Langerhans cell histiocytosis in children – a disease with many faces. Recent advances in pathogenesis, diagnostic examinations and treatment

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Abstract

Langerhans cell histiocytosis is a rare clonal disease characterized by the proliferation of CD1a-positive immature dendritic cells. The purpose of this article was to present an updated review of recent advances in the pathogenesis, clinical features, imaging and treatment of this disease. The discovery of oncogenic BRAF mutations and the presence of proinflammatory cytokines and chemokines confirmed the unusual characteristics of this disease. Currently, children with organ involvement who do not have a good response to chemotherapy and have neurodegeneration or diabetes insipidus are the most problematic patients. Further research is needed to improve the results of treatment.

Key words: Langerhans cell histiocytosis, children, pathogenesis, symptoms, treatment.

Introduction

Langerhans cell histiocytosis (LCH) is a rare clonal disease of the monocyte-macrophage system characterized by uncontrolled proliferation and accumulation of CD1a~/CD207+ dendritic cells (DCs) as a result of continuous immune stimulation [1].

Its clinical presentation is highly variable and may range from isolated, self-healing skin and bone lesions to life-threatening multi-system disease. Reaching an appropriate diagnosis in a short time is a significant challenge. Extremely wide symptomatology is associated with formations in various tissues and organ infiltrations that consist of pathological DCs that are morphologically similar to epidermal Langerhans cells.

The histiocytic disorders are generally defined by their constitutive cell type, on the basis of widely recognized pathologic and histological criteria. In light of recent insights Emile et al. proposed to parsimoniously gather the large number of categories of histiocytic disorders into 5 groups (Table 1) [1]. The nomenclature of histiocytic disorders has changed in the last 50 years. Historically LCH was called “histiocytosis X”, which is a term that includes three entities: eosinophilic granuloma, Hand-Schüller-Christian disease and Letterer-Siwe disease [2–6].

At present, LCH is classified into three distinct forms: single-system single site (SS-s), single-system multi-site (SS-m), and a multisystem type (MS) [7–10]. Approximately 65% of patients have SS disease [11–13]. An epidemiological Japanese study reported that the SS-s, SS-m, and MS forms of LCH are diagnosed at a ratio of almost 1 : 1 : 1 [12]. In SS-s LCH, 70–82% of patients have bone lesions, and only 12% have skin lesions [12, 14]. Multisystem LCH may have two forms, either with or without organ involvement. Low-risk organs include the skin, bones, lymph nodes, and the pituitary gland, whereas high-risk organs include bone marrow, liver, spleen and the lungs [2, 3, 7, 9]. The latest studies have not confirmed that the lungs are risk organs [15, 16].

The involvement of specific organs affects the course, treatment and prognosis of LCH [2, 3, 7, 17].

Epidemiology

Langerhans cell histiocytosis can occur at any age. However, peak occurrence occurs between 1 and 4 years.
of age. The incidence of LCH in children is determined as 2–9/1000 000/year and is slightly predominant in males (M/F = 1.2–1.4) [6, 8, 13, 14, 18].

Aetiology and pathogenesis

The nature of the disease still raises doubts. For many years, debates over models of LCH pathogenesis focused on functional dysregulations versus malignant transformation of epidermal DCs. The detection of some virus DNA sequences in peripheral blood and tissues and the associated Epstein-Barr virus infection with LCH suggested that LCH is reactive [19, 20]. Langerhans cell histiocytosis lesions include infiltrating pathological DCs (8%), macrophages, eosinophils and lymphocytes enriched for regulatory T cells. Dendritic cells express high levels of T-cell costimulatory molecules and proinflammatory cytokines. Cytokines, chemokines, and cytokine/chemokine receptors constitute the “cytokine storm” [5, 21–23]. Recently, it was reported that DCs abundantly express the pleiotropic cytokine osteopontin, which promoted the generation of T helper cells, recruited histiocytes/monocytes and activated osteoclasts, which in turn produced large amounts of tissue-destructive enzymes [17, 22]. Murakami et al. found that LCH patients had high serum levels of IL-17A and that IL-17A receptor expression levels helped to distinguish between LCH forms [24]. Biopsies of lesions from both SS and MS disease were found to have proliferation of DCs from a single clone [25, 26]. Despite clonality, pathological DCs do not appear dysplastic, mitoses are rarely observed, and chromosomes are typically intact [23, 27]. Clonal proliferation leading to the formation of the infiltrates and the discovery of the mutually exclusive activating somatic BRAF V600E (57%) and MAP2K1 gene mutations that occur in approximately 75% of patients suggest there is a neoplastic process [5, 28, 29]. The BRAF V600E mutations lead to constitutive activation of the downstream kinases MEK and ERK [27, 30]. The presence of circulating cells carrying BRAF V600E mutations is associated with high-risk clinical status (multisystem disease and liver, bone marrow, spleen involvement) and with a two-fold increase in the risk of recurrence [31]. Haematopoietic cell progenitors with the BRAF V600E mutation were also identified in patients with high-risk LCH. The new evidence supported a model in which LCH occurs as a consequence of a misguided differentiation programme of myeloid DC precursors. Genetic, molecular and functional data implicated activation of the ERK signalling pathway at critical stages in myeloid differentiation as an essential and universal driver of LCH pathology. The low-risk disease arises from ERK activation in tissue-restricted precursors [27, 31]. The frequency of BRAF V600E mutations in LCH depends on ethnicity [29].

Table 1. The revised classification system of histiocytoses and neoplasms of macrophage-dendritic cell lineages [1]

<table>
<thead>
<tr>
<th>No.</th>
<th>Group of histiocytic disorders</th>
<th>Histiocytosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>L group</td>
<td>LCH</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– ECD</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– Mixed LCH/ECD</td>
</tr>
<tr>
<td>2</td>
<td>C group</td>
<td>Cutaneous non-LCH:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– XG family:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– AXG, AXG, SRH, BCH, GEH, PNH</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– Non-XG family:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– cutaneous RDD, NKG, other not otherwise specified</td>
</tr>
<tr>
<td>3</td>
<td>M group</td>
<td>Primary MH</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Secondary MH (following or associated with another hematologic neoplasm)</td>
</tr>
<tr>
<td>4</td>
<td>R group</td>
<td>Familial RDD</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Sporadic RDD:</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– Classical (nodal)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– Extra-nodal</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– RDD with neoplasia or immune disease</td>
</tr>
<tr>
<td></td>
<td></td>
<td>– Other, non-C, non-L, non-M, non-H histiocytoses</td>
</tr>
<tr>
<td>5</td>
<td>H group</td>
<td>Primary HLH: Mendelian inherited conditions leading to HLH</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Secondary HLH (non-Mendelian HLH)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>HLH of unknown/uncertain origin</td>
</tr>
</tbody>
</table>


Diagnosis

The diagnosis of LCH is based on histological criteria established by the Histiocyte Society in 1987 [7, 21]. To make a diagnosis, it is necessary to perform a histological examination with immunophenotyping. The identification of characteristic cells and the presence of markers, such as CD1a and/or CD207 (Langerin), allows for a definitive diagnosis of LCH. The Birbeck granule is a LCH-specific tennis racquet-shaped organelle detected by electron microscopy. The presence of Birbeck granules, once considered the “gold standard” in the diagnostics, currently does not have to be assessed. They have been replaced by langerin, which is a cell-surface receptor that induces formation of Birbeck granules [17, 32].

Clinical presentation of LCH

The disease has different clinical courses from a mild form, which is limited to one organ, to a severe form with...
the involvement of many organs and systems that progresses very quickly [2, 8, 17]. The clinical presentation of LCH includes a wide range of symptoms, from symptoms of severe life-threatening systemic diseases, such as septic fever with accompanying haemorrhagic rash and hepatosplenomegaly, to the presence of lumps under the skin with little specificity or a single asymptomatic bone focus [2, 6, 10, 17, 33].

For this reason, LCH presents many diagnostic and therapeutic difficulties. Langerhans cell histiocytosis lesions may spontaneously regress or repeatedly reactivate, which contributes to long-term disabilities, such as diabetes insipidus (DI) and neurodegenerative disease [10, 17, 18, 33, 34]. Approximately 30–40% of patients develop lasting negative consequences [35–38].

Langerhans cell histiocytosis can be completely asymptomatic. The most frequent presenting signs and symptoms of LCH include painful bone lesions and rash. Often non-specific symptoms become prominent, such as fever, poor appetite, weight loss, fatigue, irritability and changes in behaviour [17, 18]. Finally, more characteristic symptoms associated with the involvement of bones, skin, the pituitary gland, liver, spleen, the haematopoietic system, lungs, lymph nodes, the central nervous system (CNS), thymus, and the gastrointestinal tract, may appear (Table 2) [2, 4, 5, 9, 17, 44].

In the youngest children, the disease is often an MS disease with fever and symptoms of failure in various organs [2, 10, 14]. In a Japanese study, the patients with skin lesions were younger than those with bone lesions [12]. In a previous French study, 77% of the patients with the skin lesion were less than 1 year old [14].

The prognosis of LCH varies depending on its form. The prognosis of patients with SS-s LCH is generally good [12]. However, the patients with the involvement of the craniofacial bone(s) (orbital, temporal, mastoid, sphenoid, zygomatic, ethmoidal bones, the maxilla, paranasal sinuses, or anterior or middle cranial fossa) with intracranial soft tissue extension (the so-called CNS-risk lesion(s)) had a higher risk for developing DI [39]. The LCH-III protocol study conducted by the Histiocyte Society suggested that chemotherapy should be offered to these patients, even if there was only a single lesion [40]. At present, the factors for poor prognosis are age under 2 years old, MS disease, high-risk organ involvement and dysfunction or a poor response to initial therapy [2, 3, 7, 41, 42]. It is believed that hepatic involvement at the time of the diagnosis is particularly associated with a poor prognosis [6].

### Bones

Bones are involved in approximately 80% of LCH patients [2, 14, 33, 43]. Unifocal involvement is more common than multifocal involvement [43]. Langerhans cell histiocytosis can involve any bone, but there is a predilection for the axial skeleton, with more than 50% of bone lesions occurring in the flat bones (skull, ribs, pelvis). In the long bones, the femur is the most commonly affected, followed by the humerus, tibia and vertebrae (“vertebra plana”) [17, 44]. The symptoms include pain, swelling of the affected area and pathological fractures. The location of lesions in the vertebral column may cause compression fractures and scoliosis [10, 17, 43]. There may be visual loss or exophthalmos due to orbital involve-

<table>
<thead>
<tr>
<th>No.</th>
<th>System involved</th>
<th>Organ involved</th>
<th>Symptomatology</th>
<th>Incidence</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Skeletal</td>
<td>Flat and long bones, spine/skullcap bones, femurs, humeri, spine</td>
<td>Bone pains, lumps</td>
<td>80%</td>
</tr>
<tr>
<td></td>
<td>Orbital cavity</td>
<td>Exophthalmos</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>Temporal bone</td>
<td>Discharge from the middle ear, hearing loss</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>Integumentary</td>
<td>Skin</td>
<td>Seborrhoeic erythematous rash, haemorrhagic rash</td>
<td>60%</td>
</tr>
<tr>
<td>3</td>
<td>Lymphatic</td>
<td>Lymph nodes</td>
<td>Swollen lymph nodes</td>
<td>33%</td>
</tr>
<tr>
<td></td>
<td>Thymus</td>
<td>Widened opacity of the mediastinum</td>
<td>2.6%</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Spleen</td>
<td>Splenomegaly, cytopenias</td>
<td>15%</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Liver</td>
<td>Liver</td>
<td>Hepatomegaly, liver dysfunction</td>
<td>15%</td>
</tr>
<tr>
<td>5</td>
<td>Respiratory</td>
<td>Lungs</td>
<td>Respiratory failure</td>
<td>1–15%</td>
</tr>
<tr>
<td>6</td>
<td>Bone marrow cavity</td>
<td>Bone marrow</td>
<td>Neutropenia, anaemia, thrombocytopenia</td>
<td>15–30%</td>
</tr>
<tr>
<td>7</td>
<td>CNS</td>
<td>Hypothalamic-pituitary disease</td>
<td>Diabetes insipidus, short stature, secondary hydrocephalus, cranial nerve palsies</td>
<td>25%</td>
</tr>
<tr>
<td></td>
<td>CNS</td>
<td>Neurodegenerative disease of the CNS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>Digestive</td>
<td>Gastrointestinal tract</td>
<td>Haemorrhagic diarrhoea, anaemia</td>
<td>&lt; 5%</td>
</tr>
</tbody>
</table>

Table 2. Incidence and symptomatology of systems and organs involvement in Langerhans cell histiocytosis in children [1, 2, 4, 5, 9]
The involvement of the temporal bone is manifested by symptoms imitating the inflammation of its structures – the middle ear and the mastoid process with the typical recurrent ear discharge, secondary hearing deterioration and even deafness [17, 43]. Histiocytic infiltrates in the upper and lower jaws cause swelling and looseness of gums and teeth ("loose teeth syndrome") [43, 44]. The typical radiographic findings are a solitary lytic lesion of the skull and a "vertebra plana" [17, 43, 44].

### Skin

Skin lesions are the second most-common clinical manifestation of LCH (30–60%) [17, 33], which is particularly frequent in infants [17] and the most common presentation of "non-self-regressive" cutaneous LCH in neonates and young infants [45]. However, an isolated cutaneous form of the disease is rare (the number of described cases does not exceed 12% of all patients with LCH), typically concerns only male infants and has a good prognosis [12, 46]. However, both patients born with skin-only LCH and patients who develop skin-only LCH during infancy can have a "self-healing" disease with spontaneous regression or may progress to MS-LCH with a worse prognosis [47]. The special form of congenital cutaneous LCH in neonates is uncommon Hashimoto-Pritzker disease characterized by self-regressive red-brown cutaneous nodules [45, 47]. The skin disease in patients older than 18 months of age at diagnosis is associated with the presence of MS-LCH [48]. The incidence of cutaneous involvement is significantly higher in children with MS disease (53%) [48]. Skin lesions may take different forms and different degrees of severity. The eruption may be extensive and involve the scalp, face, trunk, buttocks and intertriginous area. The trunk is the most affected site [48]. A wide spectrum of cutaneous presentations varying from crusted or scaly nodules and papules, blisters, vascular tumour-like lesions, and scaling orange to red or purpuric macules (frequently in seborrhoeic region) was documented by Morren et al. [11]. Crusted or scaly papules and papulovesicles are the most common presentation of skin LCH [11]. In early childhood, they often take the form of bumps and blisters. On the other hand, a maculopapular desquamative rash, which is located mainly within the scalp, retroauricular, inguinal or axillary area, can also appear in older children. Among other lesions occurring in the course of the disease, nodules and petechiae should be listed [46]. Sometimes there are superficial ulcerations of the lesions accompanied by oozing as the result of secondary processes leading to a bacterial superinfection. Ulcerative lesions behind the ears or involving the scalp, axillae, genitalia, or perianal region are often misdiagnosed as bacterial or fungal infections. Rare lesions include a haemangioma-like lesion, a varicella-like eruption, or purpura. "Blueberry muffin baby syndrome" has also been described as a presentation of LCH in neonates [49]. A seborrhoea-like eruption on the scalp and lesions in the large body folds were also more frequent in the group with MS-LCH, with these lesions being more erosive and ulcerative compared to the skin-only group [11]. Late onset and a protracted course of skin lesions were associated with MS LCH, whereas wild-type and mutated BRAF were found in rapidly resolving skin lesions. The involvement of the hands and feet suggests there is a self-regressive form, and necrosis might also be associated with a good prognosis [45]. Moreover, there are reports of changes in nails in LCH, but it is unclear whether the change predicts a poor prognosis, although lung involvement has been described in approximately 50% of such patients [50]. Nail involvement can occur in the form of longitudinal grooving, onycholysis, subungual hyperkeratosis, subungal pustules, purpuric striae of the nail bed, V-shaped notches, pterygium formation, and end onycholysis (oval depression in the nail plate). Paronychial erythema and swelling as well as nail-fold destruction have also been reported. The histology of nail lesions is similar to cutaneous forms (infiltrations of typical Langerhans cells) [50].

### Lymph nodes

The cervical nodes are most frequently involved and may be soft or hard-matted groups with accompanying lymphoedema [10, 51].

### Oral mucosa

The symptoms of oral LCH include gingival hypertrophy and ulcers of the soft and hard palate, buccal mucosa, or on the tongue and lips. Early eruption of teeth occurs with LCH infiltration of the mandible or maxilla [52, 53].

### Central nervous system

The involvement of the CNS is one of the rarer forms of LCH (24% of all LCH children) [54]. Central nervous system lesions based on MRI include active lesions involving hypothalamic-pituitary disease, space-occupying lesions at other sites (the meninges, the choroid plexus, the pineal gland, the cerebellar parenchyma), and neurodegenerative lesions of the cerebellum, pons, brainstem, basal ganglia, grey or white matter of the brain [55–57]. The most common MRI changes include solid enlargement of the pineal gland, a thickened enhancing stalk and signal changes of the dentate nuclei [58]. The involvement of the hypothalamus and the pituitary gland, which are manifested by DI, is the most common known form of CNS LCH (15–30% of LCH patients) [37, 39, 59]. Fifty-six percent of LCH patients with DI will develop anterior pituitary hormone deficiencies within 10 years of DI onset [60]. These patients may present growth deficiency (which may affect up to 10% of LCH patients), precocious or delayed puberty, galactorrhea and symptoms of hypothyroidism, and in the case of hypothalamus inflitrat-
tions (rare), there are typically changes in behaviour, eating, sleep, or thermoregulation disorders. Lesions in the pituitary and hypothalamus may be accompanied by foci of histiocytic infiltrates associated with the meninges and choroid plexuses. The patient may then complain of headaches, vomiting, or blurred vision, or display convulsions and other focal symptoms. DI occurs more often in patients with craniofacial bone, ear, eye and/or oral lesions, known CNS-risk lesions [39, 59, 61].

The syndrome of neurodegenerative central nervous system Langerhans cell histiocytosis (ND-CNS-LCH) is less known, develops in approximately 1–3% of LCH patients and remains a progressive and devastating complication. It is initially manifested by subtle neurological deficits that can enhance and develop years after the first LCH symptoms. The deficits include ataxia, dysarthria, tremor, vertigo, nystagmus, psychomotor development delay, behavioural disorders, and neuropsychological deficits. Patients at high risk for developing ND-CNS-LCH are usually children with MS-LCH, or DI, or have involvement of the temporal, petrous, orbital, and zygomatic bones [57, 59, 62]. The studies by Grois et al. showed that 75% of patients with DI had changes that were visible in the MRI in the bones of the facial skeleton, and as many as 76% of these patients in their check-up MRI examination conducted 5 or more years after a diagnosis of diabetes insipidus showed parenchymal, neurodegenerative changes in their encephalon [61]. The risk of LCH within the CNS in these cases is 10 times higher than in patients with SS-LCH. ND-CNS-LCH occurs more frequently in young patients who present with skull/facial lesions at LCH onset [63, 64].

The pathogenesis of these disorders is not fully explained and probably involves intracranial immune interactions and inflammatory mechanisms between LCH cells/T cells and glial cells around microvessels in the CNS. Microglia activation, gliosis, and neuronal and axonal destruction with secondary demyelination are typical [57, 63]. ND-CNS-LCH may be a consequence of undetectable active disease or paraneoplastic syndrome with DC activation of the immune system to recognize CNS antigens [62]. In recent years, MRI tests revealed the presence of pathological changes in the structures of the cerebellum – focal lesions, which are mostly periventricular in location, and hypodense foci consisting of demyelination changes and gliosis of the white matter [61]. The cerebellum is the most frequent and earliest site of involvement, and it is sometimes associated with the brainstem, basal ganglia and hemispheric white matter. MR spectroscopy showed a reduced cerebellar NAA/Cr ratio and may complement structural MRI [58, 65, 66]. Histopathology from cerebellar biopsies and autopsies revealed neuronal loss, axonal degeneration and profound T-cell inflammation without the infiltrates of cells characteristic for LCH [59, 63]. There is no correlation between the degree of neurological impairment and the topography, extent and grading of neurodegenerative lesions in most patients [65].

Both Imashuku and Krenova and Sterba underscore the necessity of cerebrospinal fluid examinations and neuropsychological/neurological/psychological assessments at the time of diagnosis in ND-CNS-LCH risk patients [67, 68].

**Liver**

The involvement of the liver in the course of LCH may be a primary or secondary process to, for example, the involvement of hepatic hilar lymph nodes. A typical symptom is hepatomegaly (in the diagnostic criteria liver > 3 cm below the costal margin in the midclavicular line in the USG). It may be accompanied by organ dysfunction, an increase of transaminases, hypoalbuminemia (< 30 g/dl) and secondary oedema or ascites, jaundice (bilirubin > 3 × N) or clotting factor deficiencies [7, 9]. Ultrasonography (USG), computed tomography (CT) and magnetic resonance (MR) images of the liver typically show hypoechoic or low signal intensity along the portal veins or biliary tracts [69]. Langerhans cell histiocytosis may also be located within the bile duct, which may cause cholestasis, sclerosing cholangitis (15–25% of LCH children), biliary cirrhosis and organ damage that require transplantation [70, 71]. Biopsies typically do not include histiocytes but demonstrate periportal lymphocytic infiltrates, bile duct inflammation, and loss of bile ducts. Seventy-five percent of children with sclerosing cholangitis will not respond to chemotherapy, and all of these patients require liver transplantation. The coexistence of liver involvement and sclerosing cholangitis or liver cirrhosis may hinder the diagnosis on account of the regression of Langerhans cells [71]. Liver involvement drastically changes a patient’s prognosis and treatment [70]. In this group of patients, the 5-year overall survival rate is only 25% [13].

**Spleen**

Spleen involvement may be the consequence of the primary or secondary process (portal hypertension as a result of periportal fibrosis). The diagnostic criterion for splenomegaly is the spleen image in the ultrasound (> 2 cm below the costal margin in the midclavicular line). Splenomegaly may be accompanied by hypersplenism [5, 7, 9].

**Mediastinum**

Mediastinal involvement is very rare in LCH (2.6% of patients). Dendritic cells infiltrate mainly the thymus and lymph nodes [72]. This location concerns primarily younger children (< 1 year old). With age, its incidence decreases, although there are reports of diagnoses made in people > 18 years old. This location predisposes to the occurrence of life-threatening conditions for the patient. The symptoms of mediastinal involvement include cough, tachypnoea, respiratory failure or superior vena cava compression.
cava syndrome. An enlarged thymus or involvement of the mediastinal lymph nodes can mimic lymphoma or an infection. Patients with LCH in the thymus and mediastinal lymph nodes usually already display MS-LCH at the time of diagnosis, and nearly half of them have one of the so-called ‘risk organs’ affected. More frequently, they have skin lesions, and the foci are rarely in the bones. The group of patients with thymus involvement and mediastinal lymph nodes is considered to be particularly vulnerable to developing serious infections during treatment of the underlying disease, which is explained by the involvement of the thymus itself as well as a younger age at the time of diagnosis [72, 73].

**Bone marrow**

The occurrence of histiocytic infiltrates within the bone marrow causes haematopoietic failure that clinically manifests as anaemia, leukopenia, or thrombocytopenia. Bone marrow involvement occurs most frequently in young children who have diffuse disease involving the liver, spleen, lymph nodes, and skin, and it may be associated with marked thrombocytopenia or neutropenia. Other occurrences have only mild cytopenias and were found to have marrow involvement with LCH through sensitive immunohistochemical or flow cytometric analysis of the bone marrow [74]. Haematopoietic system involvement is a well-known prognostic factor in LCH. The study by Raney et al. revealed that among haematological dysfunction parameters, thrombocytopenia had the most significant effect on survival [75]. Langerhans cell histiocytosis patients with very high-risk disease may present haemophagocytosis that mimics haemophagocytic lymphohistiocytosis [76].

**Lungs**

Involvement of the lungs is the most characteristic for adults, in whom smoking is a key aetioloical factor [77, 78]. In the paediatric population, the isolated form of the disease affects less than 1% of LCH children [15, 16, 79]. More often, the histiocytic lesions observed in the lungs are in the form of MS-LCH (up to 25%) [16, 80]. The most common symptoms include non-productive cough and shortness of breath. Less frequent symptoms include pain in the chest, general malaise and fatigue, weight loss, fever, and cyanosis. Extremely rare haemoptysis can be observed in children [15]. In 10% of children over the course of lung LCH, a pneumothorax typically develops, which is most frequently a unilateral one (although cases of bilateral lesions are also possible) and is often recurrent (on the same side or the opposite). Chest X-rays may show non-specific interstitial infiltrates with peribronchial thickening (100%), cystic changes (39%), and pneumothorax (11%) [16]. A high-resolution CT of the chest is needed to visualize the cystic/nodular pattern of LCH that leads to the destruction of lung tissue. Declining diffusion capacity may also herald the onset of pulmonary hypertension. In adults, as a result of a long-term disease process, pulmonary fibrosis or pulmonary hypertension is observed, and in extreme cases it requires transplantation of the lungs or both the lungs and heart [15].

Howarth et al. reported 85% disease-free survival in children and adults with isolated lung involvement [78]. Currently, isolated pulmonary involvement is not considered to be an independent prognostic factor [15, 16].

**Gastrointestinal tract**

Gastrointestinal (GI) involvement in children with LCH is rare and has been associated with systemic illness and poor prognosis [81]. The peak incidence occurs in children <2 years old with a 2-fold male predominance [82]. The reported symptoms include vomiting, abdominal pain, intractable diarrhoea, a child's failure to gain weight, malabsorption, bloody stool, protein-losing enteropathy, and even intestinal perforation [83, 84]. Through endoscopy, superficial erosions and haemorrhagic ulcerations are typically found throughout the GI tract [82]. A long-term disease process can lead to the development of fully symptomatic malabsorption or protein-losing enteropathy and complications such as intestinal perforation in extreme cases [81].

**Other lesions**

In addition to these locations of the disease, there have been unusual reports in the literature of thyroid, pancreas, kidney and salivary gland involvement [85–87].

**Recent advances in imaging evaluation**

Bone foci in LCH are lytic. X-rays typically demonstrate osteolytic “punched out” lesions with sharp margins. Radiography, which remains the gold standard for diagnostic and staging procedures, may be complemented by CT and/or MRI to delineate the extent of osseous destruction or the extent of bone marrow and soft tissue involvement [43]. Magnetic resonance imaging is necessary for the assessment of CNS-risk lesions and ND-CNS-LCH [54, 58, 61, 88].

Recently, new imaging techniques, such as positron emission tomography-computed tomography (PET-CT) and whole body MRI, have been developed, allowing improved assessment of the extent and severity of the disease [89, 90]. Positron emission ET-CT is useful for detecting and assessing LCH lesion activity and for evaluating the response to treatment [91]. There are relatively few papers regarding FDG-PET in LCH compared to other malignancies. There are mostly case and multi-case reports or reviews but not prospective, systematic evaluations. This lack of research is due to the low incidence of the disease as well as a wide range of involvement and prognosis. However, FDG-PET has been studied for usefulness in staging, therapy response assessment and diagnosis of disease recurrence.
on non-avid active LCH, the vast majority of lesions were found to have a significant increase in radiolabelled glucose uptake. The advantage of FDG-PET is that it is performed as a whole body scan with high sensitivity for lesion detection, which allows for more precise staging regardless of the location of disease active foci. With similar high diagnostic confidence, FDG-PET allows for evaluation of therapy responses. The ability to detect metabolically active LCH is crucial, especially for evaluating disease in bones where both bone scanning and conventional CT may show abnormalities due to bone remodelling in the lack of active disease. This approach helps to make management decisions in patients with LCH. Moreover, the level of glycolytic activity reflected in the tracer concentration (SUV measurement) may be considered an important factor in prognosis estimation. However, for sensitivity, it must be noted that the method is not specific and that there is no pathognomonic pattern of tracer uptake to indicate LCH, so a differential diagnosis must be considered when reading scans. Nevertheless, there is a large amount of evidence of the superiority of FDG-PET compared to conventional imaging in the LCH evaluation. With the increased availability of the method as well as the decreasing cost of an examination, the cost-benefit ratio and cost effectiveness of FDG-PET are rising. The recent introduction of PET-MR systems also resulted in reduced radiation exposure, which is especially important for paediatric patients and patients requiring multiple scans, which can be common in cases of LCH [92].

The most important imaging for LCH patient assessment is ultrasonography. The use of a new imaging technique called static elastography for assessment of changes in the skull and other bones allows for more complete differential diagnoses because the typical images reveal hyperplastic (hard changes) and fluid changes (rainbow artefact or the artefact buffalo eye) in unicameral bone cysts (Figures 1, 2) [93].

Treatment

Treatment depends on the form of LCH. Systemic therapy is indicated for all patients with SS-m and MS-LCH as well as for special localizations. A standard two-drug regimen with vinblastine and prednisone consisting of an initial intensive phase for 6–12 weeks, followed by maintenance therapy for a total treatment duration of at least 12 months, is recommended [7]. Non-responders, particularly those with progressive disease in a high-risk organ, are eligible for salvage therapy with cladribine, cytarabine, clofarabine or combinations of those medications as well as bone marrow and/or solid organ transplantation [3, 17, 48, 94].

Patients with SS-LCH limited to the skin may be observed. If the skin lesions do not spontaneously resolve, a variety of methods may be used, including topical steroids, oral methotrexate or thalidomide, topical nitrogen mustard, or psoralsens with UV light [40]. A single bone lesion involving the frontal, parietal, occipital bones and any other bones may be treated by curettage only or curettage with a local injection of methylprednisolone [95]. Other skull lesions (mastoid, temporal or orbital) should be treated with chemotherapy to reduce the risk of DI and ND-CNS-LCH. Vertebral or femoral bone lesions at risk for collapse require isolated radiotherapy. Treatment of CNS lesions (enlargement of the hypothalamus and the pituitary gland or grey and white matter, the choroid plexus, and leptomeningeal lesions) includes systemic chemotherapy [2, 40].

There is no established optimal therapy for patients who develop ND-CNS-LCH. The ND-CNS-LCH is commonly treated with intravenous immunoglobulin alone or in combination with chemotherapy regimens [40, 62, 64, 96]. Moreover, retinoic acid [97], thalidomide and its analogues [98–100], as well as high doses of steroids [55, 100], have been proposed for the treatment of ND-CNS-LCH. Cytotoxic-immunosuppressive approaches have included cladribine [101, 102] and cytarabine because of good CNS penetration [9, 17, 56].

The possible therapeutic use of target therapy, at least in the subset of patients with BRAF V600E mutation (e.g., vemurafenib), should be confirmed in the near future [103, 104]. In addition, recent prophylactic measures for preventing the development of ND-CNS-LCH were discussed by Krenova and Sterba and Imashuku et al. [67, 68]. The remaining questions include which LCH patients and what drugs and methods (immunoglobulin or leukapheresis or appropriate intrathecal/systemic chemotherapy or a molecular targeting agent) should be used.

The prognosis for LCH varies. It depends on the form of the disease (SS-LCH vs. MS-LCH) as well as its location and response to chemotherapy. In unifocal LCH involving a bone or an isolated skin lesion, the prognosis is good. There have been cases of spontaneous remission or symptoms subsiding after local treatment. In forms with multifocal bone involvement, relapses of the disease occur more often. In cases of extensive skin lesions, the likelihood of internal organ involvement increases even several years after the completion of the first-line treatment of the disease. The prognosis worsens significantly with the involvement of “risk organs” and the GI tract. Spontaneous regression of the disease located in those areas is rare. Recurrence is one of the most important problems in the management of LCH. Yagci et al. observed recurrence in nearly one-third of patients. New bone lesions were the most frequent finding at the time of recurrence [105]. Jubran et al. reported a seven-times higher risk of reactivation for patients with multiple bone lesions compared to patients who have single bone involvement [106].
Survivors may experience long-term permanent consequences that include endocrine disorders (DI and growth hormone deficiency), orthopaedic problems, decreases or loss in hearing and sight, the loss of teeth, neurological defects (which affect the CNS in up to 30% of cases), and impaired lung and liver function [2, 17, 35, 37, 38]. Among the most potentially devastating complications is ND-CNS-LCH. Patients with LCH have a higher risk for developing secondary cancers (leukaemia and non-Hodgkin lymphoma) [2].

Thus, the broad spectrum of clinical symptoms and rarity of LCH make the diagnosis difficult. The unclear nature, the
heterogeneous course of the disease from an asymptomatic form to those with various clinical symptoms, the diverse prognosis and the risk of long-term complications cause LCH to remain a major challenge for medicine. However, many years of experience indicate that both the diagnosis and treatment of this disease are becoming more successful.

**Conflict of interest**

The authors declare no conflict of interest.

**References**


55. Imashuku S. High dose immunoglobulin (IVIG) may reduce the incidence of Langerhans cell histiocytosis (LCH)-associated central nervous system involvement. CNS Neurol Disord Drug Targets 2009; 8: 380-6.


45. Imashuku S. High dose immunoglobulin (IVIG) may reduce the incidence of Langerhans cell histiocytosis (LCH)-associated central nervous system involvement. CNS Neurol Disord Drug Targets 2009; 8: 380-6.


