Spontaneous Remission of Severe Systemic Langerhans Cell Histiocytosis with Bladder Involvement: A Case Study

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Keywords
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Abstract
\textbf{Background:} The clinical presentation of Langerhans cell histiocytosis (LCH) is heterogeneous ranging from single-organ involvement to systemic disease causing substantial morbidity and mortality. We describe an unusual course of severe multisystem LCH with spontaneous remission. \textbf{Case Presentation:} We report on a 45-year-old Caucasian woman with cervical cancer, FIGO stage IVB. Five months after the end of combined radiochemotherapy and brachytherapy, the patient was readmitted because of severe dysuria. Sterile leukocyturia
was seen, and cystoscopy revealed only 3 unspecific small mucosal lesions compatible with postradiation cystitis. Incidentally, a computed tomography (CT) scan of the body showed diffuse micronodular and cystic lesions in lungs and hypodense lesions in the liver. Biopsies revealed infiltrations of CD1a and Langerin (CD207)-positive histiocytes in the lung, liver, and bladder. Additionally, positron emission tomography-CT (PET-CT) was compatible with bone involvement. Retrospective analysis revealed that the increase in alkaline phosphatase might have been a surrogate of bone marrow infiltration with osseous activity. Repeated pneumothoraces occurred, and only one course of vinblastine-prednisolone could be applied. Despite ongoing tobacco consumption and without further therapy, PET-CT showed considerable remission 2 months later. However, despite stable remission, documented by serial PET and conventional CT scans, persistent infiltration of the bladder by Langerhans histiocytes could still be demonstrated 17 months later. Unfortunately, cervical cancer recurred and progressed. **Conclusion:** Multisystem LCH may rapidly occur, may be oligosymptomatic and, even in high-risk cases, remission without specific therapy might occur. Whether alkaline phosphatase might be a surrogate to monitor osseous disease activity has to be further explored.

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**Background**

Langerhans cell histiocytosis (LCH), previously known as histiocytosis X, is a rare disease mostly affecting children. For adults, the annual prevalence of LCH can only be estimated and is about 1 case per 560,000 adults [1]. Most of the knowledge about diagnosis and therapy derives from pediatric studies [2, 3].

LCH is characterized by clonal proliferation of pathologic cells with characteristics of histiocytes with varying proportions of macrophages, T-lymphocytes, eosinophils, plasma cells, and multinucleated giant cells accumulating and forming characteristic granuloma-like lesions in various organs [4]. The main diagnostic feature is morphologic identification of the characteristic Langerhans cells. Additionally, positive staining of the lesional cells with CD1a and langerin (CD207) is required for definitive diagnosis. It has been demonstrated that the expression of Langerin strongly correlates with the presence of cytoplasmic Birbeck granules, which can be seen on electron microscopy [2, 4].

The clinical presentation is very heterogeneous ranging from a single organ or system involvement, generally benign, to a life-threatening multisystem disease [4]. Virtually every organ can be involved. Specific data about organ involvement in adults are sparse; the organs most frequently involved in children and adolescents are skeleton (80% of cases), skin (33%), and the pituitary gland (25%). Other organs involved are the liver, spleen, the hematopoietic system and the lungs (15% each), lymph nodes (5–10%), and the central nervous system (2–4%, excluding the pituitary) [5]. Patients with disease limited to one organ, usually bones, skin, lungs or lymph nodes, also called “single-system” or “low-risk disease,” have a good prognosis and usually need minimal or in some cases no treatment. In contrast, multi-system disease with risk organ involvement, defined as infiltration of bone marrow, liver, spleen, and/or central nervous system, is associated with a less favorable prognosis with increased risk of mortality (10–20%) and morbidity [3, 6]. In a mixed cohort of children and adults of the Mayo Clinic, 30% had multisystem and 70% single-system disease. In cases with single-system disease, bones were most often involved (52%), followed by pulmonary (39%), mucocutaneous (6.5%), thalamic (1%), and lymphatic LCH (0.5%). Predicting factors for progressive LCH were multisystem LCH with bone or pituitary-thalamic axis involve-
ment, multisystem LCH in 3 or more organs, single osseous LCH in 3 or more bones, single-system LCH of mucous membranes, young age (<5 years), and hepatosplenomegaly in patients at the age of 3 years or less [7].

The immune mechanisms underlying LCH are still not fully understood [4]. Cancer-associated mutations (BRAF V600E in 50% of cases, MAP2K1 in 30% of cases and MAP3K1 mutations in 10% of cases) were identified [3, 8, 9], and clonality was shown in all forms of LCH, except for smoking-associated adult pulmonary LCH, indicating that LCH might rather be neoplastic with variable behavior than a reactive disorder triggered by infections or malignancies [3, 4].

We describe a case of multisystemic LCH with involvement of the bladder, which is extremely unusual. Despite risk organ involvement, considerable regression occurred and symptoms disappeared without specific therapy.

Case Presentation

A 45-year-old Caucasian woman was diagnosed with stage IVB cervical cancer (Fig. 1). Obstruction of the right kidney was seen, and a ureter catheter was inserted. The cancer was treated with 6 courses of combined radiochemotherapy followed by brachytherapy. The patient’s past medical history did not reveal any previous cancer manifestation. She had been smoking for several years, cumulative 45 pack-years. She abused cocaine and heroin, but definitively stopped drug abuse a long time ago. Regarding previous surgical procedures, the patient had a gastric banding, abdominoplasty, and recurrent abscess excision in the breast due to mastitis non-puerperalis. She reported spontaneous vaginal deliveries when she was 18 and 20 years old.

Side-effects of radiation-chemotherapy were transient local irritations with dysuria and pollakisuria. Additionally, one pyelonephritis episode occurred 3 months after the end of therapy while ureter catheter was in situ. Eight months after cancer diagnosis and 5 months after the end of the radiation-chemotherapy, the patient complained again of heavy dysuria, pollakisuria, and additionally lower-abdominal cramps (Fig. 1). Urinary tract infections were assumed but could not be proven microbiologically. Empiric therapy with different antibiotics was not successful in pain relief. Therefore, the differential diagnosis of overactive bladder syndrome and postradiation irritation was made. Analgesic and spasmolytic therapy with oxycodon, morphine, and trospium chloride were not sufficient to control the complaints, and the patient was admitted for further diagnostic evaluation. Sterile leukocyturia was seen, and cystoscopy revealed 3 unspecific small mucosal lesions and one ulceration compatible with postradiation cystitis. To exclude a relapse of her cervical cancer, computed tomography (CT) of the thorax, abdominal wall, and pelvis was performed. Surprisingly, multiple micronodular and cystic lesions in both lungs and diffuse hypodense lesions in the liver were detected. ELISPOT assay for Mycobacterium tuberculosis, HIV screening test, and ANA/ANCA were negative. Biopsies of the liver and the lung revealed infiltrations of Langerin (CD207) and CD1a-positive histiocytes admixed with eosinophils in both organs, and LCH was diagnosed (Fig. 1 and Fig. 2a, b). Sequencing of BRAF and MAP2K1 revealed wild-type genes. At this time, LCH of the bladder was suspected and also confirmed by biopsy (Fig. 2c, d). On this occasion, the ureter-catheter could be removed. To complete the staging for LCH, a whole-body positron emission tomography-CT (PET-CT) was performed, and the final diagnosis of multisystem LCH involving lungs, liver, bone, and the bladder was established (Fig. 1 and Fig. 3a–d). Interestingly, neither clinical symptoms nor lab values re-
flected the severe involvement of lungs, liver, and bone marrow seen on PET-CT. However, retrospective analysis revealed that an increase of alkaline phosphatase might have been a surrogate of bone marrow infiltration with osseous disease activity (Fig. 1). Lung function performed after diagnosis showed capacity at the lower limit of normal (86% of predicted value), severely impaired diffusion capacity (48% of predicted value) as well as slight hypoxemia (pO$_2$ of 63 mm Hg) despite hyperventilation (pCO$_2$ of 32 mm Hg). However, the patient did not suffer from cough or dyspnea.

The patient was advised to quit tobacco use. However, despite nicotine-replacement therapy and varenicline, the patient was not able to stop smoking. Due to the multisystem presentation with risk organ involvement (lung, liver, and bone), therapy with vinblastine (6 mg/m$^2$ i.v. day 1) and steroids (40 mg/m$^2$ daily) was started (Fig. 1). However, 3 days later the patient suffered from a right-sided pneumothorax. Resection of bullous cysts and talc-pleurodesis were performed. In order not to jeopardize the healing process, vinblastine and steroids were discontinued. During the next weeks, the patient experienced 2 more pneumothoraces (one on the left side and one recurrence on the right side) in need of further interventions with pleurodesis and wedge resection of the bullous cysts (Fig. 1).

Surprisingly, despite therapy discontinuation, the patient became asymptomatic, and two months later a follow-up PET-CT showed remarkable partial remission of LCH (Fig. 1). However, despite stable remission, documented by serial PET-CT and conventional CT scans (Fig. 1 and Fig. 3e–g), persistent infiltration of the bladder by LCH could be demonstrated at the last follow-up 17 months after diagnosis. Unfortunately, cervical cancer recurred, and laparoscopic evaluation showed a noncurative condition with positive pelvic lymph nodes (Fig. 1 and Fig. 4).

**Discussion**

We describe a case of multisystem LCH with risk organ involvement. There are several teaching points that make our case particularly interesting. First, the involvement of the urogenital tract by LCH is a rarity. There are only few reports in the literature, which describe such an involvement. Besides the bladder, infiltration of the kidneys and the testes has been described [10]. In contrast, genital and oral mucosa is the most common mucous infiltration in cases of mucocutaneous LCH [7]. In our case, there was no mucocutaneous manifestation of LCH. Dysuria, pollakisuria and lower abdominal cramps were the leading symptoms, initially misjudged as infection, overactive bladder syndrome, and postradiation cystitis. Second, because the patient was asymptomatic, LCH was missed and only diagnosed by incidental findings on CT scans. In spite of extended lesions in the liver, lungs, and bone marrow, there were no specific symptoms either clinically, or in the laboratory, which could suggest manifestations in these organs. Interestingly, a transient increase in the alkaline phosphatase was retrospectively detected. We assume that this increase was caused by osseous disease activity in the context of bone marrow infiltration. Whether alkaline phosphatase could be a useful marker in patients with osseous LCH has to be further explored. In the literature, around 20% of adults with pulmonary LCH, isolated or as part of multisystem disease, have no symptoms [11]. Our case confirms that even extended multisystemic LCH can evolve subclinically, and symptoms of specific organ involvement can be misleading. Third, it is thought that the beginning of the disease process may precede symptoms by several years [7]. The median delay from first symptoms to diagnosis is around 1 year; however, it is shorter in patients with mucocutaneous, thalamic, and lymphatic manifestations [7].
Further, pneumothoraces caused by pulmonary LCH and liver manifestation are judged to be indicators for advanced and late manifestations of LCH [7]. In our case, 9 months before, a PET-CT scan was performed for cancer staging and revealed no signs of LCH. Therefore, our case demonstrates that even extended multisystem disease may evolve very fast and may occur within few months. Fourth, multisystem LCH can be associated with other malignancies. In a retrospective study of a mixed population of children and adults with biopsy-proven LCH (median follow-up 4 years, range 1–47.5 years), 27 out of 314 (8.5%) had a co-existing neoplasm [7]. LCH may precede, occur currently, or occur after the malignancy [12]. The causality between LCH and the occurrence of a second malignancy has still to be determined on an individual basis. A key part of the classification of LCH as a neoplasm has been the identification of BRAF V600E (B-Raf proto-oncogene, serine/threonine kinase) gene mutations in 35–60% of cases [13]. In our patient, gene sequencing of BRAF and MAP2K1 revealed wild-type genes, rather supporting the hypothesis of an immune-reactive disorder and not a neoplastic disorder. There is some evidence that immature dysregulation leads to an aberrant reaction between Langerhans cells and T lymphocytes. This reaction could be triggered by various stimuli including viruses or malignancies [4]. Our patient developed LCH after the diagnosis of advanced cervical cancer, which was caused by Human Papilloma Virus 16. Additionally, the affection of the bladder by LCH corresponded to the area of radiotherapy. Due to the anatomical proximity of the bladder and radiation field, one may postulate that radiotherapy was a trigger for Langerhans cell activation. Further, the patient is a smoker and smoking is a strong risk factor for pulmonary LCH involvement. Taken together, in our patient there are some potential triggers which may have led to an immune-reactive dysregulation of histiocytes and the evolution of LCH. Fifth, multisystem LCH with risk-organ involvement is associated with a poor prognosis, and experts recommend chemotherapy as treatment [2, 7, 11]. In our patient, the clinical course was astonishing. Shortly after diagnosis, the pulmonary situation worsened and the patient suffered three pneumothoraces. In the following, impressive partial remission occurred, followed by stable disease until the last follow-up 17 months after diagnosis. The role of cigarette smoking in pulmonary LCH appears very convincing, and cessation of smoking is recommended for regression [14, 15]. Surprisingly, the patient remained asymptomatic regarding LCH although smoking cessation could not be achieved, cervical cancer could not be cured and even progressed, and therapy with vinblastine and steroids was discontinued after the first course.

Conclusion

We described a rare case of adult multisystem LCH with unusual presentation and course. In this case, heavy dysuria mimicking urinary tract infection was the first symptom and not dyspnea despite advanced pulmonary LCH. Physicians should be aware of unusual organ manifestations and oligo-symptomatic presentation of LCH. Further, the case demonstrates that even in severe LCH with risk organ involvement partial remission and stabilization may occur without systemic therapy. Whether alkaline phosphate might be a useful marker to monitor osseous disease activity has to be explored in further studies.
Statement of Ethics

Hereby, we confirm that the patient has given her written consent for the case report to be published.

Disclosure Statement

All authors declare that they have no competing interests. The case study was not funded.

Author Contributions

Herein, we declare that all authors have made substantial contributions to the case study. I.M.M. drafted the manuscript, A.T. carried out the histopathology and revised the manuscript for important intellectual content, F.K. revised the oncological data, C.R. carried out the radiological studies, P.G. carried out the pneumological studies, V.H.S. revised the manuscript, M.M. and F.D.S. designed the study, have been involved in drafting the manuscript and in final approval. All authors read and approved the final manuscript.

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Fig. 1. Disease course of Langerhans cell histiocytosis (LCH). Green line: Values of the alkaline phosphatase (green dotted line: missing values); x axis: follow-up in months (dotted line: compressed time axis); PET-CT, positron emission tomography-computed tomography; Dx, diagnosis; Bx, biopsy.
**Fig. 2.** a Langerhans cell histiocytosis infiltrating the lung. HE. ×160. Note histiocytic cells with cleaved nuclei and broad amphophilic cytoplasms intermingled with eosinophilic granulocytes. b Expression of langerin in the pulmonary infiltrates. Immunoperoxidase staining. ×160. c Langerhans cell histiocytosis infiltrating the urinary bladder. HE. ×100. Note the partially desquamated urothelial surface and the eosinophilia. d Expression of CD1a in the bladder infiltrates. Immunoperoxidase staining. ×160. Note the negatively staining urothelium.
Fig. 3. a Intense, diffuse FDG accumulation in the lungs and liver, and increased accumulation in the bone marrow of both femora as a typical finding for Langerhans cell histiocytosis (maximum intensity projection images of 18F-FDG PET/CT). b Hypermetabolic hepatic lesions (fused axial PET/CT slices). c Diffuse hypermetabolism in both lungs (fused axial PET/CT slices). d Multiple small cystic and micronodular lesions in the lungs, ground glass opacities and small pneumothorax on the right-side ventral view. e Regression of the hypermetabolic lesions in the liver (fused axial PET/CT slices). f Regression of the hypermetabolism in the lungs (fused axial PET/CT slices). g Nearly complete regression of the cystic and micronodular lesions, the ground glass opacities and the pneumothorax (axial CT).

Fig. 4. Invasive endocervical squamous cell carcinoma. HE stain. ×360.