anatomic complexity. The primary MCC lesion may be detected incidentally at routine imaging for other reasons (19). Careful assessment of the surrounding cutaneous region is needed, as MCC has a tendency to “jump” discontinuously to adjacent normal-appearing skin, and in-transit and satellite cutaneous metastases can occur (20).

Although MCC has nonspecific imaging features, there are evocative findings associated with

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**Figure 4. Management of MCC.**

- Imaging is encouraged whenever metastatic or unresectable disease is suspected on the basis of the history and physical examination findings. The most reliable staging tool for identifying subclinical nodal disease is SLNB.
- Quantitation of MCPyV oncoprotein antibodies may be considered part of the initial workup. Seronegative patients may have a higher risk of recurrence; in seropositive patients, a rising titer may be an early indicator of recurrence.
- Consider observation of the primary site in cases where the primary tumor is small (eg, <1 cm) and widely excised with no other adverse risk factors such as lymphovascular invasion or immunosuppression. FNA = fine-needle aspiration, − = negative, + = positive. (Adapted and reprinted, with permission, from reference 17.)

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<table>
<thead>
<tr>
<th>Clinical N0</th>
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<tbody>
<tr>
<td>- Management of the primary tumor: Wide excision, followed by Adjuvant radiation therapy to the primary tumor site or Consider observation⁵</td>
</tr>
<tr>
<td>- Management of the draining nodal basin: Sentinel lymph node biopsy (SLNB) with appropriate immunopanel.</td>
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<td>-SLNB (+) baseline imaging if not performed</td>
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<tr>
<td>-SLNB (−) Observation or Consider radiation therapy to the nodal basin in high-risk patients</td>
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<td>- Management of the primary tumor: Follow clinical N0 pathway</td>
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<td>- Management of the draining nodal basin</td>
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<tr>
<td>FNA core biopsy Immunopanel</td>
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<tr>
<td>Node biopsy positive</td>
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<td>Node biopsy negative</td>
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<td>Follow appropriate clinical N0 pathway</td>
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<th>Clinical M1</th>
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<tr>
<td>Multidisciplinary tumor board consultation</td>
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<td>Clinical trial preferred if available</td>
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<td>Consider any of the following therapies or combinations of:</td>
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<td>- Systemic therapy</td>
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<td>- Radiation therapy</td>
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<td>- Surgery</td>
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<td>or</td>
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<td>Best supportive care</td>
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[Diagrams and flowcharts are not transcribed in this text format but are included in the original document.]
it that are helpful. Common features of a primary MCC include a cutaneous or subcutaneous nodule (sessile or fungating cutaneous nodule or mass) and focal or diffuse skin thickening (Fig 5). It can manifest as a perifascial muscular or intramuscular mass (Fig 6). Necrosis is common, whereas calcifications are not.

At CT, cutaneous fat stranding near the primary MCC may suggest the presence of engorgement and edema from lymphatic invasion (21). Often, there is large lymph node enlargement with fine compressed or retained fatty tissue (22). US can be used, especially for MCC in the head and neck (23). US evaluation of primary MCC usually demonstrates hypoechoic nodules arising from the dermis and extending into the subcutaneous fat, with variable degrees of posterior acoustic transmission (21). US enables real-time imaging with possible simultaneous fine-needle or core-needle biopsy and provides a concise workup at lower cost.

At MRI, MCC is hypo- to isointense on T1-weighted images and iso- to hyperintense on T2-weighted images or fat-saturated T2-weighted images (24,25). On gadolinium-enhanced images, the lesion usually has diffuse or heterogeneous enhancement. Large lesions can have inhomogeneous signal intensity on both T1- and T2-weighted images. Focal central increased signal intensity within large lesions on T2-weighted images has been described as being associated with central necrosis and hemorrhage (24).

At histologic analysis, the skin, subcutaneous mass, and reticular stranding are found to be involved by lymphangitic carcinomatosis and soft-tissue lymphatic metastases. MCC is a highly metabolic tumor and demonstrates intense fluorine 18 (18F) fluorodeoxyglucose (FDG) uptake at PET. The mean maximum SUV (standardized uptake value) at 18F-FDG PET for a primary MCC is reported to be 4–6.5 (26,27). Also, MCC often expresses increased somatostatin receptor (SSTR) on its surface and therefore shows increased radiotracer uptake at somatostatin-seeking scintigraphy and PET/CT (discussed later).

Sentinel Lymph Node Biopsy
SLNB is the most reliable tool for investigating subclinical nodal metastases. The NCCN guidelines recommend SLNB for all clinically node-negative patients who are fit for surgery because it is an important staging tool and in combination with subsequent treatment affects regional control for those with positive sentinel lymph nodes (28). Patients with a positive SLNB result have a higher risk of in-transit recurrence and may benefit from adjuvant radiation therapy with inclusion of the in-transit field in amenable cases.
In patients with stage I and II MCC, SLNB is more sensitive than \[^{18}F\]FDG PET/CT (30). SLNB should be performed before wide local excision or Mohs micrographic surgery because surgical excision before SLNB may alter the lymphatic drainage patterns.

At our institution, technetium 99m (\[^{99m}Tc\]) sulfur colloid (1 mCi [\(37 \times 10^6\) Bq], 0.2 \(\mu\)m filtered) is used for lymphoscintigraphic sentinel lymph node mapping (Fig 7). The dose is injected intradermally at four locations around the primary tumor or tumor biopsy site. Multiple static images of the expected lymph nodal basin are obtained to localize the sentinel lymph node. A handheld gamma probe is used to confirm the node.

For anatomically complex areas or when planar images are difficult to interpret, SPECT/CT can be performed to localize the sentinel lymph node. In addition, the higher sensitivity and spatial and contrast resolution of SPECT allow visualization of foci undetected on planar images. At our institution, SPECT/CT is routinely performed for MCC of the head and neck owing to its anatomic complexity. The sentinel lymph node is further localized with blue dye in the operating room. The combination of the two mapping methods is an accurate approach and widely performed.

**Detecting Nodal or Distant Metastasis**

MCC has a high propensity for nodal metastasis, with 27%–31% of patients presenting with clinical nodal disease. In addition, another 16%–38% have occult nodal metastasis demonstrated at SLNB (29). On the basis of various large patient databases, the most frequent distant metastasis site is a nonregional lymph node, occurring in 33%–85% of cases, followed by the skin or subcutaneous tissue, bones, liver, and lung or pleura (26,31–33). Less frequent sites include the pancreas (Fig 8), muscle (Figs 6, 9), central nervous system (Fig 10), adrenal glands, heart (Fig 11), gastrointestinal tract (Fig 12), retroperitoneum, urinary bladder, and breast.

There is no imaging feature specific for MCC metastasis with any modality, and US, CT, MRI, or PET/CT can be used as indicated. Imaging findings of MCC metastasis have been described only in several case reports, case series, or review articles. MCC metastasis can manifest as lymph adenopathy, a cutaneous or subcutaneous soft-tissue nodule, a pulmonary nodule or masses, or related chest wall invasion.

A solid visceral metastasis in the abdomen tends to be hypervascular with peripheral rim enhancement at contrast-enhanced CT or MRI and hypointense or hypointense in the portal venous phase (19,34,35) (Fig 8). Metastasis to hollow organs such as the stomach or bowel can manifest as wall thickening or a wall mass, ulceration (19), or bowel thickening with aneurysmal dilatation (Fig 12). A bone metastasis can be osteolytic or osteoblastic at radiography or CT (36). MRI can be used for detection of bone marrow involvement and extraosseous extension (37).
Cardiac metastasis is rare. Owing to its invasiveness, it is difficult to achieve pathologic confirmation with biopsy. Thus, imaging is important to establish a diagnosis of this rare manifestation (Fig 11).

$^{18}$F-FDG PET is increasingly used for detection of distant metastasis. MCC is a hypermetabolic tumor with a reported SUV$_{\text{max}}$ (maximum standardized uptake value) for distant metastasis of 7.2–11.5 (26,27). Many studies have indicated favorable performance of $^{18}$F-FDG PET/CT (discussed later). However, $^{18}$F-FDG PET has

Figure 7. MCC in a 72-year-old woman referred for sentinel lymph node mapping. (a) Anterior and lateral images of the head and neck from $^{99m}$Tc sulfur colloid scintigraphy show a large focus of intense radiotracer uptake (black arrows) within the right cheek, which corresponds to the intradermal injection site around the primary tumor. A fainter smaller focus of radiotracer uptake (yellow arrows) below the injection site suggests the sentinel lymph node. (b) Axial SPECT/CT image shows intense radiotracer uptake at the injection site along the right cheek (arrow). (c) Axial SPECT/CT image shows a focus of increased uptake in the sentinel lymph node adjacent to the right parotid gland (arrow). (d) Axial SPECT/CT image of the neck below the level of the parotid gland shows an additional focus of increased radiotracer uptake in a smaller sentinel lymph node in the submandibular region (arrowhead).

Figure 8. Pancreatic metastasis in a 30-year-old man with MCC of the right lower extremity who was referred for staging. Axial contrast-enhanced CT image shows a hypoattenuating exophytic mass (arrows) arising from the pancreas (P). No dilatation of the main pancreatic duct is noted.
Figure 9. Intramuscular metastasis in a 55-year-old man with MCC of the lower extremity. (a) Whole-body posterior image from indium 111 ($^{111}$In)–pentetreotide scintigraphy shows faint increased radiotracer uptake along the left side of the pelvis (arrow). Rt = right. (b) Coronal SPECT/CT image shows increased radiotracer uptake in a left iliacus muscle metastasis (arrows).

Figure 10. Central nervous system metastases in four patients. (a) Brain metastasis in a 67-year-old woman with MCC of the right mandible that was originally staged IIB. Axial contrast-enhanced MR image shows an enhancing lesion (arrow) in the corpus callosum. (b) Complex cystic metastasis in a 65-year-old man with MCC of the forehead. Axial contrast-enhanced MR image shows a predominantly cystic lesion (arrow) in the right frontal lobe with an enhancing solid component (arrowhead). (c) Hemorrhagic metastasis in a 63-year-old man with MCC. Axial susceptibility-weighted image shows a heterogeneous right cerebellar mass (arrow) with areas of low signal intensity indicative of hemorrhage. (d) Epidural metastasis in a 65-year-old man with MCC. Axial contrast-enhanced MR image of the spine shows an enhancing epidural nodule (arrow).

decreased sensitivity for brain metastasis owing to intense physiologic FDG activity in the brain, in which case contrast-enhanced brain MRI should be performed (38).

The reported imaging characteristics of brain metastasis from MCC include an enhancing homogeneous or heterogeneous parenchymal nodule or masses, a cystic mass with or without a mural enhancing nodule, leptomeningeal metastasis, direct parenchymal invasion from a skull metastasis, or a parenchymal brain metastasis with skull infiltration (Fig 10). Necrosis or hemorrhage can be seen. Lesions may or may
not be associated with surrounding vasogenic edema (39–41). Contrast-enhanced MRI is also preferred for diagnosis of spinal metastasis (Fig 10d), and both intradural and extradural metastases to the spine have been reported (40,42–45).

**Treatment**

Standard treatment of MCC is generally surgery, radiation therapy, or systemic therapy, depending on staging. According to the NCCN guidelines, wide local excision of the primary lesion is a component of initial management, but surgical margins should be balanced with the morbidity of surgery (17,28). After wide local resection, observation may be reasonable for patients with small primary lesions (eg, <1 cm) that have been widely excised and who present with no risk factors such as lymphovascular invasion or immunosuppression (17).

Adjuvant radiation therapy to the primary site is generally recommended for all other cases, especially for patients with microscopic or grossly positive margins or other risk factors for recurrence (17). If there is regional nodal involvement (confirmed with biopsy), node dissection and/or radiation therapy to the lymph nodes should be performed. Multidisciplinary tumor board consultation is recommended, and a clinical trial for adjuvant therapy is preferred if available (17).

For MCC patients with distant metastasis, treatment in the context of a clinical trial is preferred if available. Other options to consider include systemic therapy, radiation therapy, and/or surgery (17,28,46). Until 2016, chemotherapy (carboplatin/cisplatin and etoposide or a combination of cyclophosphamide and doxorubicin) was the only systemic treatment option listed in the NCCN guidelines for MCC. However, responses are not durable, with median progression-free survival of about 94 days, and toxic effects are considerable (47).

Recently, immunotherapy such as targeting the PD-1 (programmed cell death 1) pathway was found to activate the immune system to attack tumor cells. PD-1 or PD-L1 (programmed death ligand 1) antibody therapy has demonstrated promising long-term effects (48). On the basis of preliminary data from nonrandomized trials showing promising response rates to PD-1 or PD-L1 blockade, several of these agents (avelumab, nivolumab, and pembrolizumab) are included as recommended systemic therapy options for treatment of disseminated disease in the current NCCN guidelines (17). These agents are the preferred options for patients who select systemic therapy as part of their treatment of disseminated disease (17).

**Follow-up or Assessment of Treatment Response**

Currently, there is no consensus regarding the accuracy and utility of imaging modalities for follow-up or assessment of treatment response. For follow-up imaging, the NCCN guidelines...
recommend imaging studies “as clinically indicated,” such as in the case of emergent adenopathy, unexplained changes in liver function test results, or development of new suspicious symptoms. In addition, routine imaging should be considered for high-risk patients (eg, those stage IIIB or higher or with immunosuppression) (28).

The guideline states that “whole body PET with fused axial imaging (CT or MR) or neck/chest/abdomen/pelvis CT with contrast, with or without brain MRI, may be useful to identify and quantify regional and distant metastases” (17). In clinical practice, interpreting PET/CT or PET/MR images would require reviewing raw PET images and attenuation-corrected CT or MR images in multiple planes in addition to fused axial images. It is important for radiologists to guide the clinical team in terms of which modality is most appropriate in each case.

On the basis of the current guidelines, more MCC patients will receive immunotherapy rather than cytotoxic chemotherapy (17). If imaging is performed in MCC patients receiving immunotherapy, these patients may have an unconventional response, and immune-modified Response Evaluation Criteria in Solid Tumors (imRECIST) may better capture the treatment response (49).

For follow-up of MCC patients with bone metastasis receiving radiation therapy or systemic therapy, metabolic change at \(^{18}\text{F}-\text{FDG PET}\) better correlates with true disease status than does change at CT. New sclerotic change at CT may be due to treatment response or progressive disease (50). This may be misleading in a sensitive situation like a patient with MCC metastasis receiving radiation therapy or systemic therapy. A decrease in the extent or intensity of FDG activity in the face of increasing sclerosis at CT usually heralds healing (50).

Role of Specific Nuclear Medicine Studies

\(^{18}\text{F}-\text{FDG PET}\)

\(^{18}\text{F}-\text{FDG PET/CT}\) is a good modality for staging and is increasingly used, as MCC is usually
very FDG avid (Figs 11, 13). According to a large meta-analysis of 10 studies comprising 328 MCC patients who underwent 549 $^{18}$F-FDG PET/CT scans, $^{18}$F-FDG PET/CT has sensitivity of 90% and specificity of 98%, allowing pathologic and nonpathologic reference standards (ie, use of clinical or radiologic follow-up as a standard) (51). $^{18}$F-FDG PET/CT is useful for detection of nodal involvement and distant metastasis. $^{18}$F-FDG PET/CT demonstrates more bone metastases than CT (52).

Studies have reported that initial or baseline staging $^{18}$F-FDG PET/CT significantly influenced treatment decisions and management in up to around 40% of patients (26,27,32,53–55). A prospective study of 58 MCC patients (AJCC [American Joint Committee on Cancer] version 7 stages IIA–IIIB) demonstrated that staging $^{18}$F-FDG PET significantly influenced treatment decisions in 27.6% of patients, with disease in 25.9% of patients being upstaged whereas no disease was downstaged (54). In this study, posttreatment PET was not found to be prognostic (54). Another large study of 270 scans in 97 MCC patients reported that initial $^{18}$F-FDG PET/CT led to upstaging in 16% of patients (26).

A retrospective study of 102 consecutive MCC patients demonstrated that initial staging PET had a high clinical impact (the PET results changed the primary treatment modality or intent) in 22% of patients and a medium impact (the treatment modality was unchanged, but the radiation therapy technique or dose was altered) in 15% (53). These authors also reported that the PET stage was significantly associated with overall survival. A smaller retrospective study of 23 MCC patients found that initial PET/CT led to a change in staging in seven of 18 patients (39%) and a change in treatment in six of 18 patients (33%) (55).

A retrospective study of 62 MCC patients who were treated definitively and underwent posttreatment PET found that restaging $^{18}$F-FDG PET had a high impact (the PET results changed the primary treatment modality or intent) in 24 of 53 cases (45%) and a medium impact (the treatment modality was unchanged, but the radiation therapy technique or dose was altered) in six of 53 cases (11%) (56). Metabolic response was significantly associated with overall survival (56).

More recently, there have been several case reports suggesting the usefulness of $^{18}$F-FDG PET/CT for assessment of immunotherapy response (57,58). Caution is needed for interpretation because $^{18}$F-FDG can also accumulate in inflammation or infection, and careful correlation with CT is necessary.

**Bone Scintigraphy**

Bone scintigraphy using bone-seeking radiotracers such as $^{99m}$Tc–methylene diphosphonate (MDP) can be used to detect bone metastasis (Fig 14). It is a relatively concise imaging technique with low cost, and whole-body evaluation is possible. The tracer is a nonspecific radiotracer binding to the hydroxyapatite of the osseous matrix at a site of active bone remodeling (59).

Bone metastasis responding to treatment may cause more bone remodeling or healing and therefore more radiotracer uptake at bone scintigraphy, mimicking progression (flare phenomenon) (60). Multiple meta-analyses have shown that $^{18}$F-FDG PET has higher sensitivity than bone scanning for detecting bone metastasis in other cancers (61–63).

**Somatostatin Receptor–seeking Nuclear Medicine**

MCC is a unique cutaneous NET and exhibits SSTR on the tumor cell surface. Like other NETs, MCC has a higher affinity for SSTR types 2A and 5 (64). Owing to these characteristics, a certain radioisotope can be linked to a peptide that binds to SSTR and used for gamma camera imaging or PET or peptide receptor radiation therapy (Fig 15).

Indium 111 ($^{111}$In)–pentetreotide (OcetreScan; Curium, Maryland Heights, Mo) has long been used for scintigraphic imaging of NETs including MCC (Figs 16, 17). It has high affinity for SSTR types 2 and 5, to a lesser extent for type 3, but not for types 1 and 4 (59).

Gallium 68 ($^{68}$Ga) DOTA (tetraazacyclododecane tetraacetic acid)–Tyr$^3$-octreotate
It is reported that SSTR analog PET has higher sensitivity for bone, soft-tissue, and brain disease than CT but lower sensitivity for liver and lung disease, stressing the importance of combined imaging. Somatostatin analog peptides labeled with different radioisotopes that can be used in diagnostic imaging (\(^{111}\)In-pentetreotide, gallium 68 \(^{68}\)Ga–DOTATATE) or peptide receptor radionuclide therapy (PRRT) (lutetium 177 \(^{177}\)Lu, yttrium 90 \(^{90}\)Y).

Widespread metastases in a 69-year-old woman with MCC of the left arm. (a) Whole-body \(^{111}\)In-pentetreotide image shows several foci of increased radiotracer uptake compatible with metastasis having SSTR expression (arrows), some with higher uptake than the liver. The areas of metastasis include the right side of the calvaria (A), right scapula (B), sternum (C), left subpectoral region (D), left proximal humerus (E), right cardiophrenic angle (F), right external iliac node (G), and soft tissue around the right proximal femur (H). (b–d) Axial SPECT/CT images show the sternal metastasis (arrow in b), the right external iliac node metastasis (arrow in c), and the metastasis in the soft tissue around the right femur (arrow in d).

\(^{68}\)G DOTA–Na\(^{3}\)-octreotide (DOTANOC), and \(^{68}\)Ga DOTA–TyI\(^{-}\)-octreotide (DOTATOC) are PET tracers with high affinity for SSTR and can be used for imaging of NETs including MCC (Fig 18). In the United States, the Food and Drug Administration (FDA) recently approved \(^{68}\)Ga-DOTATATE for imaging of NETs. The advantages of \(^{68}\)Ga-labeled somatostatin analog PET over \(^{111}\)In-pentetreotide imaging are higher spatial resolution and sensitivity, shorter scanning time, and the fact that quantification of several parameters such as standardized uptake value (SUV) is possible. With \(^{68}\)Ga-labeled somatostatin analog PET, a patient can be scanned 45–60 minutes after radiotracer administration, whereas \(^{111}\)In-labeled pentetreotide imaging is typically performed 24–72 hours after radiotracer administration. It is reported that SSTR analog PET has higher sensitivity for bone, soft-tissue, and brain disease than CT but lower sensitivity for liver and lung disease, stressing the importance of combined imaging.
PET/CT (65). Physiologic radiotracer activity in the liver, spleen, or kidneys may interfere with evaluation of these sites.

Decreased sensitivity for lung lesions is likely due to the smaller size of lung lesions, which are below the resolution of PET. Evaluation of SSTR expression is important to validate targeted molecular therapy such as somatostatin analog or peptide receptor radionuclide therapy (PRRT).

18F-FDG PET versus 68Ga–Somatostatin Analog PET
Currently, 18F-FDG is the only PET tracer indicated for imaging of MCC in the NCCN guidelines. There is no consensus on use of 68Ga–somatostatin analog PET for imaging of MCC. In gastrointestinal NET, there is an inverse relationship between the World Health Organization (WHO) tumor grade and the European Neuroendocrine Tumor Society (ENETS) tumor grade based on Ki-67 and SSTR expression rate (66). Ki-67 is a marker of cellular proliferation: well-differentiated tumors have lower Ki-67 level (<3%), whereas poorly differentiated tumors have higher Ki-67 level (>20%) (67). In non-MCC NETs, 68Ga–somatostatin analog PET/CT is recommended for imaging lower-grade tumors with lower Ki-67 expression and 18F-FDG PET is recommended for imaging higher-grade more aggressive tumors (68).

Little is known regarding an association between SSTR expression, tumor grade, and Ki-67 level in MCC. A preliminary study showed that 68Ga–somatostatin analog PET/CT provides good diagnostic performance equivalent to that of 18F-FDG PET (69). These results do not suggest that 18F-FDG PET/CT should be replaced by 68Ga-SSTR imaging. However, it could be considered in select cases of SSTR-positive MCC—that is, “personalized medicine.” Further studies are needed to establish the usefulness of 68Ga–somatostatin analog PET for imaging of MCC.

Peptide Receptor Radionuclide Therapy
The SSTR binding peptide is paired with a β particle–emitting radioisotope such as yttrium 90 (90Y) or lutetium 177 (177Lu) using a chelator (bonding agent). The radiolabeled peptides are delivered directly to tumor cells via SSTR and irradiate tumor cells. In Europe, PRRT has been
used for treatment of SSTR-positive metastatic well-differentiated gastrointestinal NETs since the 1990s.

Retrospective analysis has shown promising results for treatment of gastrointestinal NETs. In the United States, the FDA approved $^{177}\text{Lu}$-DOTATATE for treatment of gastrointestinal NETs in January 2018. Currently, there are only a few case reports that have demonstrated favorable results in MCC (70–72). A larger trial is needed to further understand the efficacy of PRRT in treatment of MCC.

**Conclusion**

MCC is an aggressive skin cancer with unique characteristics both clinically and radiologically. MCPyV is causally linked to its development, and antibodies to the virus can be used as a “tumor marker” in seropositive patients. Immunotherapy is now the recommended first-line systemic therapy, but caution is needed to interpret the treatment response. MCC has neuroendocrine features with SSTR expression, which can be used for SSTR-seeking molecular imaging and potentially theranostics (PRRT). Although MCC is rare, its incidence is steadily increasing, and radiologists should be aware of its characteristics.

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