Putative Malignant Pleural Mesothelioma in situ (MPMIS) with Sequential Acquisition of Genomic Alterations on Fluorescence in situ Hybridization (FISH) Examination

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Established Facts

- Criteria for the diagnosis of malignant pleural mesothelioma in situ have been proposed to be based on clinical, radiological, and morphological features.
- Fluorescence in situ hybridization analysis is a sensitive and highly specific ancillary tool for the diagnosis of malignant mesothelial cells.
- Satellitosis is a morphological phenomenon described among others in the context of acute alcoholic hepatitis.

Novel Insights

- Fluorescence in situ hybridization analysis and BAP1 immunochemistry are a potential screening tools to detect malignant pleural mesothelioma in situ in patients presenting with recurrent effusions of unknown origin and no clinical evidence of mesothelioma.
- Genomic transition from a diploid to an aneuploid state might play a role in progression from mesothelioma in situ to invasive mesothelioma.
- Satellitosis of mesothelial cells by lymphocytes in pleural effusion cytology is a rare and not tumor-specific phenomenon with unclear relevance.

Keywords

Mesothelioma in situ · BAP1 · CDKN2A · 9p21 deletion · Satellitosis

Abstract

Malignant pleural mesothelioma (MPM) is a rare and deadly disease. A precursor in situ lesion, malignant pleural mesothelioma in situ (MPMIS), has recently been proposed. On cytological examination, the distinction between reactive and malignant mesothelial cells is often challenging, and sometimes even impossible without ancillary methods. Fluorescence in situ hybridization (FISH) for detection of 9p21 deletion is a powerful diagnostic tool in this context, both in histological and in cytological specimens. Here, we present a case of MPM with initial presentation as a putative MPMIS...
with disomic chromosomal pattern and homozygous 9p21 deletion with subsequent development of an aneuploid pattern after whole genome duplication during tumor progression.

Introduction

Malignant pleural mesothelioma (MPM) is a rare cancer associated with poor prognosis [1]. The existence of a malignant mesothelioma in situ has long been postulated [2]. Recently, the criteria for the diagnosis of malignant pleural mesothelioma in situ (MPMIS) had been proposed to be based on clinical, radiological, and morphological features [3, 4]. The prerequisite is the absence of a clinical and radiological tumor manifestation as evidence of an established malignant mesothelioma. Here, we present a case of MPM with initial presentation as a putative in situ lesion showing sequential acquisition of genomic alterations on fluorescence in situ hybridization (FISH) examination during tumor progression.

Case Presentation

A 57-year-old man, non-smoker, and electro-mechanic by profession, presented with cough and B-symptoms for more than 3 months. Clinically and radiologically, a MPM was suspected. A thoracentesis was performed, and pleural fluid was obtained. On cytological examination, there were numerous malignant mesothelial-like cells with frank atypia (shown in Fig. 1a). The mesothelial phenotype was confirmed by immunohistochemistry (IC) and showed calretinin expression and absence of BerEP4 expression. There was a complete loss of desmin expression. The diagnosis of malignant mesothelial cells was ascertained by FISH using a

Fig. 1. a Pleural effusion at the time of diagnosis showing large clusters of unequivocal malignant mesothelial cells with prominent nucleoli (Papanicolaou [Pap] stain, ×630). d Initial pleural effusion highlighting satellitosis of mesothelial cells by lymphocytes; the mesothelial cells only show mild atypia (Pap stain, ×630). b, e Initial pleural biopsy: atypical mesothelial cells on histology (b, hematoxylin and eosin stain, ×400) with loss of BAP1 expression. Internal positive control with nuclear BAP1 expression is observed in the adjacent leucocytes (e, immunoperoxidase, ×400). c Multiprobe FISH result of the initial effusion: regular disomic pattern for chromosomes 3 (red), 7 (green), and 17 (blue) and homozygous deletion of the chromosomal region 9p21 (lack of gold signal) (×630). f Multiprobe FISH result of the pleural biopsy of malignant mesothelioma 8 months later: homozygous deletion of the chromosomal region 9p21 (lack of gold signal) and polysomy of the chromosome 3 (red), 7 (green), and 17 (blue) in the cytological specimen (f, DAPI). FISH, fluorescence in situ hybridization.
multiprobe FISH assay (UroVysion™, Abbott, Chicago, IL, USA) revealing unbalanced polysomy of the chromosomes 3, 7, and 17 and homozygous loss of 9p21 (shown in Fig. 1f). The MPM diagnosis was confirmed by a subsequent pleural biopsy. Interestingly, 8 months before, we had received pleural effusion fluid and a pleural biopsy from the patient, which had been diagnosed as unspecific organizing pleuritis without signs of malignancy (shown in Fig. 1b, 2a–b). At this time, no mesothelioma was suspected, radiologically and clinically. Review of the biopsy samples showed single layered and mildly atypical mesothelial cells without infiltrative growth pattern. Also, the effusion cytology has been rendered as unsuspicious. On review, it contained lymphocytes and mesothelial cells without frank atypia. Interestingly, a substantial number of the mesothelial cells were surrounded and encircled by lymphocytes, which we refer to as satellitosis of mesothelial cells by lymphocytes (shown in Fig. 1d). The encircled mesothelial cells were positive for calretinin. Satellitosis of mesothelial cells by lymphocytes was not present in the cytological specimen with the full-blown mesothelioma. Retrospective IC and FISH performed at the initial presentation revealed a loss of desmin and BAP1 expression (shown in Fig. 1e) as well as a homozygous deletion of 9p21 (shown in Fig. 1c) but without polysomy of the chromosome 3, 7, and 17 as seen in the subsequent effusion specimen with MPM diagnosis (shown in Fig. 1f). Considering these results, we hypothesize that during the time interval between the 2 samples (8 months), the tumor cells had undergone whole genome duplication and that the lesion initially biopsied could represent an in situ form of MPM as, at this time, there was no radiological and clinical evidence of MPM. A pleural biopsy confirmed the diagnosis of mesothelioma (shown in Fig. 2c), and a loss of MTAP could be observed on retrospective IHC (shown in Fig. 2d–e). Unfortunately, there was no more tissue left for MTAP IHC of the early manifestation after various IHC and FISH analyses.

Being intrigued by the satellitosis of mesothelial cells by lymphocytes in the initial specimen, we performed a data base search for Papanicolaou-stained cytological samples containing “malignant pleural mesothelioma” and “lymphocyte-rich effusion.” These specimens were reviewed in order to determine the incidence of this phenomenon in pleural effusions. We identified 60 cases with either histological or radiological proven MPM and reviewed 60 cases with the diagnosis of “reactive mesothelial cells” and “lymphocyte-rich effusion.” We observed “satellitosis” in a total of 5 cases, 2 cases with malignant mesothelial cells including the case presented here, and 3 cases with reactive mesothelial cells (shown in Fig. 3a–d).

**Fig. 2.** a Low-power view of the initial pleural biopsy being consistent with chronic pleuritis with lymphocytic inflammation (a, HE, ×5). b High-power view of the initial pleural biopsy, highlighting a single layer of cuboidal mesothelial cells, without frank atypia at the surface of the pleura (b, HE, ×20). c Low-power view of the invasive pleural mesothelioma with infiltration of the lung parenchyma (c, HE, ×5). d, e Complete loss of MTAP in the neoplastic mesothelial cells by IHC. Retained MTAP expression in adjacent lymphocytes and stromal cells (d, e, HE, ×20).
Discussion and Conclusion

The existence of an in situ precursor of invasive malignant mesothelioma has long been disputed but is now an accepted diagnostic concept [2–4]. The main challenge in clinical practice lies in the large surface of the pleura, which makes it impossible to exclude coexistent focally invasive MPM due to limited biopsy sampling. The International Association for the Study of Lung Cancer recently proposed that MPMIS lesions should be diagnosed based on clinical, radiological, and morphological features, the prerequisite being the absence of clinical and radiological evidence of mesothelioma [3]. In this setting, an atypical mesothelial proliferation limited to the pleural surface only qualifies as MPMIS if a BRCA1 associated protein-1 (BAP1) loss and/or cyclin-dependent kinase inhibitor 2A (CDKN2A) homozygous deletion is present [3, 4]. In our case, the neoplastic mesothelial cells present in the pleura sample at the time of the initial effusion meet the International Association for the Study of Lung Cancer criteria proposed for MPMIS, as the patient did not show any radiologic evidence of mesothelioma and as the mesothelial proliferation was restricted to the pleural surface with homozygous deletion of 9p21 containing the CDKN2A gene and loss of BAP1 expression.

FISH analysis has been shown to be a sensitive and highly specific ancillary tool for the diagnosis of malignant mesothelial cells and for its distinction from reactive mesothelial cells in histological biopsies and effusion cytology, respectively [5–7]. In the present case, FISH played a key role in identifying the mesothelial cells as malignant both at the time of the first effusion and at the time of clinically manifest disease 8 months later. Using a multiprobe FISH assay also allowed us to observe genomic progression from a diploid to an aneuploid state, which was most likely driven by whole genome doubling. Whole genome doubling has recently been recognized as a macro-evolutionary event in cancer arising early in carcinogenesis after an antecedent transforming driver mutation [8]. Homozygous 9p21 deletion and/or inactivating BAP1 mutation could qualify as such early transforming alterations in case of malignant mesothelioma. The combina-
tion of BAP1 IC and FISH for detection of 9p21 deletion has been shown to be complementary to increase the sensiti-
tivity for diagnosing MPM [9]. In addition, loss of
MTAP expression by IC has emerged as a promising sur-
rrogate marker for homozygous 9p21 deletion, which is
particularly interesting for laboratories without access to
FISH analysis [7, 10]. The MTAP gene is located near to
CDKN2A/p16 and co-deleted in the vast majority of
MPM with 9p21 deletion. However, until there are more
data on the robustness of MTAP IC, FISH remains the
gold standard to detect homozygous 9p21 deletions. The
rapid progression from invisible early diploid mesothe-
lial neoplasia to an advanced aneuploid manifestation of
diffuse mesothelioma within a short time interval of 8
months would be remarkable, but possible considering
the variable progression dynamics of pleural mesothelio-
ma [8]. The interval between in situ and invasive meso-
theлиma have been as short as 12 months [8]. Neverthe-
less, we cannot exclude an undetected focus of established
aneuploid mesothelioma at the time of the first effusion
due to limited sampling.

Even if the main cytological features that indicate a me-
sothelial origin for neoplastic cells are well described, in
practice, the diagnosis of malignant mesothelioma can be
difficult [11]. Sometimes, the malignant cells are indistin-
guishable from benign, reactive mesothelial cells, as evi-
denced by the first effusion of our patient [12]. The in-
triguing satellitosis of the rare neoplastic mesothelial cells
by lymphocytes could be suspected to reflect early interac-
tion of T-cells due to neoantigen recognition. Interest-
gly, it has previously been proposed that human peritoneal
mesothelial cells are equipped with an antigen-presenting
machinery to recall antigens to T-cells [13–15]. The ob-
served satellitosis might be a morphological correlate of
such interaction between T-cells and mesothelial cells.
Satellitosis – defined as abnormal clustering of cells encir-
cling others cells – has been observed in the context of
acute alcoholic hepatitis, where ballooned, damaged he-
patocytes are surrounded by neutrophils or in certain
brain tumors, for example, in oligodendroglioma, where
the neoplastic oligodendrocytes tend to cluster around
neurons exhibiting a so called “perineuronal satellitosis”
[16, 17]. Our review of a consecutive series of benign and
malignant mesothelioma effusions showed that satellit-
osis of mesothelial cells by lymphocytes is a rare phenom-
enon that is not specific for malignant mesothelial as we
observed it in both reactive and malignant samples. We
cannot exclude that this phenomenon only represents a
physical artifact caused by unknown variables during pro-
cessing of the samples. Nevertheless, further investiga-
tions are needed to study if there are specific interactions
between lymphocytes and mesothelial cells in reactive
conditions or during early mesothelial oncogenesis that
could explain this morphological finding.

In conclusion, we present a case of putative MPMIS
with homozygous 9p21 deletion and diploid chromo-
somal pattern that progressed to a clinically manifest an-
euploid MPM as shown by FISH analysis. This report em-
phasizes the utility of ancillary testing to detect MPMIS
or early invasive MPM even in cytology or biopsy speci-
mens that would otherwise be diagnosed as benign or re-
active in patients presenting with recurrent effusions of
unknown origin. It also illustrates the potential usefulness
of ancillary testing by FISH analysis and/or BAP1 IC
as screening tools for MPMIS in patients at risk due to a
history of asbestos exposure who present with otherwise
unexplained pleural effusion. Satellitosis of mesothelial
cells by lymphocytes is a rare and not tumor-specific phe-
nomenon with unclear relevance.

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Statement of Ethics

This study was covered by the local institutional approval EK
253/08 on human research (EKBB/EKNZ). A written informed
consent could not be obtained from the patient as he passed away
10 years ago. As the report does not contain any critical data (pic-
ture or radiography of the patient) and consequently anonymity is
guaranteed, a consent from the patient is, in this particular case,
not strictly required according to the ethics approval.

Conflict of Interest Statement

The authors have no conflict of interest to declare.

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Author Contributions

L.B. and S.H. conceived and designed the project. L.B., S.S., J.R.,
and S.H. analyzed the data. H.B. obtained cytological and histo-
logical specimens. S.H., L.B., H.B., and S.S. wrote the manuscript.
All authors agreed to the content of the manuscript.
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