The Clinical Significance of the Primary Malignant Melanoma of the Lower Respiratory Tract and/or Lung Based on the Analysis of Published Case Reports and of Two Patients

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Considering the data and including two patients of the authors, there exist only 18 authentic cases of primary malignant melanoma of the lower respiratory tract and/or the lung. The tumor was localized in the endobronchial space in 7 cases and only once in the trachea. Endobronchial localization, together with the involvement of the surrounding lung tissue, was found in two cases, whereas in 8 patients, the tumor was found exclusively in the lung parenchyma. Successful resection could be performed in 14 of the 18 cases. Survival was influenced primarily by operability, and on actual size and extension. The authors question the role of the obduction in the diagnostic criteria, because most of the survivors, even up to ten years postsurgery were considered primary. Apart from the various imaging methods, diagnostic endoscopy (i.e., bronchoscopy) and the histology of the biopsy material are major aids in the diagnosis of primary character, location, and operability, as well as in the elaboration of the surgical plans, than it is usual in cases of other tumors.

KEY WORDS: melanoma malignum, primary melanoma in lung, primary melanoma in the bronchial tract

INTRODUCTION

Eighty-eight percent of the primary malignant melanomas appear in the skin or the eye (1). As primary tumors, melanomas occur considerably less frequently in the oral cavity, upper respiratory tract, esophagus, liver, gall bladder, ovaries, cervix uteri, uterus, urogenital system, and leptomeninx. The malignant melanoma usually occurs as a metastatic tumor in the lower respiratory tract and/or lung (in 7% to 9% of those of cutaneous origin and in 2% of the autopsies) (2,3), and the primary forms of the same localization are extremely rare. It is assumed that the primary melanomas derive from the benign melanocytes moving together with the mesoderm and/or entoderm (4,5). Similar cells have been isolated from the mucosa of the esophagus and larynx. The latter organs derive from the 6th branchial arch as parts of the pulmonal system. Nevertheless, the presence of even benign melanocytes in the bronchopulmonary system is not usual (6). According to some suggestions, melanocytic metaplasia also may take place under various stimuli in epithelial cells, salivary glands, and nervous structures (7). It is possible also that some precursor cells (e.g., Kultschitsky cells) may undergo both neuroendocrine and melanocytic differentiation (8,9,15).

A careful analysis of the literature reveals that the majority of the really primary malignant melanomas located in the lower respiratory tract and/or lung have been described in English-speaking countries (Table 1); we find only Danish (11), French (12), and Spanish (13) publications in Europe, in addition to those in Great Britain (5,13a–21). Thus, we describe observations obtained from our two patients and survey the relatively few publications.
Mrs. B. J. is a 43-year-old patient. In her anamnesis there was a uterus extirpation because of myomas. She recovered in our department on November 3, 1989, because of shadow seen by X-ray examination in the left thoracic region. She has been coughing for several weeks and is occasionally febrile. Physical examination did not reveal any alterations on the skin; no lymph nodes could be found. The repeated posteroanterior summation X-ray examination and tomographic imaging of the left side in planes of sagittal direction revealed a round vascular shadow about 6 to 7 cm in the left hilar region (Figure 1). Bronchoscopy revealed the presence of a tumor at the beginning of the left upper lobe bronchus; the tumor was about 8 mm with a slightly uneven surface and brown-gray color (Figure 2). Histologic and cytologic analyses of the excised sample suggested that the alteration represented an anaplastic cancer of the lung consisting of intermediary cells. Because of this diagnosis, left pneumonectomy was performed on November 22, 1989.

Pathologic, histologic and macroscopic findings were left lung, $18 \times 12 \times 3$ cm weighing 290 g. In one of the branches of the upper lobe bronchus we found a brown-gray tumor about 8 mm. The bronchial wall infiltrated by the tumor was in close proximity to a huge lymph node of the hilus; this node was $5 \times 4 \times 3.5$ cm, of tumorous appearance, part gray-white, part dark brown-gray, and soft and fragile.

Microscopic examination showed the tumor infiltrating all layers of the cartilaginous wall of the bronchus, and intruding into the bronchial lumen in polyplike form. The tumor surface is covered in part by typical ciliated respiratory epithelium, displaying in some places slightly atypical planocellular metaplasia or atypical melanocytic hyperplasia. Under the epithelium, several naevoid cell groups that were confluent with the tumor were observed.

### Table 1 Basic data of case reports recognizable as primary malignant melanoma of the lower respiratory tract and/or lung

| Authors         | No | Year | Age | Sex | Country      | Survival time | Search for primary melanoma |
|-----------------|----|------|-----|-----|--------------|---------------|-----------------------------|
| Salm            | 1  | 1963 | 45  | M   | England      | 6 months      | done                        |
| Reed and Kent   | 2  | 1964 | 71  | M   | USA          | 10 years      | done                        |
| Reid and Mehta  | 3  | 1966 | 60  | F   | New Zealand  | 11 years      | done                        |
| Allen and Drash | 6  | 1968 | 40  | F   | USA          | not published | done                        |
| Taboada et al.  | 7  | 1972 | 56  | M   | USA          | 3 years       | done                        |
|                 | 8  | 1972 | 40  | M   | USA          | 1 year        | done                        |
Table 1  Continued

| Authors         | No | Year | Age | Sex | Country | Survival time | Search for primary melanoma |
|-----------------|----|------|-----|-----|----------|---------------|----------------------------|
| Walter et al.   | 9  | 1972 | 33  | M   | France   | 22 months     | done                       |
|                 |    |      |     |     |          |               |                            |
| Adebonojo et al.| 10 | 1979 | 55  | F   | Nigeria  | 3 years       | done                       |
|                 |    |      |     |     |          |               |                            |
| Gephardt        | 11 | 1981 | 47  | M   | USA      | before interv. | done                       |
|                 |    |      |     |     |          |               |                            |
| Cagle et al.    | 12 | 1983 | 80  | M   | USA      | 5.5 months    | done                       |
|                 |    |      |     |     |          |               |                            |
| Alghanem et al. | 13 | 1987 | 42  | F   | USA      | 2.5 years     | done                       |
|                 |    |      |     |     |          |               |                            |
| Santos et al.   | 14 | 1987 | 58  | M   | Spain    | 18 months     | done                       |
|                 |    |      |     |     |          |               |                            |
| Bagwell et al.  | 15 | 1989 | 62  | M   | USA      | 2 months      | done                       |
|                 |    |      |     |     |          |               |                            |
| Jennings et al. | 16 | 1990 | 34  | F   | USA      | 19 months     | done                       |
|                 |    |      |     |     |          |               |                            |

Several small exulcerations also were found on the tumor surface. Most of the tumor mass consisted of spindle cells with hyperchromic nuclei; the cells were partly ordered in bundles of various directions or in vortexlike structures (Figure 3). Although the tumor was rich in cells, the occurrence of apparent mitoses was not high. The massive brown pigment depositions of the tumor cells proved to be melanin with silver stains; the distribution of this pigment is not uniform (Figure 4). The histologic structure of the lymph node is unrecognizable; it is filled completely with tumor tissue. Similar to other bronchial tumors in all aspects, the tumor is in part fusocellular in structure containing abundantly melanin pigment, and in part large, pale rounded or polygonal cellular in structure with large, vacuolated nuclei and mostly ballonized cytoplasm. These latter cells contain much less melanin. The HMB-45 melanoma marker procedure proved to be positive. The resection line of the left main bronchus did not contain tumorous tissue.

Diagnosis: Melanoma malignum bronchi lobi superiors pulmonis sinister. Metastasis tumoris in lymphanodu hilis pulmonis sinister.

Opinion: The tumor is considered a primary bronchial melanoma. This opinion is supported by the following morphologic signs: a single tumor of central location in the bronchial lumen; the polypoid surface of the tumor covered by an epithelium displaying atypical melanocytic proliferation and atypical papillomatous metaplasia, and epithelium naevoid cell nests confluent with the tumor (Figures 3–4).

The postoperative period was without complication. The patient received Detecin (dacarbacin) therapy in 12 cycles during the first postoperative year. The metastatic origin of this tumor in the bronchus of the left upper lobe of lung could be excluded with high probability based on the negative results of dermatologic, otolaringologic, oculistic, and gynecologic, and abdominal ultrasound examinations. Because the skin did
Mrs. K. L., age 81 years complained of diffuse chest pains and dyspnoe. X-ray examination showed an egg-sized shadow in the segment 10 of the left side and a lymph node conglomerate in the hilus (Figure 5). Bronchoscopy was performed on November 21, 1990 and revealed that the lower part of the left main bronchus was almost completely closed by a tumorous gray-brown mass infiltrating the bronchial wall and penetrating from the direction of the bronchus of the lower lobe (Figure 6). To increase the bronchial lumen and improve the drainage, small parts of the tumor were removed. This intervention was repeated after 2 weeks, because the patient reported an improvement of her condition.

Histologically, the tumor is rich in cells and already has reached the surface of the bronchial mucosa. The tumor cells form solid groups, are medium-size with partly pale, eosinophilic cytoplasm, and contain numerous, focally localized, finely distributed, brownish, melaninlike pigment granules. The cell nuclei are hyperchromic and are ovoid or elongated, and some contain large nucleoli. There is an expressed cellular atypia and polymorphism. The brownish pigment tested positive for Grimelius and Masson-Fontana silver stainings, and the immunohistochemical HMB-45 melanoma result was also positive. All these results supported the diagnosis of melanoma malignum.

The biopsy material also was analyzed electronmicroscopically; the cells containing the brownish melaninlike pigment also displayed numerous premelanosomes or melanosomes of high electron density (Figures 7 and 8).

The patient was observed for 4 months and was hospitalized for 23 days from this period. The analyses performed could not reveal the presence of malignant melanoma except in the bronchopulmonary location. The careful derma-
The first authentic primary malignant bronchopulmonal melanoma was described by Salm in 1963. In some of the cases described in later publications, one has to doubt the primary lung and/or bronchial origin. (22–26). The most serious issue is the exclusion of the metastatic origin of the tumor, because according to some opinions, skin melanomas may spontaneously disappear without any trace after they already have metastased. (2,27). However even if one accepts the possibility of the eventual, rare spontaneous regression of the primary skin melanoma,
oneshouldgenerallyexpectalatergeneralization (27). It
is also a fact that more than 15% of the metastatic
melanomas have a hidden primary tumor (28). Therefore
Jensen and Egedorf already have defined in 1967 the
mainly clinical conditions necessary for the safe diagno-
sis of the primary malignant melanoma of the lower res-
piratory tract. These conditions include:

1. No previous removal of pigmented skin tumor or
   any kind of skin tumor
2. No removal of eye tumors, especially through enu-
   cleation
3. Surgically removed specimen with a solitary tumor
4. Morphology of tumor corresponds to the primary
   origin
5. No detectable melanoma in any other organ at time
   of surgical intervention
6. No primary malignant melanoma found, especially
   in the skin and the eyes, at autopsy.

The macroscopic and histologic properties of the tumor
may offer further evidence for the origin of the tumor from
the lower respiratory tract (6,15,21,29). Primary process
is likely, if the tumor is polypoid and localized intralumi-
nally, whereas the metastases are usually disseminated.
The histologic conditions include:

1. Junctional alterations with the nestlike location or
   "dropping" of the melanoma cells
2. Invasion of melanoma cells in the bronchial epithe-
   lium at non-eroded places
3. Intraepithelial alterations that correspond unani-
   mously to melanoma.

An accessory sign is considered to be the presence of
naevoid alteration in the bronchial epithelium nearby
to the tumor and the melanoma flare that somewhat more
distant from the tumor. (6,15,21). In our first reported case,
the listed histologic criteria were present, whereas in our
second case, not all criteria could be met, mainly because
of the small size of the biotic materials.

Generally speaking, the above criteria are considered
valid, although the autopsy cannot always be performed,
as described in some published cases, because of long
postoperative survival or other reasons. On the other hand,
autopsy itself has some limitations especially in answer-
ing the dilemma of primary tumor or metastasis.
Therefore, if the obduction cannot be performed, the neg-
ative outcome of an extensive search for primary tumor
in vivo can be accepted. Using modern endoscopic and
imaging procedures, the probability of error can be re-
duced. Therefore, although the criteria defined by Jensen-
Egedorf (1967) for the diagnosis of the primary malignant
melanoma of the lower respiratory tract should be con-
sidered the standard, cases should be qualified based on
the analysis of all test results and on common sense.

In both our cases, we found a solitary tumor with metas-
tases in the hilus of the identical side. The malignant
melanoma was proved by histologic, histochemic, and
electron microscopic techniques. The tumors did not show
any morphologic signs that suggested their metastatic
origin.

According to the data (see Table 1), the primary malig-
nant melanoma of the lower respiratory tract and/or lung
involves both sexes almost equally. At time of diagnosis, the
youngest patient was 33, the oldest 81 years of age. Of the
14 operated cases (lobectomy and pulmonectomy), two and
one patients, respectively, are still alive, i.e., after 3 to 4 years,
and one patient in each group survived after 10 to 11 years.
Three patients were alive 2 to 30 months after the interven-
tion. Five patients died between 5.5 and 22 months, and one
patient died during the postoperative phase of tracheal re-
section. There was one patient who underwent successful
right lower lobectomy whose survival is not known.

The tumor was endobronchial in seven patients, and
was intratracheal in one patient. The endobronchial loca-
tion was accompanied by the involvement of the adjacent
lung area in two persons, and exclusive parenchymal loca-
tion was seen in eight cases. These facts indicate the
significance of the bronchoscopy in the diagnosis.
Operability and Survival are determined not by the
bronchial or parenchymal location but by extension and
circumscription of the tumor. The presence of a solitary,
metastatic hilar lymph node, even if it is of considerable
size, does not predict an unequivocal bad prognosis, as
was shown in our first case.

In the therapy of the primary malignant melanoma of the
lower respiratory tract, surgical removal is most important;
considerable survival can be expected only from this in-
tervention. The cytostatic treatments applied so far in the
inoperable cases did not give any result; the effects of the
adjuvant treatments cannot be measured either. Therefore,
the main task is to detect the tumor as early as possible and
if operable, surgical intervention must be performed.

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