Early B-Cell Differentiation in Merkel Cell Carcinomas: Clues to Cellular Ancestry

Axel zur Hausen¹, Dorit Rennspiess¹, Veronique Winnepenninckx¹, Ernst-Jan Speel¹, and Anna Kordelia Kurz²

Abstract

Merkel cell carcinoma (MCC) is a highly malignant neuroendocrine nonmelanoma skin cancer, which is associated with the Merkel cell polyoma virus (MCPyV). Recently, expression of the terminal deoxynucleotidyl transferase (TdT) and the paired box gene 5 (PAX 5) has been consistently reported in the majority of MCCs. We tested 21 MCCs for the expression of MCPyV, TdT, PAX5, IgG, IgM, IgA, kappa, and lambda by immunohistochemistry and assessed IgH and Igk rearrangement in all 21 MCCs. All of the MCCs revealed specific expression of PAX5 and 72.8% of the MCCs expressed TdT. In addition, most of the MCCs revealed specific expression of one or more Ig subclasses and kappa or lambda. One MCC did reveal monoclonal IgH and Igk rearrangement next to two other MCCs showing Igk rearrangement. As coexpression of TdT and PAX5 under physiologic circumstances is restricted to pro/pre- and pre-B cells we propose, on the basis of our results, that the cell of origin of MCCs is a pro/pre- or pre-B cell rather than the postmitotic Merkel cells. MCPyV infection and transformation of pro-/pre-B cells are likely to induce the expression of simple cytokeratins as has been shown for SV40 in other nonepithelial cells. This model of cellular ancestry of MCCs might impact therapy and possibly helps to understand why approximately 20% of MCCs are MCPyV-negative.

Introduction

Merkel cell carcinoma (MCC) is a highly malignant non-melanoma skin cancer characterized by the expression of neuroendocrine markers, that is, CD56, chromogranin A, and synaptophysin and a typical dot-like perinuclear expression of cytokeratin 20 (CK20; ref. 1). MCC occurs mainly in elderly or immunosuppressed patients and its clinical prognosis is very poor (1). The recent finding of the Merkel cell polyomavirus (MCPyV) in MCCs has substantially contributed to the understanding of the etiopathogenesis of this malignancy (2). Since 2008, MCPyV DNA has been detected in approximately 80% of more than 1000 MCCs yet tested. MCPyV has been shown to be integrated in the MCC genomes by diverse molecular techniques (2–6).

MCC was first introduced by Toker in 1972 as a trabecular carcinoma of the skin (7). Dense core granules were identified in 3 MCCs, which was confirmed by subsequent immunohistochemical studies testing neuroendocrine and simple epithelium markers (8–13). On the basis of the similarity of the expression patterns, MCC is considered to be most closely related to merkel cells, which are located at the epidermal dermal junction and are supposed to function as noticceptive receptors of the skin. However, serious doubts on this theory have been postulated—also based on histomorphologic growth patterns—and an alternative proposal suggesting a role for a pluripotent stem cell in the dermis as cells of origin of MCCs still remains to be proven (14).

In the past years, evidence is accumulating that MCCs express early B-cell lineage markers (15–22). For example, specific expression of terminal deoxynucleotidyl transferase (TdT) as assessed by immunohistochemistry, which normally is restricted to thymocytes and precursors of lymphocytes in the bone marrow, has been reported in 72.5% (95 of 131) of MCCs (15–19). In addition, a number of studies have reported immunostaining of the B-cell–specific activating protein (BSAP; syn: PAX5) in 83.6% (102 of 122) of MCCs (16, 19–22). Coexpression of TdT and BSAP under physiologic circumstances is restricted to pro/pre-B cells and pre-B cells. Pre-B cells exhibit immunoglobulins and in addition rearrangement of the IgH and Igk locus. Here, we examined 21 MCCs for the expression of TdT, PAX 5 (BSAP), IgG, IgM, IgA, kappa, and lambda by immunohistochemistry. In addition, clonal rearrangement of the IgH and Igk locus was tested in all MCCs to find further evidence for our assumption that MCCs possibly derive from pro/pre- or pre-B cells.

Materials and Methods

Patients and tissues

Formalin-fixed and paraffin-embedded (FFPE) tissues of resection specimens of 21 MCCs were included...
in this study. All respective tumors had been excised for diagnostic and therapeutic reasons. Eighteen specimens were obtained from the Maastricht Pathology Tissue Collection. All use of tissue and patient data was in agreement with the Dutch Code of Conduct for Observational Research with Personal Data (2004) and Tissue (2001, www.fmwv.nl). Three MCCs were collected at the Institute of Pathology, University Hospital Freiburg, Freiburg, Germany, which had been previously tested for the presence of MCPyV (23).

MCC diagnoses were previously defined by histology and immunohistochemistry for CK20, CD56, synaptophysin, and chromogranin A in routine diagnostic. The histology of all MCCs included in this study has been reviewed by 2 expert dermatopathologists (V. Winnepenninckx and A. zur Hausen). The patient group consisted of 9 men (age: range, 63–93 years; mean, 78.3 years) and 12 women (age: range, 60–93 years; mean, 76.2 years). Only MCC tissues were included in this study of patients with no history of other hematologic oncologic disease. Further details of clinicopathologic parameters are included in Table 1. Serial sections of all specimens were used for hematoxylin and eosin (H&E) staining, immunohistochemistry, and DNA isolation.

Immunohistochemistry

The following antibodies and dilutions were used in this study: anti-MCPyV (clone: CM2B4) dilution 1:50, Santa Cruz, Inc.; anti-TdT "Ready to use Antibody"; anti-PAX5 (clone: DAK-Pax5), "Ready to use Antibody"; anti-IgA dilution 1:20,000; anti-IgG dilution 1:20,000; anti-IgM dilution 1:2,000; anti-kappa dilution 1:40,000; anti-lambda dilution 1:40,000, all DAKO. All immunohistochemical stainings were conducted on a Dako Autostainer Link 48 using the EnVision FLEX Visualization Kit K8008 DAKO according to standard diagnostic routine protocols and manufacturers’ instructions.

DNA extraction and immunoglobulin light chain restriction

First, an H&E stain of the selected specimens was reviewed (V. Winnepenninckx and A. zur Hausen). Five 10 μm-thick consecutive sections were cut from each paraffin block of formalin-fixed tissue and, if needed, microdissection was conducted. After deparaffinization, the tissues were lysed by proteinase K overnight until complete tissue lysis, and DNA was extracted using QIAamp mini kit (Qiagen) or Maxwell 16 FFPE Plus LEV DNA isolation kit (AS1135), Promega. Purified DNA was measured in a spectrophotometer (Nano-drop, 2000; 

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<th>Table 1. Clinicopathologic, immunohistochemical, and molecular pathology data of 21 MCCs</th>
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NOTE: Expression of Igs and kappa and lambda is given as percentage tumor cells expressing respective Igs or kappa and lambda. Abbreviations: –, negative; +, weak expression; ++, moderate expression; ++++, strong expression.

Abbreviations: BSAP, B-cell-specific activating protein; F, female; int., intermediate; M, male; monoclonal; m, vk/intron k deleting element; n, not sufficient DNA quality; n.i., not informative; p, polyclonal; s.c., small cell; PAX5, paired box gene 5; SCS ladder, specimen control size ladder indicating whether DNA quality is sufficient in order to interpret results; y, sufficient DNA quality.
Detection of clonal immunoglobulin gene recombina-
tions was conducted according to an established and validated protocol (24) in a routine diagnostic setting in the nationally accredited diagnostic molecular pathology laboratory of the Department of Pathology, Maastricht University Medical Center, Maastricht, the Netherlands.

Results

MCPyV, PAX5, and TdT expression in MCCs

The presence of MCPyV was assessed by the expression of the large T antigen (LTAg) as tested by immunohistochemistry. Eighteen MCCs (85.7%) were tested positive for LTAg expression. Of the 21 MCCs, 13 revealed very strong nuclear expression of the LTAg, 2 moderate nuclear expression, and 3 weak nuclear expression (Fig. 1; Table 1). In these MCPyV-positive cases, LTAg expression was homogeneous and found in all tumor cells. Three MCCs were completely negative for LTAg expression.

Specific nuclear expression of PAX5 was seen in all MCCs, of which 10 MCCs revealed strong PAX5 expression, 6 moderate, and 5 weak (Table 1).

Specific nuclear TdT expression was detected in 16 MCCs (76.2%), of which 6 were strongly, 2 moderately, and 8 weakly expressing TdT. Five MCCs were completely negative for TdT expression. Although no correlation was observed between PAX5, TdT, and LTAg expression it is of interest that when PAX5 expression was high, TdT expression was low, and vice versa. MCCs could be grouped according to TdT and PAX5 expression (see Table 1): TdT high, PAX5 low expression, and
Expression of immunoglobulins and kappa and lambda in MCCs

Testing 21 MCCs for the expression of Igs by immunohistochemistry revealed that almost all MCCs did show Ig expression of one or more Ig subclasses to variable extent (Fig. 1A). In detail, 10 (47.6%) MCCs were expressing IgA, 10 (47.6%) MCCs were expressing IgG, 5 (23.8%) MCCs were tested positive for IgM expression to a varying degree. In addition, 7 (33.3%) MCCs revealed kappa and 12 (57.1%) MCCs revealed lambda expression. Expression was membranous or cytoplasmic and only strong expression intensity was scored as positive. The percentage of tumor cells expressing Ig or kappa or lambda did show marked diversity among the MCCs. Details are summarized in Table 1. Noteworthy, all of the MCPyV-negative MCC (n = 3) did not show expression of Ig or kappa or lambda.

Molecular testing of IgH and Igk rearrangement in MCCs

According to the specimen control size, ladder 18 of the 21 MCCs revealed sufficient DNA quality to proceed with molecular testing (Table 1). Analysis of IgH and Igk rearrangements in MCCs according to van Dongen and colleagues (24) identified a monoclonal IgH locus rearrangement combined with a monoclonal Igk rearrangement in one case (case no. 38–20, see Fig. 1B, Table 1). Furthermore, light chain restriction was found in 2 additional cases (case no. 15–18 and 21–11; Table 1). All MCCs revealing molecular rearrangements were restricted to the group of MCCs with low TdT and high PAX5 expression.

Discussion

At present, profound doubts are challenging the concept that the postmitotic merkel cells are the cells of origin of MCCs, which had been proposed in the early 1980s (8–13, 25, 26). The recently discovered MCPyV has been shown to play an important role in the etiopathogenesis of the majority of MCCs. Of interest, MCPyV cannot be detected in non-neoplastic merkel cells adjacent to MCPyV-positive MCCs (26). On the basis of MCC mouse models, it has been speculated that the cell of origin of MCC derives from epidermal stem cells or other primitive totipotent stem cells (reviewed in refs. 25, 26).

However, none of the previous reports on the cellular origin of MCCs took into account that in a significant number of MCCs, TDT and PAX5 are expressed consistently, that is, in 72.5% and 83.6% of cases, respectively. Here, we assessed TDT and PAX5 expression in 21 MCCs and indeed were able to confirm these previous results. Specific nuclear expression of TdT was found in 16 of 21 MCCs (76.2%) and all of the 21 MCCs revealed specific nuclear PAX5 expression. Under physiologic conditions, coexpression of TDT and PAX5 is restricted to pro/pre- and pre-B cells. In addition to TDT and PAX5 expression, pro/pre- and pre-B cells are characterized by the expression of immunoglobulins (Igs) and VJH clonal rearrangements.

Taken together, the obtained data on TdT and PAX5 expression in MCCs may either reflect an epiphenomenon in the course of malignant transformation or be of meaningful significance possibly pointing to the cellular ancestry of MCC. To assess further pro/pre- and pre-B cells differentiation in MCCs, we examined the expression of immunoglobulins (IgA, IgG, and IgM), kappa and lambda and in addition B-cell clonality in MCC tissues.

Indeed, we were able to show strong expression of immunoglobulins or kappa/lambda in a significant number of MCCs. Only strong membranous or cytoplasmic Ig and/or kappa/lambda staining of the MCC cells were considered as positive expression. Although this is the first report to show Ig expression by immunohistochemistry in FFPE tissues of MCCs, it is not the first report on Ig's and kappa/lambda expression in MCCs on the transcriptional level. Recent cDNA microarray analyses have shown that the expression of certain Igs is strongly upregulated. Comparing the transcriptional expression profiles of MCC cell lines versus that of cell lines of small cell lung cancer, Van Gele and colleagues have shown a 100-fold increase in membrane-bound and secreted immunoglobulin gamma heavy chain in MCC cells (27). In addition, Paulson and colleagues identified transcriptional expression of the IgJ polypeptide and Ig kappa constant as genes most highly upregulated in good prognosis tumors (28). Thus, the immunohistochemical expression data of Ig and kappa/lambda are in line with previous observations on the transcriptional level.

Although only a minority of the MCCs in this study did show B-cell clonality, the finding of B-cell clonality in 3 MCCs (14.3%) strongly points to pro/pre- or pre B-cells as the cells of origin of MCCs. B-lymphocyte development is a timely regulated specific process controlled by protein products of functionally rearranged Ig heavy (H) and light (L) chain genes. This process of Ig rearrangement begins in pro-B cells at the IgH locus (29).

On the basis of our results, we propose that pro/pre- or pre-B cells, which either or not have yet started the process of Ig rearrangement turn into a tumor cell as the starting point of MCCs (Fig. 2). Although other transforming events may be considered, MCPyV is the main suspect to transform pro/pre- and pre-B cells. More than 80% of MCCs harbor the recently discovered Merkel cell polyomavirus in its integrated form and in addition reveal oncogenic mutations in the LTAg of MCPyV. It is likely that MCPyV transformation leads to spontaneous loss of cytokeratin and neuroendocrine marker gene expression control and thus the expression of simple cytokeratins and neuroendocrine markers is induced in pro/pre- and pre-B-cells. At least, the induction of cytokeratin expression in cytokeratin-negative cells upon infection with another polyomavirus, that is, SV40, has been shown previously (30). Phylogenetically, MCPyV is closely related to SV40 and thus shares the same set of transformation relevant genes. Moreover, it has been shown that, for example, JC polyomaviruses preferentially infect stem cells or progenitor cells (reviewed in ref. 25). The finding of Ig and kappa and lambda expression in MCCs on the protein level, as has been described in 2 previous independent cDNA microarray studies on the transcriptional level strongly support our model. Clonal IgH and Igk rearrangement in MCCs as found in our study in addition highly substantiates our model. The relative low number of 3 MCCs revealing B-cell clonality might indicate that the transforming event, that is,
MCPyV infection, of pro/pre- or pre-B cells occurs before the start of Ig rearrangement. The fact that all MCCs carrying IgH or IgL rearrangements are in the group of MCCs with lower TdT than PAX5 expression also supports our hypothesis. In addition, the striking histologic similarity of pro/pre- and pre-B cells and the intermediate type of MCCs further supports this hypothesis. The proposed model can also help to understand why there are MCPyV-positive and MCPyV-negative MCCs and why most MCCs reveal a lymphoma-like growth pattern in the skin and only a very small minority of MCCs being connected to the epidermis. MCCs are almost always hematologically disseminated at the time of diagnosis. Identifying pro/pre- or pre-B cells as the cellular ancestors of MCCs is also expected to impact current treatment regimens of MCCs, which is of major significance in the context of the very poor prognosis for patients with MCCs after primary MCC diagnosis.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors’ Contributions

Conception and design: A. zur Hausen, A.K. Kurz
Development of methodology: D. Rennspiess, E.-J.M. Speel
Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): D. Rennspiess
Analysis and interpretation of data (e.g., statistical analysis, biostatistics, computational analysis): A. zur Hausen, D. Rennspiess, E.-J.M. Speel
Writing, review, and/or revision of the manuscript: A. zur Hausen, V. Winnepenninckx, E.-J.M. Speel, A.K. Kurz
Study supervision: A. zur Hausen

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