Abstract. Merkel cell carcinoma (MCC) is a rare, aggressive primary cutaneous neuroendocrine carcinoma. It usually appears on the face and neck of elderly Caucasian people as a flesh-colored, erythematous or violaceous dome-shaped, non-tender nodule with a smooth surface. In immunocompromised patients with T-cell dysfunction, such as patients with acquired immunodeficiency syndrome (AIDS) or solid organ transplant recipients, the incidence of this disease is markedly increased. This suggests a link between the development of MCC and the immune system. Merkel cell polyomavirus (MCPyV) is clonally integrated into the majority of MCCs, suggesting its causative role in the pathogenesis of the majority of these tumors. Despite wide local excision, sentinel lymph node biopsy, and eventually, adjuvant radiation therapy, which remains the first-line treatment for MCC, the identification of MCPyV has opened novel therapeutic insights. Novel therapeutic strategies could be to inhibit MCPyV oncoproteins and to stimulate immune responses against virus-infected tumor cells by immunostimulatory cytokines, including interferons and interleukin-2.

1. Introduction

Merkel cell carcinoma (MCC) is a rare but aggressive primary neuroendocrine skin cancer, with a strong propensity to metastasize. It is considered to be caused by the malignant transformation of neurosecretory Merkel cells, which are located in the basal layer of the epidermis and are involved in skin mechanoreception (1). However, the origin of the malignancy of these cells remains a controversial issue. It is well known that MCC originates in the dermis, and only occasionally exhibits an epidermal involvement, whereas Merkel cells lay in the basal part of the epidermis; in addition, touch-sensitive areas, such as the lips and palmoplantar surfaces, are rich in Merkel cells, but it is unusual for MCC to originate in these locations (2). These findings demonstrate that the origin of MCC may not reside in the Merkel cells but, as recent studies have suggested, the origin of MCC may reside instead in pluripotent progenitor stem cells in the dermis (3) or in precursor B cells (4).

2. Clinical presentation

The clinical presentation of MCC is often non-specific: The clinical aspect of the lesion is usually a flesh-colored, erythematous or violaceous dome-shaped, non-tender nodule with a smooth surface. A differential diagnosis of MCC based on the clinical appearance includes such characteristics as epidermoid cysts, lipoma, basal cell or squamous cell carcinoma, amelanotic melanoma, pyogenic granuloma or lymphoma cutis. This skin cancer is characterized by an extremely rapid rate of growth, from a few weeks to months, leading in certain cases to ulceration (5). Typically, MCC arises in elderly Caucasians of either sex on sun-exposed skin. The most frequently affected site is the head and neck region (50%), followed by the trunk (30%) and the limbs (10%), although MCC may arise in any body site, including the mucosae (6).

3. MCC and the immune system

MCC is an uncommon skin cancer, although its incidence is rising, probably due, on the one hand, to a higher level of detection via more advanced diagnostic techniques, such as cytokeratin 20 immunostaging (7), and on the other hand to the higher prevalence of MCC risk factors, including T cell suppression and sun exposure (8). The annual incidence
of MCC in the USA is 0.6/100,000 per year, with a median age of 76.2 years for women and 73.6 years for men (9). Immunocompromised patients with T-cell dysfunction appear to be more likely to be affected by this disease: In these patients, the incidence of MCC is dramatically increased. For example, patients with acquired immunodeficiency syndrome (AIDS) have an incidence rate that is 11-13 times greater compared with the general population, and solid organ transplant recipients are 5-10 times more likely to develop MCC (10). There is also an association with ultraviolet (UV) radiation exposure: The incidence of MCC was determined to be 100-fold greater in patients who underwent photochemotherapy (PUVA) treatment (11), and MCC usually affects sun-exposed areas, such as the head and neck region, whereas the trunk and the limbs are less commonly involved (6).

The higher susceptibility of immunocompromised patients, together with the positive association with UV exposure, suggests a link between MCC and the immune system. Furthermore, several case reports have described regression after an improvement in immune function, highlighting the importance of the immune system to the development of this cancer (12,13).

This clear link with the immune system, particularly with T-cells, suggests that MCC may be caused by an infective agent. In 2008, Chang and Moore (14), using the technique of digital transcriptome subtraction, analyzed four samples of MCC and identified the 5,387-base-pair genome of an unknown polyomavirus that they termed ‘Merkel cell polyomavirus’ (MCPyV). Furthermore, through the analysis of a further 10 samples, these authors demonstrated that MCPyV was clonally integrated in 80% of MCCs, suggesting that infection and integration occur prior to the clonal expansion of tumor cells, thereby underlining its oncogenic role (15). The remaining 20% of MCC is MCPyV-negative, and this is considered to be caused by a different oncogenic pathway that involves UV-induced DNA damage and chromosomal aberrations (16).

4. MCPyV

MCPyV is a non-enveloped, double-stranded circular DNA virus. It is found ubiquitously, and is frequently isolated from healthy subjects: The seroprevalence of this virus ranges from 9% in children under 4 years of age to 35% in teenagers (14), increasing to 80% in adults, suggesting that it may be part of the cutaneous microbiome (17). The means of transmission has yet to be fully elucidated: Assembled virions may be detected on clinically normal skin, suggesting a cutaneous transmission (18). However, MCPyV DNA has also been identified in the gastrointestinal tract, on oral and anogenital mucosa, and in respiratory secretions, supporting oro-fecal, trans-mucosal and respiratory means of transmission, respectively (19-23). MCPyV infection is asymptomatic, and this accounts for its high prevalence throughout the general population (24).

The MCPyV genome is made up of early and late gene regions separated by a non-coding regulatory region. The late gene region expresses the major capsid protein, VP1, and the minor coat proteins, VP2 and VP3, encoding the viral capsomere and capsid, whereas the early gene region encodes large T antigen (LT), 57 kT antigen (57 kT) and small T antigen (sT) (16). The early gene region targets important cell proteins involved in cell cycle regulation and tumor suppression: LT interacts with the oncosuppressors, p53 and pRB, whereas sT binds to protein phosphatase 2A (PP2A), which is involved in cell proliferation through the regulation of gene transcription. This explains why this region is termed the ‘tumor antigen locus’. In addition to this function, LT is also important in viral DNA replication: It is involved in the initiation of viral DNA synthesis through its origin binding domain (OBD), ATPase domain and helicase domain, which are localized in its carboxy-terminal part (25).

5. Pathogenesis

Following asymptomatic infection with MCPyV, which usually occurs in early childhood, the immune system normally gives an appropriate humoral and cellular response. Thus, the virus becomes a part of the microbiome of the skin, from which it is chronically shed as encapsidated virions. UV radiation or other environmental mutagens, aging and infective or drug-induced immunosuppression are able to promote reactivation of the virus via the reduction in immunosurveillance. At this point, the viral genome is able to integrate into the host chromosome, courtesy of a defect in the virus itself, or following UV or mutagen exposure. In addition, a mutation in LT renders the virus unable to replicate (25). Indeed, MCPyV isolated from MCCs, in contrast with MCPyV from non-tumor sources, has been revealed to present mutations that are responsible for the premature truncation of the MCV LT helicase. These mutations do not affect the Rb binding domain, but eliminate the capacity of the viral DNA to replicate. In this way, the virus loses its capability to replicate in MCC tumoral cells, but continues to express motifs that may potentially lead to uncontrolled proliferation (26). Furthermore, sT is required for tumor cell proliferation, since sT knockdown inhibits cell replication in MCC (27). It has been revealed that sT prevents the dephosphorylation of the cellular translation factor, eIF4E-binding protein 1 (4E-BP1); hyperphosphorylated 4E-BP1 releases eIF4E, promoting active cap-dependent translation. Therefore, viral integration into the host genome and LT truncation mutations suggest that MCPyV may be responsible for MCC carcinogenesis via a ‘hit-and-run’ mechanism of transformation (25).

6. Immunotherapy

Currently, a wide local excision with at least a 1 cm margin, sentinel lymph node biopsy and, eventually, adjuvant radiation therapy are recommended by the National Comprehensive Cancer Network (NCCN) guidelines (28) as the first-line treatment for MCC. However, the identification of MCPyV in patients with MCC opened up novel therapeutic insights: On the one hand, the possibility to develop antiviral therapies interfering with the function of the oncoproteins, and on the other hand, the stimulation of immune responses against virus-infected tumor cells by immunostimulatory cytokines, such as interferons (IFNs) or interleukin-2 (IL-2) (24).

The proapoptotic action of IFN types I and II is well established (29). In particular, type I IFN is able to induce apoptosis in MCPyV+ MCC cell lines in vitro and in vivo through the modulation of the virally encoded LTA: IFN type...
I reduces the expression of the MCV LT and increases the expression of promyelocytic leukemia (PML) protein, which interferes with the function of the LT (30). Furthermore, a multicenter study revealed that isolated hyperthermic limb perfusion with tumor necrosis factor-α, IFN-γ, and melphalan resulted in a complete or partial response of locally advanced MCC (31).

Recombinant IL-2 causes regression of solid tumors by enhancing T cell activity, and it has already been approved by the Food and Drug Administration (FDA) for the treatment of metastatic melanoma and metastatic renal cell carcinoma (32). Regarding the treatment of metastatic MCC, a phase I/II clinical trial with autologous T cells and IL-2 (aldesleukin) is currently under way (33).

An alternative method of treatment would be to inhibit the downregulation of T cell function. An example of a group of agents that would be effective for this purpose is the cytokotoxic T-lymphocyte-associated protein 4 (CTLA-4) blockers, such as ipilimumab; a randomized clinical trial has been set up that compares therapy with ipilimumab with observations made following surgical resection of MCC (34). CTLA-4 is a negative regulator of T-cell-mediated antitumor responses, and it is expressed only on T cells (35).

Other putative target therapy agents are programmed death-1 (PD1; CD279) and programmed death ligand-1 (PD-L1) blockers. PD1, expressed on T cells but also inducible in B-cells and natural killer (NK) cells, after binding with its ligand, PD-L1, is expressed only in tumor cells, thereby down-regulating T cell function (35).

Since CD56 is expressed on almost all MCC tumors, a phase I clinical trial has examined the use of the CD56-targeting antibody drug, IMGN901 (36); this molecule is a monoclonal antibody, made up of a CD56-binding domain attached to emtansine (DM1), a cytotoxic agent. Following its binding to CD56, IMGN901 is internalized into the cell and DM1 is released, thereby killing the cancer cell via inhibition of the polymerization of tubulin (36).

One difficulty that must be circumvented in this type of tumor is chemoresistance. One of the major mechanisms of MCC chemo-resistance is inhibition of apoptosis through the upregulation of survivin (a member of the family of inhibitor of apoptosis proteins) and B-cell lymphoma 2 (Bcl-2) protein (37). In this situation as well, target therapy may help in fighting chemoresistens: YM-155, a novel small-molecule survivin suppressant, appears to downregulate survivin expression, promoting apoptosis in MCC xenograft tumors (38). A Bcl-2 antisense oligonucleotide has been demonstrated to arrest tumor growth in an MCC xenograft animal model, although a different Bcl-2 antisense oligonucleotide, G3139, did not exhibit any therapeutic efficacy in MCC (39). In addition, other apoptotic inhibitors, such as ABT-263, have demonstrated a certain level of clinical efficacy (40).

Following the identification of MCPyV, a novel and promising therapeutic approach would appear to be viral antigen-directed immunotherapy, or the use of a vaccine. Zeng et al (41) developed a DNA vaccine encoding MCPyV LT (pCDNA3-LT), which contained an LT-specific CD4+ T-helper epitope. This DNA vaccine generated antitumor effects that were predominantly mediated by CD4+ T cells against LT in mice (41).

7. Conclusions

MCC is an aggressive tumor with poor prognosis. Although surgical removal with negative margins, eventually followed by radiotherapy, remains the first-line treatment, immunotherapy appears to represent a very promising alternative approach. The identification of the MCPyV-specific cellular immune response has suggested novel therapeutic targets. In this respect, it would be helpful to identify MCPyV-positive patients among all the patients with MCC in order to optimize the use of antiviral therapy and DNA vaccines encoding MCPyV LT. Furthermore, particularly in the case of immunocompromised patients, such as organ transplant recipients and AIDS patients, prevention should not be discounted: Since UV exposure appears to be associated with the etiology of MCC, these patients should consequently limit their exposure to UV radiation and adopt sun safety practices. However, one single approach is not likely to be effective for all the patients, due to the inter-individual variability of the immune system and the mechanisms of immune evasion for MCC. Therefore, further studies are required to investigate multiple target therapies and to improve our understanding of the molecular mechanisms of immune evasion.

References


