Cervical lymph node diseases in children

Abstract

The lymph nodes are an essential part of the body’s immune system and as such are affected in many infectious, autoimmune, metabolic and malignant diseases. The cervical lymph nodes are particularly important because they are the first drainage stations for key points of contact with the outside world (mouth/throat/nose/eyes/ears/respiratory system) – a critical aspect especially among children – and can represent an early clinical sign in their exposed position on a child’s slim neck.

Involvement of the lymph nodes in multiple conditions is accompanied by a correspondingly large number of available diagnostic procedures. In the interests of time, patient wellbeing and cost, a careful choice of these must be made to permit appropriate treatment. The basis of diagnostic decisions is a detailed anamnesis and clinical examination. Sonography also plays an important role in differential diagnosis of lymph node swelling in children and is useful in answering one of the critical diagnostic questions: is there a suspicion of malignancy? If so, full dissection of the most conspicuous lymph node may be necessary to obtain histological confirmation.

Diagnosis and treatment of childhood cervical lymph node disorders present the attending pediatric and ENT physicians with some particular challenges. The spectrum of differential diagnoses and the varying degrees of clinical relevance – from banal infections to malignant diseases – demand a clear and considered approach to the child’s individual clinical presentation. Such an approach is described in the following paper.
1 Introduction

Neck masses are a symptom ENT physicians in hospitals and surgeries are frequently confronted with among children. If the mass originates from the lymph nodes, the first step should be to establish whether the lymph node itself is enlarged: this is the case above a diameter of >1 cm (in the angle of the mandible >1.5 cm) and is defined as lymphadenopathy. A distinction is made between an acute (<2 weeks), subacute (2–6 weeks) and chronic (>6 weeks) course of the lymphadenopathy [1]. Enlarged lymph nodes must also be differentiated from other possible causes of neck masses, such as midline thyroglossal cysts and branchial cysts, lipoma, vascular malformations (e.g. hemangioma), paraganglioma/neurinoma, lesions of the salivary or thyroid glands, lymphangioma, teratoma, (epi-)dermoid cysts, and ectopic thyroid tissue [2] (see Figure 1, Figure 2, Figure 3, Figure 4).

As Table 1 shows, there are many possible causes of lymph node enlargement. For this reason, anamnesis and thorough clinical examination are crucial steps towards securing a diagnosis [3].

2 Anamnesis

The more varied the possible differential diagnoses are, the greater the importance of the primary diagnostic instruments, namely anamnesis and clinical examination, available to the physician. The course of the illness is one of the first aspects to be considered in diagnosis: the majority of cases of acute lymphadenitis, which subside after 2 weeks, are of infectious origin; this contrasts with chronic lymph node enlargement, which is more likely to originate from neoplastic or metabolic disorders or opportunistic infections. Subacute lymph node enlargement has various possible causes, and additional criteria therefore need to be drawn on before determining the subsequent diagnostic and therapeutic steps (Figure 5).
| Infectious diseases: | Immunologic diseases: | Metabolic diseases: | Neoplastic diseases: |
|----------------------|----------------------|-------------------|---------------------|
| **Viral:** | Granulomatous diseases: | Storage diseases: | **Malignant lymphomas:** |
| EBV, CMV, rubella, measles, varicella, HIV, adenovirus, HSV II, rhinovirus, enterovirus, parvovirus B19 | - Sarcoidosis | - Niemann-Pick disease | - Non-Hodgkin lymphoma |
| | - Common variable immunodeficiency (CVID) | - Gaucher’s disease | - Hodgkin’s lymphoma |
| | - Hyper-IgM syndrome | - Tangier disease | **Leukemias:** |
| | - Chronic granulomatous disease | - Amyloidosis | - Acute lymphoblastic leukemia |
| **Bacterial:** | Rheumatoid disorders: | Hypersensitivity: | - Acute myeloid leukemia |
| Staphylococci, streptococci, tuberculosis, non-tuberculous mycobacteria (NTM), brucellosis, Bartonella henselae, tularemia, Mycoplasma pneumoniae, Yersinia pestis, Pasteurella multocida, cervical actinomycosis | - Arthritis | - Serum sickness | **Lymph node metastasis from solid tumors:** |
| **Protozoan/parasitic:** | Systemic lupus erythematosus | - Adverse drug reaction (e.g.: antiepileptic drugs; heparin; antituberculosis drugs such as isoniazid; antibiotics such as cephalosporins, penicillins; phenytoin, hydralazine, procainamide, allopurinol, dapsone, carbamazepine, atenolol, captopril, gold, primidones, pyrimethamines, quinidines, sulfonamide, sulindac) | - Rhabdomyosarcoma |
| Toxoplasmosis, trypanosomes, toxocariasis, leishmaniasis, microfilaria | - Dermatomyositis | **Neuroblastoma** | **Nasopharyngeal carcinoma** |
| **Fungal:** | Lymphoproliferative and histiocytic disorders: | |
| Dermatophytes (tinea), coccidioidomycosis, histoplasmosis, blastomycosis | - Sinus histiocytosis with massive lymphadenopathy (Rosai-Dorfman disease) | |
| | - Castleman’s disease | |
| | - Autoimmune lymphoproliferative syndrome (ALPS) | |
| | - Langerhans cell histiocytosis | |
| | - Hemophagocytic lymphohistiocytosis (HLH) | |
| | - Kawasaki syndrome | |
| | - Kikuchi-Fujimoto disease | |
| | - PFAPA syndrome | |
The anamnesis must include any reference to a focus, including in the past, such as a sore throat, earache or toothache, insect bites or injuries. Local tenderness suggests an inflammatory component, while fever can occur equally in infections and as part of the B symptoms. Other B symptoms include night sweats (drenching) and weight loss (>10% of body weight in a period of 6 months). Positive B symptoms typically occur in malignant lymphomas (classification as A/B symptoms according to Ann Arbor), but may also accompany infectious diseases (tuberculosis, HIV, parasitosis). The same applies to symptoms such as bone/joint pain, weakness and lowered resilience or anorexia.

Details of foreign travel and possible contact with animals (in particular cats, rabbits, rodents, tick bites) and equally of all vaccinations must also be obtained in the anamnesis. Other environmental influences and noxa (including drugs) may more rarely be the cause of lymphadenopathy, but they should nevertheless always be considered. A family anamnesis not only includes acute infections but also chronic and systemic diseases such as sarcoidosis or tuberculosis.

3 Clinical examination

Clinical examination of the young patients initially comprises inspection, palpation, ENT examination and more extensive examinations as indicated, including palpation of other lymph node regions (axillary, inguinal etc.) or of the liver and spleen.

Inspection

Inspection must never focus solely on the region of the lymph node itself but also extend to the drainage regions and in particular local portals of entry (tonsils, scratches or skin lesions e.g. associated with an allergic exanthema).

Palpation

Palpation is similarly not restricted to the conspicuous lymph nodes and includes thorough examination of the head and neck region. It is advisable to work in a fixed order. The lymph nodes and diseases associated with them are listed in Table 2. One after the other, the submental and submandibular lymph nodes, the lymph nodes at the angle of the mandible, the pre- and post-auriculare lymph nodes and the (sub-)occipital lymph nodes are palpated. These are followed by the group of vertical lymph nodes of the neck along the jugular vein and the posterior cervical lymph nodes along the anterior edge of the trapezius muscle and the supraclavicular lymph nodes in the supraclavicular fossa. Examination must also include other regions, in particular the axillary and inguinal lymph nodes, liver and spleen, in order to assess...
Table 2: Overview of lymph node regions of the neck

According to the AJCC Classification of cervical lymph nodes: A = Level Ia, B = Level Ib, F = Level II-IV; G = Level V; I = Level VI

| Region         | Drainage areas |
|----------------|----------------|
| Submental      | Bottom lip, tip of tongue, floor of mouth, skin of cheeks |
| Submandibular  | Tongue, submandibular gland, lips, mouth, conjunctiva |
| Pre-auricular  | Eyelids, conjunctiva temporal region, parotid, middle ear |
| Post-auricular | Auditory canal, auricle, scalp |
| Suboccipital   | Scalp, neck |
| Jugular        | Tongue, tonsils, parotid, auricle |
| Posterior cervical | Scalp and neck |
| Supraclavicular | Lungs, esophagus, abdomen (thoracic duct) |
| Paratracheal   | Trachea, thyroid gland |

Lesion of the thyroid

- **A**: Infectious processes of the oral cavity, nose, maxillary sinus of the face
- **B**: Infectious processes of the oral cavity, nose, maxillary sinus of the face
- **C**: Infectious processes of the ears, teeth, lid infections, conjunctivitis
- **D**: Infectious processes
- **E**: Infectious processes, toxoplasmosis, rubella, lymphomas
- **F**: Pharyngitis, rubella, cancer metastases, lymphomas
- **G**: Tuberculosis, melanomas, lymphomas
- **H**: Thoracic and retroperitoneal carcinomas, lymphomas, sarcoidosis, tuberculosis
- **I**: Lesion of the thyroid
systemic involvement. Palpation of the deep cervical lymph nodes is facilitated by the patient inclining their head forwards slightly to relax the strong muscles that support the head [4].

The conspicuous lymph nodes are palpated for tenderness, consistency and mobility in relation to the surrounding tissue/adjacent lymph nodes. For example, swollen lymph nodes originating from an infection are generally painful, soft, fluctuant if abscessed, while nodes associated with lymphoma are typically firm, rubbery and painless, and lymph node metastases from carcinomas hard and seldom mobile [5], [6].

If not just a single lymph node is conspicuous and enlarged, but several – possibly also on both sides – are affected, other aspects must be taken into account for differential diagnosis. If infectious in origin, bilateral enlargement tends to suggest a viral (e.g. Epstein-Barr virus = EBV) or group A streptococcal infection (e.g. in tonsillitis), while rubella or Erythema infectiosum (fifth disease) are more rarely the cause. Table 3 provides an overview of distribution.

The location of the lymph node may play an additional role in securing diagnosis: in infections of the scalp, the (sub-)occipital lymph nodes are often affected and therefore require careful inspection of the skin (beneath the hair). The (sub-)occipital lymph nodes are also enlarged in toxoplasmosis, inflammation of the outer ear and rubella. Enlarged pre-auricular lymph nodes may indicate infections of the eyes (e.g. lid infections, conjunctivitis), ears, teeth or the parotid gland. Submental/submandibular lymph nodes are often conspicuous in infectious processes of the oral cavity, the nose, the maxillary sinus or the face. Lymphomas may appear practically anywhere but nevertheless often chiefly affect the lymph nodes along the jugular vein and in the occipital or supraclavicular regions. In relation to the supraclavicular lymph nodes, differential diagnosis should consider sarcoidosis and the – albeit rare – existence of lesions in the thorax and gastrointestinal tract [6].

The age of the patient is a further criterion to be considered alongside location and palpation. Generally, the cause of lymph node swelling in young patients – from babies to infants and adolescents – is benign in over 80% of cases. This figure decreases considerably with age, to the extent that a malignant cause is found in over 60% of patients age 50 and above [7]. However, if there is a suspicion of malignancy in a young patient, the most frequent causes in children under 6 years of age are acute leukemia, neuroblastoma, rhabdomyosarcoma and non-Hodgkin lymphoma. Between the ages of 7 and 13, non-Hodgkin lymphoma and Hodgkin’s lymphoma are roughly equal, with rhabdomyosarcoma and thyroid cancer occurring more rarely. From the age of 13, Hodgkin’s disease is the leading malignant cause of neck masses during childhood and adolescence [8].

Clarifying whether a suspected malignancy exists is crucial to the further diagnostic process: a well thought-out and considerate approach to dealing with the concerned parents and the patient is essential in order to spare them any unnecessary (additional) worry while not trivializing a manifest suspicion. The doctor should assess the parents’ and patient’s need for information and be able to justify and communicate the next steps in the process. Conclusive evidence of malignancy is only possible by histological examination. In a study by Torsiglieri et. al, histological examination of 445 neck masses in children showed a malignant disease in 11% (n=48) of the cases [9]. The following criteria can be used to distinguish between malignant and benign:

Criteria more consistent with malignancy:
- over 2/2.5 cm in size
- firm/hard consistency
- supraclavicular/axillary localization, posterior edge of the sternocleidomastoid muscle
- absence of tenderness
- low mobility
- progressive course
- B symptoms

Criteria less consistent with malignancy:
- under 1.5 cm in size
- soft consistency
- tender
- erythema
- fluctuation
- mobility
- regressive course

Conclusion

Anamnesis and clinical examination raise the following cardinal questions, the answers to which are instrumental in determining subsequent procedure:

- Is the lymph node swelling clearly pathological?
- How long has the lymphadenopathy been apparent?
- How old is the child?
- Is the course progressive, constant, regressive or intermittent?
- What is the localization?
- Is the lymph node swelling on one or both sides, uni- or multifocal?
- Apart from “tumor”, are there other signs of inflammation such as “rubor”, “calor”, “dolor”, “functio laesa”?
- Are there any additional significant clinical findings?
- Is there any suspicion of malignancy?
Table 3: Origin, localization and course of disease

| Origin      | Side    | Course          | Rare                                                                 | Frequent                                                                 |
|-------------|---------|----------------|----------------------------------------------------------------------|----------------------------------------------------------------------------|
| Infectious  | unilateral | acute          | Staphylococcus aureus [A, B, F, G], anaerobic organisms [A, B, F]    | EBV [F, G], CMV [F, G], adenovirus [A, B, F, G], enterovirus, serogroup A streptococci |
|             | bilateral | chronic        | Non-tuberculous mycobacteria (M/MBC), bartonellosis                   | Arthritis, systemic lupus erythematosus, dermatomyositis, sarcoidosis,       |
|             |          |                | Serogroup B streptococci [A, B, F, G], tularemia [E, F, G, H],      | sarcoïdosis                                                                |
|             |          |                | Pasteurella multocida, Yersinia, gram-negative bacteria [A, B]       | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,    |
|             |          |                | Toxoplasmnosis, TBC [F, G, H], actinomycosis                         | Arthritis, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,     |
|             |          |                | Parovirus B19, rubella virus [E, J], HS, mycoplasma pneumoniae       | Arthritis, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,     |
|             |          |                | HIV, toxoplasmosis, TBC [F, G, H], syphilis                          | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |
|             |          |                | Chronic, granulomatous disease                                       | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |
|             |          |                | Hyper-IgM syndrome, Kawasaki syndrome [F-H],                          | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |
|             |          |                | Common variable immunodeficiency                                      | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |
|             |          |                | HHV-8, Langerhans cell histiocytosis, Castleman’s disease, Rosai-      | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |
|             |          |                | Dorfman disease, Kikuchi-Fujimoto disease [G]                         | Adenovirus, systemic lupus erythematosus, dermatomyositis, sarcoïdosis,   |

Langetal.: Cervicallymph node diseases in children
4 Further diagnosis

If the anamnesis and clinical examination prove inconclusive, or if confirmation of a suspected diagnosis is required, further diagnostic means are available, including serological tests, sonography as the main imaging technique, and for certain special indications also MRI and CT. As a means of obtaining histological confirmation, full dissection of a suspicious lymph node should take preference over fine-needle aspiration cytology (FNAC), compare 4.4.

Clinically it is neither reasonable nor cost-effective to routinely perform every one of the available serological and microbiological tests for every patient with lymph node swelling of uncertain origin. The same is true of the imaging and histological procedures. It is therefore a question of selecting a set of diagnostic measures that best meet the individual’s clinical presentation with the highest possible sensitivity and specificity. Table 4 gives an overview of clinical indications from anamnesis and examination, together with the possible cause and accompanying further diagnosis in each case.

4.1 Laboratory

Laboratory diagnosis is widely used in lymphadenopathy of uncertain origin. Parameters such as the erythrocyte sedimentation rate (ESR), C-reactive protein (CRP) and lactatedehydrogenase (LDH) are elevated in many infectious but also neoplastic and immunologic disorders and therefore tend to be nonspecific. They are nevertheless often determined in order to add a further aspect to the diagnosis.

A blood count including microscopic differential hemogram can further help to confirm a suspected malignancy. Once the diagnosis has been established, the laboratory tests are used to establish which organs are involved in systemic diseases (uric acid, creatinine levels for renal involvement e.g. in SLE; GOT, GGT for liver involvement e.g. in mononuclearosis; LDH and muscle enzymes e.g. in dermatomyositis) and as part of the follow-up and clinical monitoring (e.g. blood count for malignant tumors).

4.2 Microbiology

The success of microbiological diagnosis depends to a critical degree on the quality of the material submitted for testing. If this kind of diagnosis is not performed routinely on a regular basis and there is any doubt as to the sampling procedure, it is advisable to first contact the experts for further information. How the samples are transported (temperature, culture medium, aerobic/anerobic milieu) likewise has a decisive influence on diagnosis.

Serology

Serology is primarily used to diagnose virus-associated diseases. Pathogens such as rubella, measles, CMV and EBV are diagnosed serologically (in combination with the corresponding clinical assessment). Here it is always necessary to obtain the patient’s vaccination anamnesis.

(Swab) samples

Samples must be placed in a transport medium. The time and site of sample removal must be recorded, together with details of any antibiotic treatment already initiated. The role of swabs in cervical lymphadenitis is primarily to diagnose abscessed lymph nodes. *Staphylococcus aureus* (S. aureus) and *Streptococcus pyogenes* (S. pyogenes) are found in the majority of cases (40–80%) [10], [11], [12], [13], [14]. The proportion of meticillin-resistant *S. aureus* (MRSA) bacteria in childhood ENT infections is 10–30% [15], [16]. In a study by Neff and colleagues, swabs were taken intraoperatively from 277 immunocompetent children and cultures set up for aerobic, anaerobic, acid-fast pathogens and fungi [17]. The results showed none of the fungal cultures, 1% of the anaerobic and 2% of the acid-fast cultures to be positive. In the study, the cost of the swab cultures was put at 402 $ US dollars per patient (aerobic 94 $, anaerobic 108 $, acid-fast 100 $, fungal 96 $). MRSA pathogens were found in 38 patients (14%), 100% of which were sensitive to clindamycin. Neff and colleagues conclude that only aerobic cultures should be employed and acid-fast pathogens only cultivated if a strong clinical suspicion exists. The authors recommend empiric (single) antibiotic therapy effective against *S. aureus* and *S. pyogenes*.

Blood culture and other means of pathogen detection

Blood cultures are considerably less reliable if antibiotic treatment has already begun, since antibiotic substances continue to work in the culture bottles and may hinder pathogen growth. Full diagnosis includes cultures in an aerobic and an anaerobic medium and should be repeated at intervals of several hours to raise their diagnostic value. Blood cultures are primarily indicated in septic patients, fever of unknown origin and serious infectious diseases such as meningitis.

Other means of detecting pathogens include indirect material sampling (e.g. bronchoalveolar lavage, BAL) or testing secretions (sputum, gastric juice etc.), where appropriate combined with a polymerase chain reaction (PCR).

Tuberculin skin test

There are various methods of tuberculin testing: The *percutaneous* test is suitable for children under 10 years of age. It involves applying a salve containing tuberculin to the skin; the test is read after 72 hours [18]. In the *Tine test*, four needles treated with tuberculin are pressed into the skin for three seconds and moved briefly. This test is also read at the earliest 72 h after administration. Both tests are approximate screening methods –
Table 4: Clinical parameters and steps in differential diagnosis

| Cause                              | Clinical Presentation                                                                 | Diagnosis                                                                 |
|------------------------------------|----------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| **Infectious**                     |                                                                                        |                                                                           |
| Streptococci, staphylococci        | Local findings, signs of inflammation                                                  | Clinical assessment, if indicated swab                                     |
| Tuberculosis                       | Painless, often dense cervical lymph nodes, immune deficiency                          | Tuberculin test, histology, culture: sputum, gastric juice, biopsy         |
| Brucellosis                        | Contact with animals, consumption of unpasteurized milk, hepatosplenomegaly, fever, night sweats | Blood culture, serology                                                    |
| Bartonellosis (cat-scratch disease) | Contact with cats                                                                       | Clinical assessment, if indicated histology                                |
| Tularemia                          | Contact with animals (rodents), sudden onset of fever, abscesses at entry site with regional, often purulent, lymph node inflammation, tick bite | Blood culture, serology                                                    |
| EBV infection                      | Painful, mainly localized lymphadenopathy, general symptoms                              | Serology, abdominal sonography                                             |
| CMV infection                      | Hepatosplenomegaly, secondary hepatitis                                                | Blood count (lymphocytosis with atypical lymphocytes) CMV antibody IgM     |
| HIV infection/AIDS                 | HIV risk constellation, recurrent colds, rash                                           | HIV serology                                                              |
| Measles                            | Characteristic skin manifestations                                                     | Clinical assessment, measles serology IgM                                  |
| Rubella                            | Retroauricular/(sub-)occipital lymphadenopathy                                        | Clinical assessment, rubella serology IgM                                  |
| Toxoplasmosis                      | Contact with cats, immune deficiency, consumption of raw meat, 90-90% are otherwise asymptomatic | Toxoplasma antibody IgM                                                   |
| Leishmaniasis                      | Tropical anaemia, fever, splenomegaly                                                  | Pancytopenia                                                              |
| Lymphatic filariasis               | Fever                                                                                   | Eosinophilia, microfilaria in peripheral blood                             |
| Yersiniosis                        | Intestinal symptoms                                                                     | Serology                                                                  |
| Borreliosis                        | Anamnesis, erythema chronicum migrans, tick bite                                        | Borrelia serology IgM                                                     |
| **Immunologic diseases**           |                                                                                        |                                                                           |
| Lupus erythematosus                | Characteristic skin manifestations                                                     | Clinical criteria, ANA, anti-dsDNS, complement                             |
| Sarcoïdosis                        | Bilateral hilar mediastinal lymphadenopathy, fever, dyspnea                              | Histology                                                                 |
| Langerhans cell histiocytosis      | Mono- or multisystemic disease                                                         | Biotic findings                                                           |
| Arthritis                          | Swelling of the joints/bone and soft tissue pain                                       | Clinical criteria, rheumatic factors                                       |
| Kawasaki syndrome                 | Exanthema, cheilitis, fever                                                            | Clinical assessment                                                       |
| Angiofollicular lymph node hyperplasia | Mediastinal lymphomas, localized or multicentred                                      | Histology                                                                 |
| Histiocytic necrotizing lymphadenitis | Fever, (cervical) lymphadenopathy, night sweats                                      | Histology                                                                 |
| Dermatomyositis                    | Characteristic skin manifestations, proximal muscle weakness                            | Muscle enzymes, electromyography                                           |
| **Metabolic diseases**             |                                                                                        |                                                                           |
| Gaucher’s disease                  | Splenomegaly and neurological symptoms                                                 | Clinical assessment, histology                                             |
| Niemann-Pick disease               | Splenomegaly and neurological symptoms                                                 | Clinical assessment, histology                                             |
| Tangier disease                    | Enlarged, yellow and grey tonsils                                                      | Clinical assessment, histology                                             |
| Drug-induced                      | Drug anamnesis                                                                         | Anamnesis, serology                                                       |
| Serum sickness                     | Fever, fatigue, skin conditions, arthralgia, drug/vaccine exposure                     | Clinical assessment, complement                                            |
| **Neoplastic diseases**            |                                                                                        |                                                                           |
| Malignant lymphomas                | Indolent lymph nodes, B symptoms                                                      | Imaging, bone marrow aspiration, histology                                |
| Leukemias                          | Indolent lymph nodes, swollen joints/bone and soft tissue pain, increased bruising    | Changes in blood count, bone marrow aspiration                             |
| Metastases                         | Indolent lymphomas, tumor anamnesis, general symptoms                                 | Imaging, histology                                                        |
the percutaneous test in particular is considered unreliable – and are therefore not recommended [18]. The intracutaneous Mendel-Mantoux test is used to screen for suspected TBC by intracutaneously injecting tuberculin. The test is read after 72 hours; the induration must have a diameter of at least 6 mm to count as positive.

What is important is that all the tuberculin tests make no distinction between an acute or past infection and only confirm exposure to the disease. The test is also positive after BCG vaccination. Infection with non-tuberculous mycobacteria (NTM) may produce a false-positive test. Another diagnostic test for tuberculosis is the quanti-feron test. It is a serological investigation and – depending on the interferon-γ levels – evaluated as positive or negative. The advantages of the quanti-feron test are its higher sensitivity (75–80%) and its independence from the vaccination status [19].

4.3 Imaging

Imaging is used in cervical lymph node swelling in children to confirm or eliminate a specific diagnosis (e.g. abscess) or to assess the location and number of affected structures and can significantly contribute to diagnosis where clinical assessment and anamnesis are inconclusive. Particular attention must be paid here to the age of the patients, some of whom are very young, especially in terms of their exposure to radiation, e.g. during CT, the practicability and difficulty of the procedure, and stress.

Sonography

Sonography is a very valuable way of assessing lymph node swelling in children. In inflamed processes, sonography is the instrument of choice for determining localization, number, size and characteristics of the affected structures [17], [20], [21]. In the ultrasound, inflamed lymph nodes appear enlarged and are hypoechoic compared to the adjoining connective and muscle tissue [22]. Both abscess-forming lymph nodes and tuberculous lymphadenitis are characterized by central cystic changes with loss of the echogenic hilum [23].

Size, form, structure, intranodal necrosis, hilum structure, calcification and edema of the surrounding connective tissue are also assessed in non-infectious processes [24]. Doppler sonography additionally permits imaging of the vascularization of the hilum structure and the node. Ultrasound is not only valuable in helping to identify inflammatory, abscess-forming conditions but also and above all in assessing malignant processes. Here it is important to differentiate between suspected metastases from a solid tumor and – particularly in children – manifestations of other entities, such as lymphomas.

Ultrasound criteria that suggest a malignant process are changes in size and form [25]. The specificity and sensitivity here essentially depend on the defined limits [26]. In general, a L/D ratio (length/depth) of <2 is seen as an indication of a malignant process [24], although this is an unreliable criterion especially in the case of small lymph nodes [27]. The situation is similar with the hilum echogenity: 84–92% of benign lymph nodes have an echogenic hilum, while the hilum structures in 76–96% of malignant lymph nodes cannot be detected on account of intranodal processes [28]. It is essential to note in sonographic assessment that this is rather to be seen as a late sign and that the negative predictive value of early involvement of the lymph nodes is low [29].

The cortex configuration of the lymph node similarly allows to draw conclusions as to the dignity of the lymph node. It is considered in relation to the hilum structure and is suspicious if the cortex diameter is greater than half the diameter of the hilum [29]. A distinction must be made between concentric (bulging into the hilum) thickening of the cortex, which is considered nonspecific and may also occur in reactive changes in the lymph nodes [29], and eccentric (bulging outwards) cortical thickening, which is more likely an early sign of malignancy with a high positive predictive value [30].

The internal structure of the lymph nodes may also help to identify origin. For example, microcalcification and necrosis can be caused by infections (e.g. tuberculosis or other inflammatory granulomatous conditions) and neoplastic diseases (e.g. metastases from thyroid carcinomas). At the same time, the existence of necrosis is less consistent with lymphoma [31].

A valuable contribution to differential diagnosis has been made with ongoing improvements in the quality of resolution by intranodal angiography using Doppler sonography: neoplastic lymph nodes often exhibit pronounced hypervascularization, i.e. formation of new vessels, chiefly in the periphery of the lymph node but in some cases also beyond the lymph node capsule, which is characterized by irregular distribution and flow properties [32]. In lymphatic neoplasms, the periphery is rarely affected on its own due to their intranodal origin [33]. In the literature, the measurement of vascular resistance and contrast agent-supported sonographic imaging are still subjects of controversial debate in relation to sensitivity and specificity in differential diagnosis [27].

Computer tomography and magnetic resonance imaging

On a CT scan, inflamed lymph nodes appear enlarged, absorb contrast agent and are oval in shape, with purulent abscesses exhibiting a ring with no contrast agent in the centre and thickening on the periphery. By contrast with tuberculous lymphadenitis, fatty deposits can often be observed in the nodes [34], [35]. This ring-shaped accumulation of contrast agent is often referred to in eliminating a suspected abscess, but since it is apparent in over 60% of lymphadenitis cases, it cannot be considered highly sensitive [17]. Alongside exposure to radiation, this is a further reason why sonography should always be the preferred method of primary examination [20], [21].
A number of authors recommend computer tomography as the diagnostic instrument of choice in deep neck infections in order to achieve the best possible picture of the spread and origin of infection, monitor its course and help to assess the endangered mediastinal compartments [36]. A CT scan provides morphological evidence to verify a suspected malignancy, primarily by size, shape and, in advanced stages, infiltration of the surrounding structures.

Magnetic resonance imaging (MRI) is a further procedure available as part of more extensive diagnosis and can above all help to identify neck masses not originating in the lymph nodes. In this case – where necessary and possible – sedation is preferable to endotracheal anesthesia in children.

4.4 Histological examination

There are two main examination techniques to obtain histological confirmation of a diagnosis: Lymph node dissection and fine-needle aspiration cytology (FNAC). Full dissection of the affected lymph node is considered the standard procedure of choice when it comes to obtaining histological confirmation of a diagnosis, as it permits assessment of the histopathological structures and their distribution (e.g. vascular growth beyond the capsule). It also supplies sufficient tissue for pathological and microbiological examination. This makes it possible to perform more extensive immunohistochemical staining, which is essential to differential diagnosis of lymphomas in particular, and more detailed analysis of the suspected pathogen DNA by PCR.

Varying levels of sensitivity and specificity are also discussed in the literature in relation to fine-needle biopsy. In combination with sonography, this method produces accurate results with a high sensitivity (89–98%) and specificity (95–100%) in diagnosing malignant conditions [37], [38], [39]. Since the negative predictive value is approx. 80%, diagnosis is incorrectly eliminated in a fifth of cases [40]. This is particularly significant if only a single lymph node is affected, and FNAC causes a hemorrhagic lesion that hinders histological examination if dissection of the lymph node becomes necessary [6]. In this case, the experience and expertise of the examining physician and the cytopathologist are crucial [41]. Performing FNAC can be difficult anyway, particularly so in small children, and this is another consideration when deciding on procedure.

A fine-needle biopsy is an option if (i) several lymph nodes are suspicious, (ii) surgical dissection is associated with a high risk that can be eliminated by FNAC, (iii) sufficient expertise is available, and (iii) the patient is in compliance (age of the patient!).

Conclusion

Further diagnosis essentially builds on the anamnesis and clinical examination. Special laboratory tests should be conducted in particular if a specific clinical suspicion exists. Sonography is the first and standard choice of imaging procedure and provides answers to a range of questions in differential diagnosis. If a condition persists or malignancy is suspected, histological examination is warranted without too much delay, in which case the gold standard is full dissection of the affected lymph node.

5 Causes

The following chapter looks at the causes of cervical lymphadenopathy, divided according to infectious, immunologic, metabolic and neoplastic origin.

5.1 Infectious

Indications of lymphadenopathy of infectious origin may be specific (pathogen entry site, concurrent tonsillitis/pharyngitis) and unspecific (signs of inflammation, fever, elevated CRP, ESR). Viral and bacterial infections are the most common causes in childhood, while parasites or fungal infections tend to play a less frequent role. Generalized/bilateral lymphadenitis is more consistent with viral infection, unilateral with bacterial.

5.1.1 Viral pathogens

Pathogenesis and etiology: Viral infection is the most frequent cause of lymphadenitis in childhood [42]. Precisely the recurring viral pathogens that are associated with infections of the upper respiratory tract (RS viruses, influenza and parainfluenza viruses, rhino-, adeno- and acute corona viruses) [43], occur repeatedly in childhood and often cause acute cervical lymphadenitis. Other common viral agents, which can also cause chronic lymphadenitis, are the Epstein Barr virus, particularly in adolescence, and cytomegalovirus (CMV) [1]. Less frequently lymphadenopathy is also caused by mumps, measles, rubella, varicella, Erythema infectiousum or fifth disease, herpes simplex and Coxsackie viruses [44]. HIV-associated lymphadenopathy is far less common among children than adults, but depending on the risk profile it should also be considered as a potential cause in children and especially adolescents.

Clinical presentation: The main symptoms are defined by the infection of the upper respiratory tract: fever, pharyngitis accompanied by odynophagia, occasionally otitis/sialadenitis. These may be attended by cutaneous symptoms typical of the relevant viral infection.

Lymph nodes: The involved lymph nodes are usually affected multiply on both sides; without erythema/ fluctuation if there is no bacterial superinfection.

Diagnostics: If a specific suspicion exists, the diagnosis may additionally include virus serology for the relevant pathogen. If viruses that also affect the spleen and liver (e.g. EBV, CMV) are suspected, an ultrasound should always be performed on the abdomen to eliminate the possibility of hepatosplenomegaly.
Therapy: Immune competent patients receive supportive therapy, while prophylactic antibiotic or antiviral therapy is only indicated for immunocompromised patients or a severe clinical course [45]. If the symptoms persist, re-evaluation is necessary.

5.1.2 Bacterial pathogens

Pathogenesis and etiology: Bacterial infections also affecting the lymph nodes can likewise be divided into acute and chronic infections. The majority (40–80%) of acute bacterial infections are caused by staphylococci and beta hemolytic serogroup A streptococci [10], [11], [12], [13], [14]. Francisella tularensis (tularemia), pasteurella, Haemophilus influenzae type B and anaerobic organisms such as propionibacteria, fusobacteria and peptostreptococci are more rarely the cause of cervical lymphadenopathy [1]. Chronic bacterial infections of the lymph nodes are chiefly caused by non-tuberculous mycobacteria (NTM) [46], followed by other agents such as bartonella (cat-scratch disease), tuberculosis bacteria and brucella (brucellosis), cf. Table 5.

Clinical symptoms: The clinical symptoms are often comparable to an acute viral infection of the upper respiratory tract, accompanied by fever, odynophagia, otitis media or cough. Anaerobes may be apparent in connection with dental/gingival infections.

Lymph nodes: The lymph nodes in acute bacterial infections are typically painful, enlarged on one side and exhibit progressive growth. The predilection sites are the submental and submandibular, jugular and occipital lymph nodes [44].

Diagnosis: Diagnosis of bacterial lymphadenitis is based on the clinical symptoms and detection of pathogens in swabs/secretion by microscope, culture or PCR, serological procedures and evidence of antigens.

Therapy: The therapy is adapted to the relevant pathogen. The majority of acute bacterial infections are sensitive to a combination of benzylpenicillins + beta lactamase inhibitors. If empiric therapy does not lead to regression, other pathogens must be considered. In very young or immunosuppressed patients in particular, therapy may initially need to take place intravenously under inpatient supervision to avoid any serious complications.

5.1.3 Parasites/protozoa

Pathogenesis and etiology: If a parasitic or protozoa-associated lymphadenopathy is suspected, an anamnesis of foreign travel and environmental factors is of primary importance. Pathogens enter the body in food (unpasteurized milk, meat, contaminated water) or through wounds and close contact with animals. The neck and throat region, as the area associated with food intake, and the draining lymph node regions are predilection sites for manifestation of the diseases.

Clinical presentation: Parasitic infections are characterized by a deterioration in the patient’s general condition, organ involvement (toxocara (roundworm) → lungs, eyes; toxoplasmosis → eyes, lymphoreticular system; trypanosoma → heart, nerve cells; leishmaniasis → kidneys, intestine, skin; microfilaria → lymphoreticular system, heart, nerve cells) [47].

Lymph nodes: As part of the lymphoreticular system, the lymph nodes may be affected in parasitic infections.

Diagnosis: Serological procedure to determine antibodies or detect pathogens in a film of blood/secretion.

Therapy: Therapy is adapted to the relevant pathogen. Prophylaxis is important, particularly in preparing food and contact with animals. The recommended drug treatment for toxoplasmosis is a combination of pyrimethamine, sulfadiazine and folinic acid as well as corticosteroids [48]. Toxocara infections may be treated with Anthelmintics, if indicated by the symptoms [49]. There are several therapy options in the treatment of leishmaniasis, including N-methylglucamine antimionate, miltefosine, allopurinol, paromomycin, although a completely curative therapy is often not possible here [50].

5.1.4 Fungi

Pathogenesis and etiology: Histoplasmosis, coccidioidomycosis, blastomycosis and tinea are fungal infections caused by environmental contamination through the skin or the respiratory tract.

Clinical presentation: Characteristic symptoms are skin manifestations and lung involvement.

Lymph nodes: The lymph nodes are frequently affected as a result of organ involvement (lungs → supravacuicular; scalp → suboccipital [51])

Diagnosis: Serological tests, microscopic examination of swab and cultures are available means of diagnosis.

Therapy: The majority of infections require no treatment; an antifungal therapy may be necessary in the event of systemic complications and in immunosuppressed patients.

5.2 Immunologic

5.2.1 Granulomatous diseases

Sarcoidosis

Pathogenesis and etiology: Sarcoidosis (also known as Boeck’s disease, Schaumann-Besnier syndrome) is a systemic disease of the connective tissue. An increase in inflammatory activity with increased cellular immune response leads to the formation of non-caseating granulomas. The age peak lies between 20 and 40 [52], although the disease can occur in childhood. The cause of the increased immune response is not yet clear, but in epidemiological terms a genetic cause (BTLN2 mutation) appears at least to play a role [53]. Incidence of the disease in Germany is 10–12:100,000 inhabitants [54].

Clinical presentation: A distinction is made between the chronic and acute form (Löfgren syndrome) of the disease. Löfgren syndrome is characterized by the typical triad of symptoms: bilateral lymphadenopathy, polyarth-
| Pathogen | Acute | Chronic | Clinical presentation | Diagnosis | Histology | Therapy |
|----------|-------|---------|-----------------------|-----------|----------|---------|
| Staphylococcus and beta-hemolytic streptococci | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Swab, if necessary | Haemorrhages, neutrophil infiltration of the lymph node parenchyma | Benzylpenicillin + beta-lactamase inhibitor |
| Haemophilus influenzae type B | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Aspergillus, examination of secretion | Areas of necrosis, surrouned by palisade cells, neutrophils and granulomatous infiltrates | see above |
| Aerobic and anaerobic gram-negative bacilli, anaerobic gram-positive bacilli, and fusobacteria | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Propionibacterium and microaerobic bacteria | Areas of necrosis, surrouned by palisade cells, neutrophils and granulomatous infiltrates | see above |
| Pasteurella multocida | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | If indicated streptomycin, doxycycline, gentamicin | Necrotizing granulomas with star-shaped microorganisms | see above |
| Francisella tularensis (tularemia) | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | If indicated cotrimoxazole, doxycycline, quinolone | Necrotizing granulomas with star-shaped microorganisms | see above |
| Non-tuberculous mycobacteria (NTM) | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Serology | Necrotizing granulomas with star-shaped microorganisms | see above |
| Tuberculosis bacteria | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Serology | Necrotizing granulomas with star-shaped microorganisms | see above |
| Bartonella henselae | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Serology | Necrotizing granulomas with star-shaped microorganisms | see above |
| Brucella abortus | X     |         | Tender on palpation, usually unilateral lymph node enlargement (2-4 cm) | Serology | Necrotizing granulomas with star-shaped microorganisms | see above |
rthritis and erythema nodosum [55]. General symptoms such as fever, fatigue and arthralgia are also typical. After the lungs (90%) and the lymph nodes (90%), the liver (60–90%) and spleen (50–60%) are the most frequently affected organs. Others include the eyes (25%), heart (5%), the skeleton (25–50%), nerve tissue and bone marrow (15–40%) [52]. The clinical course of the disease depends on which organs are affected and the pulmonary stage. Scadding [56] distinguishes between 5 different clinical stages of pulmonary sarcoidosis:

- **Stage 0:** Lungs normal, other organ possibly affected (reversible)
- **Stage I:** Symmetrical lymph node enlargement, no involvement of lung tissue (reversible)
- **Stage II:** Bilateral lymph node enlargement with diffuse granulomas in lung tissue (reversible)
- **Stage III:** Lungs affected without lymph node enlargement (reversible)
- **Stage IV:** Pulmonary fibrosis with loss of function (irreversible)

In early stages and in the acute course of the disease, patients have a very good clinical prognosis with up to 90% spontaneous remission. However, around 5% of patients experience chronicity with a potentially fatal outcome, chiefly as a result of pulmonary or cardiac involvement [57].

**Lymph nodes:** In children, peripheral lymphadenopathy is the main symptom of sarcoidosis. Typically, the lymph nodes on both sides are fixed and firm on palpation. The supraclavicular lymph nodes are affected in over 80% of cases [1].

**Diagnostics:** The clinically asymptomatic course of the disease often means that sarcoidosis is discovered as an incidental finding of a chest x-ray. Depending on the clinical symptoms, further imaging may be necessary to ascertain or eliminate organ involvement. The available procedures are sonography, CT, PET-CT, MRI and the now rarely indicated scintigraphy [58].

Most laboratory parameters are unspecific. Elevated levels of angiotensin converting enzyme (ACE), erythrocyte sedimentation rate (ESR), immunoglobulins, calcium and calcitriol and a left shift in the blood count may be detected [59].

Diagnosis of lung function is worthwhile, and if the diagnosis is unclear, bronchoalveolar lavage with cytological examination to detect lymphocytic alveolitis with an increased CD4/CD8 ratio (>5 in acute sarcoidosis; normal 2).

Histological examination following lymph node dissection or transbronchial ultrasound-controlled biopsy is relatively unspecific, typically revealing non-caseating, epitheloid cell granulomas with Langhans giant cells and periiphery (monocytes, lymphocytes and fibroblasts) [60]. Diagnosis is therefore based on a combination of clinical signs and additive examinations.

**Therapy:** Symptomatic therapy is provided if organs are affected. Corticosteroids are the first choice of immunosuppressive medication for pharmacological treatment. Should they prove ineffective, azathioprine and chloroquine may also be used [61]. Therapy should not begin too early, since there is discussion of a higher recurrence rate following immunosuppressive therapy [62]. Tetracycline can be used for cutaneous symptoms [63]. Anti-inflammatory, analgesic and in some cases antiarthritic drugs (aspirin, ibuprofen, diclofenac) are the first choice of medication in treating Löfgren syndrome [64].

**Common variable immunodeficiency**

**Pathogenesis and etiology:** Common variable immunodeficiency (CVID) is a congenital immune defect with hypogammaglobulinemia and a resulting defective antibody response leading to recurring infections [65], [66], [67]. It is thought to comprise a heterogeneous group of genetic defects with effects on dendritic cells and on B and T cell function [68], [69].

**Clinical presentation:** Patients present with recurring infections, particularly of the upper respiratory tract. Many also develop organ-granulomas, disorders of the gastrointestinal tract (diarrhea e.g. caused by Giardia lamblia infections, enteroviruses) and skin manifestations (hair loss, vitiligo) [70]. Autoimmune phenomena such as idiopathic thrombocytopenic purpura (ITP) or autoimmune hemolytic anemia (AIHA) likewise occur with greater frequency [71].

**Lymph nodes:** Around one fifth of patients with CVID develop granulomas, 42% of them in lymph nodes [72]. Histological examination reveals non-caseating granulomas with multinucleated macrophages [73]. Differential diagnosis from sarcoidosis is considerably more difficult here, and particular attention must therefore be paid in the anamnesis to recurring infections, as these are typical of CVID [74].

**Diagnostics:** A serum protein electrophoresis is indicated if CVID is suspected. In the case of hypogammaglobulinemia, other possible causes should subsequently be eliminated (e.g. nephrotic syndrome, exsudative entero-pathy).

**Therapy:** The pathogenesis of the disease is such that only symptomatic therapy options are available, e.g. substitution of immunoglobulins, antibiotics, immunosuppressive therapy and bone marrow transplantation [75].

**Hyper-IgM syndrome**

**Pathogenesis and etiology:** Hyper-IgM syndrome is a rare (incidence 1:1,000,000 [76]) immune system disorder in which a malfunction prevents IgM switching to IgG [77]. This results in elevated (or normal) IgM levels and reduced IgG levels [78]. To date, five different genetic defects have been identified [79].

**Clinical presentation:** Recurring infections of the upper respiratory tract, paranasal sinuses and the ears are frequently exhibited in affected patients. In 40% of cases, the initial symptom is an opportunistic lung infection with pneumocystis carinii [76].
Lymph nodes: Swelling of the lymph nodes occurs frequently in the course of the recurrent infections of the upper respiratory tract. Some case descriptions refer to necrotizing granulomas and affected lymph nodes [80].

Diagnostics: Serum protein electrophoresis and genetic analysis are used in diagnosis.

Therapy: Possible therapy involves treatment of the opportunistic infections, immunoglobulin substitution and bone marrow transplantation [81].

Chronic granulomatous disease

Pathogenesis and etiology: Chronic granulomatous disease (CDG) is a rare genetic immune disorder (incidence 1:200,000 [82]) with dysfunction of the neutrophil granulocytes (NADPH oxidase) [83]. Chronic granulomatous disease has so far been divided into five different forms depending on the location of the genetic defect. The majority of patients present clinically in the first two years of life [84].

Clinical presentation: Frequent bacterial infections and fungal infection of the internal organs, particularly of the lungs, osteomyelitis, pulmonary inflammation and gastrointestinal complaints frequently manifest during childhood due to granuloma formation.

Lymph nodes: Over 50% of cases result in purulent inflammation of the lymph nodes (predominantly associated with staphylococci [82]).

Diagnostics: The differential blood count is normal, diagnosis is obtained by granulocyte function testing and genetic analysis. Imaging (sonography, CT) is necessary to eliminate an abscess or assess the affected organs (lungs, abdomen) [85].

Therapy: Exposure prophylaxis is important, particularly to mold spores. Surgical intervention is necessary if abscess formation occurs, combined with a pathogen-adapted antibiotic therapy. Granulocyte transfusions are only indicated in severe infections [86].

5.2.2 Rheumatic diseases

Juvenile idiopathic arthritis

Pathogenesis and etiology: Juvenile idiopathic arthritis (JIA) is a chronic inflammatory disease of unknown origin that affects the joints and begins in childhood. There is discussion as to whether it is triggered by e.g. an acute infection that elicits a chronic inflammatory response in patients with a polygenic predisposition. There are several different forms of juvenile idiopathic arthritis [87]:

- Juvenile chronic polyarthritis, adult type (positive rheumatoid factor)
- Juvenile ankylosing spondylitis
- Juvenile chronic arthritis, systemic onset (Still’s disease)
- Juvenile chronic arthritis (negative rheumatoid factor), polyarticular onset
- Juvenile chronic arthritis, oligoarticular onset
- (Juvenile) psoriatic arthropathy
- Other juvenile arthritis

Clinical presentation: The symptoms are common to all subforms of the disease: inflammation of one or more of the joints for longer than six weeks without identifiable cause, accompanied by rubor (rare), calor, dolor, tumor (swelling, effusion) and functio laesa (restricted mobility) [88].

Lymph nodes: Although joint problems are the main symptom, lymph node enlargement occurs in the majority of cases, particularly in the systemic forms of the disease [89], [90].

Diagnostics: Clinical findings, laboratory parameters and imaging are productive in diagnosis of JIA. In 2010, a set of common criteria of the rheumatological associations in the US (American College of Rheumatology; ACR) and Europe (European League Against Rheumatism; EULAR) was established in order to simplify early detection of rheumatic diseases [91]. Depending on the number of joints affected, the acute phase reaction (ESR or CRP) and other serological parameters (rheumatoid factors, anti-citrullinated protein/peptide antibodies; ACPA) and the duration of the arthritis, points are awarded (up to a maximum of 10). Rheumatoid arthritis is diagnosed at a total of six points and above.

Therapy: Treatment of rheumatoid arthritis involves dietary measures, physiotherapy, medication and interventions. A low-meat diet and regular exercise have a positive impact on the clinical course [92]. Four different groups of therapeutic agents are used:

1. Analgesics
2. Non-steroidal anti-inflammatory drugs (NSAIDs)
3. Glucocorticoids
4. Basic therapeutic agents, disease modifying anti-rheumatic drugs (DMARD)

Intervention options include (partial) arthroectomy, endoprosthesis, arthrodesis, and radiation synoviorthesis.

Systemic lupus erythematosus

Pathogenesis and etiology: Systemic lupus erythematosus (SLE) is a subform of lupus erythematosus, a collagen disease caused by autoantibodies and immune complexes (incidence 20–70:100,000) [93]. Many patients have a genetic predisposition, which may be triggered by factors such as EBV infection [94].

Clinical presentation: The clinical signs vary considerably depending on which organ is affected (kidneys, cardiac valves, bones, vessels, PNS and ZNS, lungs). General symptoms are fatigue, joint pain and sensitivity to light [95].

Lymph nodes: Lymphadenopathy presents in around 15% of cases of childhood SLE [96]. Histological examination shows zones of necrosis with haematoxylin inclusion bodies [97].
**Langerhans cell histiocytosis**

**Pathogenesis and etiology:** Langerhans cell histiocytosis (histiocyteosis X) describes a proliferation of dendritic cells from the bone marrow.

**Clinical presentation:** A distinction is made between three different clinical forms. The first – eosinophilic granuloma – describes the localized form of the disease. It is the most frequent manifestation of histiocytosis X (70%) and is characterized by its uncomplicated course. The acute and disseminated form of histiocytosis X, Abt-Letterer-Siwe syndrome, is the most severe (10%). Patients are usually under the age of 2, and without treatment the disease has 90% mortality. The third form, Hand-Schüller-Christian syndrome, may be localized or generalized and occurs more frequently over the age of 2. The clinical course of the disease depends on which organs are affected.

**Lymph nodes:** It is not unusual for lymphadenopathy to be the initial symptom of histiocytosis X [110]. Histological examination detects the Langerhans cells, which immunohistochemically stain positive for S100, vimentin and CD1a. X-shaped granula known as Birbeck granules are often detected by electron microscope [111].

**Diagnostics:** Diagnosis is confirmed histologically.

**Therapy:** Depending on the course the disease, aggressive chemotherapy or bone marrow transplantation may be necessary in some cases to improve prognosis [112].

**Hemophagocytic syndrome**

**Pathogenesis and etiology:** Hemophagocytic syndrome (HPS), or reactive hemophagocytic syndrome (RHS); macrophage activation syndrome (MAS); or lymphohistiocytic syndrome (LHS), occurs in a familial form of genetic origin and a secondary form in reaction to an infection, neoplasia or autoimmune disorder. Hypercytokinemia with multiple organ involvement is common to all forms [113].

**Clinical presentation:** Patients exhibit bouts of high fever, hepatosplenomegaly and lymphadenopathy, cutaneous manifestations and polyserositis. The fatality rate is high at up to 50% [114].

**Lymph nodes:** Moderate lymphadenopathy is not obligatory for diagnosis, but together with icterus, edema,
pericardial or pleural effusions is a supportive clinical symptom [115].

**Diagnostics:** Diagnosis of the syndrome should consider a possible genetic origin (some forms may be hereditary) and – usually infectious – trigger factors, as these affect both course and prognosis [116]. Henter and colleagues developed a revised list of criteria. Diagnosis is considered probable if five or more of the following clinical criteria are met: fever, splenomegaly, cytopenia of at least two cell lines, decreased hemoglobin, thrombocytopenia <100×10⁹/l, neutropenia <1×10⁹/l, hypertriglyceridemia (≥3 mmol/l) or hypofibrinogenemia (<1.5 g/l), hyperferritinemia (≥500 µg/l), soluble CD25 >2400 U/ml, hemophagocytosis of the bone marrow, the spleen or in lymph nodes, low or zero cytotoxicity of NK cells [117].

**Therapy:** The type of therapy depends on the cause and manifestation of the syndrome. In the primary form, curative therapy is possible by stem cell transplantation. In the secondary, the trigger factor must be treated. A possibility of immunosuppressive therapy with corticosteroids, cyclosporin A and the cytostatic etoposide also exists [115].

**Autoimmune lymphoproliferative syndrome**

**Pathogenesis and etiology:** Autoimmune lymphoproliferative syndrome (ALPS) is a disease affecting the reticuloendothelial system (RES) with a dysfunction of the FAS receptors and consecutive defect in apoptosis [118]. This results in increased survival time of the lymphocytes, which collect in the RES and produce clinical symptoms. **Clinical presentation:** The main clinical symptoms are triggered by the increased lymphoproliferation: lymphadenopathy and hepatosplenomegaly are present in the majority of patients and manifest early. The average age of clinical presentation is approx. 1 year. ALPS is often associated with autoimmune disorders or cancer (10%). **Lymph nodes:** The lymph nodes are affected in 90% of ALPS. Histologically the lymph nodes exhibit paracortical hyperplasia with expansion of the interfollicular regions and population with CD3+CD4-CD8- (double negative) T-cells [119].

**Diagnostics:** The diagnostic criteria were established by Straus and colleagues and distinguish between obligatory and supportive criteria [120].

**Therapy:** Therapy depends on the clinical symptoms and predominantly treats the associated autoimmune or malignant disorders. The first choice of therapy is with corticosteroids or immunoglobulins, the second mycophenolate mofetil, sirolimus or mercaptopurine [118].

**Castleman’s disease**

**Pathogenesis and etiology:** Castleman’s disease (or giant lymph node hyperplasia, angiofollicular lymph node hyperplasia) is a rare (incidence <1:100,000) disorder of uncertain origin with angiofollicular lymph node hyperplasia [121].

**Clinical presentation:** A distinction is made between the localized and the multicentric form. Half of patients with the localized form of the disease are clinically asymptomatic. Possible symptoms are fever, weight loss, fatigue and pain. Multicentric forms of the disease may also exhibit hepato- and/or splenomegaly or the so-called POEMS syndrome (polyneuropathy, organomegaly, endocrinopathy, monoclonal gammapathy, skin alterations) [122].

**Lymph nodes:** The lymph nodes are affected in >80% of cases, the main manifestations are abdominal, thoracic and cervical. A distinction is made pathogenetically and histologically between two subtypes: the hyaline vascular type with the onion-skin structure of the hyaline lymph nodes, usually occurring at a single site, and the plasmacytic type (rarer, multifocal) with plasma cell proliferation [123].

**Diagnostics and Therapy:** Diagnosis is confirmed histologically. The affected regions imitate both benign and malignant entities in imaging [124]. Dissection is simultaneously the therapy.

**Kawasaki syndrome (mucocutaneous lymph node syndrome)**

**Pathogenesis and etiology:** Kawasaki syndrome (also known as mucocutaneous lymph node syndrome, MCLS) is a necrotizing vasculitis (incidence in Germany 9:100,000 [125]). It predominantly affects small children (<5 years); it is thought to be of infectious origin from an unknown pathogen and affects patients with a genetic predisposition.

**Clinical presentation:** The disease typically progresses in three phases: feverish, subacute and reconvalescent. The main symptoms are persistent high fever over a period of 5–10 days, rashes in the oral cavity and on the lips, swelling, redness, eye involvement with conjunctival injection, skin involvement (exanthema), skin rash predominantly on the trunk, painful swelling of the hands and feet, and enlargement of the cervical lymph nodes [126]. The most serious complications are caused by secondary symptoms and are associated with cardiac involvement and the formation of aneurysms and thromboses of the coronary arteries on the one hand [127], and neurological symptoms including non-infectious meningitis and central hearing loss on the other. The lungs, intestine and joints may also be affected by the vasculitis [128].

**Lymph nodes:** The lymph nodes in the anterior cervical triangle are typically affected in the majority of patients by acute, non-purulent and relatively painless swelling. **Diagnostics:** The disease is diagnosed clinically (obligatory fever + 4/5 skin symptoms) and can be confirmed serologically. Echocardiography (cardiac involvement) and beginning therapy early are important [129].

**Therapy:** Initial treatment comprises high-dose intravenous immunoglobulins combined with high-dose aspirin. Further treatment options are available in the event of chronification, e.g. with pentoxifylline, abiciximab, infliximab and cyclophosphamide [130], [131], [132].
Kikuchi-Fujimoto disease

Pathogenesis and etiology: The pathogenesis of Kikuchi Fujimoto disease (lymphadenitis) is unknown to date. It primarily affects younger women [133], [134].

Clinical presentation: The symptoms are a combination of fever, lymphocytosis and necrotizing lymphadenitis as a result of stimulated apoptosis, mainly affecting the cervical and axillary lymph nodes. The disease is self-limiting, with remission after 1–4 months [135], in isolated cases however persisting for years. Recurrence is observed in rare cases (4%) [136].

Lymph nodes: Histology following lymph node dissection is critical to diagnosis: typical paracortical lesions with confluent zones of necrosis, atypical mononuclear cells, T-lymphocytes and histiocytes without granulocytes are exhibited. The lymph nodes are usually affected bilaterally, located in the posterior cervical triangle and are hard and tender.

Therapy: Therapeutic options are limited to symptomatic treatment (lowering fever, analgesia, intake of fluids) [135].

PFAPA syndrome

Pathogenesis and etiology: PFAPA syndrome (periodic fever, aphthous stomatitis, pharyngitis and cervical (lymph)adenitis) is similarly a disorder of unknown origin. It chiefly affects children under the age of five [137].

Clinical presentation: The clinical manifestations after which the disease is named may recur periodically in a 2–9 week cycle, spontaneously subsiding again after a few days [138].

Lymph nodes: The lymph nodes are enlarged on both sides.

Diagnostics: Diagnosis is based on the blood count, the level of enzyme activity and other means (e.g. sonography to detect hepatosplenomegaly).

Therapy: Miglustat as referred to earlier or so-called chaperone therapies are employed to reduce progression [144].

Tangier disease

Pathogenesis and etiology: The rare autosomal recessive inherited enzyme defect in Tangier disease affects a cholesterol transporter protein. It results in an accumulation of cholesterol deposits in the cells of the reticuloendothelial system [145].

Clinical presentation: Typical manifestations are yellow tonsils, with cholesterol deposits also found in other organs of the reticuloendothelial system, e.g. liver, spleen, in muscles and the nerve cell system [146].

Lymph nodes: The fat-storing macrophages in the affected lymph nodes appear as foam cells.

Diagnostics: Genetic analysis can confirm a clinical suspicion.

Therapy: No curative therapy exists. A low-fat diet is recommended prophylactically to reduce cardiovascular risk [147].
Amyloidosis

Amyloidosis is an umbrella term for the pathological deposition of certain proteins in the stroma. The symptoms are very varied depending on the localization and proteins deposited [148]. The lymph nodes may be affected by these deposits; histological examination shows the typical green birefringence under polarized light [149]. The therapy is adapted to the clinical findings.

5.3.2 Hypersensitivity

Serum sickness

Serum sickness is a Type III hypersensitivity reaction (Arthus reaction), as classified by Coombs and Gell, in which immune complexes are formed against exogenous antigen proteins and lead to activation of the complement system and damage to the surrounding tissue [150]. As formation of the antibodies must first be completed, in contrast to anaphylactic shock (Type I allergic reaction) it is usual here for days and weeks to elapse between exposure to the antigens and allergic reaction. This must be taken into account in the anamnesis of lymph node swelling (e.g. vaccination in the preceding months [151]) for differential diagnosis. Possible clinical symptoms are: swelling of the lymph nodes, fever, exanthema, arthralgia, circulatory deficiency. The therapy is adapted to the symptoms, where applicable watchful waiting, NSAIDs and corticoids. Vaccinations are not the only trigger for the aforementioned reactions, which are also possible to drugs and in particular antibiotics such as amoxicillin, cefaclor, cephalaxin and trimethoprim-sulfamethoxazole [152].

Adverse drug reactions must not necessarily be accompanied by the other symptoms of serum sickness but can also cause isolated lymphadenopathy [153]. The anamnesis should therefore include the following drugs: antiepileptics; heparin; tuberculostatics such as isoniazid, antibiotics such as cephalexin, sulindac [153].

A special form of the drug-induced systemic reaction with lymphadenopathy is DRESS syndrome (drug reaction with eosinophilia and systemic symptoms). This severe form involves multiple organ failure with 20% mortality [154]. Drugs associated with DRESS syndrome are allopurinol, minocycline, hydrochloride, anticonvulsants, sulfonamides and antibiotics. Generalized lymphadenopathy is exhibited in these cases.

5.4 Neoplastic

5.4.1 Malignant lymphoma

Non-Hodgkin lymphoma

Pathogenesis and etiology: Non-Hodgkin lymphomas (NHL) are a heterogeneous group of neoplastic disorders of the lymphatic cell line which can affect both B- (approx. 80%) and T-lymphocytes (approx. 20%). Their cause is considered to be hereditary or acquired genetic defects leading to a mismatch of proliferation and apoptosis of cells in the lymphatic system [155], [156].

Clinical presentation: Lymph node swelling and general symptoms such as fever, weight loss and night sweats are the predominant manifestations. Patients additionally often exhibit fatigue, weakness and changes in blood count. The stages are:

- Stage I: A single lymph node region above or below the diaphragm is affected,
- Stage II: Two or more lymph node regions above or below the diaphragm are affected,
- Stage III: Both sides of the diaphragm are affected,
- Stage IV: Organs other than primarily lymphatic organs are affected,

and appended: A = no general symptoms, B = with general symptoms, S = spread to spleen, E = extranodal spread.

The WHO divides NHL into precursor and mature B and T-cell lymphomas.

Lymph nodes: The swollen lymph nodes are usually firm and non-tender; swelling may occur on one or both sides.

Diagnostics: Histological procedures with special immune staining and bone marrow aspiration are necessary for diagnosis. Imaging is useful in staging the disease (chest x-ray, CT and sonography of neck, thorax and abdomen) [157].

Therapy: Treatment is according to the stage and entity and comprises chemo, radiation and antibody therapy [155].

Hodgkin’s lymphoma

Pathogenesis and etiology: Hodgkin’s lymphoma is similarly a neoplastic disease of the lymphatic cell line and is distinguished histologically from NHL by its characteristic Reed-Sternberg cells. There is discussion of the disease being of multifactor origin with genetic predisposition, associated with viral oncogenes (e.g. EBV) or following immunosuppressive therapies. It is possible to differentiate histologically between five different forms of the disease (nodular sclerosing, mixed cellularity, lymphocyte rich, lymphocyte depleted and lymphocyte predominant).

Clinical presentation: Hodgkin’s lymphoma patients often present with swollen lymph nodes and B symptoms. Hepatosplenomegaly or involvement of other organs (lungs, CNS) may additionally occur. The Ann Arbor classification distinguishes between four stages:
• Stage I: A single lymph node region is affected, 
• Stage II: Two or more lymph node regions on one side of the diaphragm are affected, 
• Stage III: Two or more lymph node regions on both sides of the diaphragm are affected, 
• Stage IV: Disseminated involvement of several extra-lymphatic organs with or without involvement of the lymph nodes, 

and appended: A = no general symptoms, B = with general symptoms, S = spread to the spleen, E = extranodal spread.

**Lymph nodes:** The swollen, rubbers, non-tender lymph nodes are frequently the initial noticeable symptom. Histological examination reveals Hodgkin and multinucleated Reed-Sternberg cells and multicolor reactive lymphocytic reactions [158].

**Diagnostics:** Anamnesis and clinical examination provide diagnostic indicators; staging is performed as in NHL with imaging, bone marrow aspiration and histological evidence.

**Therapy:** Chemotherapy, and if applicable radiotherapy, according to the stage of the disease are used [159], [160], [161].

### 5.4.2 Leukemias

**Acute lymphoblastic leukemia**

**Pathogenesis and etiology:** Acute lymphoblastic leukemia (ALL) is a neoplastic disorder of the lymphatic precursor cells. This leads to suppression of the bone marrow with consecutive insufficiency of the hematopoietic system. The early age peak (four years) means that ALL is particularly relevant among children and the most common malignant disorder in childhood [162].

**Clinical presentation:** Children present with anemia, weakness and exhaustion. They additionally have an increased bleeding tendency or increased susceptibility to infections. Lymph node swelling, hepatosplenomegaly, meningeosis or bone pain may be responsible for symptoms [163].

**Lymph nodes:** Around 50% of cases present with swollen lymph nodes.

**Diagnostics:** The laboratory tests are important in establishing diagnosis. Immunophenotyping by fluorescence-activated cell sorting (FACS) makes it possible to assign different grades of maturity, with the relevant consequences for prognosis and therapy [164].

**Therapy:** Therapy for ALL is divided into different phases, the duration and intensity of which depends on certain risk factors (detection of specific genes, CNS involvement). A distinction is made between the induction therapy and the subsequent consolidation therapy and the re-induction therapy and maintenance therapy [165]. Antibody therapies and inhibitor therapies are also employed.

**Acute myeloid leukemia**

**Pathogenesis and etiology:** Acute myeloid leukemia (AML) is a neoplastic disorder of the myeloid precursor cells. As in ALL, this results in a bone marrow insufficiency. AML is relatively rare in childhood and is responsible for around 20% of leukemias in this age group [166].

**Clinical presentation:** The symptoms are chiefly caused by the bone marrow suppression and are comparable to those of ALL (anemia, weakness, exhaustion, bleeding tendency, susceptibility to infections). Lymph node swelling, hepatosplenomegaly, meningeosis, bone pain, unilateral testicular swelling and gingival hyperplasia [167] are other common signs.

**Lymph nodes:** Involvement of the lymph nodes is frequently also found in AML.

**Diagnostics:** Diagnosis is based on morphology, immunophenotyping and cytogenetics. Bone marrow aspiration is indicated in addition to laboratory testing [167].

**Therapy:** Chemotherapy (induction and post-remission phase) and, if this proves ineffective, hematopoietic stem cell transplantation are available in the treatment of AML [168].

### 5.4.3 Lymph node metastasis from solid tumors

**Rhabdomyosarcoma**

**Pathogenesis and etiology:** Rhabdomyosarcoma is a highly malignant tumor of mesenchymal origin and responsible for around 5% of childhood tumor diseases [169].

**Clinical presentation:** Swelling and pain are the cardinal symptoms; other symptoms are possible depending on localization (e.g. hematuria if the bladder is affected).

**Lymph nodes:** Lymph node metastasis must be distinguished from a primary neck mass. Characteristic histological indicators are eosinophilic cells and multinucleate giant cells [170].

**Diagnostics:** Diagnosis is established by imaging (sonography, MRI) and histology.

**Therapy:** Therapy for rhabdomyosarcoma involves surgical excision with adjuvant radiotherapy. Neoadjuvant chemotherapy may be indicated for preoperative size reduction [171].

**Neuroblastoma**

**Pathogenesis and etiology:** Neuroblastoma is the third most common malignant disease in childhood and consists of proliferating cells of the neural crest called neuroblasts. The head and neck region is a common predilection site.

**Clinical presentation:** Neurological symptoms, B symptoms or abdominal complaints may arise depending on location. Classification is according to clinical aspects, taking into account localization, spread, lymph node involvement and distant metastasis [172].
Lymph nodes: Neuroblastoma often metastasizes in regional and distant lymph nodes.

Diagnostics: Staging examinations, bone marrow aspiration and histological examination confirm diagnosis [173].

Therapy: Depending on the stage, therapy options include dissection, (neo-)adjuvant chemotherapy, radiotherapy and stem cell transplantation as well as retinoids, arsenic trioxide and antibody therapies. A number of therapy options are associated with high comorbidity/comortality [174].

Nasopharyngeal carcinoma

Pathogenesis and etiology: Nasopharyngeal carcinomas are a rare tumor entity among children of western industrial nations (approx. 1% of malignant tumors) [175]. By contrast, the incidence of the disease is rising rapidly among patients from Asiatic countries [176].

Clinical presentation: Nosebleeds and nasal breathing difficulties and neck masses from lymph node metastasis are classic symptoms of a nasopharyngeal carcinoma.

Lymph nodes: Metastasis from the carcinomas is lymphogenous, rarely hematogenous. A distinction is made between keratinizing, well-differentiated squamous cell carcinomas, non-keratinizing squamous cell carcinomas, and undifferentiated lymphoepithelial carcinomas (Schmincke-Regaud).

Diagnostics: Diagnosis is confirmed by clinical examination, imaging (MRI) and histology.

Therapy: Therapy is dependent on the localization and staging of the disease, primary radiochemotherapy is generally indicated.

6 Conclusion

Cervical lymph node swelling of uncertain origin in childhood can pose a special challenge to the attending ENT and pediatric physicians. Faced with the spectrum of possible differential diagnoses and malignant diseases, they must follow a clearly structured algorithm in their diagnostic steps and therapeutic strategies to avoid unnecessary delays in further diagnosis and safeguard against overhasty, and possibly too invasive, diagnosis and therapy. This paper is intended to support them in these efforts.

Notes

Competing interests

The authors declare that they have no competing interests.

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