Reflections on the Histopathology of Tumor-Infiltrating Lymphocytes in Melanoma and the Host Immune Response
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Abstract
In the past five decades, the role for lymphocytes in host immune response to tumors has been shown, at least in some patients, to be a critical component in disease prognosis. Also, the heterogeneity of lymphocytes has been documented, including the existence of regulatory T cells that suppress the immune response. As the functions of lymphocytes have become better defined in terms of antitumor immunity, specific targets on lymphocytes have been uncovered. The appreciation of the role of immune checkpoints has also led to therapeutic approaches that illustrate the effectiveness of blocking negative regulators of the antitumor immune response. In this Masters of Immunology article, we trace the evolution of our understanding of tumor-infiltrating lymphocytes and discuss their role in melanoma prognosis from the very basic observation of their existence to the latest manipulation of their functions with the result of improvement of the host response against the tumor. Cancer Immunol Res; 3(8); 827–35. ©2015 AACR.

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Learning Objectives
Lymphocytes comprise a critical component of the host immune response to tumors, and the presence of tumor-infiltrating lymphocytes (TIL) correlates with disease prognosis. Identification of TILs in the tumor microenvironment and characterization of their role in melanoma prognosis, as well as the strategies available for manipulating TIL functions, will facilitate the augmentation of host immune response against tumors. Upon completion of this activity, the participant should gain insight from the discovery of TILs in the tumor microenvironment to the histopathology of TILs in melanoma and their roles in host immune response.

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Introduction
The importance of lymphocytes in tumors has been long recognized. At first, it was considered that they were causative of malignancy. Their possible role in prognosis in melanoma and other tumors has only been recognized in the past 40 to 50 years. This Masters of Immunology article relates, often through the lens of personal experience, the history of the discovery and characterization of tumor-infiltrating lymphocytes (TIL) and
the elucidation of their possible role in the control of human melanoma.

This work has been accomplished by many collaborations (see Note on page 1, bottom right). We acknowledge the role of various fellows and staff of the Massachusetts General Hospital, the Brigham and Women’s Hospital, and the Moffitt Cancer Center, as well as the members of the WHO and EORTC Melanoma Pathology groups for their aid, guidance, and support. As the focus will be on the histopathology aspects of this topic, we further wish to acknowledge the successful past and ongoing immunotherapy clinical trials of the adoptive transfer of tumor-infiltrating lymphocytes in patients with advanced melanoma, first pioneered at the NCI Surgery Branch (1, 2) and expanded to other treatment institutions nationally (3, 4) and internationally (5, 6), which will not be covered in our reflections.

**Historical Aspects of Tumor-Infiltrating Lymphocytes in Melanoma**

The observation of lymphocytes and mononuclear cells associated with human cancer was first noted more than a century ago, and these infiltrates were thought to be causative of the disease (7). The concept of an immunologic response to malignant tumors in patients was probably first proposed by Paul Ehrlich in 1907 and later expanded in 1909 (8). Coley injected bacteria into tumors, so-called Coley toxin, with some tumor shrinkage recognized as a host response to tumors (9). Subsequently, studies of various cancers, including osteogenic sarcoma, neuroblastoma, carcinoma of the colon, and melanoma, have been performed and have elucidated the relationship of host immune response to tumors and patient survival. Moore and Foote hypothesized for medullary carcinoma of the breast that the ‘rather characteristic lymphoid infiltrate indicates some maladjustment between tumor and host, and that the clinical behavior is a further indication thereof’ (10). Already, by the time of the publication of “Melanoma and Skin Cancer” in the 1972 Proceedings of the International Congress on Skin Cancer in Sydney, Australia, descriptions of host immune response to melanoma in the form of antimelanoma antibodies (11) and lymphocytotoxicity (12) to human melanoma and mouse melanoma cells had been documented. The injection of minced melanoma cells into 26 patients with metastatic melanoma resulted in 2 patients with complete tumor regression and 5 patients with partial regression of tumors (13). In a similar study, Ryan and colleagues (14) reported a high index of lymphocytotoxicity in patients with metastases injected with irradiated autologous melanoma cells. They also reported regression of metastases after injections with Bacillus Calmette-Guerin (BCG). Histology of the regressing tumors exhibited marked infiltration with lymphocytes.

The term "tumor-infiltrating lymphocytes" was used for the first time in the experience of one of us (M.C. Mihm) during work with Wallace H. Clark Jr. In the detailed study of malignant melanomas that was spearheaded by Clark, the anatomic levels of invasion were proposed (15). The epidermis, the papillary dermis, and the superficial vascular plexus were considered reactive sites where cutaneous inflammatory processes occurred. Clark considered these sites the sensitizing area of the skin and used the example of poison ivy to support his hypothesis. During the course of these studies (16), Clark was struck with the prominent inflammatory infiltrate (Fig. 1A) that accompanied the superficial spreading variant of malignant melanoma in the radial growth phase (RGP), and we both invoked the possibility of a contact-like hypersensitivity reaction. He also noted that, in the lesion that we described as the precursor of melanoma, lentigo maligna, there was often no inflammation until the lesion became microinvasive. These observations and hypotheses led to the designation of level II as a microinvasive melanoma that was probably controlled by the host response through a sensitization of the host by the tumor cells (15). Another feature that was described both in superficial spreading melanoma and in lentigo maligna melanoma was regression in the RGP (17). This phenomenon was identified by the presence of partial areas of fibrosis with lymphocytes and melanophages flanked on one or both sides by the tumor (Fig. 1B). This observation was considered to be evidence of a host immune response to the tumor. Levels III, IV, and V represented a new event in tumor progression that was associated with the acquisition of metastatic potential. However, at level III, the tumor was still confined to the papillary reticular dermal junction and also often had an inflammatory infiltrate, frequently at the base of the tumor. Levels IV and V were proposed to confer the most adverse prognosis, either because the tumor could overcome the barrier of the papillary dermal interface and successfully invade through the dermis and into the fat or because the host response had failed. In 1967, a patient at the Massachusetts General Hospital (Boston, MA) Pigmented Lesion clinic presented with a metastatic nodule surrounded by a halo of hypopigmentation. A biopsy of McCarten reported that patient survival was related to the density of the lymphoid infiltrate at the advancing edge of the tumor. Favorable prognosis was associated with a dense response (19). Day and colleagues (20) reported better survival in patients with primary melanoma that had a moderate to marked infiltration of intratumoral and/or peritumoral lymphocytes compared with those with no infiltrate.

Meanwhile, in Australia, McGovern and Lane Brown began to study lymphocytic infiltration in halo nevi and melanoma (21). Thompson reported an improved prognosis in patients with a prominent infiltrate at the advancing edge of the melanoma nodule in a very limited study (22). McGovern studied the evolution of regression and considered a dense infiltrate of lymphocytes in a nodule of melanoma to represent early regression (23). A panel of 11 pathologists and a surgeon, G.W. Milton, chaired by Vincent McGovern with one of us (M.C. Mihm) as rapporteur was convened at the Sydney Conference in 1972 to formalize the classification of melanoma and its reporting. The group recognized that lymphoid infiltrates appeared to be important in primary melanoma but were not well-enough studied to be
included in the routine pathology report (24). In a subsequent
review of prognosis in melanoma, McGovern reported on varying
studies that described improved prognosis to be associated with
lymphocytes, especially in the base of the tumor nodule (25).

In the early 1970s, a great deal of interest was expressed in
vaccination of metastases. In collaboration with M. Lane Brown,
Cosimi, and T.B. Fitzpatrick, we injected vaccinia virus into several
melanoma nodules of the lower leg of a patient with multiple
metastatic lesions in the right lower limb. The injected nodules
had marked inflammatory reactions and some nodules regressed.
All melanoma nodules, with or without vaccinia injection, reacted
with at least some swelling and erythema. These changes were
possible examples of the now-recognized abscopal response. A
biopsy of one of the injected lesions revealed histologically a
striking infiltrate of mainly lymphocytes with some mononuclear
cells and focal neutrophils. Direct interactions of the lymphocytes
with melanoma cells were observed with pyknosis of melanoma
cells. Roenigk and colleagues (26) reported on a similar response
of vaccinia virus injections into metastatic melanoma nodules.
These changes certainly reinforced earlier suggestions for devel-
oping a therapy that uses the lymphocyte response to control
melanoma.

In the United States, Morton and colleagues pursued studies
showing the importance of the immune response in melano-
mas that led him to publish a rationale for immunotherapy
(27). One study showed significant sensitization of lympho-
ocytes by melanoma antigens (28), and further studies with BCG
vaccination of tumor nodules recorded the importance of the
lymphocytes in the reaction (29).

In addition to the study of melanoma, one of us (M.C.
Mihm) formed a very active collaboration with Harold
F. Dvorak in the study of delayed hypersensitivity reactions
as a component of allergic and tumor responses. Because of the
interest in hypersensitivity reactions, Dvorak and colleagues
(30) studied four examples of RGP superficial spreading mel-
anoma with electron microscopy and compared these with
prior findings in human delayed hypersensitivity responses,
including allergic contact dermatitis and human allograft
skin rejection. This investigation revealed activation of lym-
phocytes and macrophages, prominent vascular changes,
basophils, and mast cells with a partial loss of mast cell granule content, as seen in delayed-type hypersensitivity reactions. There was evidence for killing of melanoma cells through direct contact with lymphocytes. The vascular changes included endothelial cell swelling as well as vessels manifesting necrosis of the endothelium. The basal lamina showed marked reduplication of the endothelial cell swelling as well as vessels manifesting necrosis of the endothelium. The basal lamina showed marked reduplication typical of recurrent injury such as chronic inflammation. While these changes, other than the lymphocyte–melanocyte satellitosis, were not specific, they were compatible with the findings in human cutaneous hypersensitivity responses (30).

During these years, Clark, along with Elder and other collaborators, started to study the patterns of lymphocyte infiltration that resulted in the publication in 1989 of the most commonly used classification of TILs as brisk, non-brisk, and absent (31). To qualify as a TIL, there had to be evidence of lymphocyte-melanoma cell interaction (Fig. 2). The study by Clemente and colleagues (32) in the WHO Melanoma Pathology group confirmed the Clark data and demonstrated a highly significant survival advantage in patients with lesions with a brisk infiltrate, an intermediate survival for those with lesions with non-brisk infiltrates, and a poor prognosis for those with melanomas without an infiltrate (Fig. 3). However, neither study described the variation in the density of the infiltrating lymphocytes. The WHO group subsequently studied the presence of TILs in clinical stage II melanomas with positive lymph nodes from patients treated on a trial of IFNα, some of whom did very well with a prolonged survival (33). These lymph node metastases were evaluated for TILs anonymously with the use of the grading system of brisk, non-brisk, and absent as applied to the primary lesion. Fig. 3A–E are schematic diagrams illustrating the different types of TILs. To qualify as brisk, all the metastatic tumor deposits, whether in one or several lymph nodes, had to exhibit a diffuse infiltrate of lymphocytes interacting with tumor cells. The non-brisk category included those in which there was partial infiltration of the tumor by lymphocytes. The results indicated that patients with the brisk immune infiltrates had significantly better survival than those with absent inflammation or non-brisk inflammation. There was no evidence that the IFN therapy had any effect on the outcome. An additional study of these patients indicated a matched clonality of lymphocytes in the primary tumor and in the metastatic tumor with regard to T-cell receptor (TCR) usage; this finding suggested the potential specificity of the host response (34). In a later study of prognostic variables in 259 Eastern Cooperative Oncology Group interferon trials, positive TILs had a beneficial survival effect, independent of therapy (35).

Over the past 20 years, with Glenn Dranoff and F. Stephen Hodi, we have studied the histopathology of the response to vaccination with autologous melanoma cells engineered to produce GM-CSF as well as that due to antibodies blocking the immune checkpoint antigen, CTLA-4, in mice and in humans (36, 37). This work clearly demonstrated the critical role of lymphocytes as they infiltrated the tumors in the response to melanoma metastases due to vaccination or due to the release of the immune response with appropriate anti-CTLA-4 antibodies. During the same time period, several additional histopathology studies of TILs in melanoma have been published. Some supported the antitumor significance of TILs in melanoma and T cells in sentinel lymph nodes (SLN). Others failed to show significant correlation, as reviewed by Schatton and colleagues (38). While the basis for these differences remains to be established, one factor might relate to the heterogeneity of T-cell subsets. The importance of regulatory T-cell (Treg) inhibition of protective antimelanoma host responses, for example, has been documented in murine studies (39). The importance of stimulatory and costimulatory molecules in determining whether lymphocyte activation or inhibition occurs has also been revealed. A more complete definition of melanoma antigens, including mutated neoeptopes, has been achieved as well. All of this new information will lead to a more comprehensive understanding of antimelanoma immunity including the critically different types of T cells (38, 40).
Present and Future Histopathology Studies of TILs in Melanoma

A recent article by Thomas and colleagues (41) described TILs in 2,845 patients with a 5-year follow-up from the Population-Based Genes, Environment and Melanoma (GEM) study. A definite survival benefit was found when comparing melanomas with brisk and non-brisk TILs versus those with absent TILs (P < 0.001). These investigators used the criteria established by Clark and colleagues (31) in evaluating the infiltrates, and the report included information on Breslow thickness, mitoses, and ulceration, as well as the age and the site of the tumor. All cases included the American Joint Committee on Cancer (AJCC) tumor-staging features. The index cases were predominantly vertical growth-phase lesions but also included a small number of stage T1b lesions according to the AJCC tumor determinations. The rates of melanoma-specific death were 30% and 50% less in tumors with non-brisk and brisk TILs, respectively, compared with tumors with an absence of TILs. The brisk infiltrates were found more commonly in tumors from young patients, with the absence of infiltrates found in tumors with increased mitoses and ulceration. Interestingly, in a larger population of patients from the GEM study, including the index cases, the RGP tumors were consistently associated with a brisk infiltrate. This finding harkens back to the studies of Clark and Dvorak and colleagues discussed above. During these years, the Sydney Melanoma group, now formalized into the Melanoma Institute of Australia under the leadership of John Thompson, Jonathan Stretch, and Richard Scolyer, has engaged in numerous melanoma studies. A recent benchmark article on TILs and their relationship to sentinel lymph nodes and on patient survival demonstrated that the presence of dense TILs in the primary tumor was associated with 5.6% SLN positivity versus 27.8% positivity in tumors with absent TILs. Furthermore, there was 100% survival in patients with dense TIL infiltration (42).

Cintolo and colleagues (43) recently reported on 161 patients and determined the influence of TILs and tumor regression in patients evaluated by the standard parameters, including AJCC staging criteria. The tumors had a median thickness of 5.27 mm. The favorable prognostic factors included female sex, site, non-axial, positive TILs, age < 60 years, no ulceration or histologic regression, no prior elective lymph node dissection, or no evidence of any regional nodal disease. The presence of TILs was prognostically significant only when there was no RGP regression. The benefit of TILs was lost if RGP regression was present (P < 0.001). These authors hypothesized that the presence of regression indicated partial success of host response to the RGP lesions but failure as far as the deeper tumor was concerned. A recently complete study of 1,240 patients with a 30-year follow-up by the EORTC Pathology Group suggests that certain patterns of TILs have a definite survival benefit (Cook M et al., manuscript in preparation).

The Role of Chemokines, Mononuclear Dendritic Cells, and Immunosuppression in Melanoma Pathogenesis

The original concepts that lymphocytes could generate cancer have long been based on the fact that chronic inflammation can be associated with malignancy. One example is the association
of the inflammatory infiltrate triggered by Helicobacter pylori infection and gastric malignancy or the persistent inflammation of chronic prostatitis and prostate cancer. These observations pose the dilemma as to why some TILs in melanoma result in prolonged survival in some cases, whereas in other cases, the same inflammatory infiltrate is associated with immunosuppression. The answer, in part, to these observations may be found in understanding the role of inflammatory chemokines in lymphocyte function, as well as understanding the immunosuppressive aspects of certain mononuclear dendritic cells and the tumor milieu. At least 48 chemokines (in 4 groupings, i.e., CC, CXC, C, and CX3C) and at least 20 chemokine receptors are involved in the various types of response to stimuli that result in inflammation and/or are important in homeostasis (44). Taube and colleagues recently described how both melanoma cells and benign nevi, in the presence of immune host responses and secretion of IFNγ, express the programmed death ligand-1 (PD-L1), whose interaction with the PD-1 receptor leads to lymphocyte exhaustion. The lymphocyte is the apparent source of IFN that then results in rendering the TILs’ response effete. Blocking of this interaction by appropriate antibodies leads to enhancement of the immune response (45).

There are different types of dendritic cells in the skin and in the lymph nodes. On exposure to antigen, some result in sensitization and others can lead to tolerance (46). Plasmacytoid dendritic cells are abundant in positive SLN and appear to be associated with immunosuppression (47). Finally, with regard to the issue of malignant transformation in the milieu of chronic inflammation, the medley of cytokines and chemokines that are expressed lead to altered DNA, angiogenesis, and inactivation of suppressor genes among others with resultant malignant change. Among the most active of cytokines are TNFa, IL1, IL6, and IL8 with support of stromal fibroblasts. In fibroblasts and myofibroblasts, the reaction of, CXCL12 with CXCR4 results in angiogenesis associated with tumor growth (48, 49). The future lies in unraveling the complexities of tumor-host and stromal interaction to better understand the role of TILs in relation to tumor progression, control, and eradication, as well as establishing the definitive role of the monocyte macrophages in enhancing and suppressing the immune response (50).

**Ectopic Lymph Node-like Structures or Tertiary Lymphoid Structures within Melanomas**

The term “tumor-localized ectopic lymph node-like structures” (TL-ELN) was introduced to one of us (J.J. Mulé) for the first time in Seattle by Karl Erik and Ingegerd Hellstrom along with Ellsworth "Buster" Alvord during the studies of the capacity of immune lymphocytes to selectively accumulate in tumors in vivo (Fig. 4; refs. 51, 52). Years later, Messina and colleagues (53) interrogated a novel 12-chemokine gene expression signature (GES) on genomic arrays of stage IV melanoma metastases. They found this GES to accurately predict the degree and type of lymphoid infiltrate, organized as TL-ELNs (53). TL-ELNs had been identified earlier by others in human non–small cell lung cancers and in a few other solid tumors (reviewed in ref. 54). Harlin and colleagues (55) suggested that a lack of critical chemokines in a subset of melanoma metastases might limit the migration of activated T cells, which in turn could limit the effectiveness of antitumor immunity. Of interest, Hong and colleagues (56) reported that CXCR3 ligands and the CCL5 chemokine could determine T-cell infiltration into cutaneous melanomas. In addition, certain chemotherapy drugs could induce the expression of these chemokines in human melanoma cell lines. These chemotherapy-induced chemokines correlated with T-cell infiltration in melanomas, also resultant with tumor control and prolonged patient survival.

**Figure 4.**
The image depicts in light microscopy, the Ki-67 immunohistochemical staining of a section of a typical TL-ELN (center) within a stage IV (nonlocoregional), visceral melanoma metastasis. The tissue was stained using the avidin–biotin complex method with retrieval under high pH. Prediluted antibody to Ki-67 [Ventana Medical System, Inc.: anti–Ki-67 (30-9) primary antibody is a rabbit monoclonal antibody (IgG2) directed against C-terminal portion of Ki-67 antigen.] was used for the proliferative analysis of the TL-ELN. Lymphocytic proliferation within and surrounding the follicle (brown staining cells) was found to contain both B and T lymphocytes, suggesting newly formed and/or activated TL-ELNs within the tumor microenvironment. (Image generated by Jane Messina, Departments of Anatomic Pathology and Cutaneous Oncology at the Moffitt Cancer Center, Tampa, FL.)
By immunohistochemistry, Messina and colleagues (53) demonstrated that TL-ELNs uniformly contained prominent CD20+B-cell follicles surrounded by marginal zone areas of CD3+ T cells (comprising both CD4+ and CD8+ subsets). In addition, CD86+ follicular dendritic cells, but not Foxp3+ Treg cells, were present within these TL-ELNs. Stage IV melanoma metastases negative for the GES were uniformly found to have either an absence or a diffuse pattern of lymphoid infiltrate. There was a highly statistically significant (P<0.0001) and consistent association between a marked increase in overall patient survival, the value of the mean score of the GES, and the presence of TL-ELNs in stage IV melanoma metastases. These findings suggested an ongoing adaptive immune response within the melanoma microenvironment. More recently, van Baren and colleagues (57) have confirmed lymphoid neogenesis in melanoma. Moreover, Cipponi and colleagues (58) performed an analysis of the repertoire of rearranged immunoglobulin genes in B cells of the microdissected follicles within melanoma metastases and demonstrated clonal amplification, somatic mutation, and isotype switching, all of which suggested a local antigen-driven B-cell response. TL-ELNs have recently been described in primary cutaneous melanoma as well (59), but this has not been observed by others in a very limited number of specimens examined (58).

Future Directions

The recent progress in advanced melanoma therapy, both targeted and immune response related, has emphasized the importance of the relationship of melanoma and the host response as well as the tumor microenvironment. The treatment of BRAFV600-mutated melanoma with the BRAF-specific inhibitor BRAFi results not only in cessation of tumor growth but also in diminution in tumor size. This effect is also associated with an increase of CD8+ TILs in the affected tumor nodule; the treated tumor cells show an increase in MART-1 expression (60–62). Thus, targeted therapy is associated with TIL infiltration as a marker of its effect.

Furthermore, a study of the checkpoint blockade antigen PD-1 with one of its ligands, PD-L1, results in effective blocking of host response to the tumor as a result of exhaustion of the PD-1–bearing lymphocytes. PD-L1 can be found on a variety of cell types, including tumor cells, stromal cells, and macrophages. Treatment of patients with melanoma with a specific humanized mouse monoclonal antibody directed against PD-1, pembrolizumab, or other antibodies directed against PD-L1 can result in freeing of the host response and effective tumor regression. Tumeh and colleagues (63) reported that melanomas that have a great number of CD8+ lymphocytes at the advancing tumor margin are the ones that respond to this treatment. Results from their study indicate that release of the blockade leads to a proliferation of the CD8+ cells that migrate into the tumor and induce apoptosis and necrosis of tumor cells. In contrast, melanomas without CD8+ lymphocytes at the advancing tumor edge do not show response and the tumors progress. Once again, antigen-specific TILs that infiltrate the tumor show prognostic and therapeutic significance. TILs, by their presence in both of these instances, have led to better understanding of the tumor–host interaction and, at least in part, to the therapeutic response. The future lies in more intense study of the various inhibitors of TILs and of better ways to enhance their interaction not only with melanomas but also with other cancers. Special emphasis should be given to co-stimulatory and inhibitory molecules and the effect that TILs may have on enhancing or thwarting their positive or negative effects, respectively (63). Moreover, the phenomenon of the abscopal effect in patients with melanoma following treatment with the combination of anti–CTLA-4 antibody and radiotherapy, although linked to the participation of the immune system, has not yet been thoroughly examined, particularly with respect to changes in the melanoma microenvironment as it pertains to TILs (64, 65).

The existence of a functional connection between TL-ELNs and identifying and (and expanding) tumor-specific, therapeutic TILs is now being examined in melanoma. Laser-capture microdissection with DNA/RNA isolation and sequencing methodologies can also be used to address questions of clonality and functional relationships of the resident T cells within the TL-ELNs, including in-depth T-cell receptor (TCR) transcript repertoire analyses. Furthermore, the predictive nature of TL-ELNs in patients with melanoma who have participated, or are participating, in TIL-based (3, 66) and/or checkpoint antibody–based clinical immunotherapy trials is likely to be examined retrospectively. It will also be of interest to determine whether the presence of TL-ELNs reflects an underlying pathogen infection or is related to baseline gene mutational load of the particular melanoma.

In summary, we have traced TILs and their relationship to melanomas from their first identification under the microscope over a century ago to a progressive understanding of their relationship to tumor control and prognosis. In the molecular biology era, we find molecular-based therapies that directly affect patient survival. It is hoped that the great enthusiasm in the scientific community engendered by these discoveries will lead not only to melanoma control but also to a possible understanding of melanoma initiation and eradication by immunotherapy alone or in combination with other promising therapeutics.

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