Abstract. A mutual spatial and functional relationship occurs between mast cells (MCs) and endothelial cells and the density of MCs is highly correlated with the extent of tumor angiogenesis. The aim of this study was to investigate the pattern of MCs around the blood vessels in melanoma samples by means of an approach derived from spatial statistics, based on the analysis of the distribution of the distances of MCs from vessels to objectively establish if the two structures (MCs and vessels) are distributed independently over the studied area or if they displayed any kind of spatial association. Results showed that a higher number of vessels and MCs can be observed in melanoma as compared with samples from common acquired nevi (control group). The percent of area covered by vessel profiles was significantly higher in the melanoma group than the control group and the MC density was also significantly different; the melanoma group showing a number of MCs per unit area twice as high as the number measured in the control group. Furthermore, in the melanoma group, MCs were closer to each other and to the vessels. In fact, both the mean distance from vessels and the mean distance from the nearest cell profile were significantly lower than in the control group. This close association between MCs and the endothelium does not necessarily imply a participation of MCs in angiogenic processes, but might rather indicate that MCs are involved in the maintenance reaction necessary for the long lasting functional integrity of the endothelium.

Introduction

In solid tumor growth, a specific critical turning point is the transition from the avascular to the vascular phase (1). Having developed an intrinsic vascular network, the neoplastic mass is able to grow indefinitely both in situ and at distant sites (metastasis) in so far as an intrinsic vascular network enables its cells to enter the vascular bed and colonize other organs (2).

Human melanoma progresses through different steps: nevocellular nevi, dysplastic nevi, in situ melanoma, radial growth phase melanoma (Breslow index <0.75 mm), vertical growth phase melanoma (Breslow index >0.75 mm) and metastatic melanoma. In agreement with progression, it acquires a rich vascular network (3,4), whereas an increasing proportion of tumor cells express the laminin receptor, which enables their adhesion to the vascular wall (5).

Tumor cells are surrounded by an infiltrate of inflammatory cells, such as lymphocytes, neutrophils, macrophages and mast cells (MCs). These cells communicate by a complex network of intercellular signaling pathways mediated by surface adhesion molecules, cytokines and their receptors (6). It is becoming clear that stromal cells cooperate with endothelial and cancer cells in promoting angiogenesis. In particular, infiltrating inflammatory cells secrete a diverse repertoire of growth factors and proteases that enable them to enhance tumor growth by stimulating angiogenesis.

The density of MCs is highly correlated with the extent of both normal and pathological angiogenesis, such as that in chronic inflammatory diseases and tumors (7). In MC-deficient mice, tumor angiogenesis is reduced to significant extents (8). MCs release a variety of factors known to enhance angiogenesis; for example, heparin, histamine and angiogenic cytokines such as transforming growth factor ß, tumor necrosis factor a, interleukin-8, fibroblast growth factor-2 and vascular endothelial growth factor.

Two types of human mast cell (MC) have been described on the basis of the differences in their neutral protease composition (MC T cells, which contain tryptase only, and MC Tc cells, which contain both tryptase and chymase) (9). Tryptase positivity reflects the total number of MCs, whereas chymase is not present in all MCs. Blair et al (10) have shown that tryptase released by MCs at an angiogenesis site may play an important role in neovascularization. Direct addition of tryptase to microvascular endothelial cells cultured on Matrigel caused a pronounced increase of capillary growth, which was suppressed by specific tryptase inhibitors. Also,
tryptase directly induced endothelial cell proliferation in a
dose-dependent fashion.

We have previously demonstrated that, in B-cell non-
Hodgkin’s lymphoma, myelodysplastic syndromes, B-cell
chronic lymphocytic leukemia and melanoma, there is a
striking association between tryptase-positive MCs and micro-
vessel counts, and both increase in function of malignancy
(11-14). Moreover, we have further demonstrated that tumor
vascularity and tryptase-positive MCs correlate with a poor
prognosis in melanoma (15).

The aim of this study was to investigate the pattern of
tryptase-positive MCs around the blood vessels in melanoma
samples by introducing a quantitative approach to characterize
their spatial distribution.

Materials and methods

Tissue samples. Tissues were selected from the six clinical
steps of melanoma progression described by Clark et al (16).
The control group included tissue samples obtained from 15
common acquired nevi. The melanoma group included 20
advanced primary melanomas with a thickness >1.5 mm. The
cohort included 10 men and 10 women with an age range of
30-75 years. The distribution of anatomic location of the
primary tumors was as follows: trunk, ten; head and neck,
five; extremity, five.

Specimens were fixed in 10% buffered-formalin and
embedded in paraffin. Histological sections, 5 μm thick, were
prepared for immunohistochemical study.

Immunohistochemical staining. A murine monoclonal antibody
against tryptase (MAb AA1, Dako) was used. Briefly, sections
were collected on 3-amino-propyl-triethoxysilane-coated
slides, de-paraffinized by the xylene-ethanol sequence, re-
hydrated in a graded ethanol scale and in Tris-buffered saline
(TBS, pH 7.6) and incubated overnight at 4˚C with AA1
(1:1500 in TBS), after prior antigen retrieval by enzymatic
digestion with Ficin (Sigma, St. Louis, MO) for 30 min at
room temperature. The immunoreaction was performed using
the streptavidin-peroxidase complex (LSAB2, Dako) and
Fast Red as chromogen, followed by haematoxylin counter-
staining. An unrelated monoclonal IgG1 produced by the
P3x63Ag8 mouse secretory myeloma replacing the MAb
served as negative control (5).
Computer-assisted image analysis was performed to characterize the distribution of MCs around the vessel profiles. The image analysis system included a Leica DM-R microscope (Leica Microsystems, Wetzlar, Germany) and a high-resolution digital camera (DC200, Leica Microsystems) that transmits image data to a PC equipped with appropriate software for image acquisition and analysis (QWin, Leica Microsystems, Cambridge, UK). At a primary magnification of x16, 4 fields covering the whole of each of two sections per biopsy were considered and the image of each field was acquired in full colour (24-bit), processed to correct the shading and enhance the contrast, and stored as a TIFF file.

The main steps of the image analysis procedure are illustrated in Fig. 1. The study area within the image was defined as the minimum rectangle bounding MCs and vessels. Colour thresholding was then applied for the identification of the MC profiles. According to this procedure, pixel colours were represented as HSI (hue, saturation, intensity) values (17). In general, for stains used in biological samples, the ‘hue’ component identifies the location of a particular stain, while the ‘saturation’ corresponds to the amount of staining and the ‘intensity’ indicates the overall density of the stained specimen. Selecting only pixels with a red hue and saturation greater than the mean saturation level exhibited by the background plus two standard deviations, a binary image containing the MC profiles only was identified (Fig. 1B). A second binary image (Fig. 1C) corresponding to the vessel profiles present in the image was also obtained by interactively tracing their contour. From these images, the number of MC profiles, their positions (i.e. x, y-coordinates of the gravity centers), the distance of each profile from its nearest neighbour and the area fraction occupied by vessels were evaluated. To estimate the distance of each MC profile from the vessels, the binary image of the vessel profiles was further processed to calculate its ‘distance transform’ (17). This algorithm provides a map where each background pixel is labelled with a value equal to its distance from the nearest pixel belonging to a vessel profile. The distance from vessels of each MC profile was then evaluated by the value the map exhibited at the location corresponding to the x, y-coordinates of the cell profile.

Around the observed set of vessel profiles, 100 random (Poisson) point patterns were finally computer generated (Fig. 2). Each pattern had a number of points equal to the number of observed mast cell profiles. They underwent the previously described analysis in order to provide Monte Carlo estimates (18) of the distances from vessels in the case of complete spatial randomness (CSR).
Statistics. Within each sample, MC density values (number of cells per unit area of sampled tissue), distances from the nearest neighbour and vessel area fractions were averaged to provide a representative value for that sample. Differences between the two groups of samples were then statistically tested by Student's t-test. The GraphPad Prism 3.0 statistical package (GraphPad Software Inc., San Diego CA, USA) was used for the analysis and p≤0.05 was considered as the limit for statistical significance.

For each group, the cumulative frequency distribution \( G(d) \) of all the observed cell-to-vessel distances was calculated. Its expected value under CSR \( G_0(d) \) was estimated by averaging the cumulative frequency distributions of the distances from vessels obtained from the 100 simulated random point patterns. To interpret the cell-to-vessel spatial relationship statistically, the 95% confidence envelope for \( G_0(d) \) was also calculated from the Monte Carlo simulations (19). The null hypothesis is that there is no difference between the two functions, i.e. \( G(d) = G_0(d) \) for all d. Thus, if \( G(d) \) is greater than the confidence envelope around \( G_0(d) \), then the cells are clustered around the vessels, i.e. they are closer to the vessels than expected by chance. If \( G(d) \) is lower than the envelope around \( G_0(d) \), then short cell-to-vessel distances are less frequent than expected by chance, i.e. the placement of the cells close to the vessels was ‘inhibited’ (20).

Results

As shown in Fig. 3, a higher number of vessels and MCs can be observed in the tissue sections from the lesions of patients with primary melanoma as compared with samples from patients with common acquired nevi (control group).

Table I. Tissue density of vessels and tryptase-positive cells.

<table>
<thead>
<tr>
<th>Group</th>
<th>Common acquired nevi</th>
<th>Primary melanoma</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vessels (area % ± SEM)</td>
<td>1.41±0.27</td>
<td>8.14±1.8</td>
<td>0.0061</td>
</tr>
<tr>
<td>Tryptase-positive cells/mm²</td>
<td>48.5±8</td>
<td>101.7±8</td>
<td>0.0015</td>
</tr>
</tbody>
</table>

As shown in Table I, the percent of vessels and MCs is significantly higher in the primary melanoma group than in the control group. This morphological observation was confirmed by morphometric evaluation as shown in Table I. The percent of vessels and MCs is significantly higher in the primary melanoma group than in the control group.
area. To derive this reference distribution, a simulation in which the MCs are distributed randomly over the studied corresponding to the case of complete spatial randomness, i.e. observed distribution of MC-to-vessel distances with the one association. Such an analysis involved the comparison of the structures (MCs and vessels) were distributed independently vessels with the aim to objectively establish whether the two analysis of the distribution of the distances of MCs from 24,25). The method was based on the statistical we addressed this topic by an approach derived from spatial MCs and vessels, however, has never been provided. Here, the density of blood vessels in the dermis. to capillaries and lymphatic channels (21-23). Eady and under physiological conditions, numerous MCs are close occurring between MCs and endothelial cells. In many organs much evidence of the mutual spatial and functional relationship Discussion

MCs are widely distributed in connective tissue and there is much evidence of the mutual spatial and functional relationship occurring between MCs and endothelial cells. In many organs and under physiological conditions, numerous MCs are close to capillaries and lymphatic channels (21-23). Eady et al (21) found a direct correlation between the number of MCs and the density of blood vessels in the dermis.

A quantitative assessment of the spatial co-localization of MCs and vessels, however, has never been provided. Here, we addressed this topic by an approach derived from spatial statistics (24,25). The method was based on the statistical analysis of the distribution of the distances of MCs from vessels with the aim to objectively establish whether the two structures (MCs and vessels) were distributed independently over the studied area or displayed any kind of spatial association. Such an analysis involved the comparison of the observed distribution of MC-to-vessel distances with the one corresponding to the case of complete spatial randomness, i.e. in which the MCs are distributed randomly over the studied area. To derive this reference distribution, a simulation technique was applied (18), involving the computer generation of point patterns of the same size as the observed cell pattern, but placed in the area under investigation according to a random (Poisson) distribution. Analyses very similar from a statistical point of view were recently applied to characterize the degree of spatial association between cell populations (26), the co-localization of immunogold labels (20) and the spatial relationship between microdomains in the plasma membrane (27). The main difference between these approaches and the method proposed here is that, in the present study, one of the two structures (i.e. the vessels) was not modelled as a point pattern but as a set of surfaces. As a consequence, distances from each cell to the nearest vessel profile boundary (roughly corresponding to the endothelial lining) were considered for the analysis. The results showed that, in both experimental groups evaluated in the present study, the spatial distribution of MCs was characterized by a significant spatial association between MCs and vessels, as indicated by a frequency of short distances higher then expected under the hypothesis of a random spatial distribution of cells.

This anatomical association between MCs and the endothelium, however, does not necessarily imply the participation of MCs in angiogenic processes, but might rather indicate that these cells are involved in the maintenance reaction necessary for the long lasting functional integrity of the endothelium (28).

Human dermal endothelial cells express the MC growth and chemotactic factor stem cell factor (SCF) (29,30). Mierke et al (31) found that MCs survive for many weeks in the absence of any exogenous growth factor if cultured on an endothelial cell layer. Endothelial cell-dependent MC survival is mediated by SCF and adhesion molecules, such as vascular cell adhesion molecule-1 (VCAM-1). Furthermore, SCF induce uPAR-expression in MCs, and cells stimulated in this way could also chemotactically respond to uPA released by endothelial cells (32). These findings suggest that, under physiological conditions, endothelial cells play an important role in regulating MC influx as well as MC development and function in tissue.

MCs have been linked for almost three decades to neo-vascularization. Several lines of evidence have implicated MCs in the regulation of pathological or physiological examples of angiogenesis, including that associated with haemangiomas (33), tumors (7), rheumatoid arthritis (34), nasal polyps (35), wound healing (36) and ovulation (37). We have previously demonstrated that, in B-cell non-Hodgkin's lymphoma, myelodysplastic syndromes, B-cell chronic lymphocytic leukemia and melanoma, there is a striking association between tryptase-positive MCs and microvessel counts, and both increase in function of malignancy (11-15).

Here, we demonstrate that a higher number of vessels and MCs can be observed in tissue sections from the lesions of patients with primary melanoma as compared with samples from common acquired nevi (control group). In fact, the percent area covered by vessel profiles was significantly higher in the melanoma group than in the common acquired nevi group and the MC density was also significantly different; the melanoma group showing a number of MCs per unit area twice as high as the number measured in the common acquired nevi group.

Figure 5. Mean intercellular MC distance (estimated as the distance from the nearest neighbour cell) and mean distance from vessels in the common acquired nevi group (N=15) and in the melanoma group (N=20). Error bars are SEM. *p<0.05; **p<0.01 (two-sample Student's t-test).
closer to each other and to the vessels. In fact, both the mean distance from vessels and the mean distance from the nearest cell profile were significantly lower than in the common acquired nevi group of tissue samples. Shorter intercellular distances and lower cell-to-vessel distances could be a morphological condition important to the increases in the rate of signal exchange among the cells and vessels, and to the induction of higher concentrations of signalling molecules in the pericellular and periendothelial environment.

All of these findings indicate that MCs may enhance angiogenesis, at least in the tumor model, and the preferential localization of MCs along blood vessels and sites of new vessel formation sustains the suggestion for an association between MCs and angiogenesis. It is possible, however, that the spatial association between vessels and MCs simply reflects migrating MCs from the blood stream at vessel growing sites (38). Until further data are available, the question as to whether MCs are main factors or innocent bystanders in this process will remain unsolved.

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References


